**Phytophthora insolita** on Hainan Island

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**Abstract.** *Phytophthora insolita* was isolated from forest soil, streams, and ponds on southern China’s Hainan Island. This is the first documented report of its occurrence outside Taiwan, and the significance of the finding is discussed.

**Keywords:** China; Hainan Province; *Phytophthora; P. insolita.*

**Introduction**

*Phytophthora insolita* Ann & Ko is an interesting but little known fungus, distinguished from all other species of *Phytophthora* in the production of abundant oospores in single cultures without the presence of antheridia (Stamps et al., 1990). Although parthenogenic “oospores” have been reported occasionally in *P. infestans* de Bary and *P. fragariae* Hickman (Pethybridge and Murphy, 1913; Savage et al., 1968), they are rare and produced only under certain cultural conditions. *Phytophthora insolita* was first reported from citrus soil in Changhua, Taiwan (Ann and Ko, 1980) and later from diseased poinsettia (*Euphorbia pulcherrima* Willd.), also in Changhia. Later, the common occurrence of asexual isolates of *P. insolita* was discovered in soil, ditch water, and diseased plant tissues in southern and central Taiwan (Ann and Ko, 1994). In a recent survey of *Phytophthora* species on southern China’s Hainan Island we have obtained isolates of *P. insolita* and are reporting them herewith as new findings.

**Materials and Methods**

In the summers of 1998 and 1999, samples of soil, stream and pond water were collected from natural mountain forests and other localities and brought back to the Plant Pathology Laboratory in Dangzhou City for the isolation of *Phytophthora* species. Samples were placed in 9-inch specimen dishes, with the soil samples flooded with an equal amount of deionized water. Citrus leaves were used as the bait, submerged or partially submerged in water. When lesions started to develop within a few days at room temperature (27-30°C), they were removed, scrubbed slightly with domestic bleaching powder and washed clean under running tap water. Small pieces of plant tissues (ca 2×4 mm) were cut out from the margin the lesions, blotted dry with paper towels and plated on selective medium in 9-cm petri dishes. The medium was made up by mixing 10 ml Campbell’s V-8 juice, 90 ml deionized water, and 0.2 g CaCO3. This was filtered through two layers of cheesecloth, diluted with deionized water to 1000 ml, boiled to dissolve 20 gm Bacto agar, and autoclaved at 121°C for 15 min. Prior to pouring the agar medium into sterilized petri dishes, benomyl (150 ppm), hymexazol (50 ppm), rifampicin (100 ppm), nystatin (50 ppm), and ampicillin (100 ppm) were added. The inoculated plates were incubated in dark at 25°C. When mycelial colonies appeared in 2-3 days, they were examined under light microscope to determine their identity as species of *Phytophthora* based on the colony morphology and the characteristics of the hyphae and branching characteristics (Ho et al., 1995). The culture was then transferred onto 10% V-8 agar plates for further studies. To induce sporangial production, small mycelial agar discs (ca 2×2 mm) were cut from the edge of a growing colony and transferred to sterile distilled water in 6-cm petri dishes, left under regular indoor light and at room temperature.

The pathogenicity of *Phytophthora insolita* to fruits was tested by inoculating apple, avocado, cucumber, eggplant, mango, green pepper and tomato with the fungus. The fruit was washed clean, surface sterilized with 90% ethanol, and a slit was made with a flamed scalpel. A small mycelial disc from a young colony of isolate Bds 1-18 was inserted into the crevice, which was then sealed with Scotch tape. Alternatively, a small mycelial agar disc (ca 5×5 mm) was placed on an unwounded surface, kept moist with wet cotton, and held in place by Scotch tape. The inoculated fruits were kept in sealed plastic bags and incubated at 28°C for 5 days.

**Results**

Of the numerous isolates of *Phytophthora* recovered, those that have been identified as belonging to *P. insolita* are presented in Table 1. They were all similar in producing appressed and petaloid colonies on V-8 agar plates,
Table 1. Isolates of *Phytophthora insolita* from Hainan Island, South China.

<table>
<thead>
<tr>
<th>Isolate</th>
<th>Source</th>
<th>Location</th>
</tr>
</thead>
<tbody>
<tr>
<td>JFS 9954, 55, 57</td>
<td>Soil</td>
<td>Jianfengling</td>
</tr>
<tr>
<td>Bds 1-18, 9</td>
<td>Soil</td>
<td>Dongliuqu, Bawangling</td>
</tr>
<tr>
<td>Bdw 1-6, 2-6, 3-1</td>
<td>Stream water</td>
<td>Dongliuqu, Bawangling</td>
</tr>
<tr>
<td>Bdl 1-1, 1-2</td>
<td>Fallen leaves in stream water</td>
<td>Dongliuqu, Bawangling</td>
</tr>
<tr>
<td>Lmw 1-1, 1-3, 1-4, 1-5, 1-7, 1-8, 1-9, 1-11</td>
<td>Pond water</td>
<td>Tianmushahua, Limushan</td>
</tr>
<tr>
<td>Bw 1-1, 1-2</td>
<td>Pond water</td>
<td>Badui</td>
</tr>
<tr>
<td>A 14-5</td>
<td>Lotus leaves</td>
<td>Nursery, Dazhou City</td>
</tr>
</tbody>
</table>

Figures 1-3. Figure 1A-B, Sporangia of *Phytophthora insolita*. Figure 2A-C, Hyphal swellings of *Phytophthora insolita*. Figure 3A-B, Oospores of *Phytophthora insolita*. Note the thin oogonial wall (Og) and the thick oospore wall (Os). Scale bar: 20 μm. All at the same magnification.
with a maximal temperature for growth at 40°C. Sporangia were not found on agar plates but readily produced overnight in water. They were oval to obpyriform, nonpapillate, nondeciduous and terminal on unbranched or irregularly branched sporangiophere, which displayed internal proliferation to form new sporangia nested within or outside the sporangium (Figure 1A-B). Laterally bi-flagellate zoospores were produced within the sporangium proper. The sporangia measured (27-)-36-51(-62) × (22-)-27-39(-51) μm with length/breadth ratio of (1.1)-1.3-1.6(-1.9). Hyphal swellings (Figure 2A-C) were commonly found, terminal or intercalary, single or catenuous, often lateral and sessile. The shape of hyphal swellings ranged from spherical (12-)-21-30(-31) μm diam to oval or obpyriform. Oogonia with oospores were produced in 1-5 month old cultures, well sealed with paraphilm and kept in dark at 25°C. They were found only in isolates BDW 2-6, Bds 1-18, Bdl 1-2, Bds 9, JFS 9954, 9955 and 9957, especially around the inocula. The oogonia were yellowish, spherical, measuring (27-)-29-33(-37) μm diam, and the spherical oospores, (26-)-27-30(-32) μm diam almost filled up the oogonia (Figure 3A-B). The oospore wall was (2.5-)-3-4(-6) μm thick. No antheridia could be found.

By artificial inoculation, *P. insolita* could not infect unwounded fruit but was pathogenic to wounded apple, avocado, cucumber, eggplant, mango, green pepper and tomato, causing fruit rots. The fungus was re-isolated from the diseased plant tissues in all cases.

### Discussion

In addition to the absence of antheridia, *P. insolita* is also characterized by its appressed, chrysanthemum growth pattern, high maximal temperature for growth (39-40°C), nondeciduous, nonpapillate, internally proliferating sporangia with low length/breadth ratio and hyphal swellings of medium sizes (Ann and Ko, 1994; Ho et al., 1995). For comparison, the morphological characteristics of *P. insolita* reported in literature are summarized in Table 2. The production of oospores only in old cultures of some isolates of this species on Hainan Island is of special interest. Ann and Ko (1994) confirmed the identity of the asexual isolates of *P. insolita* by comparing the electrophoretic patterns of soluble proteins of these isolates with the sexual isolates. They suggested that “in nature, the asexual isolates of *P. insolita* probably originated from sexual isolates of *P. insolita* by losing their ability to produce oospores.” The discovery of *P. insolita* isolates on Hainan Island, which produce oospores only in aged cultures, tends to support their hypothesis. These sexual isolates may be in the process of losing their ability to produce oospores. The thickness of the oospore wall may be influenced by the age and the condition of the culture. Whereas the thickness of oospores from 5-day old cultures was 2.4 μm (Ann and Ko, 1980), 4-6 μm from 2-week old cultures (Ho et al., 1995), oospore wall from 1-5 month cultures in present study measured (2.5-)-3-4(-6) μm thick.

The distribution of *P. insolita* is also intriguing. Ann and Ko (1994) concluded that the “high temperature variant of *P. megasperma* Drechsler” from alfalfa in Southern California (Ribeiro et al., 1978) should be re-classified as *P. insolita*. Otherwise, all isolates of this species have been reported from Taiwan only. Taking into consideration the occurrence of *P. insolita* on Hainan Island, the fungus is now known to exist on both sides of the Pacific Basin. Whereas Hainan Island and Taiwan are similar in having tropical to subtropical maritime climate and a central mountainous region surrounded by flat agricultural plains around the edge of the islands, southern California is different in having hot and dry weather and flat topography. We should determine if *P. insolita* exists in other parts of the world before speculating on the origin of this species. Nevertheless, since most of the isolates on Hainan Island were obtained from relatively undisturbed and protected forests on high mountains, *P. insolita* is possibly indigenous to the island. It is not clear how important *P. insolita* is as a plant pathogen in nature. It is pathogenic to alfalfa root (Ribeiro et al., 1978), causes damage to poinsettia stem base (Ann and Ko, 1990), and is associated with strawberry fruit rot (Chang, 1988). The strawberry isolate was initially identified as a “high temperature variant of *P. fragariae*” but was reassigned to *P. insolita* (Ann and Ko, 1994). Although *P. insolita* was pathogenic to various wounded fruits by artificial inoculation in the present study, no report on plant diseases on Hainan Island attributable to *P. insolita* has appeared.

### Literature Cited


海南島之缺雄疫霉（Phytophthora insolita）

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從中國南方海南島森林土壤、溪河水及池塘水中分別到缺雄疫霉（Phytophthora insolita）。這是在台灣以外的地區首次報告發現缺雄疫霉菌。文中還討論缺雄疫霉發現的意義。

關鍵詞：中國；海南省；疫霉菌；缺雄疫霉。