

**Bioaccumulation of Hg in rice leaf facilitates selenium bioaccumulation in rice (*Oryza sativa L.*) leaf in the Wanshan mercury mine**

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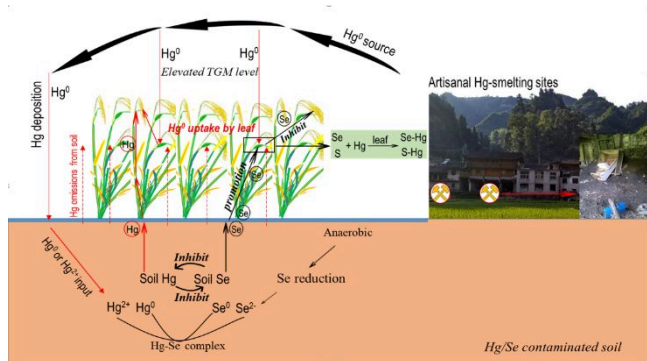
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TOC



**ABSTRACT**

Mercury (Hg) bioaccumulation in rice poses a health issue for rice consumers. In rice paddies, selenium (Se) can decrease the bioavailability of Hg through forming the less bioavailable Hg selenides (HgSe) in soil. Rice leaves can directly uptake a substantial amount of elemental Hg from the atmosphere, however, whether the bioaccumulation of Hg in rice leaves can affect the bioaccumulation of Se in rice plants is not known. Here, we conducted field and controlled studies to investigate the bioaccumulation of Hg and Se in the rice-soil system. In the field study, we observed a significantly positive correlation between Hg concentrations and BAFs of Se in rice leaves ( $r^2 = 0.60$ ,  $p < 0.01$ ) collected from the Wanshan Mercury Mine, SW China, suggesting that the bioaccumulation of atmospheric Hg in rice leaves can facilitate the uptake of soil Se, perhaps through the formation of Hg-Se complex in rice leaves. This conclusion was supported by the controlled study, which observed significantly higher concentrations and BAFs of Se in rice leaf at a high atmospheric Hg site at WMM, compared to a low atmospheric Hg site in Guiyang, SW China. 3

## 38 INTRODUCTION

39 Mercury (Hg) is a pollutant of global concern due to its long-range transport in the  
40 atmosphere, and adverse effects on ecosystems and human health<sup>1, 2</sup>. In aquatic ecosystems, a  
41 fraction of mercury can transform into methylmercury (MeHg), a potential neurotoxin that  
42 has a strong capacity to bioaccumulate along the food chain<sup>3-6</sup>. Mercury contamination in Hg  
43 mines is receiving special attention due to the extensive release of Hg into the surrounding  
44 environment (e.g., atmosphere, water, and soils) during Hg mining activities<sup>7-13</sup>. Mercury  
45 pollution is serious in southwestern China because this area has a number of large Hg mines,  
46 including the Wanshan Mercury Mine (WMM) which is the world's 3rd largest Hg mine<sup>14</sup>. To  
47 make the situation worse, around these mines, there are many rice paddies that contain a few  
48 to hundreds of  $\mu\text{g/g}$  Hg in soils<sup>14, 15</sup> and tens to thousands of  $\text{ng/m}^3$  Hg in the ambient air<sup>16, 17</sup>.  
49 Rice accumulates inorganic Hg through (1) leaf uptake of gaseous elemental Hg ( $\text{Hg}^0$ ) from  
50 the atmosphere and (2) root uptake of bioavailable Hg species from the soil<sup>18</sup>. More  
51 importantly, rice paddies, are hotspots of MeHg production<sup>19-21</sup>. High levels of MeHg are  
52 commonly found in rice near Hg mines<sup>14, 22-25</sup>. At mercury mining sites and in inland China  
53 where rice consumption is higher compared to fish consumption, rice is a major MeHg  
54 exposure source to local residents<sup>26</sup>.

55 Selenium (Se), an essential element and antioxidant, can antagonize the toxicity of Hg and  
56 many heavy metals (e.g., Cd and Cr)<sup>27-35</sup> via the formation of less bioavailable Hg-Se particles  
57 in animal and human bodies<sup>31, 33, 36, 37</sup>. An approximate daily intake of Se of 50  $\mu\text{g/day}$  has  
58 been shown to be essential and healthful for the human body<sup>38</sup>. While approximately 72% of  
59 Chinese land is in a Se-deficient state<sup>39</sup>, many Se-rich areas were recently found including

60 WMM. A recent study demonstrated that the soil in WMM contains 0.16 to 36.6  $\mu\text{g/g}$  of Se<sup>40</sup>,  
61 which is 1 to 3 orders of magnitude higher than the abundance of Se in Earth's crust (50 ng/g)  
62 and comparable with that reported in soils from other seleniferous areas<sup>41-44</sup>.

63 At high concentrations, Se has been proven to result in 8~72% of the decrease in the  
64 accumulation of Hg in rice grains through the formation of less bioavailable mercury  
65 selenides (HgSe) in soil and on the root surface<sup>32, 45, 46</sup>. In flooded paddies, the anaerobic and  
66 reducing conditions favor the interaction between Se and Hg due to the higher affinity  
67 constant of Hg to Se ( $10^{45}$ ) and lower solubility constant of HgSe ( $K_{SP} = 1.0 \times 10^{-59}$ ) than of  
68 Hg sulfides (affinity constant:  $10^{39}$ ;  $K_{SP}$ :  $1.6 \times 10^{-52}$  for  $\alpha$ -HgS and  $4.0 \times 10^{-53}$  for  $\beta$ -HgS)<sup>40, 47, 48</sup>.  
69 Mercury favors binding to thiol (-SH) functional groups over other elements in organisms<sup>49, 50</sup>.  
70 However, due to the high-affinity constant of Hg to Se, the complexation between Hg and  
71 -SeH has also been found in organisms<sup>36, 51</sup>.

72 In WMM, which is the "Capital of Mercury in China", historic large-scale mining  
73 activities and ongoing illegal artisanal Hg mining activities have resulted in extremely high  
74 Hg levels in the soil and atmosphere<sup>15, 17</sup>. In such a high Hg background, while the inhibition  
75 of Hg bioaccumulation in rice by Se has been reported<sup>40</sup>, the effects of Hg contamination on  
76 Se bioaccumulation in rice remains a mystery. As Se may antagonize the toxicity of Hg,  
77 understanding the effects of Hg contamination on Se bioaccumulation in rice is critical to  
78 evaluate the risk and toxicity of Hg in rice. A laboratory-controlled study demonstrated that  
79 adding Hg into the culture solution can promote the translocation of Se to garlic tissues, and  
80 suggested that Se can balance the Hg stress by the formation of reduced Se ( $\text{Se}^{2-}$ ) in garlic  
81 tissues<sup>52</sup>. As plant leaves mainly uptakes  $\text{Hg}^0$  from the atmosphere<sup>18</sup>, the coexistence of

82 reduction state of Se forms ( $\text{Se}^0$  or  $\text{Se}^{2-}$ ) and oxidation state of Hg ( $\text{Hg}^0$  or  $\text{Hg}^{2+}$ ) in leaves  
83 provided a possible reaction site where Hg may react with Se. Based on the garlic study, there  
84 may be an increase in Se translocation in rice plants at regions where soil and atmospheric Hg  
85 concentrations are high. In these regions, more Se is possibly needed to antagonize the  
86 toxicity of Hg in plant tissues.

87 Here, we conducted field and controlled studies to investigate the bioaccumulation of Hg  
88 and Se in the rice-soil system. In the field study, we investigated the distribution of Hg and Se  
89 in rice plants and corresponding rhizosphere soil at both artisanal mining sites and  
90 non-artisanal mining sites in WMM. In the controlled study, we conducted pot experiments  
91 regarding growing rice plants on a Se-rich soil at high TGM site in the WMM and low TGM  
92 site in Guiyang (GY). We aim to (1) test whether the excessive soil Hg could inhibit the  
93 uptake of soil Se by roots due to the formation of more HgSe in rhizosphere soil, or (2) test  
94 whether atmospheric Hg in rice leaves could facilitate Se bioaccumulation in rice leaves.

95

## 96 MATERIALS AND METHODS

97 **Field study.** To gain a first understanding of the interactions between Hg and Se in the  
98 rice-soil system, rice plants and corresponding rhizosphere soil were collected at 25 sites in  
99 the WMM area, SW China (Figure S1), in September 2017. Prior to sample collection, the  
100 TGM concentration at each site (~0.5 m above ground) was measured 3 times in July, August,  
101 and September, with > 30 min each time, using an automated Hg vapor analyzer (LUMEX,  
102 RA-915 AM, Russia). The averaged TGM data of each site was used to reflect a long-term  
103 TGM concentration. Twelve of the sampling sites were near artisanal Hg smelters (hereafter,

104 artisanal mining sites), while the remainder lacked artisanal Hg smelting activities (hereafter,  
105 non-artisanal mining sites). At each site, three  $2 \times 2$  m plots were established for sample  
106 collection. At each plot, three rice plants and corresponding rhizosphere soils (0 to 20 cm  
107 depth) were randomly sampled. The rice and soil samples from the three plots were combined  
108 to represent each site. Soil and rice samples were stored in a cooler ( $-18$  °C) and delivered to  
109 the laboratory following collection. Soil samples were freeze-dried ( $-79$  °C), crushed,  
110 homogenized, and passed through 200 mesh. Rice plants were separated into different tissues  
111 (root, stem, leaf, and grain). The hull and bran of grain samples were removed to obtain  
112 polished rice. Then, root, stem, leaf and polished rice samples were washed thoroughly with  
113 tap water followed by  $18.2$  M $\Omega$  water (Milli-Q® Integral System), freeze-dried ( $-79$  °C),  
114 weighed, and powdered using a grinding machine (IKA®A11 basic)<sup>41</sup>. All soil and rice  
115 samples were sealed in polyester plastic bags and stored at room temperature, prior to further  
116 analysis.

117 **Controlled study.** To further investigate if the bioaccumulation and translocation of Se can  
118 be affected by Hg accumulation in rice leaf, pot experiments regarding growing rice plants on  
119 Se-rich soil, were performed at high TGM site in the WMM and low TGM site in Guiyang  
120 (GY) in 2018. An active Hg related chemical plant was located nearby the WMM site.  
121 According to our study, the TGM concentration at the WMM site during the entire growing  
122 season ranges 24 to 23842 ng/m<sup>3</sup> (geomean: 1556 ng/m<sup>3</sup>), which are 1-3 magnitudes higher  
123 than that at the Guiyang site (range: 5 to 19 ng/m<sup>3</sup>; geomean: 9.6 ng/m<sup>3</sup>). The Se-rich soil,  
124 containing  $7.66 \pm 0.16$   $\mu\text{g/g}$  of Se and  $389 \pm 16$  ng/g of Hg, was collected from the Enshi  
125 seleniferous area, Hubei province, China. The soil was fully mixed before using in the pot

126 experiment. Storage boxes, 45 cm × 34 cm × 30 cm in size and each contains ~ 20 kg of the  
127 Enshi soil, were used for rice growing. Three boxes at each site and the soil layer is ~ 20 cm  
128 deep in each box. Three rice seeds (*Oryza . Sativa L*) were planted in each box. During the  
129 growing season, commercial drinking water of the same brand (Long Men drinking water Co.,  
130 Ltd.) was used for irrigation at the same time-frequency. A transparent plastic cloth was  
131 placed ~ 2 m above the box to prevent wet Hg deposition to the box. The TGM concentrations  
132 at the Guiyang site and the Wanshan site were measured 3 times in July, August, and  
133 September, with > 30 min each time, using the LUMEX automated Hg vapor analyzer. Rice  
134 samples were harvested at the end of September 2018. Soil and rice plants were sampled and  
135 processed following the method described above.

136 **Mercury concentration analysis.** For THg analysis, approximately 0.1 g of soil samples  
137 were digested in a water bath (95 °C, 6 hours) using 5 mL of aqua regia (HCl/HNO<sub>3</sub> = 3/1,  
138 v/v), and measured by F732–VJ cold vapor atomic absorption spectrometry (detection limit:  
139 0.05 ng/mL Hg) following a previous method<sup>18, 53</sup>. Approximately 0.2 g of rice tissues were  
140 digested with 5 mL of HNO<sub>3</sub> and H<sub>2</sub>SO<sub>4</sub> (4/1, v/v), and measured by Tekran 2500 cold vapor  
141 atomic fluorescence spectroscopy (detection limit: 0.1 pg Hg) following the US EPA Method  
142 1631<sup>54</sup>.

143 **Se concentration analysis.** TSe concentrations of soil and rice samples were measured  
144 following a previous method<sup>41</sup>. Briefly, approximately 0.1 g of soil samples and  
145 approximately 0.2 g of rice samples were weighed into 15 mL Teflon bombs. Soil samples  
146 were digested by 2.5 mL HNO<sub>3</sub> and 0.5 mL of HF, and rice samples were digested by 2.9 mL  
147 HNO<sub>3</sub> and 0.1 mL of HF. The Teflon bomb was placed in a steel can and heated in an oven



148 (155°C) for 36 and 18 hours, respectively, for soil and rice samples. The bombs were then  
149 screwed open, and supplemented with 1 ml of 30% (v/v) H<sub>2</sub>O<sub>2</sub> and heated on a hot plate  
150 (90°C) until the solution was evaporated to near dryness. The residual solution was added to 3  
151 mL of 6 mol/L HCl and heated in a water bath (95°C) for 2 hours, and then diluted to 15 mL  
152 with 18.2 MΩ water for hydride generation atomic fluorescence spectrometry analysis  
153 (HG-AFS 9700, BJHG, China).

154 **Quality control.** The standard reference materials GBW07405 (yellow-red soil), GBW10020  
155 (citrus leaf) and BCR-482 (lichen) and sample replicates were included during both THg and  
156 TSe analysis. The recoveries of Hg for GBW07405, GBW10020 and BCR-482 are 109 ± 6%  
157 (n=6), 101 ± 7% (n=6), and 88 ± 2% (n=3), respectively, and the recoveries of Se for  
158 GBW07405 and GBW10020 are 106 ± 5% (n=5) and 93 ± 4% (n=9), respectively. Duplicate  
159 analysis of soil and rice plant tissue samples were conducted in every ten samples, the relative  
160 standard deviations of THg and TSe of all duplicate samples were all within 5% (n=22).

161 **Bioaccumulation factors of Hg and Se in rice tissues.** Bioaccumulation factors (BAFs) of  
162 Hg and Se in rice tissues were calculated using the following equation:

$$163 \quad \text{BAF}_{\text{tissue}} = C_{\text{tissue}}/C_{\text{soil}} \quad (1)$$

164 where  $C_{\text{tissue}}$  is the THg (or TSe) concentration of rice tissue and  $C_{\text{soil}}$  is the THg (or TSe) of  
165 the corresponding rhizosphere soil.

166 **Statistical analysis.** Correlation analyses and T-test were performed using IBM SPSS 22.0  
167 software. Correlation coefficients ( $r^2$ ) and significance probabilities ( $p > 0.05$  is insignificant;  
168  $p < 0.05$  is significant;  $p < 0.01$  is very significant) were computed for regression fits. T-test  
169 was performed to compare whether Hg or Se concentrations and BAFs in rice tissues differed

170 significantly between artisanal and non-artisanal mining sites. Graphical analyses were  
171 performed by Origin 2019 and Microsoft Office 365.

172

## 173 **RESULTS**

174 **Mercury and selenium distribution in the field study.** THg and TSe concentrations in the  
175 field study are summarized in Table 1 and detailedly described in Text S1. Briefly, artisanal  
176 mining sites show significantly higher TGM levels (geomean: 369 ng/m<sup>3</sup>) and soil THg  
177 concentrations (geomean: 36.8 µg/g) than non-artisanal mining sites (geomean TGM: 38.8  
178 ng/m<sup>3</sup>; geomean soil THg: 13.5 µg/g), as shown in Figure 1. Slightly lower soil TSe can be  
179 found at artisanal mining sites (geomean: 1.65 µg/g) than non-artisanal mining sites  
180 (geomean: 2.13 µg/g). Significantly positive correlations between soil THg and soil TSe can  
181 be observed at non-artisanal mining sites ( $r^2 = 0.32, p < 0.01$ ) and at artisanal mining sites ( $r^2$   
182  $= 0.35, p < 0.05$ ), suggesting Hg and Se may share a similar source.

183 The concentrations and BAFs of Hg and Se in rice tissues in the field study are  
184 summarized in Table 1. Briefly, significantly higher geomean values of THg and TSe were  
185 observed at artisanal sites, compared to non-artisanal mining sites (Table 1). At all studied  
186 sites, root and leaf have higher THg concentrations and Hg BAFs than grain and stem (Figure  
187 S2). Similarly, at all studied sites, higher TSe concentrations and Se BAFs were also observed  
188 in root and leaf, compared to grain and stem (Figure S3).

189 **Mercury and selenium distribution in the controlled study.** THg and TSe concentrations  
190 of rice tissues in the controlled site are shown in Table 2 and Figure 2. During the growing  
191 season, the THg of soil placed at the Wanshan site increased from 389 ng/g to 535 ng/g,

192 whereas soil placed at the Guiyang site showed consistent THg concentration ( $395\pm 24$  ng/g),  
193 The increase of soil THg at the Wanshan site is thought to be caused by the intensive release  
194 of Hg from the nearby artisanal Hg sites. Soil Se showed no significant variation at both sites  
195 (Wanshan:  $7.77\pm 0.04$   $\mu\text{g/g}$ ; Guiyang:  $7.54\pm 0.14$   $\mu\text{g/g}$ ) during the growing season.

196 The concentrations and BAFs of Hg in rice tissues at the Wanshan site decrease in the  
197 following order: leaf > root > stem > grain (Figure 2A, Table 2). However, at the Guiyang  
198 site, the concentrations and BAFs of Hg decrease as follows: root > leaf > stem > grain. The  
199 concentrations and BAFs of Se at both sites show a consistently decreasing order: root >  
200 leaf > grain > stem (Figure 2B, Table 2).

201

## 202 DISCUSSION

203 **Hg distribution in rice plants in the field study.** At both artisanal and non-artisanal mining  
204 sites, roots and leaves showed much higher THg concentrations and Hg BAFs than other  
205 tissues (Table 1, Figure S2). Meanwhile, THg concentrations of leaf and stem showed  
206 significantly positive linear correlations with TGM, whereas root THg was positively  
207 correlated with soil THg (Table S1). As discussed in Text S2, these correlations are consistent  
208 with previous studies that demonstrated that rice takes up Hg by root and leaf from the soil  
209 and ambient air<sup>18</sup>, respectively, and Hg is not readily translocated among plant tissues<sup>55, 56</sup>.

210 The bioaccumulation of Hg in rice tissues may be inhibited by elevated soil TSe  
211 concentrations<sup>40</sup>. Such a hypothesis can be supported by the negative correlations ( $p < 0.01$ )  
212 between soil TSe and BAFs of Hg in rice tissues at non-artisanal mining sites (Figure 3A). At  
213 non-artisanal mining sites, a positive correlation between TGM and soil THg can be observed

214 ( $r^2 = 0.82$ ,  $p < 0.01$ ), suggesting that TGM mainly originated from the in-situ emission of  $\text{Hg}^0$   
215 from the soil. It is likely the elevated soil TSe at non-artisanal mining sites can reduce the  
216 bioavailability of soil Hg or the emission of  $\text{Hg}^0$  from the soil, through the formation of less  
217 soluble HgSe in soil<sup>57-60</sup>. At artisanal mining sites, however, we did not observe any clear  
218 correlation between soil TSe and BAFs of Hg in rice tissues (Figure 3B). Unlike non-artisanal  
219 mining sites, no significant correlation between TGM and soil THg was observed at artisanal  
220 mining sites, suggesting that the TGM was not mainly emitted from soil, but directly from  
221 artisanal mining activities. Therefore, soil Se seems not to significantly limit the  
222 bioaccumulation of Hg in rice tissues at artisanal mining sites, because of the fact that  
223 atmospheric Hg (mainly emitted from the Hg smelters) was directly uptaken by rice leaves,  
224 and there is a little chance for soil Se to complex with atmospheric Hg.

225 **Se distribution in rice plants in the field study.** The distribution patterns of TSe and Se  
226 BAFs in rice tissues in the field study, as illustrated in Table 1 and Figure S3. At non-artisanal  
227 and artisanal mining sites, the TSe concentrations of rice tissues are all positively correlated  
228 with soil TSe concentrations ( $p < 0.01$  for all), consistent with the fact that rice plant uptakes  
229 Se mainly from soil<sup>41, 44</sup>.

230 As shown in Figure 3C, the BAFs of Se in rice tissues are all negatively correlated with  
231 soil THg concentrations at non-artisanal mining sites ( $p < 0.01$  for all). Hence, we  
232 hypothesized that the uptake of soil Se by rice plant may be inhibited by high soil THg  
233 concentrations, although other mechanisms may exist. This can be explained by the formation  
234 of less soluble HgSe in paddy soil. Indeed, previous studies have demonstrated that Hg-Se  
235 interaction can occur in the rhizosphere by detecting a proportion of Hg-Se in root surface<sup>45</sup>.

236 <sup>46</sup>. Under flooded conditions, oxidized Se species ( $\text{SeO}_4^{2-}$ ,  $\text{SeO}_3^{2-}$ ) can transform to reduced Se  
237 species ( $\text{Se}^{2-}$  or  $\text{Se}^0$ )<sup>40</sup>. Reduced Se species can react with dissolved  $\text{Hg}^{2+}$  or  $\text{Hg}^0$  in soil  
238 solutions, forming Hg-Se complex in soil and rice rhizosphere. HgSe has much lower  
239 mobility and bioavailability compared to Hg-sulfides, due to their much lower  $K_{SP}$  than  $\text{HgS}^{40}$ ,  
240 <sup>47, 48</sup>.

241 At artisanal mining sites, however, we observed no clear correlation between soil THg  
242 and BAFs of Se in rice tissues (Figure 3D). However, compared to non-artisanal mining sites,  
243 relatively higher TSe concentrations and higher BAFs of Se were observed in rice leaves at  
244 artisanal sites ( $p = 0.039$ ,  $t = -2.193$ ). This is contradicting with the slightly lower soil TSe  
245 concentrations at artisanal mining sites, indicating that environmental conditions in the  
246 regions of artisanal Hg mining activities facilitated the bioaccumulation of Se in rice leaves.  
247 As mentioned above, soil Hg tends to decrease the bioavailability of Se in soils, therefore it  
248 should not be the reason for the higher BAFs of Se in rice leaves at non-artisanal mining sites.  
249 For rice leaf, significantly positive correlations were observed between Hg concentrations and  
250 Se BAFs (Figure 4A) and between TGM and Se BAFs (Figure 4B), which implies that the  
251 bioaccumulation of Hg facilitated the uptake of soil Se by leaf.

252 Artisanal Hg mining activities significantly increased the TGM levels at artisanal Hg  
253 mining sites, which resulted in elevated THg levels in rice leaves, perhaps forming Hg-Se  
254 complex in rice leaf. A substantial amount of atmospheric  $\text{Hg}^0$  can pass through stomata, and  
255 be oxidized to  $\text{Hg}^{2+}$  and accumulated by leaf of plant tissue<sup>49, 61</sup>. In leaf and other tissues, a  
256 substantial amount of  $\text{Hg}^{2+}$  binds with sulfur-containing groups (e.g., -SH) to form less  
257 soluble Hg sulfides (e.g.,  $\beta\text{-HgS}$ ), which combat the toxicity of  $\text{Hg}^{49}$ . It should be noted that

258 the leaf is also the site where the transformation of inorganic Se (e.g.,  $\text{Se}^{6+}$  and  $\text{Se}^{4+}$ ) to  
259 organic Se species occurs. Inorganic  $\text{Se}^{6+}$  and  $\text{Se}^{4+}$  are transported from the root to leaf by  
260 sulfate and phosphate transporters and are transformed to organic Se (Se-Met, Se-Cys,  
261 Se-MeSeCys, DMSe, DMDS<sub>2</sub>, etc.) and reduced Se ( $\text{Se}^{2-}$  or  $\text{Se}^0$ ) species, with the  
262 involvement of many enzymes<sup>62-64</sup>. Here, we speculate that the formation of Hg-Se may occur  
263 in rice leaf due to the presence of reduced Se ( $\text{Se}^{2-}$  or  $\text{Se}^0$ ) species, considering the stronger  
264 chemical bonding ability of Se-Hg ( $10^{45}$ ) than of S-Hg ( $10^{39}$ ). The free functional groups of  
265 -SeH are preferentially bound with  $\text{Hg}^{2+}$  over -SH<sup>36,37</sup>, even capturing the Hg that has formed  
266 Hg-(SR)<sub>2</sub> by ligand exchange reaction<sup>65</sup>. The sulfur in  $\beta$ -HgS is readily replaced by Se  
267 through isomorphism<sup>66</sup>. To test our hypothesis, more studies are needed in the future.

268 **Se distribution in rice plants in the controlled study.** Compared to the field study,  
269 significantly higher Se BAFs were observed in the controlled study. This is owing to the use  
270 of Enshi soil in the controlled study. The Enshi soil has previously been shown to have high  
271 Se bioavailability, as indicated by the high concentrations of bioavailable Se species such as  
272 water-soluble Se (0.008-0.175  $\mu\text{g/g}$ ), ligand-exchangeable Se (0.10-1.45  $\mu\text{g/g}$ ), and  
273 organically bound Se (0.61-8.11  $\mu\text{g/g}$ )<sup>41</sup>. The controlled study further supported our  
274 hypothesis that the high TGM, which is the major source of Hg in rice leaves, facilitates the  
275 uptake of soil Se by rice leaves. As shown in Figure 2B, the concentrations and BAFs of Se in  
276 root at the Wanshan site was much lower ( $p = 0.006$ ,  $t = 5.274$ ) than that at the Guiyang site,  
277 which may be due to intensive Hg deposition that decreased Se bioavailability by forming less  
278 bioavailable HgSe species in soil. However, the concentrations and BAFs of Se in rice leaves  
279 at the Wanshan site were surprisingly higher than those at the Guiyang site. In particular,

280 statistic tests suggested the concentrations and BAFs of Se in leaf at the Wanshan site are  
281 significantly higher ( $p = 0.035$ ,  $t = -3.148$ ) than that at the Guiyang site. Rice leaves receive  
282 the majority of Hg from the atmosphere<sup>18</sup>. As the same soil and water were used throughout  
283 the controlled study at both sites, we suggest that the high TGM is an important driver for the  
284 relatively higher concentrations and BAFs of Se in leaves at the Wanshan site.

285

286 **Environmental implications.** The Hg-Se interaction in the rice-soil system, as demonstrated  
287 in this study, provided some new insights into the biogeochemical cycling of both Hg and Se  
288 in the environment. As a toxin, the bioaccumulation of Hg in plants is a critical step for Hg  
289 entering the food web<sup>67</sup>. Selenium in soil has a great potential to limit the bioavailability of  
290 Hg in soil, through the formation of less soluble HgSe, especially in Se-rich regions<sup>40, 53</sup>. In  
291 highly Hg-polluted regions, mercury, in turn, has an undeniable effect on the biogeochemical  
292 fate of Se. According to our study, elevated Hg concentrations could decrease the mobility of  
293 Se in soil, due to the formation of HgSe. Meanwhile, plants receive a substantial amount of  
294 Hg<sup>0</sup> from ambient air, and the uptake of atmospheric Hg by plant leaves can facilitate the  
295 uptake of Se by rice, especially in areas associated with high TGM levels. Although the  
296 mechanism behind this phenomenon remains not well explained by this study, we  
297 hypothesized that interactions between Hg and Se may readily form HgSe in leaf, where  
298 atmospheric Hg<sup>0</sup> and soil Se are transformed to oxidation and reduction states, respectively.  
299 The formation of less soluble HgSe exhausts the available Se species (Se<sup>6+</sup> and Se<sup>4+</sup>) in rice  
300 leaves, which, in turn, facilitates the uptake of soil Se by leaves. The complex of Hg to Se has  
301 been observed in many animal tissues, and such kind of complex has been assumed to prevent

302 negative effects of Hg in animals<sup>68</sup>. The present study implies the same mechanism may also  
303 occur in plant leaves that exposed in a high TGM environment. However, it worthies  
304 mentioning that this study failed to detect the *in situ* presence of HgSe. To verify the  
305 possibility of HgSe formation in rice plant and other plant species, we encourage researchers  
306 to conduct further studies using relevant techniques (e.g., XANES). It also has been reported  
307 that the intervention of massive Hg can promote the uptake of soil Se by garlic plant in which  
308 a substantial amount of HgSe was detected by the XANES<sup>52</sup>.  
309



310 **Supporting Information**

311 **Text S1** THg, TSe concentrations of the WMM soil in the field study; **Text S2** Understand  
312 correlations among soil Hg, TGM, and Hg levels in rice tissues; **Table S1** Pearson's  
313 correlation matrix ( $r$ ) among the Hg levels in paddy soils, air, and tissues of rice plants at  
314 non-artisanal mining sites and artisanal mining sites; **Figure S1** Study area and sampling  
315 sites; **Figure S2** Distribution of Hg in soil and rice tissues; **Figure S3** Distribution of Se in  
316 soil and rice tissues.

317

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324 **Notes**

325 The authors declare no competing financial interest.

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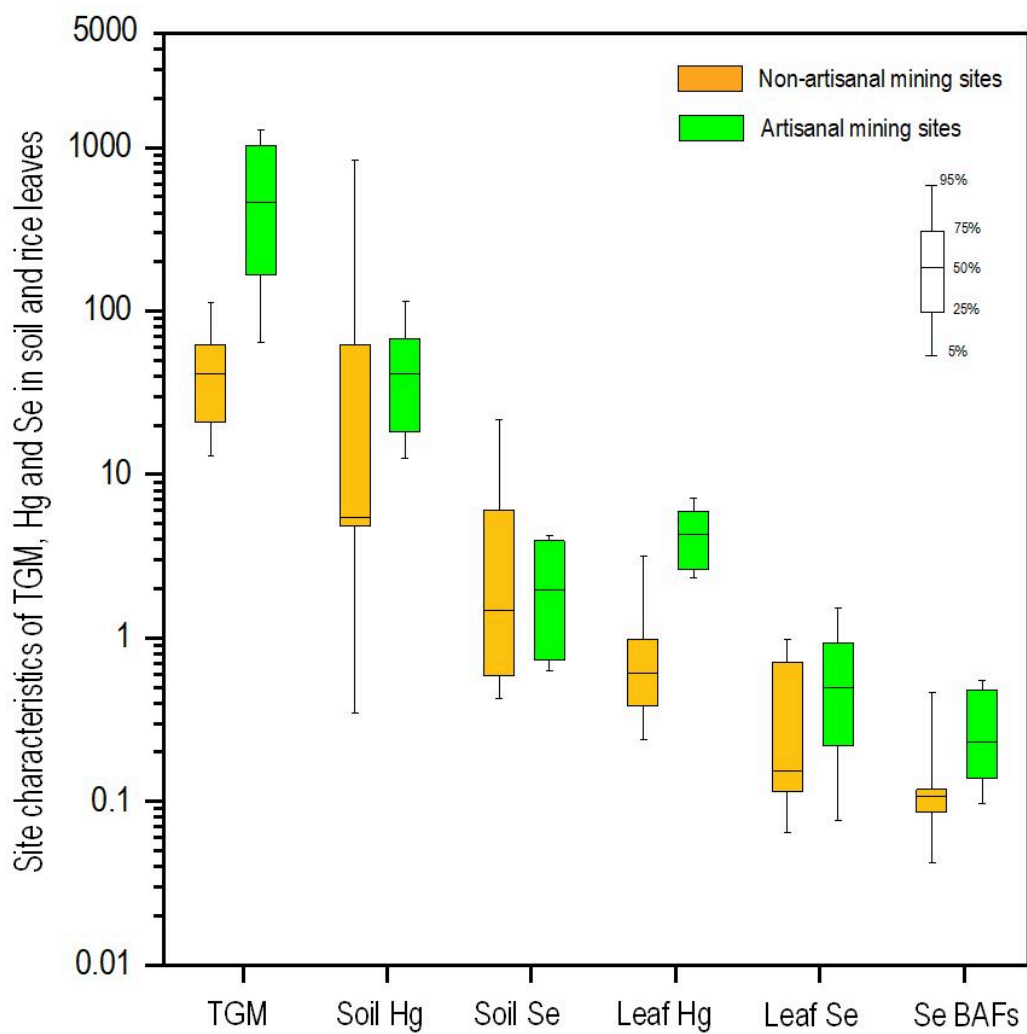
327 **Table 1** Concentrations and BAFs of Hg and Se in soil and rice tissues at non-artisanal mining sites (n=13) and artisanal mining sites (n=12).

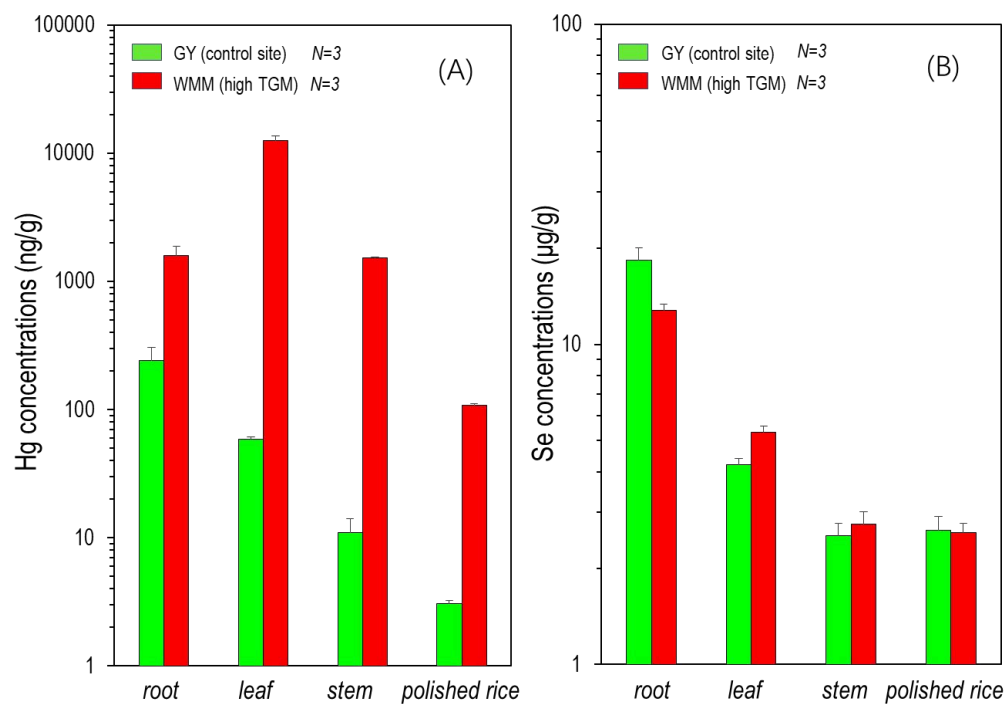
	Non-artisanal mining sites				Artisanal mining sites			
	THg ( $\mu\text{g/g}$ )		Hg BAFs		THg ( $\mu\text{g/g}$ )		Hg BAFs	
	range	mean	range	mean	range	mean	range	mean
TGM ( $\text{ng/m}^3$ )	13~113	38.8			64~1287	369		
Soil	0.35~833.7	13.5			12.5~115	36.8		
Root	0.12~20.1	1.13	0.02~0.69	0.08	0.96~20.6	2.94	0.04~0.20	0.08
Stem	0.018~0.64	0.058	$1 \times 10^{-4}$ ~0.06	0.0043	0.09~1.50	0.34	0.002~0.02	0.0093
Leaf	0.24~3.16	0.65	0.004~0.87	0.048	2.31~7.19	4.09	0.029~0.27	0.11
Polished rice	0.015~0.081	0.031	$2.04 \times 10^{-5}$ ~0.12	0.0023	0.065~0.45	0.14	0.0007~0.0081	0.0038
	TSe ( $\mu\text{g/g}$ )		Se BAFs		TSe ( $\mu\text{g/g}$ )		Se BAFs	
	range	mean	range	mean	range	mean	range	mean
	Soil	0.43~21.7	2.13			0.63~4.20	1.65	
Root	0.19~5.79	0.99	0.04~2.02	0.40	0.18~3.58	1.02	0.23~1.11	0.62
Stem	0.022~0.56	0.11	0.02~0.22	0.05	0.023~0.82	0.16	0.03~0.20	0.10
Leaf	0.065~0.98	0.24	0.04~0.46	0.12	0.077~1.53	0.39	0.10~0.55	0.24
Polished rice	0.029~0.91	0.12	0.016~0.18	0.055	0.026~0.68	0.15	0.032~0.19	0.091

328 **Table 2** Hg and Se concentrations and BAFs of pot rice plants grown in GY (low TGM, n=3) and WMM (High TGM, n=3).

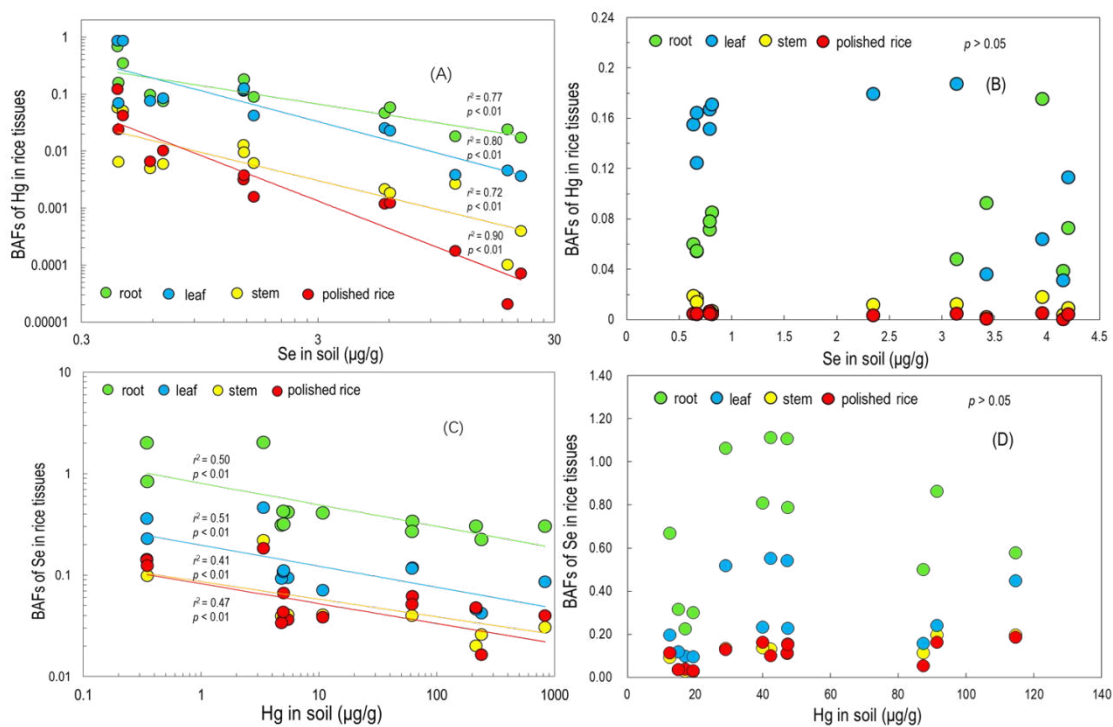
Sites	GY				WMM			
	THg ( $\mu\text{g/g}$ )		Hg BAFs		THg ( $\mu\text{g/g}$ )		Hg BAFs	
	range	mean	range	mean	range	mean	range	mean
TGM ( $\text{ng/m}^3$ )	5~19	9.6			24~23842	1556		
Soil	0.37~0.42	0.40			0.44~0.59	0.54		
Root	0.17~0.30	0.24	0.45~0.71	0.60	1.26~1.86	1.58	2.13~3.68	3.02
Stem	0.009~0.012	0.011	0.02~0.03	0.028	1.51~1.53	1.53	2.60~3.41	2.89
Leaf	0.056~0.062	0.059	0.14~0.16	0.15	11.8~13.7	12.54	21.1~26.7	23.7
Polished rice	0.003~0.003	0.003	0.007~0.009	0.0078	0.095~0.12	0.11	0.17~0.24	0.20
	TSe ( $\mu\text{g/g}$ )		Se BAFs		TSe ( $\mu\text{g/g}$ )		Se BAFs	
	range	mean	range	mean	range	mean	range	mean
Soil	7.42~7.70	7.54			7.73~7.81	7.77		
Root	16.4~19.5	18.3	2.14~2.55	2.40	12.1~13.4	12.8	1.58~1.74	1.67
Stem	2.21~2.87	2.53	0.29~0.37	0.33	2.34~3.37	2.74	0.31~0.44	0.36
Leaf	3.83~4.60	4.22	0.50~0.60	0.55	4.78~5.59	5.33	0.62~0.73	0.70
Polished rice	2.40~2.92	2.63	0.31~0.38	0.34	2.40~2.75	2.58	0.31~0.36	0.34

**Figure 1** Site characteristics of TGM, Hg and Se in soil and rice leaves in artisanal mining sites and non-artisanal mining site (TGM: ng/m<sup>3</sup>, Hg and Se in soil and rice leaves: μg/g).

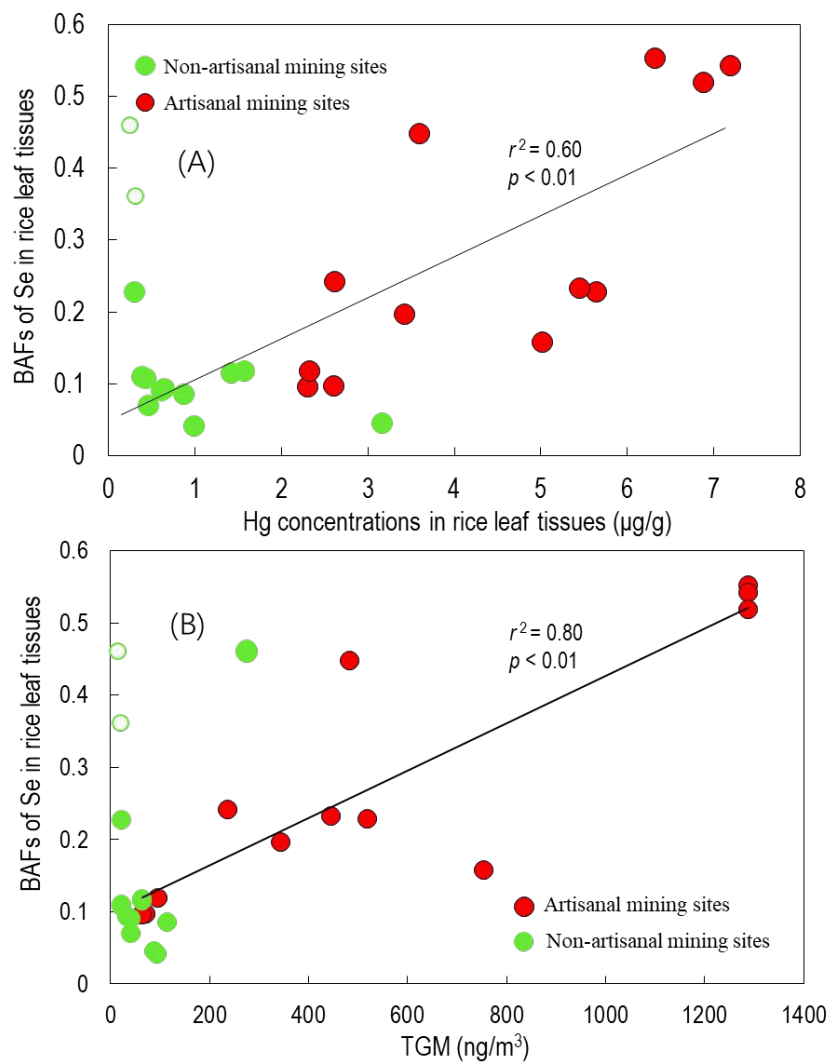


**Figure 2** Distribution of Hg (A) and Se (B) of pot rice plants placed in GY and WMM.

**Figure 3** (A) Correlations between soil TSe and BAFs of Hg in rice tissues collected from non-artisanal mining sites of the Wanshan Mercury Mine; (B) Correlations between soil TSe and BAFs of Hg in rice tissues collected from artisanal mining sites of the Wanshan Mercury Mine; (C) Correlations between soil THg and BAFs of Se in rice tissues collected from non-artisanal mining sites of the Wanshan Mercury Mine; (D) Correlations between soil THg and BAFs of Se in rice tissues collected from artisanal mining sites of the Wanshan Mercury Mine



**Figure 4** (A) Correlation between Hg concentrations and BAFs of Se in rice leaves collected from the Wanshan Mercury Mine; (B) Correlation between TGM and BAFs of Se in rice leaves collected from the Wanshan Mercury Mine.



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