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Evaluation of effectivity of phytoextracts on radial growth of *Sclerotinia sclerotiorum*

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Abstract

Sclerotinia stem rot caused by *Sclerotinia sclerotiorum* (Lib.) de Bary is the most damaging disease and has been recorded in rapeseed-mustard cultivating nations of the world. It is more common and severe in temperate and sub-tropical regions during cool and wet seasons, although it may also be found in some semi-arid regions where conditions seem unfavourable for disease development. Botanical extracts are used as fungicides to control fungal growth and detection of new antifungal compounds which have no side effects on the environment or animal health. The evaluation of effectiveness of different phytoextracts were carried out through food poisoning techniques. Application of phytoextracts through food poisoning technique significantly reduced the myceliogenic growth of *Sclerotinia sclerotiorum*. *Allium sativum* registered maximum percent of mycelial inhibition of *S. sclerotiorum*, followed by *Citrus lemon*, *A. indica*, *M. koenigii* and *A. cepa*. Concentration wise, greater myceliogenic inhibition was observed in 15% fallowed by 10% and 5% phytoextract containing petri plates. Absolute mycelial inhibition was observed in *A. sativum* (15 and 10% concentrations) treated plates. On the other hand, least efficacy of the phytoextract was observed in 5% *A. cepa* treated plate. The current *in vitro* studies of botanicals mycelial growth of *S. sclerotiorum* revealed that phytoextracts are an important component in crop disease management.

Keywords: Sclerotinia sclerotiorum, phytoextracts, radial growth

Introduction

Sclerotinia stem rot caused by Sclerotinia sclerotiorum (Lib.) de Bary is the most damaging disease and has been recorded in rapeseed-mustard cultivating nations of the world (Sharma 2015) [15]. It is more common and severe in temperate and sub-tropical regions during cool and wet seasons, although it may also be found in some semi-arid regions where conditions seem unfavourable for disease development. Disease outbreaks in drier areas occur in irrigated fields because irrigation provides favourable conditions for disease development (Sharma, 2014) [11].

In India, Sclerotinia stem rot was first reported by Shaw and Ajreker (1915) [18] from Pusa (Bihar). Later on, the occurrence of this disease has also been observed in other parts of the country (Roy and Saikia 1976) [27]. This disease is more severe in high land areas in northeastern states and certain areas of Rajasthan (Shivpuri, et al., 2000) [19]. During the eighties and nineties, this disease was of minor importance but over the years due to the adoption of newer crop husbandry practices and lack of host resistance, it has become a serious threat to rapeseed-mustard cultivation (Kolte, 2018) [20]. In Raya (B. juncea) growing areas of Rajasthan and Haryana states where farmers perform mono-cropping and cultivate the crops under irrigated conditions. This pathogen become very destructive and results some time total failure to crop yield. Sclerotinia stem rot is a serious problem in major rapeseedmustard growing areas of the country, such as Uttar Pradesh, Rajasthan, Haryana, Punjab, Madhya Pradesh, West Bengal, Bihar, and Assam (Ghasolia et al., 2004; Sharma et al., 2015) [4, 12]. This disease has now attained next position to Alternaria blight in terms of its economic importance (Parveen et al., 2007) [8]. The yield losses vary considerably with the growth stage of the plant and the number of plants infected. If plants are infected at the early flowering stage produce little or no seed, but those infected at the late flowering stage results in no yield (Kolte, 1985) [6]. Generally, the yield losses due to stem rot in rapeseed-mustard have been recorded from 40-80% in different growing regions of the country, and as high as up to 100% in severe outbreaks in Raya growing areas of Rajasthan and Haryana (Sharma et al., 2001; Singh et al., 2002; Ghasolia et al., 2004) [21, 22, 4].

S. sclerotiorum is a necrotrophic pathogen and infects the plant tissues severely and appears

to be a non-specific pathogen, producing white rot or mold or soft rot symptoms on different plant parts such as leaf, stem, root, and fruit (Sharma *et al.*, 2015) [12]. The disease is monocyclic and symptoms usually appear 4-6 weeks after sowing or at the post-flowering stage. Sudden drooping of leaves followed by drying of plants are characteristic features of the disease. Initial symptoms include the appearance of elongated water-soaked lesions at the base of the stem which usually expands rapidly, becomes bleached, necrotic, and subsequently develops patches of fluffy white mycelium (Sanogo and Puppala 2007) [23].

When the stem is completely girdled and covered by a cottony mycelium growth, it breaks from where it shows rotting and drying (Bolton *et al.*, 2006) ^[3]. However, infection is restricted to a smaller area of pith, resulting in slow stunting of the plant and premature ripening (Kolte, 1988) ^[7]. The fungus initially produces white-coloured small melanized resting mycelial aggregates on the collar or crown region of the stem, later turning into black colour hardened sclerotia. When such infected stem of the host is split large cavities are lined by fluffy mycelium and numerous black-coloured sclerotia are observed (Tripathi *et al.*, 2017) ^[16].

The sclerotia serves as overwintering or over summering structures of the pathogen and are germinated directly with new mycelial growth in presence of nutrients (myceliogenic germination) and are devoid of nutrients they germinate to form apothecia and liberate ascospores (Huang, and Kozub, 1991) [5]. Hyphae from myceliogenically germinated sclerotia infect the basal portion of the stem. During the periods of high moisture and favourable temperature, the sclerotia are germinated carpogenically and produce ascospores which are germinated on the petals deposited on the leaves and act as a nutrient substrate for developing germ tube and infection. These spores act as the main source of primary inoculum and may travel a long-distance field for causing infection and epidemics (Adams and Ayers, 1979; Sun and Yang, 2000) [1, 15]. The devastating nature of S. sclerotiorum is also attributed to its prolonged survival, structure which is often facilitated by the production of vegetative sclerotia that provide primary inoculum for subsequent growing seasons (Saharan and Mehta, 2008; Sharma et al., 2015) [10, 12]. The mono-cropping of rapeseed mustard is reported to favour the disease severity in a specific geographical area, because sclerotia of the pathogen may survive for 4 to 8 years in soil under favourable conditions (Willets and Wong, 1980; Bardin and Huang, 2001) [17, 2].

Materials and Methods

All experiments were carried out under laboratory conditions in the Department of Plant Protection, Faculty of Agricultural Sciences, Aligarh Muslim University, Aligarh (U.P.) India.

Phytoextracts

Antifungal activity of five plant extracts such as Garlic extract (*Allium sativum*), Onion extract (*Allium cepa*), Neem extract (*Azadirachta indica*), Curry leaves (*Murraya koenigii*), lemon leaves (*Citrus lemon*) was evaluated

against S. sclerotiorum, using poison food technique (Dubey and Patel, 2001) [24]. The aforementioned plant parts were collected from AMU, Campus. About 100 gm of fresh plant leaves/rhizome/bulbs were taken and thoroughly washed in distilled water. Such plant parts were cut into small pieces and then ground in mortar and pestle by adding 100 ml of sterilized distilled water. The crude material was then filtered through double layer muslin cloth and then the filtrate was filtered through Whatman no. 1 filter paper. The plant extracts so prepared were heated at 400 °C for 5 minutes to avoid contamination (Jagnathan and Narsimhan, 1988) [25]. The requisite amount of plant extracts was incorporated into sterilized non-solidified potato dextrose agar medium (PDA) and shaken well to make it homogenous. Thereafter, 20 ml of amended medium was then poured into 90 mm Petri plates.

Sterilization of glassware

The sterilization of glassware (wrapped in butter paper/brown paper) was done by autoclaving at 15 *psi* pressure for 30 minutes. Autoclaved glassware was dried in a hot air oven at 80 °C for 45 to 60 minutes. All *in vitro* experiments were carried out under aseptic conditions.

Effect of phytoextracts on radial growth of S. sclerotiorum

The efficacy of five plant extracts was evaluated for their effectiveness in inhibiting the radial growth and sclerotial formation of *S. sclerotiorum* on potato dextrose agar medium. Each treatment was replicated thrice with a suitable check, wherein un-poisoned Petri plates (without plant extracts) were centrally inoculated with the test fungus. Each treatment was replicated thrice with a suitable check, wherein un-poisoned Petri plates (without nanoparticles, dual culture, and plant extracts) were centrally inoculated with the test fungus.

Per cent inhibition (T) = $[(C-T) / C] \times 100$ Where, C = Colony growth diameter (mm) of fungus in check (in the unamended medium). T = Colony growth diameter (mm) of fungus in treatment (in fungicide amended medium).

Results and Discussion

This experiment revealed that plant extracts are important and may be used in sustainable management of plant pathogens. Application of phytoextracts through food poisoning technique significantly reduced the myceliogenic growth of Sclerotinia sclerotiorum. Allium sativum registered maximum percent of mycelial inhibition of S. sclerotiorum, followed by Citrus lemon, A. indica, M. koenigii and A. cepa. Concentration wise, greater myceliogenic inhibition was observed in 15% fallowed by 10% and 5% phytoextract containing petri plates. Absolute mycelial inhibition was observed in A. sativum (15 and 10%) concentrations) treated plates. On the other hand, least efficacy of the phytoextract was observed in 5% A. cepa treated plate (Table 1). The current in vitro studies of botanicals mycelial growth of S. sclerotiorum revealed that phytoextracts are an important component in crop disease management.

Table 1: Effect of phytoextract on radial growth of *Sclerotinia sclerotiorum*

		Concentrations						
5%		10%			15%			
Radial growth (mm)	% Inhibition	Radial growth (mi	m) % Inhibit	ion	Radial growth (mm)		% Inhibition	
Azadirachta indica	55.5	38.33	27.4	(69.55	16.7	81.44	
Allium sativum	15	83.33	0		100	0	100	
Allium cepa	72	20	65	1	27.77	60	33.33	
Murraya konini	34	62.22	33	(63.33	19.6	78.22	
Citrus limon	22.5	75	20	,	77.77	12.5	86.11	
control		90		90			90	
LSD P ≤0.05		6.75		4.32			3.85	



Fig 1: Effect of plant extracts on radial growth of S. sclerotiorum

Conclusion

Mustard plants were found severely suffering from stem rot disease caused by *Sclerotinia sclerotiorum* which is a very devastating pathogen. The effectiveness of phytoextracts was also evaluated and it was observed that plant-based products may be used for the sustainable management of sclerotinia stem rot. The evaluation of effectiveness of different phytoextracts were carried out through food poisoning techniques. Among the phytoextracts, *Allium sativum* showed absolute inhibition in the myceliogenic

growth of S. sclerotiorum followed by Citrus lemon, Azadirachta indica, Murraya koenigii and Allium cepa.

References

- Adams PB, Ayers WA. Ecology of Sclerotinia species; c1979.
- 2. Bardin SD, Huang HC. Research on biology and control of Sclerotinia diseases in Canada. Can. J Plant Pathol. 2001;23:88-98.
- 3. Bolton DM, Thomma PHJB, Nelson DB. *Sclerotinia sclerotiorum* (Lib.) de Bary: biology and molecular

- traits of a cosmopolitan pathogen. Mol. J Plant Pathol. 2006;7:1-16.
- 4. Ghasolia RP, Shivpuri A, Bhargava AK. Sclerotinia rot of Indian mustard in Rajasthan. Indian Phytopathol. 2004;57(1):76-79.
- 5. Huang HC, Kozub GC. Temperature requirements for carpogenic germination of sclerotia. Bot Bull Acad. Sin; c1991. p. 32.
- 6. Kolte SJ. Diseases of Annual Edible Oilseed Crops, Rapeseed-Mustard and Sesame Diseases. Boca Raton: CRC Press Inc.; c1985.
- Kolte SJ. Diseases of Annual Edible Oilseed Crops. Rapeseed Mustard and Sesame Diseases. Florida: CRC Press Inc. 1988;2:35-39.
- 8. Parveen K, Haseeb A, Shukla PK. Management of *Sclerotinia sclerotiorum* on *Mentha arvensis* C.V. Gomti. J Mycol Pl Pathol. 2007;37(1):33-36.
- Yadav SK, Singh RB, Trivedi H, Singh R. Studies on insect-pests complex in mustard (*Brassica juncea* L.).
 Int. J Biol. Sci. 2022;4(1):106-109. DOI: 10.33545/26649926.2022.v4.i1b.96
- Saharan GS, Mehta N. The disease and symptoms. In: Saharan GS, Mehta N, editors. Sclerotinia Diseases of Crop Plants: Biology, Ecology and Disease Management; c2008. p. 47-70.
- 11. Sharma P, Meena PD, Singh D. Effect of *Sclerotinia sclerotiorum* culture filtrate on seed germination and seedling vigour of Indian mustard (*Brassica juncea* cv. Rohini). J Oilseed Brassica. 2014;5:158-161.
- Sharma P, Meena PD, Verma PR, Saharan GS, Mehta N, Singh D, et al. Sclerotinia sclerotiorum (Lib) de Bary causing Sclerotinia rot in oilseed Brassicas: A review. J Oilseed Brassica. 2015;1(2):1-44.
- 13. Shivpuri A, Sharma KB, Chhipa HP. Some studies on the stem rot *Sclerotinia sclerotiorum* disease of rapeseed-mustard in Rajasthan. J Mycol. Pl Pathol. 2002;30(20):268.
- Kutte MM. Effect of garlic (*Allium sativum*) and Ginger (*Zingiber officinale*) on the microbial and sensorial quality of smoked mackerel fish (*Scomber scombrus*).
 Int. J Biol. Sci. 2022;4(1):188-191.
 DOI: 10.33545/26649926.2022.v4.i1c.105
- 15. Sun P, Yang XB. Light, temperature, and moisture effects on apothecium production of *Sclerotinia sclerotiorum*. Plant Dis. 2000;84:1287-1293.
- Tripathi AN, Pandey KK, Manjunath M, Meena BR, Rai AB, Singh B, et al. Morphological characterization, cross infectivity and chemo-sensitivity of Sclerotinia sclerotiorum isolates towards bio-agent and new molecules of fungicides. Veg. Sci. 2017;44(1):103-106.
- 17. Willetts HJ, Wong JAL. The biology of *Sclerotinia* sclerotiorum, S. Trifolium, and S. minor with an emphasis on specific nomenclature. Bot Rev. 1980;46:100-165.
- 18. Shaw FJW, Ajrekar SL. The genus Rhizoctonia in India. Dep. Agric. Indian Bot. Surv. 1915;7:177-194.
- 19. Zimmerman JJ, Gabbert D, Shivpuri C, Kayata S, Miller J, Ciesielski W, *et al.* Meter-dosed, inhaled beclomethasone initiated at birth to prevent bronchopulmonary dysplasia. Pediatric Critical Care Medicine. 2000 Oct 1;1(2):140-5.
- 20. Kolte SJ. Diseases of annual edible oilseed crops: rapeseed-mustard and sesame diseases. CRC press. 2018 Jan 18, 2.

- 21. Sharma R, Chisti Y, Banerjee UC. Production, purification, characterization, and applications of lipases. Biotechnology advances. 2001 Dec 1;19(8):627-62.
- 22. Singh HP, Batish DR, Kaur S, Ramezani H, Kohli RK. Comparative phytotoxicity of four monoterpenes against *Cassia occidentalis*. Annals of Applied Biology. 2002 Oct;141(2):111-6.
- 23. Sanogo S, Puppala N. Characterization of a darkly pigmented mycelial isolate of *Sclerotinia sclerotiorum* on Valencia peanut in New Mexico. Plant Disease. 2007 Sep;91(9):1077-1082.
- 24. Dubey SC, Patel B. Evaluation of fungal antagonists against *Thanatephorus* causing web blight of URD and mung bean cucumens. Indian Phytopathology. 2001;54(2):206-209.
- 25. Jagannathan R, Narasimhan V. Effect of plant extracts/products on two fungal pathogens of finger millet. Indian Journal of Mycology and Plant Pathology. 1988;18(3):250-254.
- 26. Singh RS, Singh HV, Singh P, Kaur J. A comparison of different substrates for the mass production of Trichoderma. Ann Plant Prot. Sci.; c2001.
- 27. Roy AK, Saikia UN. White blight of mustard and its control. Indian J Agric. Sci. 1976;46:197.