Steaming Inactivates Phytophthora ramorum, Causal Agent of Sudden Oak Death and Ramorum Blight, from Infested Nursery Soils in California

Wolfgang Schweigkofler, Department of Natural Sciences and Mathematics, Dominican University of California, 50 Acacia Avenue, San Rafael, CA 94901; Kathleen Kosta, California Department of Food and Agriculture, 1220 N Street, Sacramento, CA 95814; Vernon Huffman, Supriya Sharma, and Karen Suslow, Department of Natural Sciences and Mathematics, Dominican University of California, 50 Acacia Avenue, San Rafael, CA 94901; and Sibdas Ghosh, Department of Natural Sciences and Mathematics, Dominican University of California, 50 Acacia Avenue, San Rafael, CA 94901 and School of Arts and Science, Iona College, 715 North Avenue, New Rochelle, NY 10801

Accepted for publication 21 January 2014. Published 13 March 2014.

Nursery trade plays a major role in the long-distance spread of Phytophthora ramorum, the causal agent of Sudden Oak Death and ramorum blight, from infested nursery soils in California. Plant Health Progress doi:10.1094/PHP-RS-13-0111.

ABSTRACT


Nursery trade plays a major role in the long-distance spread of Phytophthora ramorum, the causal agent of Sudden Oak Death (SOD) and ramorum blight of ornamental plants. Under federal regulations, nurseries found positive for P. ramorum must destroy infected plants and treat infested soils. The use of steam is an effective method to thermally inactivate P. ramorum from nursery soils as demonstrated at the National Ornamental Research Site at Dominican University (NORS-DUC) and one commercial nursery in the Central Valley of California. Heating up the top soil layer (0-30 cm) to 50°C for 120 minutes resulted in complete thermal inactivation of P. ramorum. Consequently, the commercial nursery was released from federal quarantine. Steaming can be a fast, reliable and sustainable option for treating nursery soils.

INTRODUCTION

Phytophthora ramorum Werres, De Cock & Main’in’t Veld, is a new invasive plant pathogen that causes Sudden Oak Death (SOD), a devastating forest disease in California and Southwestern Oregon, characterized by extensive mortality of tanoak (Notholithocarpus densiflorus) and coast live oak (Quercus agrifolia) (11). Recently, a massive outbreak of disease in Japanese Larch (Larix kaempfieri) plantations in the United Kingdom was associated with P. ramorum (15). In addition, P. ramorum causes ramorum blight on several economically important ornamental nursery plants, resulting in relatively minor leaf symptoms without killing the host. The nursery trade of symptomless but infected plant material plays an important role in the long-distance spread of the pathogen (6). In infested nurseries, P. ramorum can survive on plant debris, potting mix, soil in container fields, nursery equipment, and in water for extended periods of time. According to the federal quarantine program implemented by USDA APHIS (1), nursery soil which tests positive for P. ramorum has to be treated to eliminate the pathogen and minimize the risk for further dispersal.

The National Ornamental Research Site at Dominican University of California (NORS-DUC), is focused on the development and implementation of environmentally friendly solutions to stop the spread of pathogens of ornamental plants. The research site contains a mock nursery, which was developed to study quarantine pathogens under conditions similar to those typical of commercial nurseries (8).

Methods for eradication of pathogens include chemical, biological, and physical treatments. The aim of this study was to test (i) if steaming is a suitable method to inactivate P. ramorum from infested soils in a research nursery, (ii) the time frame and costs for a steaming event, and (iii) if steaming is an option for treating soil of a commercial nursery which is under federal quarantine.

EFFECT OF WET HEATING AND DRY HEATING ON THE SURVIVAL OF P. RAMORUM

Heat treatment is a well-established method to inactivate plant pathogens in nursery soils (2); however, because P. ramorum is a federally quarantined pest, soil disinfection procedures must be planned and executed with the utmost caution as USDA has established a zero-tolerance management strategy. P. ramorum is a plant pathogen found in regions with temperate climate, and grows at temperatures between 2°C and 30°C with an optimum around 20°C (16). In order to simulate the effect of soil heating, we tested the impact of high temperatures on the growth of two P. ramorum isolates (Pr-1418886 and Pr-1418983-2) in the laboratory. Both strains originate from nurseries in Marin County, belong to clonal lineage NA1 and are of A2 mating type. Two heat sources were used: wet heating, which resembles the conditions in steamed soil; and dry heating. Leaves of Rhododendron catawbiense cv. English Roseum were inoculated with P. ramorum, incubated for three weeks at 20°C, and leaf disks (diameter 5 mm) overgrown with mycelium and chlamydoospores were used for the thermal experiment. In order to test the effects of dry heating against wet heating, leaf disks were incubated at 30°C, 40°C, 50°C, or 60°C, respectively, for 30, 60,
and 120 min in a dry incubator (Fisher Scientific, Hampton, NH; RH values varying from 31.3% at 30°C to 8.5% at 60°C) or a water bath (Thermo Scientific, Waltham, MA). After treatment, leaf disks were plated on PARPH-V8 medium (5), incubated at 20°C for a week and the percentage of disks that yielded a colony of \textit{P. ramorum} recorded. Experiments were done in triplicate; each replicate contained ten leaf disks. SPSS 12.0 (IBM Corp., Armonk, NY) for Windows was used for statistical analysis. The chi-square test was used to compare differences in survival rates between two isolates and wet vs. dry heating.

**SURVIVAL OF \textit{P. ramorum} IN THE LABORATORY**

Effect of thermal treatment on the growth of \textit{P. ramorum} is shown in Figure 1 (A, wet heating; B, dry heating). Incubation at 30°C for up to 120 min showed almost no effect on the survival rate. At 40°C, growth rates were reduced in most cases except one isolate under wet heat. At 50°C, wet heat treatment for 30 min inactivated \textit{P. ramorum} completely (survival rate = 0%). Dry heating at 50°C also lowered the growth rate, but was slightly less effective than wet heating (chi-square test, \( P = 0.00 \)). At 60°C, \textit{P. ramorum} was completely inactivated after 30 min of either wet or dry heat. Similar results have been reported for other plant pathogens (12). The two \textit{P. ramorum} isolates differed in their reaction to heating, with Pr-1418886 showing slightly higher heat tolerance than Pr-1418983-2 (chi-square test; \( P = 0.06 \)). The morphological structures of \textit{Phytophthora} spp. differ in their ability to resist adverse environmental conditions, with chlamydospores and oospores generally being more heat-stable than hyphae (14). Microscopic observations confirmed that our leaf disks contained both hyphae and chlamydospores. Oospore production of \textit{P. ramorum}, which is a heterothallic species, has not been observed in the field (6). These results are in agreement with earlier reports by Linderman and Davis (9), who found that chlamydospores of \textit{P. ramorum} mixed with plant potting medium were inactivated by aerated steam heat treatment of 50°C for 30 min; and Browning et al. (3), who reported that \textit{P. ramorum} hyphae grown on V8 medium were inactivated after exposure to temperatures of 42.5 to 50°C at a thermal gradient plate for several minutes; whereas Hamik et al. (7) reported a 60% recovery rate of \textit{P. ramorum} from infected California Bay Laurel (\textit{Umbellularia californica}) leaves incubated at 55°C for one week.

**STEAMING AT A RESEARCH NURSERY AT NORS-DUC**

Dominican University of California is located in San Rafael, CA, an area with a Mediterranean climate, with warm, dry summers (average highs 27°C in August) and mild, rainy winters (average lows around 6°C in December) and total annual precipitation averages 817 mm. The steaming experiments were carried out in research beds with a surface area of 3.7 m \times 4.6 m (17 m\(^2\)) which were lined with water-proof liners. The beds were filled with a 30-cm-deep layer of gravelly loam soil with a pH of 5.8 and 5.9% organic matter; which was originally excavated

\[\text{FIGURE 1}\]

Effect of thermal treatment on the growth of \textit{P. ramorum} isolates Pr-1418886 and Pr-1418983-2. Infected Rhododendron leaf disks were incubated at 30, 40, 50, and 60°C for 30, 60, and 120 min, respectively, in (A) a water bath (wet heat) or (B) an incubator (dry heat). Error bars indicate 95% confidence intervals.
from the Dominican campus. A steamer unit SIOUX Steam-Flo SF-11 (Sioux Corp., Beresford, SD, USA) with a boiler horsepower of 10.67 and a steaming output of 168 liters/h was used with an attached soaker hose (length 31 m) laid out on the soil surface, which then was covered with a waterproof, pond liner-type tarp. A total of five soil steaming events were carried out at NORS-DUC between July 2012 and February 2013. In addition, nine steaming events were carried out on research beds filled with potted plants (Rhododendron sp. and Viburnum sp.), which were used previously for P. ramorum infection studies. For all steaming experiments, the minimum soil temperature of 50°C was kept for 120 min (‘inactivation period’) before turning off the steamer.

STEAMING AT A COMMERCIAL NURSERY IN THE CENTRAL VALLEY, CA

Preliminary work at a commercial nursery in Stanislaus County, CA, which had been found to have P. ramorum infested soil in the container field, indicated that steam could inactivate the pathogen from nursery soil (unpublished data, S. Johnson-Brousseau). Another nursery, also found with P. ramorum infested soil, was selected for further steam treatment studies. The commercial retail nursery was located in Sacramento, CA, an area with Mediterranean climate, with hot, dry summers (average highs 33°C in August) and mild, rainy winters (average lows around 3°C in December); total annual precipitation averages 470 mm. Loropetalum sp. plants infected with P. ramorum were detected during an annual nursery inspection in the spring of 2012. Baiting of the soil on which the potted plants were kept, conducted by the California Department of Food and Agriculture (CDFA) found the soil to be infested with P. ramorum. The area put under quarantine was 20.5 m × 5.3 m (108.7 m²), composed of a layer of coarse gravel (approximately 30 cm deep, consisting of approximately ¾-inch rocks) over sandy loam. The area to be steam-treated was completely saturated with water from previous irrigations. The nursery bed was divided into two areas of 54.3 m² each, and treated on two subsequent days in August 2012.

MEASUREMENT OF TEMPERATURE AND SURVIVAL RATES

Soil temperatures were measured using HOBO-data loggers U12 (Onset Corp., Cape Cod, MA, USA) with sensors positioned at 5, 15, and 30 cm, respectively, below surface. Temperature data were collected at a 15-min interval. To serve as the positive control at the contained research site, 20 Rhododendron leaf disks overgrown with mycelium and chlamydospores of P. ramorum isolate Pr-1418886 were placed in sachets and buried at the deepest soil layer prior to steaming. After steaming, the leaf disks were plated on PARPH-V8 plates, incubated in the dark at 20°C for two weeks and then checked for growth. As a control, leaf disks infected with P. ramorum, but not-heat treated, were plated onto PARPH-V8. Phytophthora ramorum infected leaf disks could not be used at the commercial nursery due to regulations. Soil samples from the commercial nursery were tested for P. ramorum before and after treatment by a soil-baiting technique using Rhododendron leaves at both the NORS-DUC and the CDFA labs.

THERMAL INACTIVATION OF SOIL BORNE P. RAMORUM

Soil contained in a research bed used for studies on P. ramorum in a mock nursery was treated with aerated steam using a commercial steaming unit (Fig. 2). The soil temperature was measured from sensors strategically positioned at different areas within the research beds as well as at different soil depths. Three representative steaming experiments are described in more detail: the experiments 1 and 2 were carried out at NORS-DUC in different seasons (1 in July 2012, 2 in February 2013), and experiment 3 was carried out at a retail nursery in Sacramento in August 2012. Temperature profiles show different dynamics depending on soil depths. Experiment 1 (Fig. 3A) started from an ambient air temperature of 18°C; the target temperature of 50°C at 5-cm soil depth was reached after approximately 1 h, at 15 cm after 2 h, and at 30 cm after 6 h. The heat source was turned off 2 h after reaching the target temperature, after which temperatures close to the surface dropped quickly, whereas those at 30 cm rose for several degrees due to continued heat transfer from the upper layers, before declining steadily. Maximum temperatures are difficult to control with this approach; close to the soil surface temperatures above 90°C can be achieved. Soil temperatures remain well above ambient temperatures for up to three days post-steaming (data not shown).

Experiment 2 (Fig. 3B) started from an ambient air temperature of approximately 10°C; the target temperature of 50°C at 5-cm soil depth was reached after approximately 4 h, at 15 cm after 9 h and at 30 cm after 12 h. Experiment 3 (Fig. 3C) started from an ambient air temperature of approximately 20°C; the target temperature of 50°C at 5-cm soil depth was reached after approximately 2 h, at 15 cm after 5 h, and at 30 cm after 9.5 h.

The dynamics of the temperature increase in different soil depths and the time needed to reach the target temperature of 50°C are influenced by several parameters, among them ambient temperature and soil water content. Experiment 1, conducted on 10 July 2012 in San Rafael, CA, with an average air temperature of 18°C (maximum 25.5°C) and dry soil conditions resulted in a steep increase of the soil temperatures. Experiment 2, conducted on 6 February 2013 using the same soil but at lower ambient air temperatures (average 9.7°C, maximum 12.8°C), showed a much
slower temperature increase. Soils at the commercial nursery (experiment 3), which were treated at on 28 August 2012 with a average air temperature of 23.6°C (maximum 34ºC), heated up slowly because of high water content.

Steaming of research beds filled with remains of infection studies on potted plants, including pots, root balls, etc., was characterized by very fast temperature increase. Temperatures reached the target temperature of 50ºC at the center of a root ball in a one-gallon pot after one hour and rose to >80ºC (data not shown).

The heat treatment successfully inactivated *P. ramorum* in the soil and on potted plants (Table 1); no growth was detected after incubation of the leaf disks which were steamed at NORS-DUC (survival rate = 0%), whereas *P. ramorum* cultures developed from each control leaf disk (survival rate = 100%). Soil samples from the commercial nursery post-treatment were tested negative using a soil-baiting method; Rhododendron leaves were incubated with a soil-water mix at 20°C for two days and subsequently leaf disks plated on PARPH-V8 plates. The nursery was consequently released from quarantine.

The maximum soil depth in our experiments (30 cm) was limited by the dimensions of the lined research beds at NORS-DUC. However, the presence of *P. ramorum* in infected nursery soil is largely limited to the top soil layer. In a study on *P. ramorum* in nursery soils in Washington State, Dart et al. (4) discovered 86% of soil inoculum in the organic layer or the top 0-5 cm of soil, with no detection occurring below 10 cm. In our experiments, temperatures at the 5-cm soil layer reached between 84.7°C and 98.7°C, and at the 15-cm soil between 67.5°C and 83.4°C (Fig. 3). Provided that distribution of *P. ramorum* is similar in nursery soils in different geographic locations, heat treatment of the top 30 cm soil layer should result in complete deactivation of *P. ramorum*. Further support for the usefulness of thermal inactivation of *P. ramorum* comes from a recent study by Parke and Funahashi (10) at NORS-DUC, who used solarization to heat up soil to a maximum temperature of approximately 50ºC to inactivate inoculum grown on leaf disks.

FIGURE 3

Temperature profiles of three steaming events: (A) research nursery at NORS-DUC, July 2012; (B) research nursery at NORS-DUC, February 2013; (C) commercial nursery in the Central Valley, California, August 2012. Temperatures were measured at depths of 5 cm (red line), 15 cm (black line), and 30 cm (blue line). The horizontal line shows the target temperature of 50°C. The “inactivation period” (defined as the time between reaching the target temperature and turning off the steamer) is shown in red.

ACKNOWLEDGMENTS

We thank S. Johnson-Brousseau, G. Copeland, and M. Henkes for their efforts to develop and initiate the early work at NORS-DUC. We especially would like to thank the owners and managers of the nurseries who agreed to participate in the study. Our thanks to Drs. Suzanne Rooney Latham and Cheryl Blomquist from the CDFA Plant Pest Diagnostic Laboratory for their continued support and collaboration. CDFA completed the mandatory soil testing required at the commercial nurseries, as per the federal regulations. NORS-DUC is funded by a grant from the 2008 Farm Bill, Section 10201, and administrated through the United States Department of Agriculture (USDA) Animal and plant Health Inspection Service (APHIS) Plant Protection and Quarantine (PPQ) Centre for Plant Health Science and Technology (CPHST).
TABLE 1
Effect of steaming on the growth of P. ramorum on Rhododendron leaf disks.

<table>
<thead>
<tr>
<th>Experiment</th>
<th>Date</th>
<th>Location</th>
<th>Substrate</th>
<th>Control</th>
<th>Heat-treated</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>7/10/2012</td>
<td>NORS-DUC</td>
<td>soil</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>2</td>
<td>2/6/2013</td>
<td>NORS-DUC</td>
<td>soil</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>3*</td>
<td>8/28/2012</td>
<td>Sacramento</td>
<td>soil</td>
<td>positive</td>
<td>negative</td>
</tr>
<tr>
<td>4</td>
<td>8/14/2012</td>
<td>NORS-DUC</td>
<td>soil</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>5</td>
<td>8/16/2012</td>
<td>NORS-DUC</td>
<td>soil</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>6</td>
<td>2/4/2013</td>
<td>NORS-DUC</td>
<td>soil</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>7</td>
<td>7/23/2012</td>
<td>NORS-DUC</td>
<td>potted plants</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>8</td>
<td>7/24/2012</td>
<td>NORS-DUC</td>
<td>potted plants</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>9</td>
<td>9/5/2012</td>
<td>NORS-DUC</td>
<td>potted plants</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>10</td>
<td>9/7/2012</td>
<td>NORS-DUC</td>
<td>potted plants</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>11</td>
<td>9/11/2012</td>
<td>NORS-DUC</td>
<td>potted plants</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>12</td>
<td>9/13/2012</td>
<td>NORS-DUC</td>
<td>potted plants</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>13</td>
<td>1/8/2013</td>
<td>NORS-DUC</td>
<td>potted plants</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>14</td>
<td>1/9/2013</td>
<td>NORS-DUC</td>
<td>potted plants</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>15</td>
<td>4/25/2013</td>
<td>NORS-DUC</td>
<td>potted plants</td>
<td>100</td>
<td>0</td>
</tr>
</tbody>
</table>

* Soil at the commercial nursery (experiment 3) was tested using a non-quantitative soil-baiting technique.

LITERATURE CITED


