

Growth, Carbon Assimilation and Quality of Kesum (*Persicaria Minor*) as Exposed to Zinc Oxide Nanoparticles

ABSTRACT

Aims: This study was conducted to investigate the effect of zinc oxide nanoparticles towards the *Persicaria minor* that can be used as a guidance for further toxicity investigation of ZnO-NPs.

Study design: A Completely Randomized Block Design (RCBD) was used with three replication. Each unit was consisted with eight plants and the total of 96 plants were used in this study.

Place and Duration of Study: This study was conducted in plot 1, Vegetables Field plot for Teaching and Research, Taman Pertanian Universiti, Universiti Putra Malaysia (UPM) Selangor, Malaysia, from May 2018 until August 2018.

Methodology: *Persicaria minor* were exposed to four different concentration of zinc oxide nanoparticles (ZnO-NPs) which were (50,100 and 150 mg/L) and 0 mg/L as a control. The ZnO-NPs was dissolved in distilled water before being applied to plants. 40 mL of ZnO-NPs solution was applied to each plant. The growth, carbon assimilation and also secondary metabolites were measured in this experiment.

Results: The results showed that the treatment of zinc oxide nanoparticles enhanced growth of the *Persicaria minor* as the plant treated with zinc oxide nanoparticles had higher plant height and total biomass when compared to control treatment. However, the analysis revealed that the treatment of zinc oxide nanoparticles highly and significantly influenced the carbon assimilation and quality of this plant as the treated plants showed reduction in chlorophyll content, photosynthesis rate, stomatal conductance and transpiration rate but increased in production of secondary metabolites. The increased in production of plant secondary metabolites may be attributed by the plant protection mechanism due to metabolic stress caused by high concentration of zinc oxide nanoparticles.

Conclusion: This research will progressively help in contributing some reliable and valid data on the effect of zinc oxide nanoparticles (ZnO-NPs), towards the *Persicaria minor* that can be used as guidance for further experimental investigation regarding this field.

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Keywords: *Persicaria minor*, zinc- oxide nanoparticles, growth, carbon assimilation, secondary metabolites, toxicity

1. INTRODUCTION ()

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According to National Health Portal [1], the World Health Organization (WHO) estimated that 80 percent of the world's population consume herbal medicine for their health care and there are about 21000 plants species with the potential to be utilized as medicinal plants. *Persicaria minor* is one of the plants that gained great attention in this field of study. According to Christopher et al. [2], *Persicaria minor* have gained great attention in scientific

23 study due to its high content of antioxidant. This plant possesses variety of pharmacological
24 properties such as antioxidant activity, antiulcer activity, anti-inflammatory activity,
25 antimicrobial activity, anticancer activity and can enhance the digestive properties and
26 cytotoxic activity [2]. According to Rusdi et al. [3], *P. minor* is locally known as a 'kesum', and
27 is commonly used as a food additive and flavouring agent. Many studies have been carried
28 out because of the popularity of the *P. minor* as a potential medicinal plant with high
29 antioxidant and antimicrobial activities and strong anti-inflammatory properties.

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31 Recently, nanoparticles (NPs) are widely studied because its beneficial properties in
32 agriculture and allied sector [4]. According to them, zinc oxide nanoparticles (ZnO-NPs) is
33 one of NPs used widely as it has been utilized in variety of industrial sector including
34 medication, cosmetic materials, opposed microorganisms and textile industries. As it has
35 been commercially used, the toxicity effect of these ZnO-NPs to the environment and also
36 soil ecosystem are of main concern [5]. Sabir et al. [6] stated that ZnO-NPs possess
37 significant characteristics which have antimicrobial, optical and physical properties therefore
38 it has great potential to enhance agriculture. The presence of ZnO-NPs has shown to
39 enhance the antioxidant mechanism that helps to stabilize the plants and improve the
40 photosynthetic efficiency [7]. However, the effect depends on the concentration of ZnO-NPs
41 and varies from plant to plant [8-10].

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43 Secondary metabolites are the natural compound that produced by the plants and some of
44 the compound are utilized as medicines, flavoring and drugs. According to Biology
45 Reference [11] the simple classification of plants' secondary metabolites includes three main
46 groups which are terpenes, phenolics and nitrogen- containing compounds. Secondary
47 metabolites do not involve in the plant growth and development but required for plant to
48 survive in the environment because they protect plants from other organisms such as
49 pathogen and herbivores that can harm the plants [10]. These compounds possess
50 significant biological properties and also medicinal importance that can improve
51 pharmaceuticals field [10]. The application of ZnO nanoparticles might enhances the
52 production of plant secondary metabolites and thus enhanced the medicinal properties of
53 this plant.

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55 Currently, there is no study conducted on the interaction effect of ZnO-NPs on physical and
56 biochemical response of plant especially in medicinal plants like *Persicaria minor* and only
57 few research being carried out to discover about the effect of ZnO-NPs on carbon
58 assimilation and production of secondary metabolite of this plant, Hence, the objectives of
59 this study were to study the growth, carbon assimilation and quality of *Persicaria minor* as
60 affected by zinc oxide nanoparticles, to determine the optimum concentration dose of zinc
61 oxide nanoparticles that can enhance the optimum growth and secondary metabolites of
62 *Persicaria minor* and to recognize the relationship between secondary metabolites and
63 growth of *Persicaria minor* as exposed by zinc oxide nanoparticles application

64 65 **2. MATERIAL AND METHODS**

66 67 68 **2.1 Experimental site**

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70 This study was conducted in Plot 1, Vegetables Field Plot for Teaching and Research,
71 Taman Pertanian Universiti, Universiti Putra Malaysia (UPM) Selangor. The research site
72 was set up with net shading and black plastic to reduce the absorption of water by sunlight
73 since *Persicaria minor* require high amount of water and also to reduce the competition with
74 grasses and other plants. This experiment was conducted from the month of May 2018 until
75 August 2018. The microclimatic parameters during the experiment are presented in Table 1.

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Table 1. The microclimate data during the experiment

Microclimate parameters	Quantification
Relative humidity	57.14-68.23%
Light intensity	320 -860 $\mu\text{mol}/\text{m}^2/\text{s}$
Day temperature	26-34°C
Night temperature	16-23°C
Ambient CO ₂	380.23 ppm

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2.2 Planting material

82 *Persicaria minor* was obtained from Jabatan Pertanian, Serdang. The first internodes at the
83 bottom of the shoots with five number of leaves was cut in 5cm in height for the shoot stem
84 cutting preparation. The shoot stem cuttings were immersed in the tap water overnight to
85 increase their turgidity which increases the speed of the germination. Then, the propagation
86 step was done in the trays with peat moss as the medium and the shoots stem cutting were
87 left for two weeks for the development of the root. Then, the plants were transferred into
88 standard polybag (16cm x 30cm) which was filled with top soil as the medium [12].

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2.3 Soil preparation

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92 The soil medium was obtained from Unit Herba, Taman Pertanian Universiti and top soil was
93 used as the medium for the *Persicaria minor* planting. The top soil was transferred into the
94 polybag until it filled three- quarter of the polybag

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2.4 Synthesis and properties of Zinc-Oxide Nanoparticles (ZnO-NPs)

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98 The ZnO-NPs was synthesized chemically in the Laboratory of BioPhysics of Physics
99 Department, Faculty of Science UPM by using sol gel method. In this process in room
100 temperature, 200ml of ethanol was added to 0.2M of zinc acetate and then, the mixture was
101 stirred for two hours to obtain clear solution. Then, 1.0 M sodium hydroxide (NaOH) was
102 titrated into the mixture until the pH 9 is reached. After that, the mixture was stirred for one
103 hour and then left for 24 hours to allow the complete hydrolysis and gelation. The sample
104 was then filtrated to obtain white precipitate. The precipitate is dried in an oven for 48 hours
105 at 100°C to dry. And lastly, the dried sample was grinded by mortar and pestle to yield ZnO
106 powder to be used in the experiment.

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2.5 Experimental design

109 A Completely Randomized Block Design (RCBD) was used in this study with three
110 replications. Each unit consisted eight plants and the totals of 96 plants were used in this
111 study. After a month, *Persicaria minor* was exposed to four different concentration of zinc
112 oxide nanoparticles (ZnO-NPs) which were (50,100 and 150 mg/L) and 0 mg/L as a control.
113 The ZnO-NPs was dissolved in distilled water before being applied to plants. 40 mL of ZnO-
114 NPs solution was applied to each plant.

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2.6 Plant maintenance

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118 The maintenance steps are very crucial to ensure the plant develops healthy and to avoid
119 the plants from wilting or attacked by any disease that can cause the plants to die. At the
120 early phase of cultivation, *Persicaria minor* were watered two times daily. The watering was

121 unnecessary only when the heavy rain occur. This to avoid the over watering to the plants
122 that can interfere with the plants growth. The common insects that could interfere with the
123 plants growth were removed quickly from the planting area.

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125 **2.7 Collection of data**

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127 The growth data collection was done once a week after the application of treatment for the
128 plant's growth parameter. The destructive analysis and leaf gas exchange of the experiment
129 were conducted at the end of the experiment.

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131 **2.7.1 Plant growth measurements**

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133 The plant growth measurements were conducted to obtain data on plant height, number of
134 leaf and stem, diameter of stem, root to shoot ratio and the chlorophyll content. The plant
135 height was measured starting from the stem on the soil surface until the highest shoot
136 growth using a measuring tape. The plant basal diameter was measured by using vernier
137 caliper at the base of the plants and the leaves of the *Persicaria minor* were counted
138 manually in every three weeks.

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140 **2.7.1.4 Chlorophyll content measurement**

141 The total chlorophyll content of the leaves was measured by using chlorophyll meter (SPAD
142 502). The leaves of the plants in each treatment for each replication were clipped by
143 chlorophyll meter clipper to obtain the reading.

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145 **2.7.1.5 Plant fresh weight measurement**

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147 The plants were removed first from the soil and all the dirt were removed under the flowing
148 tap water. Then, the shoot and the root parts were separated for further analysis and all the
149 plants parts were weighted separately using analytical balance.

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151 **2.7.1.6 Dry weight (biomass) measurement**

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153 The plants were dried in the oven at 60°C for 48 hours. Then, the measurements were
154 recorded as observed using electronic weighing scale.

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156 **2.7.1.7 Root to shoot ratio**

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158 The root to shoot ratio was determined by dividing the weight of the roots part to the shoot
159 part after the oven drying process.

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161 **3.7.1.8 Plant leaf temperature determination**

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163 The Infrared (IR) thermometer was used to measure the plant leaves temperature. This was
164 to indicate whether the plant under stressful condition or not. The upper leaf part was chosen
165 in the determination [13].

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167 **2.7.2 Leaf gas exchange measurement**

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169 A portable photosynthesis system LI-6400XT (Li-COR Inc; Nebraska; USA) was used to
170 measure the leaf gas exchange. This equipment was warmed and was calibrated with ZERO
171 IRGA mode for 30 minutes. The measurement was set at optimum condition which were 400

172 $\mu\text{mol mol}^{-1} \text{CO}_2$, 30°C cuvette temperature, 60% relative humidity with the rate of air flow set
173 at $500 \text{ cm}^3 \text{ min}^{-1}$ and then the cuvette condition was modified at $800 \mu\text{mol m}^{-2} \text{ s}^{-1}$
174 photosynthetically photon flux density (PPFD). The measurement process of gas exchange
175 was carried out between 9.00 am to 11.00 am by using fully expanded young leaves that
176 give the measurement of net photosynthesis (A), stomata conductance (gs) and transpiration
177 rate (E). Water use efficiency (WUE) was measured by using the formula of net
178 photosynthesis dividing with transpiration rate. This is automatic operation and the results
179 were saved in the LI-6400XT console and Photosyn Assistant Software (Dundee Scientific,
180 Dundee, UK) was used to analyze it. Precautions were taken to avoid mistakes during taking
181 the measurements [14].

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184 **2.7.3 Total Phenolics and Flavonoids Quantification**

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186 The methods used for extraction and quantification of total phenolics and flavonoids contents
187 followed that described in Ibrahim et al. [15]. A fixed amount of ground tissue samples (0.1
188 g) was extracted with 80% ethanol (10 mL) on an orbital shaker for 120 min at 50°C. The
189 mixture was subsequently filtered (Whatman™ No.1), and the filtrate was used for the
190 quantification of total phenolics and total flavonoids. Folin–Ciocalteu reagent (diluted 10-
191 fold) was used to determine total phenolics content of the leaf samples. The sample extract
192 at 200 μL was mixed with Folin–Ciocalteu reagent (1.5 mL) and allowed to stand at 22°C
193 for 5 min before adding NaNO_3 solution (1.5 mL, 60 g L^{-1}). After two hours at 22°C,
194 absorbance was measured at 725 nm. The results were expressed as mg g^{-1} gallic acid
195 equivalent (mg GAE g^{-1} dry sample). For total flavonoids determination, samples (1 mL)
196 were mixed with NaNO_3 (0.3 mL) in a test tube covered with aluminium foil, and left for 5
197 min. Then 10% AlCl_3 (0.3 mL) was added followed by addition of 1 M NaOH (2 mL). The
198 absorbance was measured at 510 nm using a spectrophotometer with rutin as a standard
199 (results expressed as mg/g rutin dry sample).

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201 **2.7.4 Chlorophyll fluorescence determination**

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203 The chlorophyll fluorometer was used to measure the chlorophyll fluorescence of the
204 *Persicaria minor*. The mature leaf tissue was obtained from the *Persicaria minor* plant that
205 cultivated at 20°C in glasshouse exposed with artificial light to give minimum photon flux
206 density of $550 \mu\text{mol m}^{-2} \text{ s}^{-1}$ for 16 h photoperiod and photosynthetically active radiation were
207 supplied at $250 \mu\text{mol m}^{-2} \text{ s}^{-1}$ during 16 h photoperiod [16].

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210 **2.8 Statistical Analysis**

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212 Statistical Package for Social Sciences (SPSS) version 24 was used to analyze the recorded
213 data. A two-way ANOVA Test was conducted to analyze data for all the parameters used in
214 the experiment. Results were significant if the p-value level was ≤ 0.05 .

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216 **3. RESULTS AND DISCUSSION**

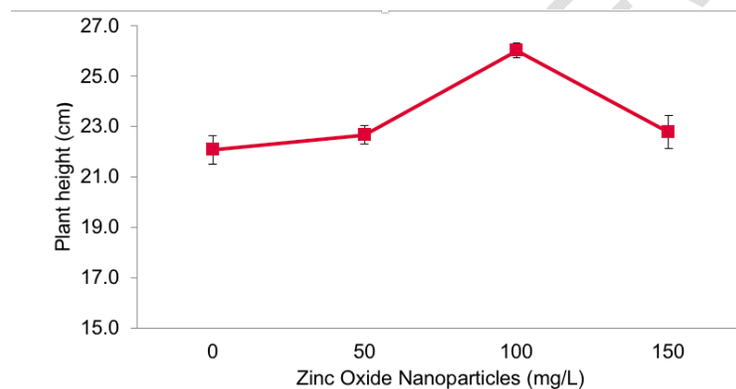
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218 **3.1 Plant Height**

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220 Fig.1 depicts the effect of three months treatment of zinc oxide nanoparticles (ZnO-NPs) on
221 the plant height of *Persicaria minor*. The result from analysis of variance showed that there
222 was a significant effect between different concentration of zinc oxide nanoparticles treatment
223 toward the height of *Persicaria minor* ($P \leq 0.05$). From the figure, the increasing of
224 concentration of zinc oxide nanoparticles treatment increased the plant height of *Persicaria*

225 *minor*. The highest plant height was recorded in 100 mg/L on 12 weeks after treatment with
226 mean 26.01 cm that might indicates the optimum concentration for the plant. Meanwhile, the
227 control treatment recorded the shortest plant height with mean 22.10 cm. The appropriate
228 concentration of zinc oxide nanoparticles plays significant role in plant growth and promotion
229 [15]. From the results, on twelfth week after harvesting, the plants treated with zinc oxide
230 nanoparticles have higher plant height as compared to plants in control treatment. This
231 finding indicates that the application of zinc oxide nanoparticles can induced the growth of
232 the plants. Kouhi et al. [17] explained that the zinc oxide nanoparticles possess plant growth
233 promoting effects and were used as micronutrient fertilizer. The presence of these
234 nanoparticles triggered the physiological processes, acting as growth regulating compound
235 that increased the plant growth such as the plant height and biomass. In addition, Prasad et
236 al. [18] reported that zinc oxide nanoparticles possess beneficial effects in enhancing plant
237 growth and development. The presence of zinc can enhance the biochemical, physiological
238 and anatomical responses of the plants thus increased the plant growth parameters such as
239 plant height and biomass [19]. Therefore, it can be concluded that the zinc oxide
240 nanoparticles treatment induced the plant growth and 100 mg/L can be considered as the
241 best concentration among the treatments rates in promoting the height of *Persicaria minor*
242 plant.

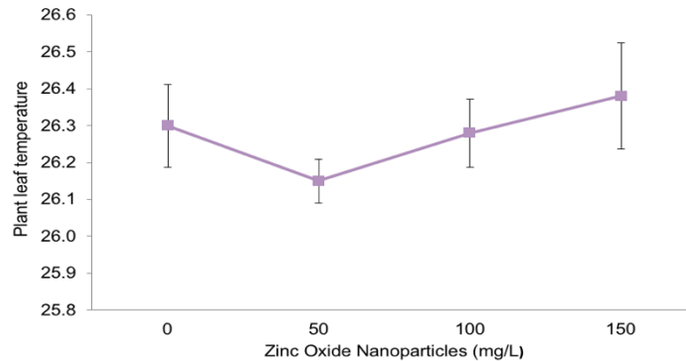


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258 **Fig.1. The effect of zinc oxide nanoparticles treatment (0, 50, 100 and 150) mg/L on**
259 **plant height of *Persicaria minor*. Data are means with standard error of mean (SEM) of**
260 **24 replicates.**

261 262 263 **3.2 Plant Leaf Temperature**

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265 Fig. 2 highlights the effect of three months treatment of zinc oxide nanoparticles (ZnO-NPs)
266 on the total leaf temperature of *Persicaria minor*. The result from the analysis variance
267 showed that there was a significant effect between different concentration of zinc oxide
268 nanoparticles treatment toward the total leaf temperature of *Persicaria minor* ($P \leq 0.05$).
269 Based on the figure, the trend shows that the plant leaf temperature increases linearly with
270 the concentration of zinc oxide nanoparticles treatment except for the control treatment. This
271 showed that application of Zinc oxide nanoparticles induces stress response to the plants.
272 The plants maintain their most important physiological process (photosynthesis) by
273 maintaining their average leaf temperature at around 21 degrees Celsius [20]. The plants
274 leaf temperature depends on the stomatal conductance and transpiration rates of the plants
275 [21]. Transpiration is one of the best mechanisms used by plants to cool themselves by
276 'pumping' out water from leaves through stomata [22]. From this study, the increasing of
277 plant leaves temperature can be explained through the reduction of the stomatal

278 conductance and transpiration rates of the plants due to the increase the concentration of
279 the treatment. The reduction in transpiration rate and stomata conductance might be due to
280 the closure of plant stomata under the exposure to the zinc oxide nanoparticles [21]. This
281 high temperature in turns will give negative effect to the photosynthesis process thus affect
282 the plant yields. Therefore, it can be concluded that the zinc oxide nanoparticles treatment
283 increased the plant leaf temperature due to the reduction of stomatal conductance and
284 transpiration rate of *Persicaria minor*.



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301 **Fig.2.The effect of zinc oxide nanoparticles treatment (0, 50, 100 and 150) mg/L on**
302 **plant leaf temperature of *Persicaria minor*. Data are means with standard error of**
303 **mean (SEM) of 24 replicates.**

304 305 306 307 **3.3 Total Biomass**

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309 Fig.3 shows the effect of three months treatment of zinc oxide nanoparticles (ZnO-NPs) on
310 the total biomass of *Persicaria minor*. The result from the analysis of variance showed that
311 there was a significant effect between different concentration of zinc oxide nanoparticles
312 treatment toward the total biomass of *Persicaria minor* ($P \leq 0.05$). From the results, it showed
313 that higher concentration of zinc oxide nanoparticles increased the total biomass of
314 *Persicaria minor* as plants treated with zinc oxide nanoparticles treatment have higher total
315 biomass as compared to the control treatment and 100 mg/L gave the highest value of total
316 biomass with mean 2.67g. The highest value of total biomass indicates that 100 mg/L was
317 the optimum concentration of zinc oxide nanoparticles for *Persicaria minor* despite 150 mg/L
318 that reduced the total biomass of the plant. This study showed that the treatment of zinc
319 oxide nanoparticles increased the plant biomass so the treatment might be effective in
320 boosting the plant growth and yield. Similar finding was observed from study conducted by
321 Venkatachalam et al. [23] that revealed total biomass significantly increased in the zinc oxide
322 nanoparticles treated plants as compared to control. This is also supported by findings of
323 Munir et al. [24] that stated the treatment of zinc oxide nanoparticles increased the shoot and
324 root dry weight thus increased the plant biomass. The presence of zinc can enhance the
325 biochemical, physiological and anatomical responds of the plants thus increased the plant
326 growth such as plant height and biomass [15]. Hence, it can be concluded that the presence
327 of zinc oxide nanoparticles can boost the *Persicaria minor* growth resulting in increasing of
328 the plant total biomass.

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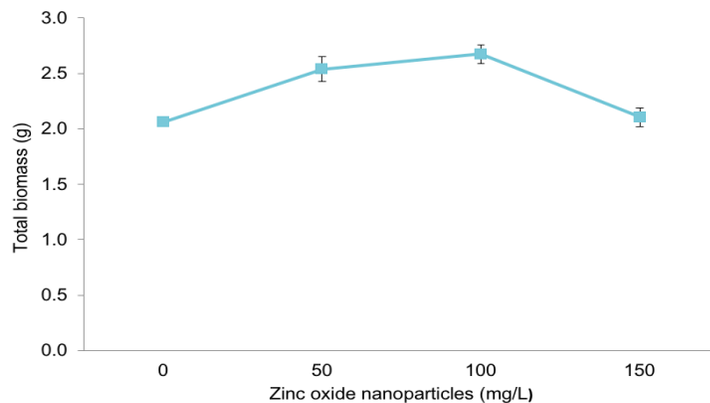


Fig.3.The effect of zinc oxide nanoparticles treatment (0, 50, 100 and 150) mg/L on total biomass of *Persicaria minor*. Data are means with standard error of mean (SEM) of 24 replicates.

3.4 Net photosynthesis rate (A)

Fig.4 illustrated the effect of three months treatment of zinc oxide nanoparticles (ZnO-NPs) on the net photosynthesis rate of *Persicaria minor*. The exposure of zinc oxide nanoparticles toward the *Persicaria minor* was highly and significantly affected the net photosynthesis rate of the plant ($P \leq 0.05$). From the result, the highest photosynthesis rate was recorded in control treatment while the lowest photosynthesis rate was recorded in 150 mg/L of zinc oxide nanoparticles treatment with mean $6.32 \mu\text{mol/m}^2/\text{s}$ and $4.00 \mu\text{mol/m}^2/\text{s}$ respectively. From the figure, the photosynthesis rate of the plant reduced with the increasing concentration of zinc oxide nanoparticles treatment. Photosynthesis is the perfect measurement to access plant performance. From this study, the net photosynthesis of *Persicaria minor* was reduced with the increasing concentration of zinc oxide nanoparticles treatment. Photosynthesis is highly affected in plants that exposed to excess heavy metal where higher level of zinc oxide nanoparticles inhibits the photosynthetic apparatus and caused critical changes to chlorophyll structure and amount [25]. This finding is also similar to that of Wang et al. [26] that revealed the presence of zinc oxide nanoparticles reduced the chlorophyll content in leaves thus reduced the photosynthetic efficiency in plants. In addition, plants exposed to high concentration of zinc oxide nanoparticles have low photosynthetic efficiency might be due to the reduction of chlorophyll content and also excess zinc oxide might damage to the photochemical system [21]. The reduction in photosynthesis with the increased application of zinc oxide was contradicting with the accumulation of total biomass suggesting that *P. minor* adapt to increased level of Zn-O nanoparticles by storing more photosynthate for the reduced photosynthetic efficiency. Therefore, it can be concluded that the presence of zinc oxide nanoparticles reduced the photosynthetic efficiency of *Persicaria minor* plants [25,26].

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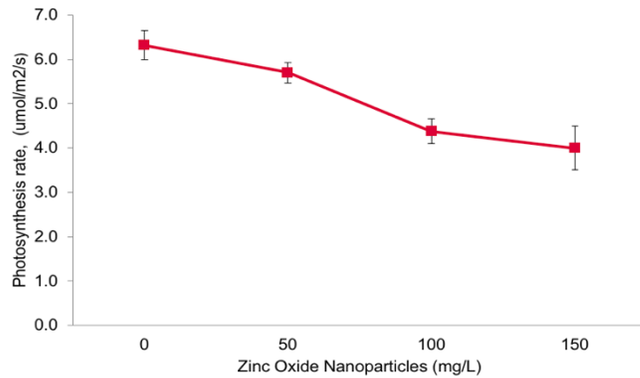
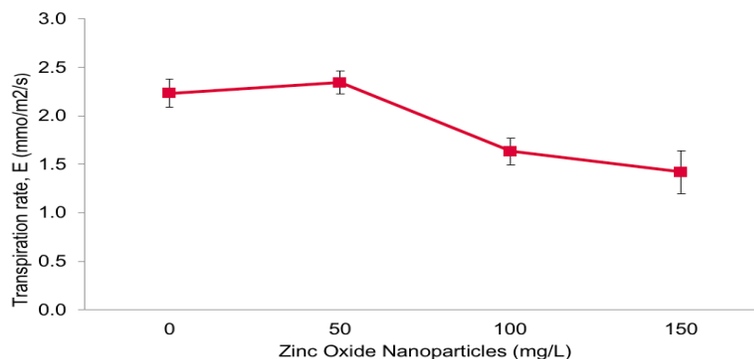


Fig.4: The effect of zinc oxide nanoparticles treatment (0, 50, 100 and 150) mg/L net photosynthesis rate of *Persicaria minor*. Data are means with standard error of mean (SEM) of 24 replicates.

3.5 Transpiration rate (E)

Fig.5 depicts the effect of three months treatment of zinc oxide nanoparticles (ZnO-NPs) on the transpiration rate of *Persicaria minor*. The exposure of different concentration of zinc oxide nanoparticles toward the *Persicaria minor* highly and significantly affected the transpiration rate of the plant ($P \leq 0.05$). From the figure, the transpiration rate of the plant reduced with the increasing concentration of zinc oxide nanoparticles treatment. The transpiration rate of 50 mg/L treatment was significantly higher with mean 2.34 mmol/m²/s while the lowest transpiration rate of *Persicaria minor* was observed in 150 mg/L of zinc oxide nanoparticles treatment with mean 1.42 mmol/2/s. Transpiration is a process of the movement of water vapors through plant and this process mainly take place in leaves. Transpiration process is mainly controlled by the opening and closing of the stomata. From the result, the transpiration rate reduced with the increasing concentration of zinc oxide nanoparticles treatment. This finding is similar with Xiaoping et al. [27] that revealed both transpiration rate and stomatal conductance reduced in plants treated with zinc oxide nanoparticles. The stomatal closure reduced the transpiration rate of the plants. According to Vankova et al. [28], the presence of zinc oxide nanoparticles induced the production of plant stress hormone, abscisic acid (ABA) and this hormone mainly accumulated in leaves. The higher level of abscisic acid triggered the stomatal closure which in turn reduced the transpiration rate in plants [29]. Hence, the application of zinc oxide nanoparticles reduced transpiration rate of *Persicaria minor* might be due to the accumulation of ABA stress hormone that cause the closure of stomata

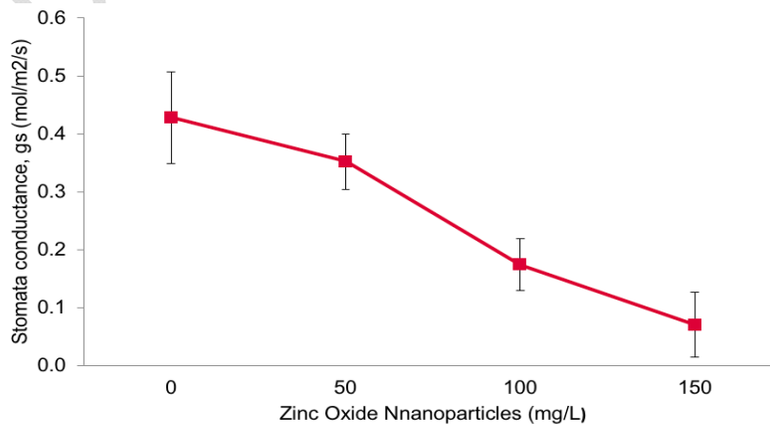


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Fig.5. The effect of zinc oxide nanoparticles treatment (0, 50, 100 and 150) mg/L transpiration rate of *Persicaria minor*. Data are means with standard error of mean (SEM) of 24 replicates.

3.6 Stomatal conductance (gs)

Stomatal conductance is a measure of the degree of stomatal opening and a good indicator in accessing plant water status [25]. Fig.6 shows the effect of three months treatment of zinc oxide nanoparticles (ZnO-NPs) on the transpiration rate of *Persicaria minor*. The exposure of different concentration of zinc oxide nanoparticles toward the *Polygonum minus* was highly and significantly affected the stomatal conductance of the plant ($P \leq 0.05$). From the figure, the stomatal conductance of the plant reduced as the concentration of the zinc oxide nanoparticles treatment increasing. The highest stomatal conductance was recorded in control treatment while the lowest stomatal conductance was observed in 150 mg/L of zinc oxide nanoparticles treatment with mean 0.43 and 0.07 mmol/m²/s respectively. The finding showed reduction in stomatal conductance might due to the increasing of zinc oxide nanoparticles concentration is similar with Xiaoping et al. [27] that proved the higher concentration of zinc oxide nanoparticles reduced the stomatal conductance resulting in low photosynthetic efficiency of the plants. Singh and Bhati [30] also stated that high amounts of zinc oxide nanoparticles can restrict the stomatal conductance. This might due to the toxicity of the treatment disturbed the cell mechanism thus alters the stomatal function. Tsonev and Lidon [31] explained that the stomatal response to high concentration of zinc oxide nanoparticles is related to the changes in carbonic anhydrase (CA) activity. Carbonic anhydrase is an enzyme that is responsible for the stomatal activity and the presence of zinc oxide nanoparticles influenced the CA activity that triggered the stomatal closure thus reduced the stomatal conductance of the plants. Hence, it can be concluded that the presence of zinc oxide nanoparticles alter the stomatal mechanism thus reducing the stomatal conductance of *Persicaria minor* plants.



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Fig.6. The effect of zinc oxide nanoparticles treatment (0, 50, 100 and 150) mg/L on stomatal conductance of *Persicaria minor*. Data are means with standard error of mean (SEM) of 24 replicates.

3.7 Maximum efficiency of photosystem II

Fig.7 depicted the effect of three months treatment of zinc oxide nanoparticles (ZnO-NPs) on the maximum efficiency of photosystem II of *Persicaria minor*. The exposure of different concentration of zinc oxide nanoparticles toward the *Persicaria minor* was highly and significantly affected the maximum efficiency of photosystem II of the plant ($P \leq 0.05$). Based on the figure, increasing the concentration of zinc oxide nanoparticles caused the maximum efficiency of photosystem II to decrease. The highest value of maximum efficiency of photosystem II was observed in control treatment while the lowest value of maximum efficiency of photosystem II was recorded in 150 mg/L treatment with mean 0.758 and 0.522 respectively. The phytotoxicity of zinc oxide nanoparticles can be accessed through the efficiency of photosynthetic mechanism (chlorophyll florescence) that act as indicator in phytotoxicity assays. The finding of this study revealed that increasing the zinc oxide nanoparticles concentration resulting in lower maximum efficiency of photosystem II of *Polygonum minus*. Wang et al. [26] stated that the treatment of zinc oxide nanoparticles reduced the chlorophyll florescence parameter and damaged the photochemical system. This finding can be explained further that the presence of zinc oxide nanoparticles induced the oxidative stress in plants and increase the production of reactive oxygen species (ROS) which alter the gene expression pathway thus reduced the chlorophyll florescence in plants. Therefore, it can be concluded that the zinc oxide nanoparticles treatment reduced the chlorophyll florescence parameters of *Persicaria minor* plants.

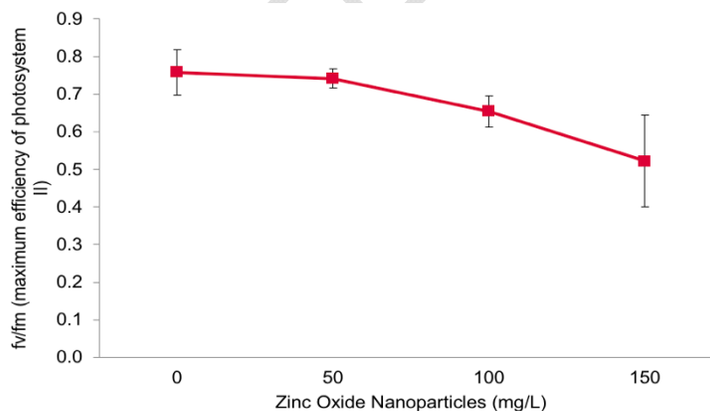
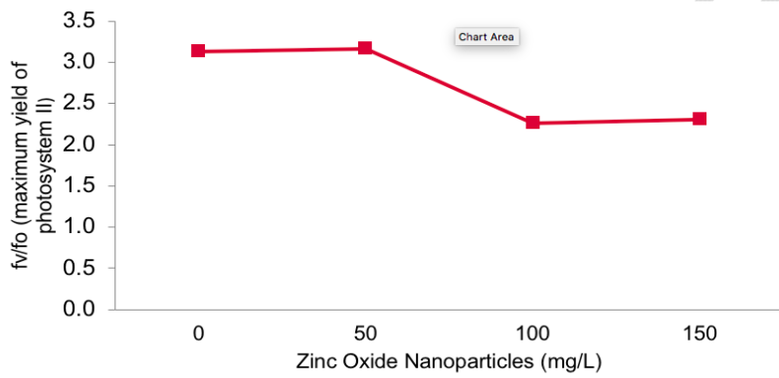


Fig.7. The effect of zinc oxide nanoparticles treatment (0, 50, 100 and 150) mg/L on maximum efficiency of photosystem II of *Persicaria minor*. Data are means with standard error of mean (SEM) of 24 replicates.

3.8 Maximum yield of photosystem II

Fig.8 shows the effect of three months treatment of zinc oxide nanoparticles (ZnO-NPs) on the maximum yield of photosystem II of *Persicaria minor*. The exposure of different concentration of zinc oxide nanoparticles toward the *Persicaria minor* highly and significantly affected the maximum yield of photosystem II of the plant ($P \leq 0.05$). The trend shows that

543 increasing the concentration of zinc oxide nanoparticles caused the maximum yield of
 544 photosystem II to decrease. Based on the figure, 50 mg/L of zinc oxide nanoparticles
 545 treatment shows the highest value of maximum yield of photosystem II with mean 3.16 when
 546 compared with other treatment. From this study, it was observed that the treatment of zinc
 547 oxide nanoparticles reduced the maximum efficiency of photosystem II which in turn reduced
 548 the maximum yield of photosystem II of *Persicaria minor*. According to Tsonev and Lidon
 549 [31], inside the chloroplast lamellae, the presence of zinc oxide nanoparticles caused the
 550 inhibition of photosynthetic electron transport and implicates the water evolving complex of
 551 photosystem II thus inhibits the photolysis and oxygen emission that disturb the
 552 conformation of photosystem II core complex. This mechanism explained how the zinc oxide
 553 nanoparticles treatment reduced the efficiency and yield of photosystem II in plants. Hence,
 554 the treatment of zinc oxide nanoparticles reduced the maximum yield and efficiency of
 555 photosystem II which in turn disturbed the photosynthetic process of *Persicaria minor* plant.
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Fig.8. The effect of zinc oxide nanoparticles treatment (0, 50, 100 and 150) mg/L on maximum yield of photosystem II of *Persicaria minor*. Data are means with standard error of mean (SEM) of 24 replicates.

3.9 Minimal florescence

579 Fig.9 shows the effect of three months treatment of zinc oxide nanoparticles (ZnO-NPs) on
 580 the minimal florescence of *Persicaria minor*. The exposure of different concentration of zinc
 581 oxide nanoparticles toward the Polygonum minus highly and significantly affected the
 582 maximum yield of photosystem II of the plant ($P \leq 0.05$). The result showed that higher
 583 concentration of zinc oxide nanoparticles treatment resulted in higher value of minimal
 584 florescence of *Persicaria minor*. The highest value of minimal inflorescence was recorded in
 585 150 mg/L with mean 627.23 while the lowest value was observed in 50 mg/L with mean
 586 462.67. Higher minimal florescence indicates higher heat dissipation of plants. This might
 587 due to the presence of zinc oxide nanoparticles that induced stress in plants thus caused
 588 plants to produce high amount of heat. From this study, the treatment of zinc oxide
 589 nanoparticles reduced the transpiration rate of *Persicaria minor*. This reduction might be
 590 related with the increasing of minimal florescence of the plants [32]. The high minimal
 591 florescence can cause heat stress to the plants. Heat stress is defined as the increase
 592 temperature beyond the threshold level that cause damage to plant growth and development
 593 [27]. Therefore, it can be deduced that the treatment of zinc oxide nanoparticles increased
 594 the minimal florescence of *Persicaria minor* due to the reduction in transpiration rate of the
 595 plants.

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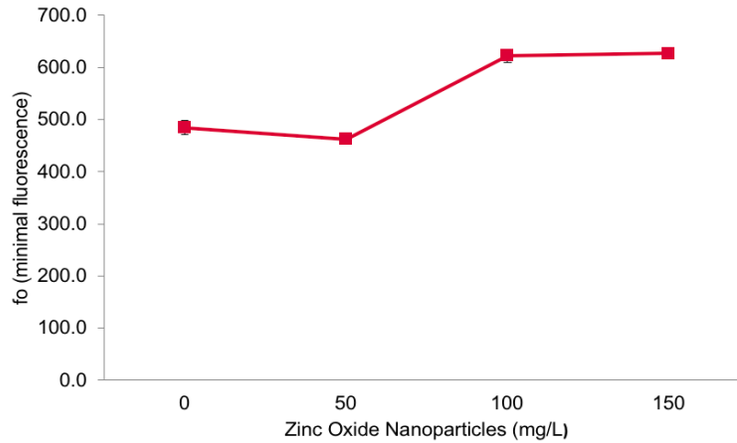
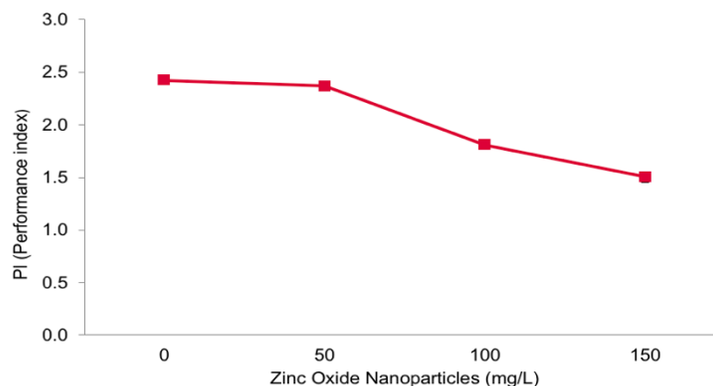


Fig.9. The effect of zinc oxide nanoparticles treatment (0, 50, 100 and 150) mg/L on minimal fluorescence of *Persicaria minor*. Data are means with standard error of mean (SEM) of 24 replicates.

3.10 Performance index (PI)

Fig.10 indicated the effect of three months treatment of zinc oxide nanoparticles (ZnO-NPs) on the performance index of *Persicaria minor*. The exposure of different concentration of zinc oxide nanoparticles toward the *Polygonum minus* was highly and significantly affected the performance index of the plant ($P \leq 0.05$). Based on the figure, the highest and lowest performance index were observed in control treatment and 150 mg/L of zinc oxide nanoparticles treatment with mean 2.42 and 1.50 respectively. The performance index of *Persicaria minor* reduced when treated with higher concentration of zinc oxide nanoparticles that indicates zinc oxide nanoparticles increased the plant stress. Performance index is an indicator of PSII functioning and informs about efficiency of assimilating apparatus. It is correlated with water accessibility for plants. High PI value implies favorable plant condition, and, by analogy, plant have less stress [32]. Nanoparticles such as zinc oxide and silver were located on the surface of plants cells and induced the oxidative stress to the cells by the activation of oxidative stress signaling [28]. From this study, it was observed that the treatment of zinc oxide nanoparticles reduced the plants performance index. Zahed et al. [33] stated that the generation of reactive oxygen species (ROS) due to the zinc oxide nanoparticles treatment alter the gene expression and cell mechanism which in turn reduced the performance index of the plants. Wang et al. [26] explained that the toxicity of zinc oxide nanoparticles reduced chlorophyll content plants, resulted in low photosynthesis efficiency thus reduced the plants performance. Hence, it can be concluded that the presence of zinc oxide nanoparticles induced stress in *Persicaria minor* resulting in low performance index of the plants.

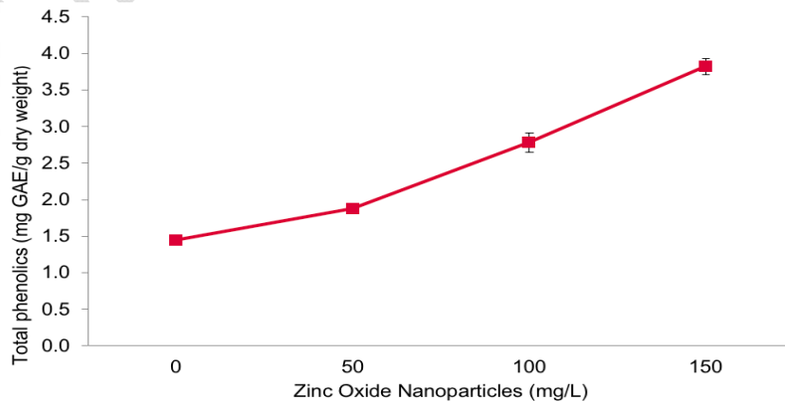


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Fig.10. The effect of zinc oxide nanoparticles treatment (0, 50, 100 and 150) mg/L on performance index of *Persicaria minor*. Data are means with standard error of mean (SEM) of 24 replicates.

3.11 Total phenolics content

Phenolics are compound that produced by plants to protect plants against stress. This compound plays significant role in plant development (lignin and pigment biosynthesis) and also provided structural integrity for plant's support [34]. Fig.11 shows the effect of three months treatment of zinc oxide nanoparticles (ZnO-NPs) on the total phenolics production of *Persicaria minor*. The exposure of different concentration of zinc oxide nanoparticles toward the *Polygonum minus* highly and significantly affected the total phenolics production of the plant ($P \leq 0.05$). Based on the figure, the total phenolics production of the plant was directly proportional with the concentration of zinc oxide nanoparticles treatment. The lowest total phenolics production was recorded in control treatment while the highest total phenolics production was recorded in 150 mg/L treatment with mean 1.44 and 3.82 GAE/g dry weight respectively. This result indicates that zinc oxide nanoparticles treatment induced stress and increased the secondary metabolites production of *Persicaria minor*. From this study, the greater production of total phenolics content in *Polygonum minus* with the increasing of concentration treatment revealed that the presence of zinc oxide nanoparticles induced stress towards the plants. This finding is supported by Rastogi et al. [35] stated that zinc oxide nanoparticles treatment induced the Reactive Oxygen Species (ROS) production in plants thus increased plants stress. They also stated that higher concentration of zinc oxide nanoparticles leads to the damage of plant cell wall and plasma membrane thus induced the production of plant secondary metabolites for plants defense against disease and threat [36]. Therefore, it can be concluded that higher concentration of zinc oxide nanoparticles lead to plant stress and boost the plants secondary metabolites production which in turn can enhanced the defense response of *Persicaria minor* plants.



701 **Fig.11. The effect of zinc oxide nanoparticles treatment (0, 50, 100 and 150) mg/L on**
702 **total phenolics production of *Persicaria minor*. Data are means with standard error of**
703 **mean (SEM) of 24 replicates**

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705 **3.12 Total flavonoids content**

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707 Flavonoids are a wide group of plants chemicals (phytonutrients) that found mostly in fruits
708 and vegetables. Flavonoids plays significant role in pharmacological field since this
709 compound is a good source of antioxidant and anti-inflammatory, protect skin, enhanced
710 brain function and also good for blood pressure regulation [33]. Fig.12 shows the effect of
711 three months treatment of zinc oxide nanoparticles (ZnO-NPs) on the total flavonoids
712 production of *Persicaria minor*. The exposure of different concentration of zinc oxide
713 nanoparticles toward the *Persicaria minor* highly and significantly affected the total
714 flavonoids production of the plant ($P \leq 0.05$). Based on the figure, the total flavonoids
715 production of the plant was directly proportional with the concentration of zinc oxide
716 nanoparticles treatment. The lowest total flavonoids production was recorded in control
717 treatment while the highest total flavonoids production was recorded in 150 mg/L treatment
718 with mean 0.43 and 1.47 mg rutin/g dry weight respectively. This result indicates that zinc
719 oxide nanoparticles treatment induced stress and increased the secondary metabolites
720 production of *Persicaria minor*. From this study, the greater production of total flavonoids
721 content in *Polygonum minus* with the increasing of concentration treatment revealed that the
722 presence of zinc oxide nanoparticles induced stress towards the plants. This finding is
723 similar with that of Zafar et al. [37] where the higher treatment concentration of zinc oxide
724 nanoparticles generates oxidative stress of plants thus increasing the plant secondary
725 metabolites production to protect plants against stress. The initial response of plants towards
726 the presence of nanoparticles involved the increasing level of reactive oxygen species
727 (ROS), cytoplasmic Ca^{2+} and up regulation of nitrogen activated protein kinase (MAPK)
728 cascades thus activates the plants secondary metabolites that act against stress to protect
729 the plants [38]. In addition, the presence of zinc oxide nanoparticles enhanced the
730 expression of genes related to antioxidant capacity thus boost the defense mechanism of the
731 plants by enhancing the production of plants secondary metabolites. Hence, it can be
732 concluded that the presence of zinc oxide nanoparticles enhanced the *Persicaria minor*
733 secondary metabolites production by increasing the total flavonoids production of the plants.

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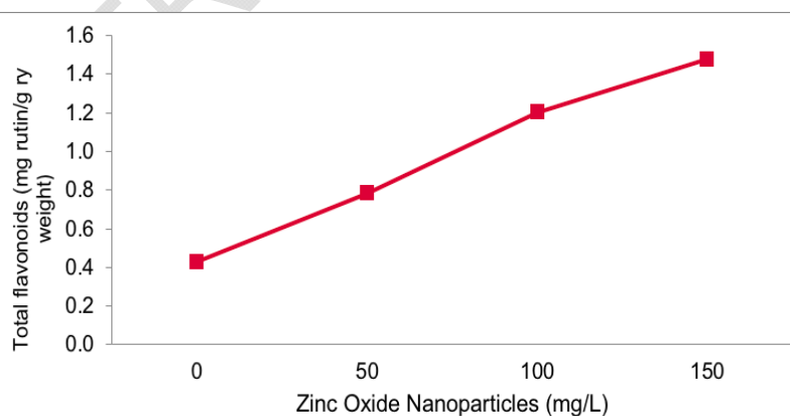
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750 **Fig.12. The effect of zinc oxide nanoparticles treatment (0, 50, 100 and 150) mg/L on**
751 **total flavonoids production of *Persicaria minor*. Data are means with standard error of**
752 **mean (SEM) of 24 replicates.**

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754 **4. CONCLUSION**

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From this study, it can be concluded that the optimum concentration of zinc oxide nanoparticles for *Persicaria minor* growth was at 100 mg/L because it recorded the highest value in the growth parameters. Overall, the treatment of zinc oxide nanoparticles increased the growth parameter of the plants as the treated plants showed higher value of plant height and total biomass when compared to plants in control treatment. However, the treatment of zinc oxide reduces the leaf gas exchange and chlorophyll fluorescence properties of *Persicaria minor* plants. Despite reduction on the leaf gas exchange and chlorophyll fluorescence the production of secondary metabolites (total phenolics and flavonoids production) were enhanced with increased levels of Zinc oxide application.

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