Population diversity among Brazilian isolates of *Cryphonectria cubensis*

L.M. van Zyl\(^a\)*, M.J. Wingfield\(^a\), A.C. Alfenas\(^b\), P.W. Crous\(^c\)

\(^a\) Tree Pathology Cooperative Programme, Department of Microbiology and Biochemistry, University of the Free State, P.O. Box 339, Bloemfontein, South Africa

\(^b\) Departamento de Fitopatologia/Bioagrio, Universidade Federal de Viçosa, 36570 Viçosa MG., Brazil

\(^c\) Department of Plant Pathology, University of Stellenbosch, Stellenbosch 7600, South Africa

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Abstract

The devastating effect of *Cryphonectria* canker on *Eucalyptus* spp. in Brazil has necessitated the initiation of a strategy to control the disease. A potential biological control strategy for *Cryphonectria cubensis* would be to make use of dsRNA, capable of debilitating the fungus. However, success would depend on the diversity of the pathogen population. In this study, the presence of vegetative compatibility groups (VCGs) in *C. cubensis* was demonstrated by pairing isolates and observing the reaction along the line of contact between the expanding colonies. Isolates (1092) of *C. cubensis*, from 182 *Eucalyptus* trees located in five regions in Brazil, were examined using this method. On a tree, one or at most two VCGs were present. However, isolates from different but closely spaced trees, represented numerous VCGs. It is probable that the multiplicity of VCGs in the Brazilian population of *C. cubensis* will limit the natural spread of dsRNA and, hence, limit the potential for biocontrol via hypovirulence. The large number of VCGs is also indicative of a well-established, and possibly endemic population of the pathogen. © 1998 Elsevier Science B.V. All rights reserved.

Keywords: dsRNA; Hypovirulence; Vegetative compatibility groups

1. Introduction

*Cryphonectria cubensis* (Bruner) Hodges is a notorious stem canker pathogen of *Eucalyptus* trees (Bruner, 1916; Boerboom and Maas, 1970; Hodges, 1980; Wingfield et al., 1989). It is prevalent in many tropical areas of the world and, particularly, between the latitudes 30\(^\circ\) north and south of the equator (Hodges, 1980; Old et al., 1986; Conradie et al., 1990). *Cryphonectria cubensis* has limited the establishment of seedling plantations of susceptible *Eucalyptus* spp. in areas where climatic conditions favour disease development (Hodges et al., 1979; Alfenas et al., 1982; Sharma et al., 1985). Humid conditions, high rainfall (>1200 mm year\(^{-1}\)), and temperatures greater than 23\(^\circ\)C or higher (Hodges et al., 1976; Sharma et al., 1985) are needed for the growth and spread of the pathogen. In cooler or drier conditions, the infection rates and the extent of canker development are much lower (Hodges et al., 1979).
C. cubensis was reported for the first time in South Africa in 1989 (Wingfield et al., 1989). Clonal forestry, which is now widely practised with Eucalyptus spp. in this country, gives rise to large, genetically uniform stands of Eucalyptus trees that will be at risk from pests and diseases, including C. cubensis. (Conradie et al., 1992). Current means of reducing losses due to the disease in the forestry areas of South Africa and Brazil, is through the plantings of disease-tolerant Eucalyptus species and clones (Conradie et al., 1990). A search for alternative methods of controlling this disease are thus being sought.

One means to reduce the impact of a tree pathogen is through the use of viruses or virus-like agents, which are capable of debilitating it. Such hypovirulence has been used successfully for limiting canker expansion and mortality of the American chestnut [Castanea dentata (Marsh.) Borkh.], due to infection by C. parasitica (Biraghi, 1950a, b; Anagnostakis and Day, 1979; Anagnostakis and Waggoner, 1981; Choi and Nuss, 1992; Chen et al., 1996). This system involves the conversion of a virulent (V) strain to hypovirulence (H) when the cytoplasmic determinants (dsRNA) of hypovirulence are transferred during hyphal anastomosis. The dsRNA is associated with the down-regulation of specific fungal genes (Powell and Van Alfen, 1987a, b), reduced sporulation and virulence, and changes in colony morphology (Anagnostakis and Day, 1979).

Transfer of dsRNA from hypovirulent to virulent isolates of C. parasitica is favoured when strains are of the same VCG (Anagnostakis, 1977; Glass and Kuldau, 1992; Leslie, 1993). Spread of hypovirulence could, therefore, be limited by vegetative incompatibility, although H strains that can convert V strains belonging to several already defined VCGs, termed networks (Anagnostakis, 1983), or conversion groups (Kuhlman et al., 1984) have been identified. Vegetative incompatibility is the inability of genetically different strains of a fungal species to fuse and form heterokaryons (Puhalla and Anagnostakis, 1970; Anagnostakis, 1988), and it results when different alleles are present at one or more vegetative incompatibility loci in the hyphae attempting to fuse (Anagnostakis, 1988).

The presence of a large number of VCGs in C. parasitica, within small geographical areas in the United States, has severely limited the natural spread of hypovirulence factors and, therefore, reduced their usefulness as biological control agents for chestnut blight (Anagnostakis et al., 1986; MacDonald and Fulbright, 1991). In a preliminary study, Van der Westhuizen et al. (1993) suggested that isolates of C. cubensis in South Africa represent a uniform population structure which is indicative of an introduced pathogen. This would facilitate spread of dsRNA which has also recently been discovered in the population (Van der Westhuizen et al., 1994).

Although C. cubensis has been known in Brazil for a considerable time, almost no information is available concerning the diversity of its population in that country. It has, however, previously been suggested that the fungus is native to Indonesia (Bruner, 1916; Hodges et al., 1986), thus implying that it was introduced into Brazil. The aim of this study was to determine the genetic variability amongst isolates of C. cubensis from Brazil. This would then provide clues as to the origin of the fungus in Brazil, and also whether dsRNA-mediated hypovirulence might be a means to reduce the impact of the disease in that country.

2. Materials and methods

2.1. Isolates and cultures

A survey of C. cubensis in Eucalyptus growing areas of Brazil (Table 1) was conducted during April 1993. One piece of bark per tree, showing characteristic disease symptoms, was collected. All bark pieces were surface-disinfested with 90% ethanol. These

<table>
<thead>
<tr>
<th>Region</th>
<th>Total number</th>
<th>Isolates</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aracruz</td>
<td>12</td>
<td>72</td>
</tr>
<tr>
<td>Teixeira de Freitas Nursery</td>
<td>113</td>
<td>678</td>
</tr>
<tr>
<td>Teixeira de Freitas Plantation</td>
<td>32</td>
<td>192</td>
</tr>
<tr>
<td>Mogi Guacu Nursery</td>
<td>22</td>
<td>132</td>
</tr>
<tr>
<td>Mogi Guacu Plantation</td>
<td>3</td>
<td>18</td>
</tr>
</tbody>
</table>
bark samples were kept moist in petri dishes containing wet filter paper and incubated at 25–27°C for 3 weeks. This was done to induce production of pycnidia and perithecia. Most of the perithecia on bark samples began to produce ascospores within 7–14 days after incubation. Ascospore masses from six perithecia per bark sample were transferred to sterile 9 cm diameter petri dishes containing 15 ml of 2% malt extract agar (MEA) (20 g malt extract, Biolab; 20 g agar, Biolab; 1 l distilled water) and incubated at 25–27°C. Isolates were stored on MEA slants in screw-capped tubes at 4°C.

2.2. VCGs per perithecium

Ascospore masses from 10 perithecia per piece of bark from each tree in the five regions were selected. Ascospores were washed onto petri dishes containing 2% water agar (20 g agar, Biolab; 1 l distilled water) and spread across the medium surface. Plates were incubated for 19–24 h at 25°C. Twenty single-germinated ascospores were then isolated from each plate with the aid of a dissecting microscope and sterile syringe needle, and incubated at 25°C in petri dishes containing 2% MEA.

Tests for determining VCGs were made by transferring small pieces of agar containing mycelium (<3 mm in diameter) from the edges of rapidly growing single ascospore cultures, less than 7 days old, onto oatmeal agar in square, sterile assay plates (30 cm²). Oatmeal agar was prepared by adding 100 g oatmeal to 1 l distilled water, steaming for 2 h and then sieving through cheese cloth before sterilizing at 120°C for 20 min. Each isolate was paired with itself and with each of the other isolates. Mycelial plugs were positioned, not greater than 5 mm apart on the surface of plates, which were then sealed with Parafilm and incubated at 25–27°C in the dark for 5–7 days. Two replicates of each plate were made.

The presence of VCGs was determined using the method described by Anagnostakis (1977). At the end of the incubation period, mycelia that were vegetatively compatible had merged, forming a confluent mycelium. Incompatible mycelia had grown to a meeting point on the agar, but remained separated by a 'barrage-like' reaction formed along the line of contact between paired colonies (Fig. 1).

2.3. VCGs per tree

The number of VCGs present on each tree was determined using ascospore masses from six perithecia per bark sample. The technique of Anagnostakis (1977) was used as described previously. In this study, however, 9 cm diameter, sterile petri dishes were used rather than the assay plates utilized previously.

2.4. VCGs per region

In regions, where more than 20 trees were sampled, isolates from six randomly selected trees were paired against each other. Initial screening of isolates was done, and those with the same VCG were excluded in order to reduce the number of isolates used. Twenty isolates were then paired against each other using the 30 cm² assay plates, in order to determine the total number of VCGs for each region.

3. Results

3.1. VCGs per perithecium

At the end of the incubation period, mycelia from cultures originating from ascospores from single perithecium had merged, forming a confluent lawn of mycelium. This indicates the existence of only one VCG amongst all ascospores in a perithecium. None of the 20 single ascospore cultures from each of the
Table 2
Genetic variability per C. cubensis canker in Brazil

<table>
<thead>
<tr>
<th>Region</th>
<th>Total number</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Trees</td>
<td>Isolates</td>
<td>Trees with single VCG</td>
<td>Trees with &gt;1 VCG</td>
</tr>
<tr>
<td>Aracruz</td>
<td>12</td>
<td>72</td>
<td>12</td>
<td>0</td>
</tr>
<tr>
<td>Teixeira de Freitas Nursery</td>
<td>113</td>
<td>678</td>
<td>103</td>
<td>10</td>
</tr>
<tr>
<td>Teixeira de Freitas Plantation</td>
<td>32</td>
<td>192</td>
<td>24</td>
<td>8</td>
</tr>
<tr>
<td>Mogi Guacu Nursery</td>
<td>22</td>
<td>132</td>
<td>22</td>
<td>0</td>
</tr>
<tr>
<td>Mogi Guacu Plantation</td>
<td>3</td>
<td>18</td>
<td>3</td>
<td>0</td>
</tr>
</tbody>
</table>

Vegetative compatibility groups (VCGs) were determined by placing isolates in close proximity on freshly prepared oatmeal agar plates.

Table 3
Total number of vegetative compatibility groups per region in Brazil

<table>
<thead>
<tr>
<th>Region</th>
<th>Total number</th>
<th>VCG/region</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aracruz</td>
<td>12</td>
<td>7</td>
</tr>
<tr>
<td>Teixeira de Freitas Nursery</td>
<td>113</td>
<td>86</td>
</tr>
<tr>
<td>Teixeira de Freitas Plantation</td>
<td>32</td>
<td>28</td>
</tr>
<tr>
<td>Mogi Guacu Nursery</td>
<td>22</td>
<td>18</td>
</tr>
<tr>
<td>Mogi Guacu Plantation</td>
<td>3</td>
<td>3</td>
</tr>
</tbody>
</table>

Vegetative compatibility groups (VCGs) were determined by placing isolate inoculum in close proximity on freshly prepared oatmeal agar plates.

3.2. VCGs per tree

Trees in the Aracruz Plantation, Mogi Guacu Nursery and Mogi Guacu Forest regions, possessed only one VCG per tree (Table 2). Most of the trees in the Teixeira de Freitas Nursery and Teixeira de Freitas Plantation regions, had been colonized by a single VCG of C. cubensis, with the exception of 10 trees from the Teixeira de Freitas Nursery and eight from the Teixeira de Freitas Plantation. However, none of these 18 trees were colonized by more than two VCGs. It would thus appear that most of the isolates from a given canker were in the same VCG. Of the 182 isolates from the same number of trees screened in this experiment, there were only 18 trees from which isolates of differing VCGs were recovered.

3.3. VCGs per region

Based on results of the study to determine the number of VCGs per tree, it was reasonable to use a single isolate from each tree to determine the number of VCGs per region. Results (Table 3) showed that at Aracruz, seven VCGs were present amongst 12 isolates sampled from the same number of trees. In the Teixeira de Freitas Nursery, 86 VCGs were identified among 113 isolates sampled from the same number of trees and very few VCGs occurred more than once in this region. In the Teixeira de Freitas Plantation, 30 VCGs were identified among 32 isolates sampled; 18 VCGs were identified from 22 isolates at Mogi Guacu Nursery and each of the three isolates from the Mogi Guacu Plantation area possessed different VCGs.

4. Discussion

Results of this study have clearly shown that C. cubensis is represented by a large number of VCGs in Brazil and, thus, have a very diverse population. These results might imply that the fungus undergoes a high level of sexual outcrossing which is known to give rise to new VCGs (Leslie, 1993). Certainly, the evidence available suggests that the fungus has been present in the area for an extended period of time.

Hodges et al. (1976) and Hodges et al. (1979) have provided conclusive evidence for the fact that C. cubensis is homothallic and thus able to give rise to perithecia from single ascospores or conidia. The fact that we found that all ascospores arising from single perithecia represent the same VCG, confirms these findings. These authors further suggested that most perithecia on stem cankers have arisen without sexual outcrossing. This result is in contrast to our findings that the population of the fungus in Brazil is extremely diverse, and that this diversity has most probably
arisen from sexual outcrossing. It is possible that sexual outcrossing is a relatively rare event but that it is at least sufficient to ensure a great diversity in the population of the pathogen. In the case of *C. parasitica*, Anagnostakis (1977) showed that the fungus is homothallic but preferentially outcrossed. The fact that all perithecia of *C. cubensis*, tested in this study, had ascospores representing single VCGs indicates that homothallism is common. A degree of outcrossing must, however, clearly be occurring to give rise to the great diversity of VCGs detected. The fact that these outcrossing events were not detected in this study is probably due to the fact that they are not as common as homothallism and may also be due to the relatively limited sample that was studied.

Almost every canker on a tree was caused by a single VCG of *C. cubensis*. This indicates that these infections arose from single propagules, which were probably ascospores. Ascospores in *Cryphonectria* are windborne and would presumably be dispersed much greater distances than conidia that are associated with moist conditions (Rankin, 1914; Heald et al., 1915; Newhouse and MacDonald, 1991). The predominance of perithecia on cankers also suggests that ascospores are the major propagule of dispersal in Brazil. This is in contrast to the situation in South Africa where perithecia of *C. cubensis* are almost never seen (Van der Westhuizen et al., 1993).

The relative number of VCGs in populations of Ascomycetes has been used in various studies to give an indication as to whether a pathogen has been recently introduced into an area, or whether it has been present for an extended period of time (Adams et al., 1990; Glass and Kulda, 1992). A great diversity of VCGs can presumably also arise in exotic situations from multiple introductions and sexual outcrossing, and results of such studies must be interpreted with caution (Perkins, 1987; Kies and Casselton, 1992). It has been suggested in the past that *C. cubensis* is most likely native on cloves [*Syzygium aromaticum* (L.) Merr. and Perry] in Indonesia, and that it was then introduced into Brazil (Hodges et al., 1986). This hypothesis is largely based on the assumption that the clove pathogen *Endothia eugeniae* (Nutman and Roberts) Reid and Booth is conspecific with *C. cubensis* (Hodges et al., 1986). Results of the current study at least show that *C. cubensis* is very well established in Brazil, and that it has very likely been present in the country for an extended period of time. Our view is that insufficient evidence exists to suggest that the fungus originated in Asia, and that further studies on the population diversity of the pathogen in both areas are required before this matter can be resolved.

The great number of VCGs in populations of fungi, and the lack of predominance of one particular VCG, as found in the Brazilian population of *C. cubensis*, may reduce the opportunities of biological control through hypovirulence. Work done on *C. parasitica* in North America showed that vegetative incompatibility reduces the frequency of transmission of dsRNA associated with hypovirulence from infected hypovirulent isolates to non-infected virulent isolates (Caten, 1972; Jaynes and Elliston, 1980; Anagnostakis et al., 1986; Anagnostakis and Kranz, 1987; MacDonald and Fulbright, 1991). The situation in Europe differs from that in North America in that biological control of *C. parasitica* through hypovirulence is successful, due to minimal diversity in the pathogen population (Anagnostakis et al., 1986; Heiniger and Rigling, 1994). Occasional conversion between some hypovirulent and virulent isolates that are of different VCGs is, however, possible (Anagnostakis, 1977; Jaynes and Elliston, 1980).

The considerable genetic diversity amongst the Brazilian population of *C. cubensis* suggests that the selection of resistant *Eucalyptus* clones will be complicated. An ongoing screening of the fungal population will have to be undertaken regularly, in order to identify the most virulent isolates. These virulent isolates will have to be integrated into an ongoing selection programme for disease tolerant *Eucalyptus* clones.

5. Conclusion

Results of this study have shown that *C. cubensis* in Brazil represents a diverse population. There is certainly good evidence to suggest that the population is endemic in that part of the world, although this matter deserves further investigation. Here, we would favour more extensive sampling, increase the sample size and the application of additional techniques to measure genetic diversity in isolates. The great diversity in the population of *C. cubensis* in Brazil also suggests that natural biological control through hypovirulence is not
likely to be effective in reducing the impact of the pathogen in that part of the world.

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References


