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# Effects of phosphorus on chemical forms of Cd in plants of four spinach (*Spinacia oleracea* L.) cultivars differing in Cd accumulation

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1 **Effects of phosphorus on chemical forms of Cd in plants of four spinach (*Spinacia oleracea***  
2 **L.) cultivars differing in Cd accumulation**

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14 **Abstract**

15 In order to clarify how cadmium (Cd) chemical forms *in planta* relate to the genotype difference  
16 in Cd accumulation of spinach (*Spinacia oleracea* L.), two low-Cd and two high-Cd cultivars  
17 were compared under a hydroponic experiment with two concentrations of Cd (1 or 5 mg Cd L<sup>-1</sup>).  
18 The concentrations of phosphorus in the hydroponic system were also adjusted to two levels  
19 (half and full P in standard Hoaglund's solutions) to investigate the influence of phosphorus on  
20 the forms and accumulation of Cd in the tested cultivars. Average Cd concentrations in shoots  
21 were 8.50-10.06 mg kg<sup>-1</sup> for high-Cd cultivars and 6.11-6.64 mg kg<sup>-1</sup> for low-Cd cultivars under  
22 lower Cd treatment, and were as high as 24.41-31.35 mg kg<sup>-1</sup> and 19.65-25.76 mg kg<sup>-1</sup>,  
23 respectively, under higher treatment. Phosphorus significantly decreased Cd accumulation in the  
24 tested cultivars and the effect had superiority over the cultivar alternation under higher Cd stress.  
25 Cadmium in the NaCl-extractable fraction of the plant tissues showed the greatest relationship to  
26 genotype difference of Cd accumulation. The difference in the capacity to binding Cd into F<sub>HAC</sub>,  
27 F<sub>HCl</sub> or F<sub>Residue</sub> was another important mechanism involving in the genotype difference in Cd  
28 accumulation of spinach. Among them, average proportion of Cd in F<sub>HAC</sub> in low-Cd cultivars was  
29 higher than that in high-Cd cultivars in association with the effect of phosphorus.

30 **Key Words:** spinach; cadmium (Cd); phosphorus; chemical form; genotype difference; food  
31 safety

32 **Introduction**

33 Contamination of agricultural soil by heavy metals such as Cu, Zn, Cd, and Pb is a  
34 substantial problem globally, especially in China (Murtaza et al. 2008; Nicholson et al. 2003).  
35 These heavy metals present a threat to human health when they enter the food chain (Satarug et  
36 al. 2003). The contamination is mainly caused by pollutant discharges from industrial and  
37 mining processes as well as a result of overuse or improper use of pesticides, insecticides and  
38 chemical fertilizers in agriculture. In China, millions of acres of agricultural lands and over 12  
39 million tons of grain are contaminated by heavy metals. Ten percent of rice in China contains  
40 excessive cadmium, a heavy metal known to cause cancer, osteoporosis, cardiovascular disease,  
41 and renal dysfunction (Nawrot et al. 2010; Wu and Zhu 2014). Various soil clean-up techniques  
42 have been proposed and proven effective (Mulligan et al. 2001). However, it is a challenge to  
43 employ these techniques in many developing countries because of their high costs (Ebbs et al.  
44 1997; Salt et al. 1995). Furthermore, in China, farmers cannot afford to leave agricultural soils  
45 long-term fallow for the remediation process due to the high demand for food products.

46 One of the alternative strategies for reducing the entrance of Cd into the human food chain is  
47 to select cultivars that accumulate low levels of Cd in their edible parts (Grant et al. 2008; Huang  
48 et al. 2015; McLaughlin et al. 1994; Wang et al. 2009; Xin et al. 2013; Yu et al. 2006; Zhu et al.  
49 2007). This cultivar selection strategy is feasible because, for a number of agronomic plant  
50 species, significant differences exist among cultivars in Cd uptake and accumulation (Grant et al.  
51 2008). A wide variation in Cd accumulation among current cultivars has been reported for some  
52 staple crops (Clarke et al. 2002; Dai et al. 2010; Liu et al. 2010; McLaughlin et al. 1994; Yu et al.  
53 2006) and leafy vegetables (Dai et al. 2012; Huang et al. 2014; Liu et al. 2010; Qiu et al. 2011a;

54 Wang et al. 2007; Wang et al. 2009; Xue et al. 2014; Zhang et al. 2013a; Zhang et al. 2013b;  
55 Zhou et al. 2013; Zhu et al. 2007).

56 There has been considerable research seeking to understand the underlying genetic,  
57 molecular, biochemical, and physiological processes that contribute to the low Cd accumulation  
58 phenotype and to lower the risk of Cd entering the food chain (Clarke, 1997; Grant et al. 2008;  
59 Huang et al. 2009; Ishikawa et al. 2012; Ishimaru et al. 2012; Li et al. 2007; Penner et al. 1995;  
60 Tanhuanpää et al. 2007). For example, Grant et al. (2008) has succeeded in breeding of a low-Cd  
61 durum wheat cultivar named AC Napoleon in Canada. Xin et al. (2010) has reported a new  
62 cultivar of water spinach (*Ipomoea aquatica* Forsk.) with high shoot biomass and low shoot Cd  
63 and Pb concentrations.

64 Phosphorus (P) is a macronutrient that accounts for ~0.2% of plant dry weight and when  
65 limiting, can reduce plant growth and yield. This element is essential for the synthesis of nucleic  
66 acids, phospholipids, and ATP. It has also been reported that addition of P-based materials to  
67 soils can influence the bio-availability of heavy metals such as Pb, Cd and Zn. The amendment  
68 of P to soils reduced the accumulation of Cd in both low-Cd and high-Cd cultivars of Chinese  
69 flowering cabbage (*Brassica parachinensis* L.) (Qiu et al. 2011b).

70 Spinach (*Spinacia oleracea* L.) is an important leafy vegetable that is cultivated and  
71 consumed all over the world, particularly in Southeast Asia during the majority of the year.  
72 Spinach has been described as a Cd accumulating species (Alexander et al. 2006; Chunilall et al.  
73 2004; Kuboi et al. 1986). A strong influence of cultivar on shoot Cu, Zn and Cd concentrations  
74 was observed in a previous study in Germany with 11 spinach cultivars (Römer et al. 2002).  
75 However, a similar study carried out in England found no significant variations in Cd, Cu, Pb or  
76 Zn concentrations among five spinach cultivars grown on metal-spiked soil (Alexander et al.

2006). There is little available information about the mechanisms affecting the genotype differences of Cd uptake, translocation and accumulation in spinach. In our previous study, the maximum difference in shoot Cd concentration varied by 7.2-fold among 29 spinach cultivars (unpublished data). We identified two low-Cd accumulation cultivars (low-Cd group) and two high-Cd accumulation cultivars (high-Cd group) in the study. These four spinach cultivars allow for a further investigation of the mechanisms associated with the genotype differences. In the present study, the chemical forms of Cd in plant tissues between the low-Cd cultivars and the high-Cd cultivars were compared in order to provide insight into the relevant biochemical mechanisms. Due to the previous researches reporting the effects of soil phosphorus on Cd accumulation of spinach (Dheri et al. 2007; Keller et al. 2001; Römer et al. 2002;), phosphorus concentration was also altered to investigate how the interaction between Cd and P contributes to the genotype difference. We hypothesize that the genotype-dependent Cd accumulation of spinach is related to chemical forms of Cd, and phosphorus is a crucial factor that interacts with Cd to influence the chemical form of Cd within the plant tissues, and therefore the extent of Cd accumulation.

92

## 93 **Material and methods**

### 94 **Spinach cultivars**

95 The four tested cultivars of spinach used in the present study were DMMNKS and CY (low  
96 Cd accumulating cultivars) and CJQNDH and CJQLDY (high Cd accumulating cultivars). Prior  
97 study (unpublished data) established the characteristics of these lines. Shoot Cd concentrations of  
98 DMMNKS, CY, CJQNDH and CJQLDY grown in Cd contaminated soil (Cd concentration up

99 to 0.79 mg kg<sup>-1</sup>) were 0.49, 0.44, 1.72 and 1.40 mg kg<sup>-1</sup>. The high-Cd group had tissue  
100 concentrations generally 3.4-fold higher than that of the low-Cd group.

### 101 **Preparation of plant samples and experimental treatments**

102 Seeds of the tested cultivars were sterilized by 2% (v/v) H<sub>2</sub>O<sub>2</sub> for 10 min and then sown  
103 into a cuboid pot (60 × 40 × 8 cm) filled with vermiculite at a rate of 80 seeds pot<sup>-1</sup>. Hoagland's  
104 nutrient solution solution was applied every day to maintain the moisture content of the culture  
105 media and provide the necessary nutrients. The Hoagland solution containing 5 mmol·L<sup>-1</sup>  
106 Ca(NO<sub>3</sub>)<sub>2</sub>·2H<sub>2</sub>O, 5 mmol·L<sup>-1</sup> KNO<sub>3</sub>, 2 mmol·L<sup>-1</sup> MgSO<sub>4</sub>·7H<sub>2</sub>O, 1 mmol·L<sup>-1</sup> KH<sub>2</sub>PO<sub>4</sub>,  
107 0.1mmol·L<sup>-1</sup> EDTA-Fe, 47 μmol·L<sup>-1</sup> H<sub>3</sub>BO<sub>3</sub>, 1 μmol·L<sup>-1</sup> MnCl<sub>2</sub>·4H<sub>2</sub>O, 1 μmol·L<sup>-1</sup>  
108 ZnSO<sub>4</sub>·7H<sub>2</sub>O, 0.01 μmol·L<sup>-1</sup> H<sub>2</sub>MoO<sub>4</sub>, and 0.25 μmol·L<sup>-1</sup> CuSO<sub>4</sub>·5H<sub>2</sub>O. The pots were placed in  
109 a greenhouse at the Guangdong University of Petrochemical Technology (Maoming City, China)  
110 with light intensity of 500-800 μmol m<sup>-2</sup> s<sup>-1</sup>, relative humidity of 40% - 45%, and day / night  
111 temperatures of 30°C / 25°C.

112 A separate hydroponic experiment using 500 mL containers was conducted to test the genotype  
113 differences in Cd chemical forms in the plant tissues. Each container was filled with 400 mL  
114 Hoagland solution with different concentrations of Cd and P as treatments. Control (with no Cd  
115 added, designated as Cd0) and two Cd treatments (adding 1 mg L<sup>-1</sup> and 5 mg L<sup>-1</sup> of Cd in form of  
116 Cd(NO<sub>3</sub>)<sub>2</sub> into the culture solution, designated respectively as Cd1 and Cd2) were conducted.  
117 The P treatments were established by reducing the concentration of KH<sub>2</sub>PO<sub>4</sub> to half of the typical  
118 Hoagland solution concentration (P1) or left at the typical solution concentration (P2). Overall,  
119 there were six Cd-P levels, which were assigned as Cd0P1, Cd0P2, Cd1P1, Cd1P2, Cd2P1 and  
120 Cd2P2. Concentrations of free Cd<sup>2+</sup> in the solutions calculated using Geochem-EZ program



121 (Shaff et al., 2010) for Cd1P1, Cd1P2, Cd2P1 and Cd2P2 were 1.1, 1.0, 23.0, and 21.5  $\mu\text{g L}^{-1}$ ,  
122 respectively.

123 On Jan. 4, 2012, the 20<sup>th</sup> day after the seeding of spinach, four seedlings with uniform size  
124 and with four leaves were identified. The seedlings were transplanted to each 500 mL plastic  
125 container covered by a cap that allowed four plants to be established in each container. The  
126 seedlings were passed through a hole (15 mm diameter) in the cap and held in place with sterile  
127 cottons. The experiment used a completely randomized design with three replicates per treatment.  
128 Thus, there are a total of 72 containers (4 cultivars  $\times$  6 Cd-P levels  $\times$  3 replicates) in the  
129 hydroponic experiment.

130 Sampling of both shoots and roots was carried out on Jan. 19, 2012 after a 15-day growth  
131 period. All shoot samples were thoroughly rinsed with deionized water and roots were with a 0.5  
132 mM  $\text{CaCl}_2$  solution for 30 min and then rinsed with deionized water. Each tissue sample was  
133 weighed, frozen in liquid nitrogen, and stored at  $-80^\circ\text{C}$  until use.

#### 134 **Extraction of Cd in different chemical forms**

135 Cadmium associated with various chemical forms in the plant tissues was determined by  
136 successively extraction tissues with the following sequence of solutions (Wu et al. 2005):

137 (1) 80% ethanol ( $F_E$ ), extracting inorganic Cd associated with nitrate, chloride, or  
138 aminophenol Cd;

139 (2) distilled water ( $F_D$ ), extracting water-soluble Cd associated with organic acids or as  
140  $\text{Cd}(\text{H}_2\text{PO}_4)_2$ ;

141 (3) 1 M NaCl ( $F_{\text{NaCl}}$ ), extracting pectate- and protein-associated Cd;

142 (4) 2% acetic acid ( $\text{HAc}$ ,  $F_{\text{HAc}}$ ), extracting insoluble  $\text{CdHPO}_4$ ,  $\text{Cd}_3(\text{PO}_4)_2$ , and other Cd-  
143 phosphate complexes;

144 (5) 0.6M HCl ( $F_{\text{HCl}}$ ), extracting Cd in oxalate;

145 (6) Cd in residues ( $F_{\text{residue}}$ ).

146 Frozen plant materials were cut into small pieces of 1–2 mm<sup>2</sup>, mixed with 37.5 mL of the  
147 appropriate extraction solution and incubated at 30°C for 18 h. The extraction solution was then  
148 separated and the residual material was re-extracted an additional volume of the same extraction  
149 solution (37.5 mL) under the same conditions for another 6 h. The two extracts were combined.  
150 This double extraction procedure was repeated a second time for the plant tissue. The residual  
151 plant material was extracted with the next extraction solution in the sequence, using the same  
152 procedure described above. All of the extracts (150 mL for each) were evaporated to constant  
153 mass and digested in a microwave digester (WX-8000, Shanghai Xinyi) with an oxidizing  
154 mixture of acids ( $\text{HNO}_3$ – $\text{HClO}_4$ , 5:1, v/v). The digests were used for analysis of Cd  
155 concentration.

#### 156 **Analysis for Cd**

157 Cadmium concentrations in the digests were determined by FAAS (Hitachi Z-5300, Japan).  
158 The precision of the analytical procedures for plant material was assessed using a Certified  
159 Reference Material (CRM) (GBW-07603) provided by the National Research Center for CRM,  
160 China. Total Cd concentrations in shoot and root samples were determined with the same method  
161 following acid digestion with  $\text{HNO}_3$ – $\text{HClO}_4$  (4:1, v/v). The Cd concentrations were based on the  
162 fresh weights of samples before separation or extraction.

#### 163 **Data statistics**

164 Total Cd concentration of each tissue was obtained by summing Cd concentrations in the 6  
165 fractions for shoots or roots. The Brown and Forsythe test and O'Brien's test were applied to the  
166 data prior to further analysis to examine homogeneity in the data. Homogeneity was confirmed

167 for all data sets so a three-way ANOVA (full model and reduced models) was performed on the  
168 data followed by so the least significant difference (LSD) was performed. Analysis were carried  
169 out in SAS 9.3 (Cary, NC).

170

## 171 **Results**

172

### 173 **Total Cd concentration**

174 Total shoot and root Cd concentrations in the tested cultivars of spinach under different Cd-  
175 P treatments are shown in Table 1. The control was undetectable, hence the data are not shown.  
176 Results from three-way ANOVA for the data of shoot Cd concentrations (Table 2) indicated that  
177 the effect of cultivar, Cd concentration, and P concentration were all significant ( $P<0.05$ ).  
178 Significance was also observed in Cd×P interaction ( $P<0.05$ ), but no significance was  
179 determined in cultivar×Cd, cultivar×P and cultivar×Cd×P interactions ( $P>0.05$ ).

180 Low-Cd cultivars (DMMNKS and CY) generally had significantly lower shoot Cd  
181 concentrations ( $p<0.05$ ) than the high-Cd cultivars (CJQNDH and CJQLDY) except for the  
182 Cd1P2 treatment (Table 1). Average shoot Cd concentrations in the low-Cd cultivars were only  
183 66.0% (P1) and 71.0% (P2) of those in the high-Cd cultivars under Cd1, while under Cd2, the  
184 differences were higher at 82.1% (P1) and 80.52% (P2). This indicated that higher Cd exposure  
185 would lead to a decrease of genotype difference in shoot Cd accumulation. Considering effects  
186 of both cultivar×P and cultivar×Cd×P interactions were not significant, level of phosphorus in  
187 cultivating solution might be less related to the genotype difference in shoot Cd accumulation in  
188 spinach, although higher level of phosphorus, compared to lower level, declined shoot Cd  
189 accumulation in both high-Cd and low-Cd cultivars.

190 The mean reduction in shoot Cd concentration as a function of cultivars (-39.0 to -51.5%,  
191 (average shoot Cd concentration in low-Cd cultivars - average shoot Cd concentration in high-Cd  
192 cultivars) / average shoot Cd concentration in high-Cd cultivars  $\times$  100) were greater than those  
193 by P supplement (-8.7 to -18.4%), (average shoot Cd concentration under P2 treatments - average  
194 shoot Cd concentration under P1 treatments) / average shoot Cd concentration under P2  
195 treatments  $\times$  100) under Cd1. Under Cd2, however, those mean reductions from P treatment (-  
196 28.4 to -31.1%) were greater than those from the different cultivar (-21.7 to -24.2%). These  
197 results illustrate why there was a significant variation in Cd $\times$ P interaction according to the three-  
198 way ANOVA.

199 For total root Cd concentrations, it was found that the effect of cultivar was not significant  
200 ( $P>0.05$ ) according to three-way ANOVA (Table 2), although the concentrations of low-Cd  
201 cultivars were all lower than those of high-Cd cultivars. The effect of Cd and P concentrations  
202 were each significant ( $P<0.05$ ). Similar to shoots, variation in root Cd concentrations derived  
203 from Cd $\times$ P interaction were significant ( $P<0.05$ ), but insignificant for those from cultivar $\times$ Cd,  
204 cultivar $\times$ P and cultivar $\times$ Cd $\times$ P interactions ( $P>0.05$ ).

205 More intense differences in Cd concentration in roots in response to the P treatment were  
206 observed. The mean decreased in response to the P treatment (-66.3 to -71.5% under Cd1 and -  
207 30.0 to -51.6% under Cd2) were generally greater than for the cultivar effect (-12 to -51.6%  
208 under Cd1 and -13.8 to -32.8% under Cd2). Different from the shoot Cd concentrations, the  
209 decrease in root Cd concentration in response to the P treatment was smaller under the Cd2  
210 treatment than the Cd1 treatment. Consistent to that in shoot, level of phosphorus in cultivating  
211 solution seemed no significant influence on genotype difference in root Cd accumulation.

212

### 213 **Cd concentrations in different chemical forms**

214 Shoot and root Cd concentrations in different chemical forms of the tested cultivars as well  
215 as results of two-way ANOVA are shown in Table 3 and 4. For the shoots, the most obvious  
216 genotype associated responses were observed in  $F_{\text{NaCl}}$ , and the differences of shoot Cd in the  
217 fraction between low-Cd and high-Cd cultivars were significant under Cd1P1, Cd2P1 and Cd2P2  
218 ( $p < 0.05$ ). The Cd in  $F_{\text{NaCl}}$ ,  $F_{\text{HAc}}$  and  $F_{\text{HCl}}$  revealed a consistent change pattern that P2 treatment  
219 significantly decreased their concentrations unrelated to cultivar under Cd1P1, Cd2P1 and  
220 Cd2P2. For the roots, there was no any Cd fraction exhibited significant variation derived from  
221 cultivar under all of the Cd-P treatments. However, P2 treatment significantly increased Cd  
222 concentrations in  $F_{\text{D}}$  under Cd2 treatment and significantly lowered Cd concentrations in  $F_{\text{NaCl}}$   
223 and  $F_{\text{HCl}}$  under both Cd treatments ( $p < 0.05$ ). These results indicated that the P treatment affected  
224 Cd speciation in spinach more effectively than the cultivar alternation did, which is consistent  
225 with those observed in the total Cd accumulation.

226

### 227 **Proportions of Cd in different chemical forms**

228 Proportions of Cd in different chemical forms in shoots and roots are shown in Figure 1 and  
229 Figure 2. In both shoots and roots, the proportions exhibited a general trend of  $F_{\text{NaCl}} > F_{\text{HAc}} >$   
230  $F_{\text{HCl}} > F_{\text{D}} > F_{\text{E}} > F_{\text{Residue}}$ . This result indicated that Cd in  $F_{\text{NaCl}}$ , which accounted for more than  
231 50% of total Cd in both shoots and roots, played the most important role in Cd accumulation and  
232 detoxification in spinach. Differences in the proportions of Cd in  $F_{\text{NaCl}}$  between low-Cd and  
233 high-Cd groups were not obvious, and the proportions for the low-Cd group were generally  
234 lower than or similar to those of the high-Cd group in both shoots and roots. Cd-P treatments did

235 not consistently influenced the proportion of Cd in  $F_{NaCl}$  in both shoots and roots, but Cd2  
236 treatments increased the proportion in roots when compared to Cd1 treatments.

237 The sums of proportions of Cd in  $F_{HAc}$ ,  $F_{HCl}$  and  $F_{Residue}$ , which were presumed to be forms  
238 with lower mobility within the plant, were 24%-36% in shoots and 24-40% in roots, and were  
239 generally higher in low-Cd cultivars than in high-Cd cultivars especially for those under Cd2.  
240 The average proportions of Cd in  $F_{HAc}$  in shoots were 19.88% (P1) and 16.81% (P2) for low-Cd  
241 cultivars, higher than those of high-Cd cultivars (17.94% under P1 and 15.09% under P2). The  
242 average proportions in roots were 17.40% (P1) and 23.64% (P2) for low-Cd cultivars and also  
243 higher than those of high-Cd cultivars (14.16% under P1 and 21.80% under P2). The average  
244 proportions displayed higher value under P2 than under P1 for both low-Cd and high-Cd groups,  
245 indicating that higher level of phosphorus can enhance formation of Cd-phosphates.

246 Under Cd2, the total proportions of Cd in  $F_{HAc}$ ,  $F_{HCl}$  and  $F_{Residue}$  greatly decreased in both  
247 shoots and roots compared to those under Cd1, indicating that the capacity to chemically  
248 deactivate Cd *in vivo* were restrained when Cd stress increased from Cd1 to Cd2. The sums of  
249 the proportions generally decreased in shoots but increased in roots with the P concentration was  
250 increased from P1 to P2, implying different effects of P on Cd chemical forms between the  
251 shoots and roots.

252 For the proportions of Cd in  $F_E$  and  $F_D$ , the fractions with higher activity, the sums were  
253 11%-20% in shoots and 6%-11% in roots, and were not consistently different between the low-  
254 Cd and the high-Cd cultivars for either shoots or roots. This demonstrated that these two  
255 fractions did not differ as a function of cultivar. The sums under Cd2 were generally higher than  
256 those under Cd1 in shoots, but were reversed in roots, indicating perhaps that roots of spinach  
257 could more effectively deactivate Cd under higher Cd exposure than shoots. The sums of

258 proportions in both tissues of all the tested cultivars (except cv. CJQNDH) were higher under P2  
259 than under P1.

## 260 **Discussion**

### 261 **Genotype-dependent Cd accumulation in spinach**

262 In the present study, differences in total Cd concentrations in shoots and roots between low-  
263 Cd and high-Cd cultivars of spinach under hydroponic condition were consistent with the results  
264 obtained under soil culture condition in our previous unpublished study. Hence, the specific  
265 genotype differences in Cd accumulation are stable, reproducible traits and not specifically  
266 dependent on the growth conditions. Similar results have been obtained in many crops such as  
267 rice (*Oryza sativa* L.) (Yu et al. 2006), asparagus bean (*Vigna unguiculata* subsp. *Sesquipedalis*  
268 L.) (Zhu et al. 2007), hot pepper (*Capsicum annuum* L.) (Xin et al. 2014), water spinach (Wang  
269 et al. 2009), Chinese flowering cabbage (Qiu et al. 2011a), small Chinese cabbage or pakchoi  
270 (*Brassica chinensis* L.) (Xue et al. 2014), Chinese leaf mustard (*Brassica juncea* L. Czern. et  
271 cross. var. *juncea*) (Dai et al. 2012), and amaranth (*Amaranthus* spp.) (Zhou et al. 2013). Some  
272 researchers have investigated the genetic mechanisms regulating Cd accumulation and  
273 detoxification, and special attention has been given to phytochelatins (PCs), a type of Cd-  
274 induced metal-binding proteins (peptides) in plants. Phytochelatins are a class of glutathione-  
275 derived peptides which can help to transport Cd into vacuole in the form of a Cd-PC complex  
276 (Clemens, 2006). RNAi-mediated silencing of *OsPCS1* had been attempted and resulted in  
277 reduction of Cd accumulation in the RNAi rice seeds approximately by half (Li et al. 2007). It  
278 was found that Cd-sensitive barley genotype had less Cd integrated with proteins/pectates as  
279 compared with Cd-resistant genotypes (Wu et al. 2005). Beside Cd tolerance, Cd accumulation  
280 was also found to be associated with proteins/pectates-bound Cd in certain vegetable crops such  
281 as Chinese flowering cabbage (Qiu et al. 2011a) and amaranth (Zhou et al. 2013). These results



282 established the relationship between PC-Cd complexes and certain Cd chemical form i.e. the  
283 NaCl extractable fraction.

284 Much high Cd accumulations were found in the tested cultivars of spinach in both the  
285 previous and the present study. According to our previous study, the maximum shoot Cd  
286 concentration among the 29 tested cultivars was 145.4 mg kg<sup>-1</sup> (dry weight basis) in soil  
287 containing 14.1 mg kg<sup>-1</sup> Cd (unpublished data). According to the water content in shoots (about  
288 90%) of spinach under soil culture conditions in the previous study, shoot Cd concentration of  
289 the high-Cd cultivars under Cd1 (1 mg L<sup>-1</sup>) in the present study would be >100 mg kg<sup>-1</sup> (dry  
290 weight basis), exceeding the critical level for Cd hyperaccumulator (Baker et al. 1989), and it  
291 would be >300 mg kg<sup>-1</sup> under Cd2 treatment (5 mg L<sup>-1</sup>). Hence, spinach is a crop with high Cd  
292 pollution risk once cultivated under Cd contaminated soils and identification and popularization  
293 of low-Cd cultivars are crucial way for ensuring food safety in spinach production. Based on the  
294 genotype-dependent Cd accumulation of spinach verified in the present study, breeding of low-  
295 Cd cultivars of the species should be considered.

#### 296 **Chemical mechanisms related to genotype difference in Cd accumulation of spinach**

297 The profile of Cd chemical forms in shoots and roots of spinach was characterized by a high  
298 proportion of F<sub>NaCl</sub>. It was found that the greatest amount of Cd was extracted by 1 M NaCl and  
299 this accounted for >50% of the Cd in both shoots and roots. This result has been observed in  
300 several vegetable crops. Qiu et al. (2011b) found that proportions of Cd in F<sub>NaCl</sub> in shoots and  
301 roots of Chinese flowering cabbage grown under Cd contaminated soils were close to or  
302 exceeded 50%. Dai et al. (2012) reported that proportions of Cd in F<sub>NaCl</sub> in shoots of Chinese leaf  
303 mustard were > 40%. For vegetable amaranth, proportions of Cd in F<sub>NaCl</sub> in stems and roots

304 were also predominated (40%-60%) when plants were grown in Cd contaminated soils (Zhou et  
305 al., 2013).

306 Similar to studies mentioned above, the proportion of Cd in the  $F_{NaCl}$  fraction of shoots from  
307 spinach were generally lower in low-Cd cultivars than in high-Cd cultivars. Under Cd1 treatment,  
308 significant genotype differences of shoot Cd were only appeared in the  $F_{NaCl}$  according to 2-way  
309 ANOVA. This may be related to the higher capacity in the high-Cd cultivars to resist the toxic  
310 effects involving in phytochelatins (PCs). As has been mentioned above, the majority of Cd in  
311  $F_{NaCl}$  is integrated with proteins/pectates, including Cd bound to PCs (Wu et al. 2005). The PC-  
312 Cd complex could pass through vacuole membrane and the Cd could precipitate within the  
313 vacuole as insoluble phosphates. This has been recognized as a major Cd detoxification  
314 mechanism of plants (Clemens, 2006). In roots of spinach, however, proportions of Cd in  $F_{NaCl}$   
315 were similar between low-Cd and high-Cd cultivars, which implied that the Cd in  $F_{NaCl}$  might be  
316 less related to the genotype difference in Cd detoxification and translocation of spinach.

317 Total proportions of Cd in the insoluble fractions ( $F_{HAc}$ ,  $F_{HCl}$  and  $F_{Residue}$ ) became generally  
318 higher in low-Cd cultivars than in high-Cd cultivars. This could be considered as one of the  
319 mechanisms involving in the genotype difference in shoot Cd accumulation of spinach. In some  
320 crops such as pakchoi (Xue et al. 2014) and watercress (Wang, 2013), the proportion of Cd in  
321  $F_{HAc}$  was the greatest for both shoots and roots when the plants were grown under Cd stresses.  
322 As a mechanism associated with Cd accumulation and detoxification, it relies on the formation  
323 of insoluble  $CdHPO_4$ ,  $Cd_3(PO_4)_2$ , and other Cd-phosphates within plant tissues (Clemens, 2006).  
324 For spinach, average proportions of Cd in  $F_{HAc}$  were also higher in low-Cd than in high-Cd  
325 cultivars, indicating that the mechanism of Cd detoxification involving  $F_{HAc}$  was relevant to the  
326 genotype difference in Cd accumulation of spinach.

## 327 **Effect of phosphorus on Cd accumulation of spinach**

328       The change in the P concentration in the culture solution resulted generally in significant  
329 decrease of total Cd concentrations in both shoots and roots of spinach. Similar results were  
330 obtained by Keller et al. (2001), Römer et al. (2002) and Dheri et al. (2007). It was worth noting  
331 that the effect of P concentration on the reduction of shoot Cd accumulation was more significant  
332 than the cultivar effect under higher Cd stress. These results were consistent with those obtained  
333 by Qiu et al. (2011b) in Chinese flowering cabbage. The effect of phosphorus correspond to the  
334 variation in Cd in  $F_{\text{HAc}}$ , which is mainly composed of  $\text{CdHPO}_4$ ,  $\text{Cd}_3(\text{PO}_4)_2$  and other Cd-  
335 phosphate complexes. Qiu et al. (2011b) reported that the proportions of Cd in  $F_{\text{HAc}}$  of the tested  
336 cultivars of Chinese flowering cabbage increased with soil P level, in consistency with an  
337 investigation by Jiang et al. (2007), who found that increased P in soil caused substantial  
338 precipitation of P-Cd complexes in cell wall and vacuoles in corn. A similar finding was reported  
339 in strawberry (*Fragaria ananassa* D.) by Nuzahath et al. (2013). In this study here, the results  
340 obtained for spinach were similar to the above-mentioned studies. Considering that increased P  
341 concentration decreased the proportions of Cd in  $F_{\text{NaCl}}$  and  $F_{\text{HCl}}$  in both shoots and roots, the  
342 lowered Cd accumulation under higher P might be attributed to the elevated Cd precipitation as  
343 insoluble Cd-P complexes.

344       As to the decrease of proportion of Cd in  $F_{\text{NaCl}}$  and  $F_{\text{HCl}}$  in spinach caused by P supply,  
345 similar results was also reported in strawberry (Nuzahath et al. 2013). Since studies on the  
346 relationship between P behavior and Cd chemical forms are, thus far, insufficient, no reasonable  
347 explanation for the phenomenon could be currently given and further investigations are required.

## 348 **Conclusion**

349 Verification of genotype-dependence in Cd accumulation of spinach is provided by  
350 comparing the results from our previous and the present study. Spinach has prominent ability to  
351 accumulate Cd, and shall thus receive more attention in identification and breeding of its low-Cd  
352 accumulating genotypes. The obvious differences in the concentrations of different chemical  
353 forms of Cd between low-Cd and high-Cd cultivars indicated that the hypothesis in the present  
354 study is partly acceptable. That is, there is a genotype-dependent effect on Cd accumulation,  
355 translocation, and detoxification that is likely related to distribution of Cd across the various  
356 chemical forms. An increased supply of phosphorus decreased significantly Cd accumulations in  
357 both high-Cd and low-Cd cultivars without significant difference between the high-Cd and low-  
358 Cd cultivars. Therefore, the external concentration of phosphorus influenced Cd accumulation of  
359 spinach, but might not be a crucial factor that affects genotype difference in Cd accumulation of  
360 the species.

361

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### 366 **Ethical Statement**

367 The authors declare that they have no conflict of interest.

368

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