A PATHOGENIC FUNGUS ASSOCIATED WITH PLATYPUS ATTACK ON NEW ZEALAND NOTHOFAGUS SPECIES

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ABSTRACT
The hypothesis that Platypus-associated mortality of Nothofagus spp. is a direct result of the invasion of the sapwood by a fungal pathogen was tested by drilling holes in living red beech (Nothofagus fusca (Hook. f.) Oerst.) trees to simulate Platypus tunnels, and inoculating these with the suspected pathogen — a Sporothrix species. Other trees were inoculated with sterile distilled water and with a Platypus ambrosia fungus, Endemycopsis platypodis Baker et Kregervan Rij. All the Sporothrix-inoculated trees and one E. platypodis-inoculated tree wilted and died. No trees inoculated with sterile water died. Sporothrix sp. was recovered from well above the inoculated zone in all the dead trees, including the E. platypodis inoculated tree.

INTRODUCTION
Mortality in New Zealand beech (Nothofagus spp.) was initially suspected to be caused by the beech buprestid Nascoides enysi Sharp (Cockayne, 1926; Morgan, 1966). However Dugdale (1965) observed that pinhole borers (species of Platypus Herbst) attacked weakened beech 1-2 years before N. enysi and suggested that a pathogenic fungus might be introduced by Platypus. More recently Milligan (1972) ruled out N. enysi as an agency in beech mortality and showed that Platypus attack can kill healthy Nothofagus fusca. On the basis of his experiments and observations, Milligan (1972; 1974) suggested that a pathogenic sapstain fungus is transmitted by Platypus beetles and becomes established initially in the innermost sapwood where the moisture content of the wood is lowest; he stated "Nothofagus mortality previously attributed to N. enysi is now more convincingly interpreted as a consequence of Platypus attack" (Milligan, 1974 p. 35).

The first step in testing these hypotheses and observations was to isolate suspected pathogens. Isolations from adults and tunnels of Platypus apicalis White, and from around fresh holes in living trees, included yeasts, a Ceratocystis sp., and other fungi (W. Faulds, unpubl.). Tests of the pathogenicity of Ceratocystis sp., yeast, Trichoderma sp., and a species of bacteria failed to show any such pathogenicity.

Inoculation tests (Faulds, 1973) to determine whether stains associated with Platypus attack in N. fusca were a response by the tree to mechanical wounds only or a response
to micro-organisms which invade these wounds clearly showed the stains were due to the presence of micro-organisms and were probably a response to toxic substances produced by them. One of the fungi used in these tests was that most frequently isolated from near Platypus tunnels in one of Milligan's experimental trees killed by induced Platypus-attack. Trees showed greater response around wounds inoculated with this fungus than with any other treatment and the fungus was recovered further from inoculation wounds than other micro-organisms. It appeared to be capable of, and well adapted to, invading live tree tissue. The conidial stage of the fungus has been identified as a Sporothrix sp. (P. Gadgil, pers. comm.). This fungus became the primary pathogen suspect and inoculation tests were undertaken to determine its pathogenicity. The locality for these tests was Kaimanawa State Forest Park (N.Z.M.S. 1. Rangitaiki, N103. 700907 (Department of Lands and Survey, 1969)).

**EXPERIMENT 1**

**Materials and Methods**

Seven apparently healthy *N. fusca*, with no visible signs of *Platypus* attack, were selected for this experiment. These trees were marked A, B . . . G and their diameters were recorded (Table 1). During March and April 1972, holes drilled in A, B, E, F, and G were inoculated with *Sporothrix*, and in C and D with sterile distilled water. The inoculum was prepared and stored as described in an earlier paper (Faulds, 1973).

The experimental treatment for each tree was as follows. A work platform was erected around the tree (Fig. 1) then a band approximately 30 cm wide completely encircling the tree was marked at a height of 1-2 m from the base of the stem (*Platypus*

![FIG. 1—Drilling and inoculation operation on work platform.](image-url)
attacks are concentrated on the lower 6 m of stem; Milligan, 1974). Loose bark was
removed from this marked area with a bark scraper. A small area in which two or three
holes were to be drilled was swabbed with 95% alcohol. The drill was cleaned with a
wire bush, dipped into a jar of 95% alcohol and flamed, and a 2.5-mm hole was drilled
until a change in the noise of the drill indicated the heartwood had been reached. A
hypodermic syringe, washed with alcohol and rinsed with sterile distilled water, was
used to inject about 1 ml of inoculum into the hole. Finally the hole was plugged with
a sterile cotton wool bung. After 2-3 holes had been drilled in the swabbed area, a
fresh area was swabbed and the process repeated. Holes were drilled randomly within
the marked area with a drill powered by a portable generator. Holes of this diameter
and depth are similar to those made by Platypus beetles (Faulds, 1973). The total
number of holes drilled and inoculated in each tree, and the number of holes per
100 cm² in the treated area are shown in Table 1.

<table>
<thead>
<tr>
<th>Tree Treatment</th>
<th>Diameter of tree at drilled zone (cm)</th>
<th>Width of treated band (cm)</th>
<th>Total holes drilled</th>
<th>Holes/100 cm² of treated band (average)</th>
<th>Time from treatment to complete wilt (months)</th>
<th>Condition of tree at felling</th>
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<tbody>
<tr>
<td>EXPERIMENT 1</td>
<td></td>
<td></td>
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<tr>
<td>A Sporothrix</td>
<td>31</td>
<td>34</td>
<td>617</td>
<td>18</td>
<td>9</td>
<td>Completely wilted</td>
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<tr>
<td>B Sporothrix</td>
<td>43</td>
<td>34</td>
<td>974</td>
<td>21</td>
<td>9</td>
<td>Completely wilted</td>
</tr>
<tr>
<td>C Sterile water</td>
<td>43</td>
<td>36</td>
<td>1069</td>
<td>22</td>
<td>—</td>
<td>Healthy</td>
</tr>
<tr>
<td>D Sterile water</td>
<td>40.5</td>
<td>32.5</td>
<td>1226</td>
<td>30</td>
<td>—</td>
<td>Healthy</td>
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<tr>
<td>E Sporothrix</td>
<td>38</td>
<td>32.5</td>
<td>366</td>
<td>9</td>
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<tr>
<td>F Sporothrix</td>
<td>32.5</td>
<td>23</td>
<td>310</td>
<td>13</td>
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<tr>
<td>G Sporothrix</td>
<td>47</td>
<td>28</td>
<td>440</td>
<td>11</td>
<td>13</td>
<td>Completely wilted except for some epicormics</td>
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<tr>
<td>EXPERIMENT 2</td>
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<tr>
<td>1 Sterile water</td>
<td>41.5</td>
<td>30</td>
<td>664</td>
<td>17</td>
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<tr>
<td>2 Sterile water</td>
<td>33.5</td>
<td>30</td>
<td>525</td>
<td>17</td>
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<tr>
<td>3 Sterile water</td>
<td>38.5</td>
<td>30</td>
<td>606</td>
<td>17</td>
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<tr>
<td>4 Sterile water</td>
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<td>30</td>
<td>561</td>
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<tr>
<td>5 Sterile water</td>
<td>33</td>
<td>30</td>
<td>523</td>
<td>17</td>
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<td>30</td>
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<td>30</td>
<td>688</td>
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<td>9 Sporothrix</td>
<td>31</td>
<td>30</td>
<td>495</td>
<td>17</td>
<td>3</td>
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<td>44.5</td>
<td>30</td>
<td>708</td>
<td>17</td>
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<tr>
<td>11 E. platypodis</td>
<td>27.5</td>
<td>30</td>
<td>438</td>
<td>17</td>
<td>22</td>
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<tr>
<td>12 E. platypodis</td>
<td>32</td>
<td>30</td>
<td>510</td>
<td>17</td>
<td>—</td>
<td>Healthy</td>
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<tr>
<td>13 E. platypodis</td>
<td>39</td>
<td>30</td>
<td>627</td>
<td>17</td>
<td>—</td>
<td>Healthy</td>
</tr>
<tr>
<td>14 E. platypodis</td>
<td>34</td>
<td>30</td>
<td>549</td>
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<td>—</td>
<td>Healthy</td>
</tr>
<tr>
<td>15 E. platypodis</td>
<td>44.5</td>
<td>30</td>
<td>709</td>
<td>17</td>
<td>—</td>
<td>Healthy</td>
</tr>
</tbody>
</table>
The trees were felled between October 1972 and September 1975 (Appendix 1), either after death or when it was considered they would remain alive indefinitely. Microorganisms were isolated from the stems by cutting discs approximately 15 cm thick from the freshly felled trees from above, below, and within the treated area, e.g., for tree A, 11 discs were taken at distances ranging from 1.2 to 17 m from the base of the stem. Slabs were chopped from the discs and split with secateurs; slivers of wood (called "isolation chips") approximately 3 X 3 mm were then taken from the freshly exposed surface, placed on 3% malt agar slopes, and incubated at 20°C. As the purpose of these isolations was to recover Sporothrix, they were not always taken at random, but often from stained areas which, in proportion to clear wood, made up a small part of the total wood in the disc. Altogether 1396 isolations were taken from 59 discs.

Results

The time from treatment to complete wilt is shown in Table 1. For trees A, B, E, F, and G the time from the first sign of wilt to complete wilt was less than 3 months. Tree G had some epicormics in the lower crown which were still green when it was felled, but all these brown and wilted trees were almost certainly dead (see Experiment 2 — Results, trees 6-8). Trees C and D (inoculated with sterile water) had new foliage flushes for three successive seasons after treatment and their foliage was still green and healthy when they were felled. Notable changes in the condition of each tree, as observed on the monthly inspections, are shown in Appendix 1.

Sporothrix was recovered from the treated zone to 1.5 m above in tree A, 3.8 m in B, 1.4 m in E, 2.4 m in F, and 5.5 m in G. It was recovered only once from tree C (from the drilled zone) and not at all from D. Most of the isolation chips from which micro-organisms grew were taken from stained or discoloured wood. Clean wood usually proved sterile.

Figure 2 (a, b, c) shows a sample of discs from tree B demonstrating stained wood in discs from which Sporothrix was recovered. It was not recovered from the clean disc (Fig. 2d). Stains associated with the fungus-inoculated holes (Fig. 3a) were similar to those recorded by Faulds (1973). Stained wood was uncommon above the treated zone in trees C and D (Fig. 3b).

No new wood was formed after treatment in trees A, B, and G (Fig. 2a). Although all the other trees formed new wood, more wood was formed in trees treated with sterile water (Fig. 3c) than in the fungus-inoculated trees (Fig. 3d).

EXPERIMENT 2

Materials and Methods

Fifteen trees, similar in diameter to those used in Experiment 1, were selected for this experiment (Table 1). During January and February 1974 five trees (No. 1-5) were inoculated with sterile distilled water, five (6-10) with Sporothrix, and five (11-15) with Endomycopsis platypodis Baker et Kreger-van Rij (Ascomycetes, Endomycetales), the suspected main Platypus ambrosia fungus. Seventeen holes per 100 cm² were drilled in a band 30 cm wide at a height of approximately 1.5 m from the base of the stem of each tree. The total number of holes per tree is shown in Table 1. Drilling and inoculation techniques were the same as in Experiment 1. These trees were felled between
FIG. 2—Discs from *Sporothrix*-inoculated tree (a) from treated zone, (b) from 1.4 m above treated zone, (c) from 3.7 m above treated zone, (d) from 6.8 m above treated zone. *Sporothrix* was recovered from stained sapwood in (a), (b), and (c), but not from clean disc (d). Note that in (a) there was no band of new sapwood produced subsequent to inoculation.

FIG. 3 (opposite)—(a) Longitudinal tangential section from sapwood of treated zone of tree B showing stains associated with *Sporothrix*-inoculated holes.  
(b) Disc from 1.7 m above treated zone of tree D inoculated with sterile water, showing clean sapwood compared with stained sapwood in Fig. 2 b, c. The finger-like intrusion of heartwood is a typical sign of earlier *Platypus* attack, i.e., attack during the year corresponding with the annual ring at the apex of the intrusion.  
(c) Disc from treated zone of tree D inoculated with sterile water. Note the wide band of sapwood formed after inoculation compared with the new sapwood formed in the *Sporothrix*-inoculated tree in (d).
(d) Disc from treated zone of *Sporothrix*-inoculated tree F, showing the small amount of new sapwood formed since inoculation. The light areas of sapwood with no new sapwood around their outer edge were dead.
March 1974 and May 1976, and micro-organisms were isolated from them by the methods used in Experiment 1. In all, 1983 isolations were made from 111 discs. In both experiments a sleeve of linen gauze was used to protect the drilled zone of each tree against insect attack, and the basal 5 m of the stem were sprayed with 0.25% dieldrin on the dates shown in Appendices 1 and 2. Also, the condition of the crowns of the trees and any other relevant observations were recorded at approximately monthly intervals.

Results

The time from treatment to complete wilt is shown in Table 1, and notable changes in the condition of each tree in Appendix 2. Most trees inoculated with *Sporothrix* were completely wilted within 4 months. They were felled only when it was certain they would not flush and recover in the following spring. Trees 1-5 and 12-15 had new foliage flushes for two successive seasons after treatment and their foliage was green and healthy when they were felled.

*Sporothrix* was recovered from the treated zone to 1.2 m above in tree 6, 2.2 m in tree 7, 2.4 m in tree 8, 1.5 m in tree 9, and 3.5 m in tree 10. It was also recovered six, four, and four times from the treated zone only in trees 12, 13, and 14 respectively. In tree 11 it was recovered from up to 1.2 m above the treated zone, but it was not recovered from trees 1-5 or tree 15. The recovery of micro-organisms in relation to wood-staining was the same as for Experiment 1.

After treatment no new wood was formed on trees which wilted and died, but new wood was formed on trees which remained alive.

DISCUSSION AND CONCLUSIONS

Most ambrosia beetles rear brood only in recently dead woody plants and will not attack living trees. Those which do attack apparently healthy trees include *Dendroplatypus impar* Schedl which attacks the red meranti group of *Shorea* in Malaysia, *Trachyostus ghanaensis* Schedl which attacks *Triplochiton scleroxylon* K. Schum, and *Doliopygus dubius* Samps. on *Terminalia superba* Engl. et Diels (Browne, 1965); *Xyleborus fornicatus fornicator* Eggers which attacks tea bushes in Ceylon (Browne, 1961); *Platypus mutatus* Chapuis which attacks a wide range of hosts (poplar, willow, she-oak, eucalypt, plane, *Ailanthus*, apple, and pear) in Argentina and Brazil (Santoro, 1957; 1962a; b; 1963); *Corthylus columbianus* Hopk. which attacks several North American hardwood species (Giese and McManus, 1965); *Austroplatypus incompertus* Schedl infesting eucalypts in Australia (Browne, 1971); and *Xyleborus truncatus* Erichson which attacks eucalypts (chiefly the branches) in Australia (Moore, 1959; 1962). *Dendroplatypus impar* apparently does no harm to susceptible *Shorea* trees; *T. ghanaensis* attack does not prevent continued growth; *X. fornicatus fornicator* has not been shown to be the vector of any fungal disease to which the tea plant is subject; *P. mutatus* tunnels weaken trees so that they snap off in high winds, but no vector relationship with tree pathogens has been demonstrated. Only *X. truncatus* is known as a disease vector, being associated with a fungus which Stahl (pers. comm. to R. H. Milligan, 1964) has identified as a species of *Ceratocystis* and considers identical with the ambrosia fungus on which broods are reared. Moore (1962) states that the fungus causes brown stains in the sapwood around
X. truncatus entries and that attacked trees exhibit terminal dieback and produce epicormic shoots; he also found that, at least in the coastal regions and the highlands of New South Wales, Eucalyptus saligna Smith deaths have been widespread.

All three New Zealand Platypus species (P. apicalis, Platypus caviceps Broun, and Platypus gracilis Broun) attack living and apparently healthy Nothofagus trees (Milligan, 1974).

The results of these experiments clearly show that Sporothrix is a pathogen of N. fusca. Even in tree 11 which died after inoculation with E. platypodis, Sporothrix was recovered from within and well above the treated zone and probably played a role in its death.

Initial establishment of the pathogen in the host tissues probably depends on its introduction into the inner sapwood (Milligan, 1972) and Platypus spp. are the only insects boring in larger-diameter living beech trees which make tunnels in this zone. Whether under natural conditions Platypus is a vector of the pathogen or whether the pathogen incidentally invades Platypus wounds is not known. However, as the fungus was recovered from trees into which it had not been inoculated it can obviously invade Platypus-like wounds in the absence of beetles. Also, it is reasonable to assume that Platypus adults emerging from infected material would be carrying fragments or spores of the fungus.

Since its original isolation Sporothrix has been isolated from other wilting Platypus-attacked beech, including Nothofagus solandri var. cliffortioides (Hook.f.) Poole from Mount Ruapehu and Nothofagus truncata (Col.) Ckn. from the Clevedon area (approximately 30 km south-east of Auckland). Milligan (1974) reported that Platypus and the associated pathogen were involved in the deaths of mature trees of the other New Zealand Nothofagus species, and that although susceptibility to this sort of mortality is not necessarily equal in the various species, none is immune. Nowhere else in the world have species of Platypodidae been implicated as vectors of, or associated with, tree-killing pathogens.

The density of inoculations used in these experiments and the diameter class of experimental trees was based on Milligan’s observations of tree mortality after induced Platypus attack. In his experiment up to 13 attacks per 100 cm² were counted in the most severely attacked parts of the trees which died; but trees smaller than 30 cm diameter did not die even though subject to comparable attack densities. Although some of the drilled trees received a maximum density of drilled holes slightly greater than this, the following differences probably more than offset this. Firstly, many of the real Platypus attacks succeeded to the stage where tangential arms off the radial tunnel were constructed (therefore exposing a far greater area of sapwood to infection); secondly, the area of maximum density of attack was larger; thirdly, there were many Platypus attacks above and below the area of maximum density of attack.

It is known that many trees survive successive annual Platypus attacks (Kershaw, 1969; Litchwark, in prep.) and that much of the stem defect in Nothofagus forests arises directly from sublethal Platypus attack (Milligan, 1974). In contrast Milligan (1974) found that trees only lightly and abortively attacked succumbed to the fungal pathogen when a drought occurred in the following summer, even without a second attack in the
drought year. Climatic and other conditions affecting tree health obviously play an important role in the successful establishment of the pathogen in the tree. The central plateau of the North Island, including Kaimanawa State Forest Park, suffered from three successive summer droughts in 1971-72, 1972-73, and 1973-74 (New Zealand Meteorological Service, 1972; 1973, 1974).

How did these droughts affect the experiments? One effect might have been to change a possible sublethal inoculation into a lethal inoculation. For example, in comparison with trees A and B which wilted and died rapidly, trees E and F, both of which had approximately half the number of inoculations that A and B had, took a long time to die. These slow-dying trees also produced new wood after treatment. Would they have received only a sublethal inoculation but for the droughts? The fact that trees C and D treated with sterile water survived the drought, in spite of having more holes and a greater density of holes drilled in them than any of the trees inoculated with Sporothrix, implies that the drill wounding was not a significant factor in mortality.

In Experiment 2 the Sporothrix-inoculated trees all wilted and died quickly. Probably these trees were under severe water stress at the time of inoculation.

Wilt caused by infective agents is thought to involve insufficient water resulting from blockage of the transpiration stream (Smith, 1970). Clearly the experimental trees were killed by interruption of the xylem and hence the water supply to the crown. Death of phloem only would not produce such rapid wilt as occurred in some trees.

The mechanism of wilt has not been studied. Probably the most intensively studied comparable fungal pathogen is Dutch elm disease. Several different possibilities have been suggested as the wilt mechanism for this disease. Many investigators thought plugging of vessels by gums, tyloses, or cytoplasm from parenchyma cells were causes of wilting (Wollenweber, 1927; Broekhuizen, 1929; Clinton and McCormick, 1936; Pope, 1943), and others have postulated systemic toxaemia (Zentmyer, 1942; Zentmyer et al., 1946; Kerling, 1955). On the other hand, Schwarz (1922) stated that degradation of cell walls might be an important factor in the wilt mechanism and Ouellette (1962) said that acute symptoms of the disease could result from the plugging of the smaller vessels by the spores and mycelium of the pathogen alone or in combination with cytoplasm and residues from adjoining cells, and particles arising from the deterioration of cell walls. He also suggested that the modification of vessel walls by fungal action and changes occurring in living parenchyma, which accumulating evidence suggests are involved in translocation (Greenidge, 1955; 1957; Postlethwait and Rogers, 1958), might independently contribute to wilting.

Although systemic toxaemia might be a factor in the wilt mechanism of Dutch elm disease, it seems unlikely to be involved in the wilt of Platypus-attacked Nothofagus. Infection of healthy elms is usually caused by bark beetles feeding in crotches of twigs and eating out a small tunnel or groove, thus exposing the xylem to the fungal spores carried internally and on the bodies of the beetles, i.e., infection occurs near the foliage (Forestry Commission, 1958). In contrast there is no such feeding by Platypus adults, so the point of infection must be the Platypus tunnels in the main stem — well away from the foliage. The wilt mechanism is more likely to involve one or more of the other factors mentioned.
ACKNOWLEDGMENTS

The author thanks Mr R. H. Milligan for helpful advice and Dr P. D. Gadgil for the identification of Sporothrix sp.

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APPENDIX 1—Notable changes and observations in Experiment 1

Tree A
March 1972 Inoculated with Sporothrix; crown healthy until 30 August 1972.
30 Aug 1972 Lower branches wilting; upper crown healthy.
16 Sept 1972 Lower branches to approximately 10 m dead or wilting; terminals in mid crown wilting; upper crown wilting.
29 Sept 1972 Lower crown wilted except for some epicormics with green foliage; mid crown wilting but still a little yellow and green foliage; upper crown wilted.
27 Nov 1972 Completely wilted—felled.

Tree B
March 1972 Inoculated with Sporothrix; crown healthy until 30 August 1972.
30 Aug 1972 Lots of foliage with a brown appearance.
29 Sept 1972 Whole crown wilted or wilting except for a few green branches in mid crown.
27 Nov 1972 Completely wilted.
13 Dec 1972 Felled.

Tree C
March 1972 Inoculated with sterile distilled water; some dead branches; foliage generally sparse.
27 Nov 1972 New foliage flushed.
22 Feb 1973 Some Platypus attack.
27 Feb 1973 Stem to 5 m sprayed with 0.25% dieldrin.
15 Nov 1973 New foliage flushed.
27 Mar 1974 New foliage flushed; stem to 5 m sprayed with 0.25% dieldrin.
10 Apr 1974 Some Psepholax (Curculionidae) attack.
14 Nov 1974 New foliage flushed.

Tree D
April 1972 Inoculated with sterile distilled water.
27 Nov 1972 New foliage flushed.
7 Feb 1973 Some gum bleed from drilled holes.
22 Feb 1973 A few Platypus attacks.
27 Feb 1973 Stem to 5 m sprayed with 0.25% dieldrin.
15 Nov 1973 New foliage flushed; a few dead branches present.
27 Mar 1974 Stem to 5 m sprayed with 0.25% dieldrin.
6 Dec 1974 New foliage flushed.

Tree E
April 1972 Inoculated with Sporothrix.
27 Nov 1972 New foliage flushed.
15 Feb 1973 Some Platypus attack.
27 Feb 1973 A few dead branches present; some Psepholax attack; stem to 5 m sprayed with 0.25% dieldrin.
27 Mar 1974 Quite a few dead branches; stem to 5 m sprayed with 0.25% dieldrin.
6 Dec 1974 New foliage flushed.
5 May 1975 Lots of dead branches in upper crown, mid, and lower crown, with wilted foliage on some branches.
27 Jun 1975 Lower crown wilted and dead; still some green leaves on upper crown.
14 July 1975 Lower crown wilted; upper crown foliage going yellow.

Tree F
May 1972 Inoculated with Sporothrix.
27 Nov 1972 New foliage flushed.
22 Feb 1973 Platypus attack.
27 Feb 1973 Stem to 5 m sprayed with 0.25% dieldrin; Psepholax attack.
15 Nov 1973 New foliage flushed.
27 Mar 1974 Stem to 5 m sprayed with 0.25% dieldrin.
6 Dec 1974 New foliage flushed.
16 Jan 1975 Small branches in upper crown with very few leaves; rest of crown healthy.
7 Apr 1975 Foliage yellow and wilting, only a few green leaves left.
5 May 1975 Completely wilted—felled.

Tree G
May 1972 Inoculated with Sporothrix.
27 Nov 1972 Crown healthy but new foliage not yet flushed.
18 Jan 1973 Lots of dead branches and terminals in upper and mid crown; lower crown green.
25 Jan 1973 Many dead branches and terminals; few remaining leaves, some of these yellow.
1 Feb 1973 Over 50% of foliage in mid and upper crown gone.
7 Feb 1973 Some Platypus attack.
15 Feb 1973 Some foliage wilting in upper crown.
27 Feb 1973 Stem to 5 m sprayed with 0.25% dieldrin; most of upper crown without foliage; a little green foliage on small branches.
7 May 1973 Upper crown dead; epicormics in lower crown still green.
23 May 1973 Felled.
APPENDIX 2—Notable changes and observations in Experiment 2.

**Trees 1-5**

- Jan 1974: Inoculated with sterile distilled water.
- 22 Mar 1974: A few *Platypus* attacks on Tree 2.
- 27 Mar 1974: Stem to 5 m sprayed with 0.25% dieldrin.
- 24 Oct 1974: Stem to 5 m sprayed with 0.25% dieldrin.
- 6 Nov 1975: Stem to 5 m sprayed with 0.25% dieldrin.
- 16 Jan 1976: Tree 3 felled — healthy when felled.
- 27 Jan 1976: Tree 1 felled — healthy when felled.
- 23 Feb 1976: Tree 2 felled — healthy when felled.
- 1 Mar 1976: Tree 4 felled — healthy when felled.
- 8 Mar 1976: Tree 5 felled — healthy when felled.

**Tree 6**

- Jan 1974: Inoculated with *Sporothrix*.
- 22 Mar 1974: Lower branches wilted; foliage in upper crown yellowish; some *Platypus* and *Psepholax* attack.
- 27 Mar 1974: Stem to 5 m sprayed with 0.25% dieldrin.
- 10 Apr 1974: Foliage wilting or yellow; only a few green leaves left in upper crown.
- 23 Apr 1974: Crown wilted except for a few leaves in upper crown.
- 17 May 1974: Completely wilted.

**Tree 7**

- Jan 1974: Inoculated with *Sporothrix*.
- 22 Mar 1974: Foliage in lower crown wilting; foliage in upper crown unhealthy; some *Platypus* and *Psepholax* attack.
- 27 Mar 1974: Stem to 5 m sprayed with 0.25% dieldrin.
- 10 Apr 1974: Foliage wilting or wilting except for patches of green leaves.
- 23 Apr 1974: Foliage wilting except for a few green leaves.
- 17 May 1974: Completely wilted.
- 12 Dec 1974: Felled.

**Tree 8**

- Jan 1974: Inoculated with *Sporothrix*.
- 22 Mar 1974: Some wilting foliage in lower crown; a few *Platypus* attacks.
- 27 Mar 1974: Stem to 5 m sprayed with 0.25% dieldrin.
- 10 Apr 1974: Lots of wilted foliage and yellow leaves throughout crown; a few branches still with green foliage.
- 23 Apr 1974: Crown wilted except for some green and yellow leaves on one side of tree and epicormics in lower crown.
- 17 May 1974: Completely wilted except for epicormics.
- 24 Jul 1974: Epicormics in lower crown starting to wilt.
- 6 Dec 1974: Felled.

**Tree 9**

- Jan 1974: Inoculated with *Sporothrix*.
- 22 Mar 1974: Some foliage on lower branches going brown; ten *Platypus* attacks and two *Psepholax* attacks seen.
- 27 Mar 1974: Stem to 5 m sprayed with 0.25% dieldrin.
- 10 Apr 1974: Most of foliage wilted; only a few small branches with green leaves.
- 23 Apr 1974: Completely wilted.

**Tree 10**

- Jan 1974: Inoculated with *Sporothrix*.
- 23 Mar 1974: Crown yellowing and some foliage wilting; foliage sparse; a few *Platypus* attacks.
- 27 Mar 1974: Stem to 5 m sprayed with 0.25% dieldrin.
- 10 Apr 1974: Completely wilted.

**Tree 11**

- Feb 1974: Inoculated with *E. platypodis*.
- 22 Mar 1974: Some *Platypus* and *Psepholax* attack.
- 27 Mar 1974: Stem to 5 m sprayed with 0.25% dieldrin.
- 24 Oct 1974: Stem to 5 m sprayed with 0.25% dieldrin.
- 20 Oct 1975: Lots of yellow foliage throughout crown and some small branches in lower crown wilted.
- 3 Nov 1975: Lower branches wilted; mid crown yellow or wilted.
- 24 Nov 1975: Completely wilted.

**Trees 12-15**

- Feb 1974: Inoculated with *E. platypodis*.
- 27 Mar 1974: Stem to 5 m sprayed with 0.25% dieldrin.
- 24 Oct 1974: Stem to 5 m sprayed with 0.25% dieldrin.
- 6 Nov 1975: Stem to 5 m sprayed with 0.25% dieldrin.
- 12 Jan 1976: Tree 12 felled — healthy when felled.
- 29 Mar 1976: Tree 14 felled — healthy when felled.
- 5 Apr 1976: Tree 15 felled — healthy when felled.