The rediscovery of *Phytophthora polygoni* Saw.

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(Received January 14, 1998; Accepted March 16, 1998)

Abstract. *Phytophthora polygoni* Saw. was first described as a new species by Sawada in 1922, causing leaf spot disease of *Polygonum japonicum* Meisn. in Taiwan. However, there has been no report of the fungus since its first discovery. Recently, a similar disease occurred on *Rumex dentatus* L. in Nanjing of Jiangsu Province in mainland China and the pathogen was identified as *P. polygoni* based on the similar biological characteristics.

Keywords: *Phytophthora polygoni*; *Rumex dentatus*.

Introduction

*Phytophthora polygoni* Saw. was first discovered and described on the leaves of *Polygonum japonicum* Meisn. (Family Polygonaceae) in Taipei and Taichung of Taiwan causing leaf spot disease from the late winter/early spring to early summer (Sawada, 1922). The disease first appeared as small, round yellow green spots which became dull yellow brown with a tint of purple. Subsequently, the lesions enlarged and turned brown and diffuse, resulting in the death of the lower leaves.

The hyphae of the pathogen were intercellular within the leaf spots, aseptate, colorless and branching with hyphal diameter 5–15 µm (usually 12–14 µm) wide. Appresoria in host cells were mostly obovate in shape. Intercellular hyphae grew through the stomata to form clusters of slender, unbranched or sympodially branched sporangiophores, 15–75 µm long and 1.5–5 µm wide (4–20 sporangiophores per stoma) each terminating in a sporangium with a conical papillum. The sporangia were mostly elongated ellipsoidal to fusiform or pyriform, 17–42 × 4–20 µm. When placed in water, a sporangium germinated either by a germ tube or by releasing motile laterally biflagellated ovoid zoospores with obtuse apex and rounded base, 11–15 × 6–9 µm, encysting to become 10–15 µm in diameter. Neither chlamydospores nor resting sporangia were observed, and no sexual structures were found. Attempts to grow the fungus on culture media failed.

Unfortunately, type specimen of *P. polygoni* is no longer in existence for re-evaluation, and the fungus has not been reported again since its first description. Efforts were thus made to locate *Polygonum* and related genera within the family Polygonaceae, with the symptoms as described by Sawada, hoping that *P. polygoni* could be rediscovered.

Materials and Methods

One genus of the family Polygonaceae, *Rumex* is a common weed in the Nanjing region of Jiangsu Province in mainland China. Field trips were made to detect and collect *Rumex* plants whose leaves showed leaf spot disease as described by Sawada (1922).

Freshly collected leaf specimens were prevented from desiccation by keeping them in a petri dish with two pieces of moist filter paper on the bottom. They were exposed to natural daylight for 28–48 h at 18–20°C to induce sporangial formation from leaf lesions. The diseased leaf was then examined under microscope at ×150 for the presence of sporangia, which were then carefully scraped off on a glass slide and mounted in a drop of lactophenol, with or without 0.05% cotton blue.

In order to isolate the fungus, two kinds of culture media were used: Lima bean agar (Zhou et al., 1997) and V-8 agar (Ann et al., 1990). Prior to pouring agar medium into sterilized petri dishes, 50 µg/ml each of penicillin, rifampicin, and PCNB (Pentachloronitrobenzene) were added. Diseased leaf was not surface-sterilized, but washed clean with running tap water and allowed to air-dry. Six small pieces of diseased tissue (about 4 × 4 mm) were cut out with a pair of scissors and placed along the edge of an agar plate, at equal distance from one another. At least three agar plates were used for each diseased leaf. In addition, diseased leaf tissues were also surface-sterilized for 5 minutes in 2% chlorine solution prepared by dissolving 1 bleach tablet in 10 ml distilled water. The bleach tablets are produced by Xuhang Chemical Company in Shanghai and marketed by Shanghai Pharmaceutical Company. The sterilized plant material was placed on Lima bean agar and V-8 agar plates without the addition of the above antimicrobial chemicals. The inoculated plates were incubated in dark at 18°C or 25°C for at least one week.
Results

Leaf spot disease was commonly seen on *Rumex dentatus* L. in Nanjing City in the spring (late March to Mid-April) when the outdoor temperature was about 5–20°C. Diseased specimens were collected from early spring till as late as early June when the temperature rose to 33°C.

The lesion spots on the leaf were roundish to irregular in shape with diffuse edges. Initially, the lesion appeared as blurry, pinkish brown spots on the leaf surface (Figure 1A), but the color was not altered or light pink when viewed from the leaf bottom (Figure 1B). No significant senescence of leaf cells was observed in the lesion. The leaf spots then changed to reddish brown in color and eventually turned light brown to greyish black. Under humid conditions downy mildew-like fungus could be vaguely observed on the surface of a fresh leaf specimen’s lesions. Under the microscope, clusters of ten to more than thirty sporangia could be seen emerging from a single stoma on the leaf surface. Fewer sporangia were found on old lesions with mostly dead leaf cells.

The fungal hyphae (8–12 µm wide) were colorless, aseptate, branching freely between leaf cells within the lesions sending oval haustoria into host cells. The sporangia were small, mostly ellipsoidal to elongated ellipsoidal, fusiform or obpyriform, each with a tapered base and a slightly elongated beak (Figure 2A–C). Sporangia measured (18.9-) 37.1 (-28.8) × (11.3-) 17.7 (-23.8) µm with length/breadth ratio of (1.86-) 2.1 (-2.8) and were semi-papillate with the apical thickening (0.5-) 1.7 (-2.7) µm deep. Zoospores were formed within the sporangium and expelled through the sporangial apex via an exit pore (5.0-) 6.4 (-7.5) µm in width. The sporangiophores were (1.8-) 2.2 (-2.5) wide, unbranched or occasionally branched sympodially in later stages. Mature sporangia were often deciduous each with a medium pedicel 20–45 µm in length, from which it was cut off by a basal cross wall.

Sexual structures were not found in diseased tissues, cleared by immersing in lactophenol or 2% potassium hydroxide solution and then examined under microscope, stained or unstained with 0.05% cotton blue. No fungal growth was detected on Lima bean agar or V-8 juice agar medium, despite numerous attempts.

![Figure 1](image_url). Leaf spot disease symptoms of *Rumex dentatus* caused by *Phytophthora polygoni*, (from left to right, early, intermediate, and final stages). (A) Upper surface of diseased leaf; (B) Lower surface of diseased leaf.
spite numerous attempts. However, the pathogenicity of *P. polygoni* to *Rumex dentatus* still needs to be tested by artificial inoculation. The absence of sex organs within the host plant can be attributable to the possible heterothallism of the fungus. The inoculating of zoospores from sporangia collected from different localities on the same leaf may increase the chance of finding the sexual stage. On the other hand, the production of sex organs may be induced by manipulating the environmental conditions. The absence of sex organ description has not prevented Stamps et al. (1990) from including *P. gonapodyides* (Peterson) Buisman, *P. japonica* Waterh., *P. undulata* (Petersen) Dick and *P. palmivora* var. *heterocystica* Babacauh in their revised tabular key to the species of *Phytophthora*. Although *P. botryosa* Chee produces small, narrowly elongated sporangia it can be easily distinguished from *P. polygoni* due to its ease of growth on common agar plates (Chee, 1969). Of those few *Phytophthora* species which are not known in culture, and produce semipapillate sporangia on the host plant (Stamps et al., 1990; Waterhouse, 1963), *P. cyperi* (Ideta) S. Ito, *P. cyperi-bulbosa* Seethal. & Ramak., and *P. lepironiae* Saw. all produce sex organs and significantly larger sporangia, distinct from *P. polygoni*. Thus, *P. polygoni* should be accepted as an identifiable *Phytophthora* species. Further attempts will be made to search for the sexual stage of *P. polygoni* in the field and under experimental conditions in the laboratory or greenhouse, in order to elucidate the life cycle of this obscure fungus.

Dried diseased specimens of *Rumex dentatus* with *P. polygoni* sporangia have been deposited in the herbarium of the Department of Plant Protection at Nanjing Agricultural University.

**Acknowledgements.** The work was supported in part by a grant from the National Geographic Society to H.H. Ho.

**Literature Cited**


**Figure 2.** Detached sporangia of *Phytophthora polygoni*. (A) Shallow apical thickening (a); (B) and (C). Beaked papillum (b) with pedicel of medium length (p) and basal septum (s). Scale bar equals 10 µm. All at same magnification.