**POLYALTHIA LONGIFOLIA** a new host record of *Nigrospora sphaerica* causing leaf blight in Pakistan and efficacy determination of various fungicides against this pathogen

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### A B S T R A C T

Severe leaf and stem necrosis of *Polyalthia longifolia* (Ulta-Ashoka) was observed in Pakistan during summer 2012. Symptoms on leaves were initiated with drying followed by whole leaf and plant drying. The causal pathogen was identified as *Nigrospora sphaerica* based on cultural characteristics and conidial morphology. Its pathogenicity was proven on healthy *P. longifolia* plants. This is the first report of *Nigrospora sphaerica* causing leaf blight on *P. longifolia* in Pakistan. Furthermore, for the selection of most effective fungicide that can be used for the chemical management of the disease under field conditions three fungicides viz., Antracol (ChloroThalonil), Halonil (Propineb) and Aerosil (Thiophenate methyl) were evaluated against mycelia growth of *Nigrospora sphaerica* under controlled conditions. Two concentrations were used for every fungicide as it was 500 and 1000 ppm. It was concluded from the results obtained that Halonil was found to be effective fungicide by reducing the mycelia growth of *Nigrospora sphaerica* by 78.4%. followed by Antracol and Aerosol respectively.

**Keywords:** *Nigrospora sphaerica*, Ulta-Ashoka, ornamental plant, first disease report and fungicides.

### INTRODUCTION

*Polyalthia longifolia* (also known as mast tree, cemetery tree, Ulta Ashoka) is a tall, ornamental and evergreen tree (Mitra, 1993; Warrier *et al.*, 2002). The genus of *P. longifolia* comprises of about 120 species and belongs to family Annonaceae (Mitra. 1993). *Polyalthia longifolia* is cultivated widely throughout tropical and subtropical asia as an ornamental and has use as an anthelmintic, germicide, and as a folk medicine for the treatment of pyrexia. Previous studies on its leaves, bark, roots, root bark, and seeds have revealed that various types of diterpenoids and alkaloids are present which plays important role in numerous biological activities such as anti-inflammatory, antihypertensive, antimicrobial and cytotoxic effects (Chen *et al.*, 2000). Its bark has medicinal properties to cure fever, skin diseases, diabetes, hypertension, helminthiasis and vitiated conditions of vata and pitta (Warrier *et al.*, 2002). The fungicidal effect of *P. longifolia* has also been reported by many workers (Nair *et al.*, 2007). In May 2012, *P. longifolia* plants growing in avenue and in landscape areas of University of Agriculture, Faisalabad- Pakistan, along roads and ornamentals plant nurseries of district Faisalabad, Pakistan were found to be heavily affected by leaf blight disease. (Haq *et al.*, 2014) Initial symptoms on leaves started with drying of leaves from margins. Severely infected leaves turn purple or reddish color, giving the leave an overall purple cast followed by complete leaf necrosis on whole plant, resulting in severe defoliation. Under severe infections branches become dried and occasionally cause plant death (Figure 1&2). The samples were collected from different localities of University of Agriculture, Faisalabad. Isolation of the causal fungus was made from diseased tissue. On isolation, different conidia that were black, single celled, globose (14-18μm), borne on a hyaline vesicle at the tip of the conidiophore were obtained and...
these characteristics were matched with the characteristics of *Nigrospora sphaerica* and hence it was identified as conidia of *Nigrospora sphaerica* on the basis of these morphological characteristics (Figure 3). To confirm and fulfil the Koch’s postulates, the pathogenicity test was conducted and *N. sphaerica* was re isolated from infected leaves. Furthermore for the selection of most effective fungicide that can be used for the chemical management of the disease under field conditions three fungicides viz., Antracol (Chlorothalonil), Halonil (Propineb) and Aerosil (Thiophenate methyl) were evaluated against mycelia growth of *Nigrospora sphaerica* under controlled conditions.

Figure 1. *P. longifolia* leaves representing drying of margin

Figure 2. Naturally infected plants *P. longifolia* showing necrosis on whole plant

Figure 3. Spores of *N. sphaerica*

Figure 4. Artificially inoculated plants of *P. longifolia* representing severe leaf necrosis
MATERIALS AND METHODS

For the isolation and identification of the involved pathogen symptomatic *P. longifolia* leaves were collected from naturally infected plants. These leaves were cut into small pieces (5mm) and surface sterilized in 0.25% sodium hypochlorite solution for 3 min, rinsed 3 times in sterilized water and dried on blotter paper under sterile air flow conditions. These small segments were then transferred into petri plates containing 20% potato dextrose agar medium and incubated at 25°C under a 12-h light and dark regime. Different fungicides (Antracol, Halonil and Aerosol) was used in vitro to evaluate their effect on colony growth of the fungus following poison food technique (Dhingra and Sinclair, 1985). A representative specimen of *N. sphaerica* from *P. longifolia* was deposited in the Herbarium of First Fungal Culture Bank of Pakistan (FCBP), Institute of Agricultural Sciences (IAGS), Punjab University (PU) Lahore, on living leaves of *A. philoxeroides* with Accession No. of FCBP#1209.

To confirm the pathogenicity of obtained isolate, *N. sphaerica* was multiplied on potato dextrose agar medium at 25°C under a cool-white florescent diurnal light with a 12 h dark and light cycle. After 10 days, conidia were harvested by applying sterilized deionized water (containing 0.01% Tween 20) and gently scraping the culture surface with a microscopic glass slide. The conidial suspension was filtered through a 0.5 mm2 pore strainer to remove mycelia debris (Akhtar et al., 2011; 2012). The resulting spore suspension was adjusted to 106 conidia per ml using a 59. haemocytometer (Farr and Rossman, 2010). Koch’s postulates were fulfilled using the whole plant assay. For this purpose, one month old, 15 plants of *P. longifolia* were sprayed with the prepared conidial suspension (106 conidia /ml) and were kept in net-house under natural conditions (average maximum 45°C, minimum 41°C temperature, and relative humidity 65%). A similar set sprayed with distilled water only was included as a negative control. Fungicial suspensions of different concentrations were prepared by dissolving requisite quantities of each fungicide in warm PDA. About 15 ml of sterilized medium was poured in each 9 cm sterilized Petri dish. After solidification, the plates were inoculated by placing 5 mm discs of 3 days old PDA cultures of the fungus. Three replicated plates were used for each concentration of every fungicide. Three replicated PDA plates received no fungicides was served as control. The inoculated plates were incubated at 28°C and data on the radial colony diameters was recorded after 4-5 days of incubation when the growth of the control plates completely covered the plate. Inhibition of radial growth was computed based on colony diameters on control plate using the following formula. (Sunder et al., 1995).

\[
\text{% Inhibition} = \frac{X - Y}{X} \times 100
\]

Where,

- \(X\) = Growth of control plate
- \(Y\) = Growth of fungicide treated plate.

The collected data was analyzed statistically by employing the Fisher analysis of variance technique (Steel et al., 1997) and treatment means will be compared by using Least Significance Difference (LSD) test at 5% probability level.

RESULTS AND DISCUSSIONS

Initially colonies of fungus were white, then changing to brown to black due to the onset and abundance of sporulation. Conidiogenous cells on superficial hyphae, lateral or terminal, swollen, ampulliform, 8–11 μm in diameter, hyaline, with a single conidium at the attenuated apex. Conidia were black, single celled, globose (14-18μm), borne on a hyaline vesicle at the isolated pathogen coincided with the description of *Nigrospora sphaerica* (Ellis, 1971) therefore, the pathogen was identified as *N. sphaerica*. Disease symptoms were started on inoculated plants as drying of leaves from margin after 3 weeks of inoculation followed by complete leaf necrosis 20 days after inoculation (Figure 4) and *N. sphaerica* was re-isolated from infected leaves. However, plants sprayed with distilled water (negative control) remained green till the end of experiment. *Nigrospora sphaerica* is a cosmopolitan species and has been reported as a leaf blight/pathogen on several hosts worldwide including mulathi (*Glycyrrhiza glabra*) in India (Verma and Gupta, 2008), Cucuma (*Cucuma wenyujin*) in China (Zhang et al., 2011), Chinese wisteria (*Wisteria sinensis*) in Turkey. This is the first report of *N. sphaerica* as a causal pathogen of leaf and stem blight of *P. longifolia* in Pakistan. Identification of *N. sphaerica* as causal agent of *P. longifolia* leaf blight in Pakistan will be helpful to suggest effective control measures to manage this disease in future.
Effect of different fungicides (at concentration of 500 ppm) on mycelia growth of *N. sphaerica* after third week: All fungicides with concentration of 500 ppm inhibit the fungal growth of *Nigrospora sphaerica* presented in Figure 5. There were three fungicides named Antracol, Halonil, and Aerosol were evaluated against *Nigrospora sphaerica*. There was variation in effectiveness among test fungicides in accordance of reducing the fungal growth considerably. It was noted that Halonil was statistically significant in reducing growth of fungus *Nigrospora sphaerica* and showed about 3.0 cm colony diameters, followed by Antracol with 4.9 cm, at concentration of 500 ppm. Aerosol at concentration of 500 ppm was statistically least significant with 5.7 cm.

![Figure 5](image)

Figure 5. Effect of different fungicides (at concentration of 500 ppm) on mycelia growth of *N. sphaerica* after second week. All fungicides with concentration of 1000 ppm inhibited the fungal growth of *Nigrospora sphaerica* presented in Figure 6. There were three fungicides named Antracol, Halonil, and Aerosol were evaluated against *Nigrospora sphaerica*. There was variation in effectiveness among test fungicides in accordance of reducing the fungal growth considerably. It was noted that Halonil was statistically significant in reducing growth of fungus *Nigrospora sphaerica* and showed about 1.7 cm colony diameters, followed by Antracol with 4.9 cm, at concentration of 1000 ppm. Aerosol at concentration of 1000 ppm was statistically least significant with 5.4 cm.

![Figure 6](image)

Figure 6. Effect of different fungicides (at concentration of 1000 ppm) on mycelia growth of *N. sphaerica* after second week.
Three different fungicides (Antracol, Halonil and Aerosil) were used in *in vitro* to evaluate their effect on colony growth of the fungus following poison food technique. Two concentrations viz: 500 ppm 1000 ppm was prepared for each fungicide. Three replications were made for each treatment. Out of three fungicides used, Halonil found to be most effective fungicide by reducing the mycelia growth of *Nigrospora sphaerica* by 80% and followed by Antracol and Aerosil by 47%, 40% respectively.

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