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1 ***In vitro* and *in vivo* antifungal properties of silver nanoparticles against**
2 ***Rhizoctonia solani*, a common agent of rice sheath blight disease**

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12

13 **Abstract**

14 Sheath blight disease in rice has caused major crop losses worldwide. Managing the causal agent of
15 disease *Rhizoctonia solani* Kühn is difficult because of its broad host range and formation of
16 sclerotia which can survive in harsh environmental conditions; therefore developing innovative
17 disease management methods without application of hazardous chemicals has been considered as
18 the main concern to maintain sustainable agriculture. This presented research has revealed the
19 negative impact of Silver nanoparticles (SNPs) on *R. solani* and disease progress both *in vitro* and
20 *in vivo*. The adverse effects of the SNPs on *R. solani* are significantly dependent on the quantity of
21 SNPs, sprayed at different concentrations *in vitro*. The highest inhibition level against sclerotia
22 formation and mycelia growth are 92 and 85%, respectively, at a SNPs concentration of 50 ppm. *In*
23 *vivo* glasshouse experiments also showed that SNPs at the same concentration favorably affects

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24 both the fresh and dry weight of rice plants with a remarkable suppressive effect on the lesion
25 development in leaves.

26
27
28 **Keywords:** Nanotechnology, silver nanoparticles, rice sheath blight disease, sclerotia formation, lesion
29 development

30

31 **1. Introduction**

32 Application of nano-materials has widely influenced drug delivery, cancer therapy [1], energy [2],
33 biomedical [3], agriculture [4] and many other high-tech industries over recent years [5].
34 Nanotechnology has led to the new ways to control diseases using atomic-scale materials [6, 7].
35 The extremely small-scale particles have emerged as modern agents owing to their large surface to
36 volume ratio which provides a large contact surface with pathogen sources [8]. Nanotechnology
37 applications in agriculture can be successful if natural processes are simulated in greater scientific
38 for successful implementation and examples of successful tools at small scale [9]; plant protection
39 products [10]; Fertilizers [11]; Water purification and pollutant remediation [12]; Nanosensors and
40 diagnostic devices [13]; plant genetic modification [14].

41 Among nanoparticles, silver NPs (SNPs) can attack microorganisms, including the cell membrane
42 structure in large-scale biological processes [15, 16]. The antibacterial activity of silver ions has
43 been well established and attributed to the ability of ionized SNPs to penetrate into the bacterial cell
44 wall and to modulate cellular signaling [17]. SNPs with fungistatic, bacteriostatic and plasmonic
45 properties are among the eco-friendly inhibitors against plant-pathogens compared to synthetic
46 fungicides [18]; however the antifungal ability of SNPs has received less attention compared to
47 medical and pharmaceutical sciences with only few studies undertaken against phytopathogenic
48 fungi such as *Alternaria alternata*, *Botrytis cinerea* [19], and *Colletotrichum gloeosporioides* [20].

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49 Rice (*Oryza sativa* L.) is a major food for a large proportion of the world's population, and is an
50 important primary crop in muddy farmlands [21]. Sheath blight disease caused by *Rhizoctonia*
51 *solani* Kühn AG1 (Teleomorph: *Thanatephorus cucumeris*; anastomosis group 1 IA, AG1 IA), is a
52 common destructive disease of rice in all rice-growing regions in the world. Sclerotia germination
53 is a key factor in the dispersion of rice sheath blight disease, hence any potential inhibitor of
54 sclerotia germination, i.e., SNPs, would be essential in order to decrease the inoculum. This impels
55 rice farmers to use a large amount of anti-nature and harmful chemicals annually to control sheath
56 blight disease, which not only adds further costs in the short term but increases devastative damages
57 in the long term to the human health and environment.

58 In this study, in order to control rice sheath blight disease with emphasis on the cleaner production
59 at a lower cost, different concentrations of SNPs were examined as a new antifungal substance to
60 suppress the pathogenic activity of *R. solani* under *in vitro* (to evaluate the inhibitory effects of
61 SNPs on sclerotia formation and mycelia growth) and *in vivo* (to investigate the effects of
62 antifungal activity of SNPs on the rice plant in a glasshouse trial) conditions.

63

64 **2. Materials and Methods**

65 *2.1. Reagent, rice seeds and fungal pathogen source*

66 SNPs suspension was obtained from Nanocide Co., Tehran, with a concentration of 4,000 ppm and
67 an average particle size of 5~10 nm, in dark brown colloid physical form. Rice seeds of *Oryza*
68 *sativa* L. var Hashemi and pure culture of *R. solani* AG-1 IA were obtained from Iran Rice
69 Research Institute (IRRI), Rasht [22]. The *Oryza sativa* L. var Hashemi potentially has a high yield,
70 and is susceptible to sheath blight disease. The fungus was maintained on potato dextrose agar
71 (PDA, Merck Co.) at room temperature.

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72 2.2. *In vitro* examination of inhibitory effects of silver nanoparticles on mycelia and sclerotia of *R.*
73 *solani*

74 To evaluate the *in vitro* antifungal effects of SNPs against *R. solani* AG1, four different
75 concentrations of SNPs suspension (5, 10, 25 and 50 ppm) were added to Petri dishes before
76 pouring plates with PDA. Uniform agar plugs with a diameter of 6 mm containing fungal mycelia
77 were inoculated simultaneously at the center of each Petri dish containing SNPs, followed by
78 incubation at $28 \pm 1^\circ\text{C}$ for three days. The mycelia growth inhibition rate was calculated using Eq.
79 1 [4]:

$$80 \text{ Inhibition rate (RH) \%} = \frac{(R-r)}{R} \times 100 \quad (1)$$

81
82 The parameter *RH* is the inhibition rate, *R* for the mycelium inhibition growth is the expansion in
83 diameter of the mycelial fungus in the control dish (cm), and for the sclerotia formation growth
84 under the inhibition process, *R* is the weight of the sclerotia in the control dish (mg). The parameter
85 *r* for mycelium inhibition growth is the expansion in diameter of the fungus mycelial when treated
86 by SNPs (cm), and for the sclerotia formation growth under the inhibition process, *r* is the weight of
87 sclerotia when treated by SNPs (mg).

88 The antifungal effect of SNPs against *R. solani* sclerotia formation was measured after adding the
89 four concentrations of SNPs to the PDA media content. Inoculated *R. solani* plates were maintained
90 at room temperature for two weeks to manifest sclerotia formation, and the sclerotia formation
91 inhibition rate was calculated using Eq. 1. All tests were carried out in triplicate.

92 The effect of SNPs on the germination of sclerotia was assayed using the following procedure [23].
93 The sclerotia of *R. solani* were formed on PDA at 15°C through incubating the inoculated plates
94 for a week. Uniform sclerotia were collected from PDA plates, and the surface was sterilized in

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95 1.5% sodium hypochlorite solution for 3 minutes. Then, three surface sterilized sclerotia were
96 treated by SNPs of various concentrations and placed in a petri dish, and then they were incubated
97 for a week at 25°C in the dark. The germination rates of sclerotia were measured and compared with
98 the control (without SNPs). The percentage of inhibition against sclerotia was calculated using Eq.
99 1.

100
101 *2.3. In vivo examination of silver nanoparticles on sheath blight disease under glasshouse*
102 *conditions*

103 Rice seeds were sown 3-4 cm below the soil surface of the pots (1 L) and they were separated into
104 six groups with four pots in each group as follows: (a) pathogen alone, (b) pathogen + SNPs (5
105 ppm), (c) pathogen + SNPs (10 ppm), (d) pathogen + SNPs (25 ppm), (e) pathogen + SNPs (50
106 ppm), and (f) control (without SNPs). Rice plants were grown in pots under glasshouse conditions
107 at 30°C and 85-95% relative humidity. As the plants reached their late tiller stage (three-week-old
108 plants) were treated by the inoculation process with *R. solani*. To achieve this, mycelia suspension
109 of *R. solani* (5×10^8 CFU/ml) was evenly sprayed using a hand sprayer on the rice plants [24]. To
110 maintain a fair coverage of SNPs on the foliage throughout the evaluation period, two sprays
111 applied including 24 hrs post inoculation and seven days, subsequently. After inoculating with
112 pathogen and SNPs spraying process, the seedlings were covered with plastic bags for three days to
113 maintain the high humidity. After 15 days, the disease severity was recorded via measuring the
114 fresh weight, dry weight and relative lesion height (*RLH*), according to the 1996 IRRI standard. The
115 relative lesion height of each tiller was calculated using Eq. 2 [25]:

$$116 \quad RLH\% = \frac{\text{Lesion height (cm)}}{\text{Plant height (cm)}} \times 100 \quad (2)$$

117

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118 2.4. *Statistical analysis*

119 Recorded data were subjected to analysis of variance with SAS software (SAS Institute, version 9).

120 Duncan's Multiple Range Test was utilized to compare means.

121

122 **3. Results**

123 3.1. *In vitro* examination of inhibitory effects of silver nanoparticles on mycelia and sclerotia of *R.*

124 *solani*

125 The effects of tested SNPS concentrations on mycelium growth, sclerotia formation and

126 germination are presented in Fig. 1. Plates treated with 50 ppm SNPs revealed the minimum

127 number of sclerotia. For the, the *RHs* of 12, 23, 56 and 92 % were found for the 5, 10, 25 and 50 of

128 SNPs concentrations, respectively. With regards to the mycelia growth, *RHs* of 8, 35, 67 and 85

129 were recorded for the SNPs of concentrations of 5, 10, 25 and 50 ppm, respectively. The *RHs* of 15,

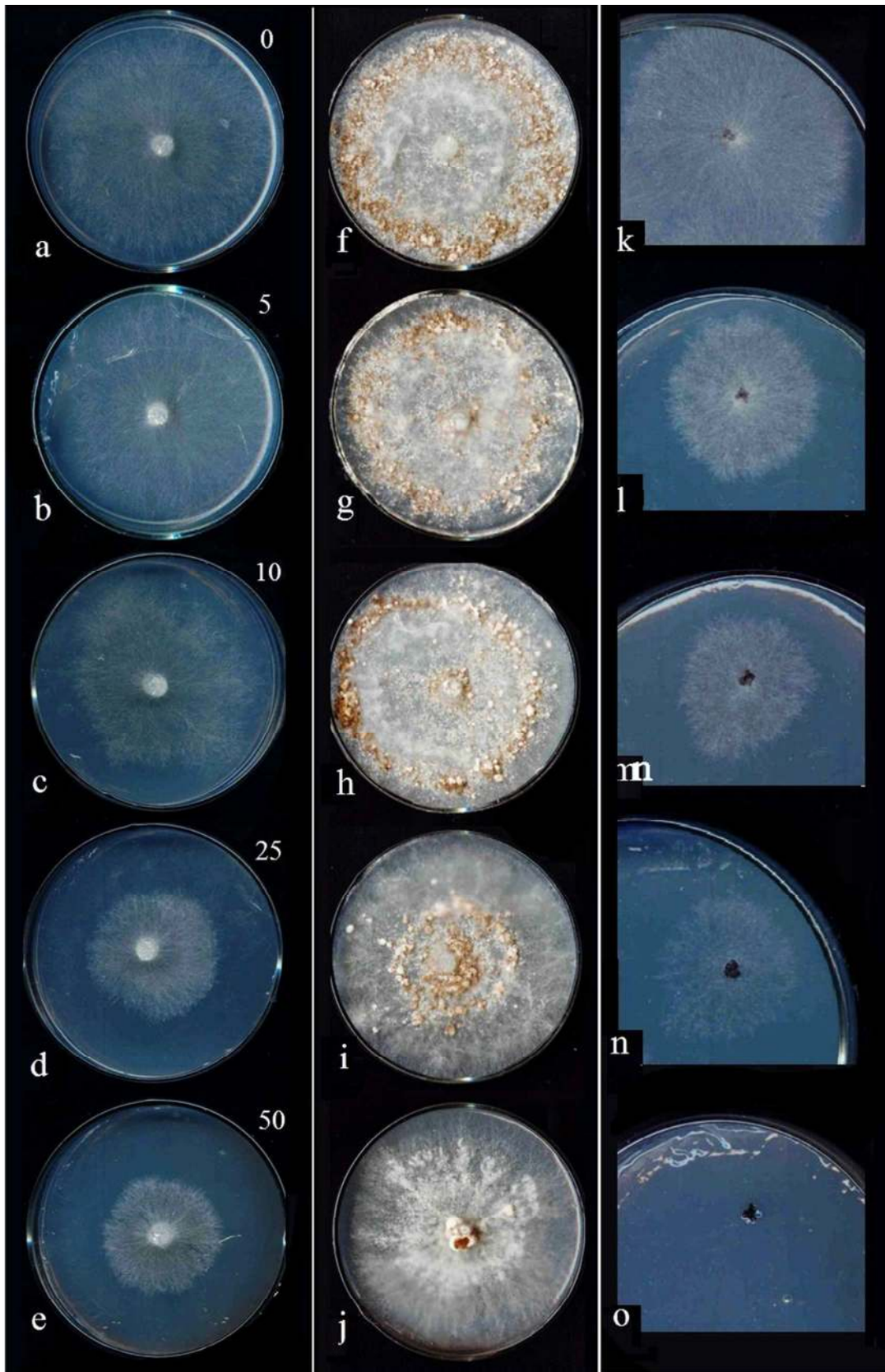
130 26, 57 and 98 % were found for the SNPs concentrations of 5, 10, 25 and 50 ppm, respectively

131 related to sclerotia germination. These results indicate that SNPs has strongly suppressed *R. solani*

132 under *in vitro* condition (Fig. 2).

133

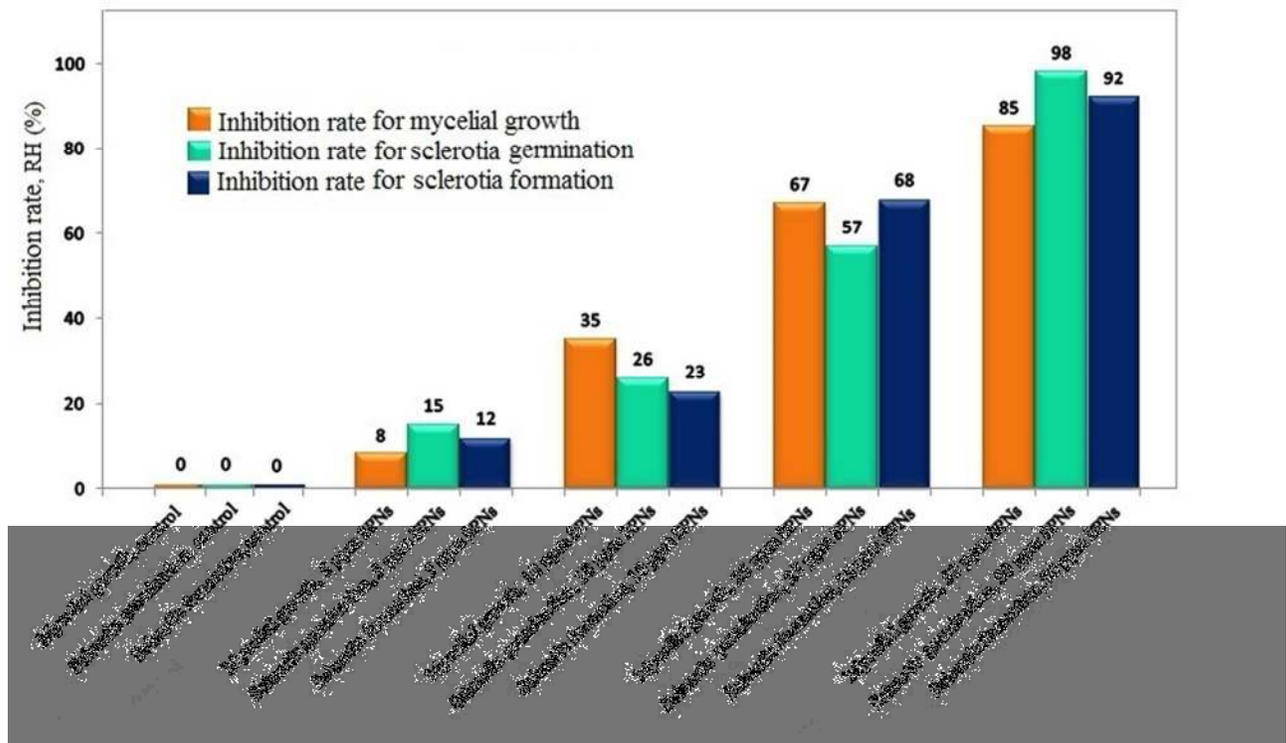
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Fig. 1. *In vitro* inhibitory effects of different concentrations of SNPs (indicated in the top right of left column) on *Rhizoctonia solani* AG1. Mycelia growth stage (left column, a-e); sclerotia formation (middle column, f-j) and sclerotia germination (right column, k-o).

134



135

136 Fig. 2. *In vitro* RH of SNPs effects on mycelia growth, sclerotia germination and sclerotia
137 formation of *Rhizoctonia solani* AG1-IA.

138
139 3.2. *In vivo* examination of silver nanoparticles on sheath blight disease under glasshouse
140 conditions

141 Fig. 3 shows the rice seedlings at the late tiller stage of 90 days under the greenhouse conditions
142 (30°C, 85-95% humidity). The *in vivo* results of SNPs against *R. solani*, the causal agent of sheath
143 blight disease, are presented in Fig. 4. In this figure, A indicates the leaf symptoms resulting from

8

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144 infection by *R. solani* alone, and b-e indicate the pathogen plus the applied 5, 10, 25 and 50 ppm of
 145 SNPs and their different degrees of inhibition in the leave-lesion development. The treatment of
 146 plants with the pathogen without SNPs resulted in typical sheath blight symptoms, but the treated
 147 plants with pathogen + SNPs show different levels of inhibitory effects. According to the results of
 148 ANOVA for different traits, all traits were different at significance level at $P \leq 0.05$. There are
 149 significant reductions in the symptoms of pathogen in pots treated with SNPs. The 5 ppm SNPs
 150 concentration has a small effect on the dry weight of the rice plants (Table 1); by increasing the
 151 concentration of SNPs to 50 ppm, the fresh weight and dry weight increased significantly. At 50
 152 ppm SNPs concentration, the *RLH* of each tiller decreased which evidence that applying SNPs has a
 153 strong antifungal influence on and minimizes lesion in the rice tillers. The comparative results of
 154 the inhibition activity of SNPs against sheath blight revealed significant reduction of lesions on the
 155 rice sheath (Fig. 5).

156
 157 Table 1. The *in vivo* inhibitory effect of SNPs on rice sheath blight under glasshouse conditions

Treatment	Fresh Weight (g)	Dry Weight (g)	RLH %
Control	22.6 ^{*a}	5.8 ^a	-
Pathogen	12.5 ^b	2.3 ^b	95%
Pathogen + SNPs (5 ppm)	13.7 ^b	2.5 ^b	90%
Pathogen + SNPs (10 ppm)	14.2 ^{bc}	2.7 ^{bc}	80%
Pathogen + SNPs (25 ppm)	18.3 ^d	3.8 ^d	55%
Pathogen + SNPs (50 ppm)	20.3 ^{ad}	4.4 ^{ad}	15%

158
 159 *The presented data are the means of four replications, and they are subjected to the analysis with the variance $n=5$
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Fig 3. Rice seedlings after 90 days under greenhouse conditions, with a temperature of 30 °C and constant humidity of 85-95%

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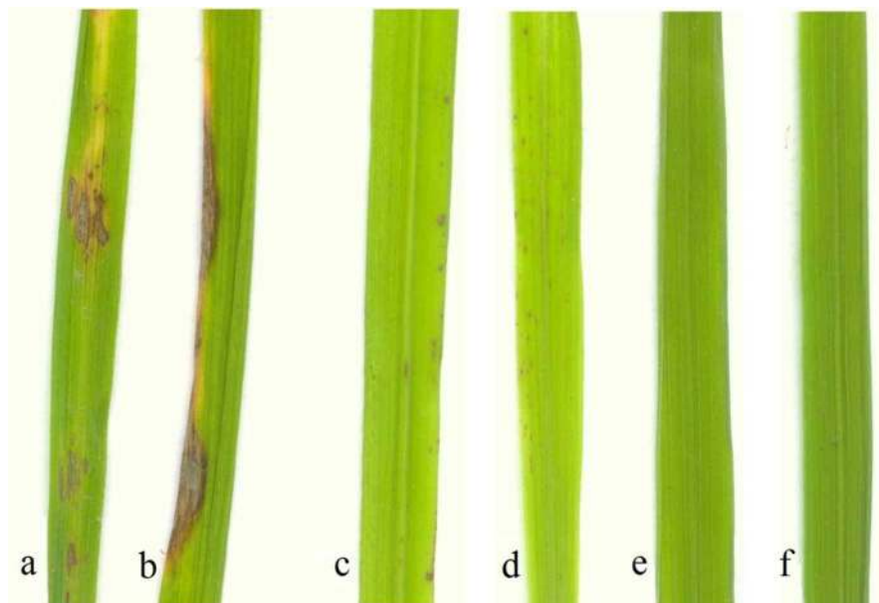
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Fig 4. Effect of SNPs treatment on the lesion development by *R. solani* on rice leaves: a) pathogen alone; b) pathogen + SNPs (5 ppm); c) pathogen + SNPs (10 ppm); d) pathogen + SNPs (25 ppm); e) pathogen + SNPs (50 ppm); f) untreated control.

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Fig 5. Effect of SNPs on the lesion development by *R. solani* on rice sheath:

a) pathogen alone; b) pathogen + SNPs (5 ppm); c) pathogen + SNPs (10 ppm); d) pathogen + SNPs (25 ppm); e) pathogen + SNPs (50 ppm); f) untreated control.

188
189 *3.3. Discussion*
190 SNPs can denature cells by attacking their membranes and structures. A previous research found
191 that SNPs disrupts transport systems, including ion efflux [26]. The dysfunction of ion efflux causes
192 a rapid accumulation of silver ions, interrupting cellular processes such as respiration and
193 metabolism by reacting with the molecules. Ji Seon et al. (2009) showed that upon the treatment by
194 SNPs spray, the hyphal walls were seriously damaged and resulted in the plasmolysis of hyphae.
195 Considering the cellular effects of silver ions, SNPs mediated the collapse in *Sclerotinia*
196 *sclerotiorum* hyphae which damaged the hyphal walls [27].
197 Silver ions are known to deactivate cellular enzymes and DNA by coordinating with electron-
198 donating groups such as thiols, carboxylates, indoles, amides, hydroxyls, etc. [28, 29] According to
199 past studies on SNPs, the smaller the SNPs, the more Ag⁺ ions they release which affects the

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200 performance of microorganisms [30, 31]. As discovered in our investigation, SNPs in an aqueous
201 solution with small sizes can penetrate into the cells of microorganisms and destroy their membrane
202 integrity [32-34].

203 NPs have a vast surface to volume ratio which significantly enhances their property of cell
204 membrane permeability compared with non-NPs forms of the same material [35-37]. NPs are able
205 to penetrate the membranes of microorganisms, leading to cell deformation [38]. NPs, with their
206 large surface to volume ratio, exhibit active antimicrobial properties due to their higher ability to
207 interact with cellular membranes through disruption of the cell wall structure, affecting the
208 respiratory chain and cell division in DNA and proteins as a microorganism [39]. It is likely that the
209 size of SNPs similarly plays a key role in their permeability and antifungal activity. In short, SNPs
210 have an active antifungal activity with great biocide properties, thus, they have the potential to be
211 considered as an economical and eco-friendly pesticide. The application of chemical fungicides
212 adds additional indirect long-term and hidden costs as it causes dangerous side effects in both
213 human health and the environment [40]. The editors of Nature estimated that any technology takes
214 some 20 years to emerge from the laboratory and be commercialized [41]. Application of
215 nanotechnology in agriculture might take a few decades to move from laboratory to land however
216 reasonable expectations would be crucial for this nascent field to blossom [42].

217 The efficacy of SNPs is increased by conjugating the antifungal drug miconazole with SNPs which
218 exhibits significant fungicidal activity [43]. SNPs with chitin inhibits the spore germination of the
219 examined pathogens [44, 45]. Moreover, bioactive capped SNPs was found to be able to control the
220 endophytic fungus of *Colletotrichum gloeosporioides*, *in vitro* [46]. The antifungal activity of SNPs
221 is comparable to those of ionic silver NPs; however, ionic silver remains cytotoxic at the
222 concentrations that inhibit the growth of the examined yeasts [47]. According previous research

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223 both positive and negative effects on plant growth and development was suggested [48]. In some
224 plants SNPs can be increase growth as shoot and root length and biochemical attributes such as:
225 chlorophyll, carbohydrate and protein contents, antioxidant enzymes [49].

226 4. Conclusion

227 *In vitro* and *in vivo* study of the antifungal activity of SNPs at concentrations of 5, 10, 25, and 50
228 ppm was conducted against fungal pathogen *R. solani* to reduce and prevent the sheath blight in rice
229 seedlings. The *in vitro* results showed the *RHs* for mycelial growth, sclerotia formation and
230 sclerotia germination were respectively (8, 35, 67, 85), (12, 23, 56, 92) and (15, 26, 57, 98) for their
231 corresponding SNPs concentrations (5, 10, 25 and 50 ppm). The results clearly show that the *RHs*
232 strongly depend on SNPs concentration, and substantially increase upon an increase in SNPs
233 concentration. In the *in vitro* examination part, we can conclude an increasing trend in the inhibition
234 rate for mycelial growth, sclerotia germination and sclerotia formation with the increasing amount
235 of SPNs. By spraying SNPs on the rice plants, the sheath disease's symptoms on the leaves
236 decreased, and at 50 ppm SNPs concentration, the symptoms completely vanished. In the *in vitro*
237 examination part, we can conclude an increasing trend in the inhibition rate for mycelial growth,
238 sclerotia germination and sclerotia formation with the increasing amount of SPNs. However the
239 author haven't give the results under condition of more than 50 ppm, which can be used to confirm
240 the relationship between inhibition rate and SPNs concentration

241 The *in vivo* results show that the SNPs solution created an antimicrobial layer around the rice plants
242 which protected the plants from pathogens. It was also demonstrated that SNPs highly affect
243 sclerotia formation and germination. SNPs can penetrate into the fungal cell membrane and cell
244 wall, killing microorganism cells. This investigation suggests SNPs can replace chemical pesticides
245 in controlling and inhibiting sheath blight, a common disease in rice.

246

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