

1 **The SLIM1 transcription factor regulates arsenic sensitivity in *Arabidopsis thaliana***

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31 Highlight: We identify a critical function of the SLIM1 transcription factor in regulating
32 arsenic transcriptional responses and propose that SLIM1 acts as both a transcriptional
33 activator and repressor.

34 **Abstract**

35 The transcriptional regulators of arsenic-induced gene expression remain largely
36 unknown; however, arsenic exposure rapidly depletes cellular glutathione levels
37 increasing demand for thiol compounds from the sulfur assimilation pathway. Thus,
38 sulfur assimilation is tightly linked with arsenic detoxification. To explore the
39 hypothesis that the key transcriptional regulator of sulfur assimilation, SLIM1, is
40 involved in arsenic-induced gene expression, we evaluated the response of *slim1*
41 mutants to arsenic treatments. We found that *slim1* mutants were sensitive to arsenic in
42 root growth assays. Furthermore, arsenic treatment caused high levels of oxidative stress
43 in the *slim1* mutants, and *slim1* mutants were impaired in both thiol and sulfate
44 accumulation. We also found enhanced arsenic accumulation in the roots of *slim1*
45 mutants. Furthermore, microarray analyses identified genes from a highly co-regulated
46 gene cluster (the O-acetylserine gene cluster), as being significantly upregulated in the
47 *slim1-1* mutant background in response to arsenic exposure. The present study identified
48 the SLIM1 transcription factor as an important component in arsenic-induced gene
49 expression and arsenic tolerance. Our results suggest that the severe arsenic sensitivity
50 of the *slim1* mutants is a result of both altered redox status as well as mis-regulation of
51 key genes.

52

53 **Key words**

54 arsenic, sulfur limitation, transcription factor, *Arabidopsis thaliana*

55

56 **Introduction**

57 Many advanced technologies used by our modern society rely on heavy metals and
58 metalloids. Some of these elements are toxic and pose a significant risk to the
59 environment and human health if consumed. However, unlike animals, plants are often
60 partially tolerant to the toxic effects of these elements and can accumulate large amounts
61 in diverse tissues (Clemens, 2006). Arsenic is a highly toxic substance commonly found
62 in anthropogenic waste (electronics and fertilizers), and natural deposits can also be
63 found at high levels in certain rocks, soils, and waters globally (Ogunseitan et al., 2009;
64 Satarug et al., 2009; Han et al., 2002; Larison et al., 2000). While this toxic metalloid
65 has no recognized role in plant or animal nutrition, plant-based products are the main
66 entry point for arsenic into the food chain (Mendoza-Cózatl et al., 2014). Thus,
67 understanding the molecular mechanisms underlying plant uptake, transport,
68 detoxification, and accumulation of arsenic is vital for enhancing the nutritional value
69 and safety of our food.

70

71 We previously described the development of a plant genetic reporter line that fused the
72 promoter of a cadmium and arsenic-inducible high-affinity sulfate transporter to firefly
73 luciferase (*pSULTRI;2::LUC*) to identify mutants in signaling (Jobe et al., 2012). A
74 major goal of this work was to identify the transcriptional regulators mediating rapid
75 arsenic-induced gene expression in Arabidopsis. This approach was successful in
76 identifying new alleles of the glutathione biosynthesis genes gamma-glutamylcysteine
77 synthetase (γ -ECS) and glutathione synthetase (GS), as being required for cadmium and
78 arsenic-induced gene expression (Jobe et al., 2012). Glutathione is required for the

79 synthesis of phytochelatins, which detoxify many toxic compounds, including cadmium
80 and arsenic, by chelation and sequestration in the vacuole (Clemens, 2006; Mendoza-
81 Cózatl et al., 2010b; Verbruggen et al., 2009; Rea, 2007; Mendoza-Cózatl and Moreno-
82 Sánchez, 2005; Song et al., 2010). Phytochelatins are short polymers of glutathione
83 synthesized in the cytosol in response to toxic metal(loid)s. Thus, arsenic exposure can
84 rapidly deplete glutathione levels, creating an extremely high demand for glutathione in
85 plant cells.

86

87 Because the tripeptide glutathione (Glu-Cys-Gly) contains the sulfur-containing amino
88 acid cysteine, the sulfate assimilation pathway is inextricably linked to glutathione
89 biosynthesis. Sulfate assimilation takes oxidized sulfur in the form of sulfate and,
90 through a series of energy-dependent reducing steps, produces sulfide. Due to the
91 toxicity of sulfide, this intermediate quickly reacts with O-acetylserine to produce the
92 amino acid cysteine (Jobe et al., 2019). Thus, unlike animals, plants do not require
93 exogenous sulfur-containing amino acids and proteins for survival (Jobe & Kopriva,
94 2018). More importantly, this creates a direct link between the sulfate assimilation
95 pathway and the ability of plants to detoxify arsenic.

96

97 While our luciferase genetic reporter approach has not identified transcriptional
98 regulators of arsenic-induced gene induction to date, a similar reporter gene approach
99 was successfully used to identify a transcriptional regulator of the sulfur deficiency
100 response in Arabidopsis. This screen used the same high-affinity sulfate transporter
101 promoter element fused to the green fluorescent protein (*pSULTRI;2::GFP*) and

102 identified four allelic mutants in an ethylene insensitive-like transcription factor called
103 Sulfur Limitation 1 (SLIM1) that failed to induce the reporter construct under sulfur
104 limiting conditions (Maruyama-Nakashita et al., 2006). All of the allelic *slim1* mutants
105 identified in this screen resulted in missense mutations altering single amino acid
106 residues (Maruyama-Nakashita et al., 2006). In *slim1-1* and *slim1-2*, high-affinity sulfate
107 uptake was decreased by ~60%, and sulfur-dependent microarray analyses on *slim1-1*
108 and *slim1-2* showed a decrease in the induction of many sulfur limitation-induced
109 transcripts compared to controls suggesting that SLIM1 is a positive regulator of sulfate
110 uptake and assimilation (Maruyama-Nakashita et al., 2006).

111

112 While the transcription factor(s) that control arsenic-induced gene expression remain
113 largely unknown, arsenic exposure is known to rapidly deplete cellular glutathione
114 levels, increasing the demand for reduced sulfur compounds from the sulfur assimilation
115 pathway (Jobe et al., 2012; Foyer and Noctor, 2011; Noctor et al., 2012). A similar
116 situation occurs under sulfur deficiency. As sulfate supply decreases, cellular levels of
117 cysteine and glutathione become depleted. Thus, because of the similarities in
118 glutathione depletion and subsequent upregulation of the high-affinity sulfate transporter
119 *SULTR1;2* under arsenic stress (Jobe et al., 2012) and sulfur limitation (Maruyama-
120 Nakashita et al., 2006), we investigated the hypothesis that SLIM1 plays a role in
121 arsenic-induced transcriptional responses. Interestingly, we found that *slim1-1* and
122 *slim1-2* seedlings were highly sensitive to arsenic. Here, we show that under arsenic
123 treatment, *slim1* mutants accumulate arsenic, experience high levels of oxidative stress,
124 and fail to induce sulfate uptake and assimilation. Microarray analyses of arsenic-treated

125 *slim1-1* seedlings identified a small number of transcripts differentially regulated by
126 arsenic in the mutant. Our results suggest that SLIM1 appears to play an important role
127 in arsenic sensitivity due primarily to its role in regulating sulfur metabolism and
128 cellular redox.

129

130 **Results**

131 ***slim1* mutants are sensitive to arsenic in root growth assays**

132 In a previous screen for regulators of cadmium and arsenic-induced gene expression
133 using a *pSULTR1;2::LUC* reporter construct, we identified new alleles in well-
134 characterized glutathione biosynthesis genes that play an important role in cadmium and
135 arsenic detoxification (Jobe et al., 2012). Because glutathione is a major sink of reduced
136 sulfur in plants, we hypothesized that the transcriptional regulator of sulfur deficiency,
137 SLIM1, might also play a role in the regulation of cadmium and arsenic sensitivity in
138 plants. To test this hypothesis, we performed root growth assays to evaluate the
139 sensitivity of the *slim1-1* and *slim1-2* mutant alleles (Maruyama-Nakashita et al., 2006)
140 to cadmium and arsenic (Figure 1A-1D).

141

142 The root lengths of wild-type (WT) (3.06 ± 0.09 cm, n=22), *slim1-1* (3.19 ± 0.07 cm,
143 n=19), and *slim1-2* (3.10 ± 0.11 cm, n=21) were not different in the control nutrient
144 media (see Methods) (Lee et al., 2003) without addition of cadmium or arsenic (Figure
145 1; $p=0.99997$ *slim1-1* & $p=1.0$ *slim1-2*, one-way ANOVA). When grown on plates
146 containing 30 μ M cadmium, WT root growth was inhibited growing only 1.88 ± 0.15
147 cm (n=10). This inhibition was similar to that observed for *slim1-1* with a final root

length of 1.86 ± 0.13 cm (n=12) and *slim1-2* having a root length of 1.82 ± 0.11 cm (p=1.0 for *slim1-1* & p=0.99998 for *slim1-2*, n=13) (Figure 1A-1D, Table S1). However, when grown on minimal media plates containing 10 μ M arsenite (As (III)), the root length of WT (1.72 ± 0.12 cm, n=14) was longer than both *slim1-1* (0.68 ± 0.06 cm, p= 7×10^{-9} , n=14) and *slim1-2* (0.75 ± 0.06 cm, p= 1.2×10^{-6} , n=10). These observations suggested that SLIM1 is involved in arsenic signaling. Thus, we further investigated possible mechanisms underlying *slim1* sensitivity to arsenic.

155

156 **Arsenic accumulation and antioxidant responses of *slim1* mutants**

To determine if arsenic accumulates in the *slim1* mutants, we measured root and shoot arsenic levels using ICP-MS. In As(III) treated seedlings, we observed no significant increase in the accumulation of arsenic in the shoots of *slim1-1* (174.8 ± 1.97 mg/Kg DW, n=3) or *slim1-2* (189.7 ± 3.78 mg/Kg DW, n=3) compared to WT (173.9 ± 4.32 mg/Kg DW, n=3, p=1) (Figure 2A). However, in As(V) treated seedlings, both *slim1-1* (309.0 ± 47.5 mg/Kg DW, n=3, p=0.01) and *slim1-2* (253.9 ± 19.9 mg/Kg DW, n=3, p=0.6) accumulated more arsenic in shoots than WT (205.6 ± 11.6 mg/Kg DW, n=3), although the difference was only significant in *slim1-1* (Figure 2A). These results suggest an increased root to shoot translocation of As(V) in the *slim1* mutants.

166

In the roots, we found arsenic accumulation in *slim1-1* (1420.0 ± 281.3 mg/Kg DW, n=3, p= 6.77×10^{-3}) and *slim1-2* (1473.0 ± 187.9 mg/Kg DW, n=3, p= 3.69×10^{-3}) compared to WT (420.1 ± 17.1 mg/Kg DW, n=3) in As(III) treated seedlings (Figure 2B, Table

170 S3). In comparison, there was no difference in root arsenic accumulation in As(V)
 171 treated seedlings (Figure 2B, Table S3).
 172

173 Because arsenic is known to cause oxidative stress and induce reactive oxygen species
 174 (ROS) production, we also tested the activity of the key antioxidant enzymes peroxidase
 175 (POD) and superoxide dismutase (SOD) in the *slim1-1* and *slim1-2* mutants. Basal
 176 superoxide dismutase activity in seedlings was similar between WT (70.9 ± 9.65 units/g
 177 FW, n=3), *slim1-1* (77.7 ± 9.44 units/g FW, p=1.0, n=3), and *slim1-2* (Figure 2C; $75.8 \pm$
 178 4.20 units/g FW, p=1.0, n=3, p=1.0 *slim1-1* and p=1.0 *slim1-2*) grown under control
 179 conditions. When exposed to arsenite (As(III)), WT superoxide dismutase increased to
 180 306.2 ± 7.78 units/g FW (n=3) while the superoxide dismutase activity in the *slim1*
 181 mutants increased dramatically to 709.5 ± 4.85 units/g FW (p=3.6x10⁻⁶, n=3) in *slim1-1*
 182 and 621.1 ± 17.7 units/g FW in *slim1-2* (p=2.0x10⁻⁸, n=3). Similarly, arsenate (As(V))
 183 treatment increased the WT superoxide dismutase activity to 234.8 ± 27.2 units/g FW
 184 while the *slim1-1* superoxide dismutase activity increased to 543.2 ± 39.4 units/g FW
 185 (p=2.9x10⁻⁸, n=3) and the *slim1-2* superoxide dismutase activity increased to $492.1 \pm$
 186 17.7 units/g FW (p=4.7x10⁻⁷, n=3) (Figure 2C, Table S4).
 187

188 In seedlings, the peroxidase activity was higher under control conditions in *slim1-1*
 189 (94.8 ± 5.95 units/g FW, p=0.009, n=3) and *slim1-2* (120.6 ± 6.04 units/g FW,
 190 p=5.4x10⁻⁵, n=3) compared to WT (50.7 ± 1.51 units/g FW, n=3) (Figure 2D). As (III)
 191 exposure increased the peroxidase activity in WT to 107.5 ± 8.91 units/g FW (n=3)
 192 (Figure 2D), while the peroxidase activity in *slim1-1* seedlings increased to 173.6 ± 1.79

193 units/g FW ($p=0.0001$, $n=3$) (Figure 2D). Similar values were observed for As (III)-
194 treated *slim1-2* seedlings (139.3 ± 4.49 units/g FW, $p=0.10$, $n=3$) (Figure 2D, Table S2).
195 Peroxidase activities showed similar trends under As (V) treatment (Figure 2D, Table
196 S5).

197

198 **Decreased shoot glutathione in arsenic-treated *slim1-1* and *slim1-2***

199 To determine if thiol production might also be altered by arsenic treatment in the *slim1*
200 mutants, we measured root and shoot cysteine and glutathione levels using fluorescence
201 HPLC of seedlings exposed to arsenite (As(III)) or arsenate (As(V)) for 48 hours
202 (Figure 3A-3D). Shoot cysteine levels were lower in *slim1-1* (10.2 ± 0.5 pmol/mg FW,
203 $p=0.04$, $n=3$) and *slim1-2* (12.2 ± 1.0 pmol/mg FW, $p=0.19$, $n=3$) than WT (23.7 ± 2.7
204 pmol/mg FW, $n=3$) in control conditions (Figure 3A, Table S6). No clear reduction in
205 cysteine levels were observed in response to As(III) or As(V) treatment (Figure 3A,
206 Table S6).

207

208 Root cysteine levels were statistically similar for WT, *slim1-1*, and *slim1-2* in control
209 conditions and were not significantly changed by As(III) or As(V) treatments (Figure
210 3B, Table S7; One-way ANOVA, Tukey HSD).

211

212 Under control conditions, shoot glutathione levels were lower in *slim1-1* (163.6 ± 24.2
213 pmol/mg FW, $n=8$) and *slim1-2* (190.2 ± 34.4 pmol/mg FW, $n=8$) than in WT ($382.4 \pm$
214 36.2 pmol/mg FW, $n=8$) (Figure 3C; $p=4.99 \times 10^{-5}$ for *slim1-1* and $p=3.9 \times 10^{-4}$ for *slim1-*
215 *2*). Shoot glutathione levels decreased in WT from 382.4 ± 36.2 pmol/mg FW ($n=8$) in

216 control conditions to 278.1 ± 23.2 pmol/mg FW (n=3) in the As(III) treatment and 269.4
217 ± 12.2 pmol/mg FW (n=3) in the As(V) treatment (Figure 3C). Similarly, shoot
218 glutathione decreased in the *slim1* mutants under As(III) and As(V) treatments with
219 *slim1-1* having only 110.4 ± 17.5 pmol/mg FW of glutathione in As(III) and 31.4 ± 1.62
220 pmol/mg FW of glutathione in As(V). Furthermore, *slim1-2* had 83.8 ± 30.9 pmol/mg
221 FW (n=3) shoot glutathione in As(III) treatment and 43.5 ± 14.7 pmol/mg FW (n=3) in
222 As(V) treatment – an 80% decrease compared to control (Figure 3C, Table S8).

223

224 Root glutathione levels decreased under both As(III) and As(V) treatments for all
225 genotypes. However, glutathione levels in roots showed no differences between
226 genotypes within each treatment (Figure 3D, Table S9; One-way ANOVA, Tukey
227 HSD). In summary, thiol measurements showed that while cysteine and glutathione
228 levels were not dramatically decreased in roots of the *slim1* mutant alleles compared to
229 WT (Figure 3B and D), glutathione levels were clearly decreased in shoots of *slim1-1*
230 and *slim1-2* compared to WT plants (Figure 3C).

231

232 **Shoot sulfate and phosphate accumulation in *slim1* mutants**

233 Arsenic is thought to be actively taken up by phosphate transporters as As(V); however,
234 once inside plant cells, it is reduced to As(III) and can move within plants through
235 aquaporins (Catarcha et al., 2007; Bienert et al., 2008). Mutants in SLIM1 were
236 previously shown to be impaired in root-to-shoot translocation of sulfate (Maruyama-
237 Nakashita et al., 2006), but the translocation of other anions, including phosphate, was
238 not reported. Thus, based on the slight arsenic accumulation in shoots of As(V) treated

239 plants noted by ICP-MS (Figure 2 A and B), we hypothesized that phosphate transport
240 might also be impaired in the *slim1* mutants.

241

242 To determine if phosphate and sulfate translocation are impaired in the *slim1* mutants
243 under arsenic treatment, we measured sulfate and phosphate accumulation in both roots
244 and shoots of plants treated with As (V) for 48 hours. Interestingly, shoot phosphate
245 accumulation was higher in *slim1-1* and *slim1-2* than WT in all treatments (Figure 4A,
246 Table S10; $p=5 \times 10^{-6}$ for *slim1-1* and $p=0.004$ for *slim1-2*; One-way ANOVA, Tukey
247 HSD).

248

249 Root phosphate accumulation was similar for WT (5.03 ± 0.27 nmol/mg FW, $n=5$),
250 *slim1-1* (4.95 ± 1.07 nmol/mg FW, $n=3$), and *slim1-2* (6.33 ± 0.80 nmol/mg FW, $n=4$) in
251 control conditions and was not different under As(V) treatment (Figure 4B). Thus, the
252 enhanced root-vs.-shoot phosphate accumulation observed in *slim1-1* and *slim1-2*
253 suggests an indirect role for SLIM1 in the regulation of phosphate and arsenate transport
254 (Figure 4A, 4B, Table S11).

255

256 Furthermore, sulfate accumulation in shoots was impaired in *slim1-1* (0.73 ± 0.10 nmol/
257 mg FW, $p=1.7 \times 10^{-6}$, $n=5$) and *slim1-2* (0.61 ± 0.20 nmol/mg FW, $p=7.3 \times 10^{-7}$, $n=5$)
258 relative to WT (3.09 ± 0.18 nmol/mg FW, $n=5$) in control conditions (Figure 4A),
259 consistent with previous findings (Maruyama-Nakashita et al., 2006). WT seedlings
260 showed a decrease in shoot sulfate upon As(V) treatment decreasing to 1.86 ± 0.44
261 nmol/mg FW ($n=5$) (Figure 4C, Table S12, $p=0.008$, One-way ANOVA, Tukey HSD).

262

263 Root sulfate accumulation was similar between WT (2.11 ± 0.14 nmol/mg FW, n=5),
264 *slim1-1* (1.95 ± 0.43 nmol/mg FW, n=3), and *slim1-2* (1.45 ± 0.21 nmol/mg FW, n=4) in
265 control conditions. Furthermore, WT (2.04 ± 0.16 nmol/mg FW, n=4), *slim1-1* ($1.72 \pm$
266 0.13 nmol/mg FW, n=5), and *slim1-2* (1.85 ± 0.15 nmol/mg FW, n=3) root sulfate were
267 not different in the As(V) treatment (Figure 4D, Table S13).

268

269 **Microarray analyses of *slim1* mutants under As treatment**

270 The current model for arsenic uptake and tolerance in plants suggests that arsenic is
271 taken up from the soil in the form of arsenate (As(V)). Once it has entered the plant, it is
272 rapidly reduced to arsenite (As(III)) by the arsenate reductase HAC1(Shi et al., 2016). It
273 has been proposed that As(III) can be removed from the root by an unidentified efflux
274 transporter(Chen et al., 2017). In rice, the aquaporin LSI1 is known to mediate As(III)
275 efflux; however, additional efflux transporters remain elusive(Chen et al., 2017). A
276 recent RNA-seq experiment using a T-DNA mutant allele of SLIM1 (*eil3*) did not find
277 misregulation of any aquaporin genes in the roots of the *slim1* mutant under control or
278 sulfur deficiency conditions(Dietzen et al., 2020). Thus, due to the observed arsenic
279 accumulation in the roots of *slim1* mutants, we hypothesized that the elusive As(III)
280 efflux transporter, or alternatively an As(III) uptake transporter, might be disrupted in an
281 arsenic-dependent manner in the *slim1* mutant background.

282

283 To test this hypothesis and uncover genes disrupted in an arsenic-dependent manner in
 284 the *slim1-1* mutant, we performed microarray analyses on WT and *slim1-1* seedlings
 285 exposed to arsenic for 48 hours. Raw expression values were normalized via the R ‘affy’
 286 package using the Robust Multi-Array Average (RMA) Expression Measure.
 287 Differential gene expression was evaluated using the R package ‘limma’, including a
 288 multiple test correction. We then performed a significance analysis to identify genes
 289 disrupted under arsenic treatment and compared these to previously published putative
 290 targets of SLIM1 obtained by DNA affinity purification sequencing (DAP-seq)
 291 (O’Malley et al., 2016).
 292
 293 From the microarray analyses, we identified 11 genes significantly differentially
 294 upregulated by arsenic (WT +As vs. *slim1-1* + As) (Supplemental Table S14). Ten of
 295 the 11 genes (AT3G49580, AT1G04770, AT1G12030, AT4G04610, AT4G21990,
 296 AT5G24660, AT5G26220, AT5G48850, AT4G20820, AT1G36370) were identified as
 297 putative targets of SLIM1 by DNA affinity purification sequencing (DAP-Seq)
 298 (Supplemental Table S14). Many of the 11 upregulated genes in *slim1-1* are associated
 299 with sulfur metabolism. For example, GAMMA-GLUTAMYL
 300 CYCLOTRANSFERASE 2;1 (GGCT2;1, AT5G26220), an enzyme involved in the
 301 break down of glutathione, was previously reported to be induced under sulfur
 302 deficiency conditions in a SLIM1-dependent manner. However, under arsenic treatment
 303 (As vs. *slim1-1* + As), this gene is expressed at significantly higher levels in the *slim1-1*
 304 mutant compared to WT, suggesting SLIM1 might be a negative regulator of GGCT2;1
 305 under arsenic treatment. Other genes that appear to be negatively regulated by SLIM1

306 include APR1 (AT4G04610) and APR3 (AT4G21990), which were upregulated in
307 *slim1-1* compared to WT in the presence of arsenic (WT +As vs. *slim1-1* + As). APR1
308 and APR3 are involved in the reduction of sulfate into sulfide (Lee et al., 2011) and
309 have been shown to be induced by toxic metal stress (Jobe et al., 2012). Similarly, the
310 LOW SULFUR 1 (LSU1, AT3G49580) and LOW SULFUR 2 (LSU2, AT5G24660)
311 genes were expressed at higher levels in *slim1-1* than WT under arsenic treatment (WT
312 +As vs. *slim1-1* + As). Interestingly, Six of the 11 genes (GGCT2;1, APR3, LSU1,
313 LSU2, SDI1, & SHM7) belong to a highly co-regulated cluster of genes that respond to
314 O-acetylserine treatment (Hubberten et al., 2012).

315

316 Microarray analyses also identified 10 significantly down-regulated genes under arsenic
317 treatment compared to WT (WT +As vs. *slim1-1* +As) (Figure 5) ($p < 0.05$, Fold Change
318 > 2). Only one gene - SULTR1; 2 (At1G78000) - was identified as a putative target of
319 SLIM1 by DAP-Seq (Supplemental Table S14). Thus, our analyses confirm the reported
320 function of SLIM1 as a transcriptional activator of *SULTR1;2* and show that this role is
321 conserved under arsenic treatment and sulfur deficiency. The remaining ten genes are
322 involved in hormone signaling (AT1G63030, AT5G13220 & AT5G52050), redox
323 regulation (AT3G06590 & AT1G03020), iron homeostasis (AT3G25190 &
324 AT5G01600), glucosinolate biosynthesis (AT5G23020), ubiquitination (AT1G24330),
325 and uncharacterized protein (AT2G17660). Based on their putative functions, these
326 general stress response genes are responding in a SLIM1 independent manner, or they
327 are previously unidentified conditional targets of SLIM1 regulation. More experiments

328 are needed to determine if SLIM1 is a direct transcriptional regulator of these genes
329 under arsenic stress.

330

331 The present transcriptome data suggest that SLIM1 can function as both a transcriptional
332 enhancer as well as a transcriptional repressor of certain genes in a condition-specific
333 manner. Furthermore, the present study provides evidence that SLIM1 plays an essential
334 role in the regulation of gene expression in response to arsenic.

335

336 **Discussion**

337 Arsenic exposure to plants causes rapid changes in gene expression (Jobe et al., 2012,
338 Fu et al., 2014, Castrillo et al., 2013). However, the transcription factors that function in
339 arsenic-induced gene expression remain largely unknown. The few transcriptional
340 regulators that have been identified, such as WRKY6, WRKY45, and OsARM1
341 (Arsenite-Responsive Myb1) (Castrillo et al., 2013; Wang et al., 2014, 2017), have been
342 implicated in the regulation of arsenic transporters while regulators of arsenic
343 detoxification remain unknown. To test the hypothesis that the SLIM1 transcription
344 factor is involved in arsenic resistance and signaling, we evaluated the sensitivity of
345 *slim1-1* and *slim1-2* to arsenic exposure. We found the *slim1* mutants were more
346 sensitive to arsenic than control plants. Arsenic treatment caused high levels of
347 oxidative stress in the *slim1* mutant alleles based on superoxide dismutase and
348 peroxidase activities. Furthermore, thiol and sulfate measurements show that *slim1*
349 mutants are impaired in both thiol and sulfate accumulation. Arsenic treatment did not
350 further decrease sulfate levels in roots. In contrast, the concentration of the thiol GSH

351 was greatly decreased in *slim1* mutant alleles. Furthermore, the peroxidase and
352 superoxide dismutase measurements show that arsenic treatments cause high levels of
353 oxidative stress in the *slim1* mutants.

354

355 We also observed a slight increase in arsenic accumulation in the shoots of *slim1*
356 mutants treated with arsenate (As(V)). This corresponded with a significant increase in
357 shoot phosphate translocation in the *slim1* mutants. Because of the chemical similarity
358 between phosphate and As(V), future research could investigate the hypothesis that the
359 misregulation of phosphate transporters may contribute to this observed increase in
360 shoot arsenic in the *slim1* mutants under As(V) treatment. We also found significant
361 arsenic accumulation in the roots of *slim1* mutants treated with As(III). This
362 accumulation could be caused by the disruption of an As(III) uptake transporter or an
363 As(III) efflux transporter.

364

365 Thiol measurements confirmed the role of SLIM1 in sulfate metabolism and thiol
366 production (Dietzen et al., 2020; Maruyama-Nakashita et al., 2006; Yamaguchi et al.,
367 2020), as *slim1* mutants contained lower cysteine and glutathione levels in shoots than
368 WT. As described previously, glutathione is essential for the production of
369 phytochelatins –arsenic chelating compounds necessary for detoxification and storage.
370 The heavy metal cadmium also binds to phytochelatins. Interestingly, recent research
371 has shown a less dramatic effect of cadmium exposure in *slim1* mutants compared to
372 wild-type controls (Yamaguchi et al., 2020), which we have also observed (Figure
373 1C,D). Thus, the present study and a recent study together show that the SLIM1

transcription factor plays a more central role in mediating arsenic resistance relative to cadmium resistance. A possible hypothesis that may contribute to this observation is that cadmium can be sequestered in vacuoles via two independent transport pathways, via phytochelatin transport(Song et al., 2010; Mendoza-Cózatl et al., 2010a) and via HMA3-mediated cadmium transport(Takahashi et al., 2012).

Sulfate measurements confirmed that SLIM1 is a major transcriptional regulator of sulfate uptake and translocation(Maruyama-Nakashita et al., 2006). Our microarray analyses also identified 11 genes significantly differentially upregulated by arsenic (Supplemental Table S14), of which ten of the 11 genes were identified as putative targets of SLIM1 by DNA affinity purification sequencing (DAP-Seq). Interestingly, nine of these genes are involved in sulfur assimilation or redox signaling. Furthermore, six of these sulfur metabolism genes belong to a highly co-regulated cluster of genes that respond to O-acetylserine treatment(Hubberten et al., 2012). While previous studies show these genes can regulate sulfur assimilation in a SLIM1 independent manner (Aarabi et al., 2020; Hubberten et al., 2012), results from DAP-Seq and the microarray results from the current study suggest SLIM1 might also act as a negative regulator of these genes during arsenic stress. The disruption of these genes in the *slim1* mutants may also contribute to their increased sensitivity to arsenic.

Shoot sulfate accumulation was significantly lower in the *slim1* mutants in all conditions tested. Decreased shoot sulfate was accompanied by an increase in shoot phosphate in the *slim1* mutants. Similar anion compensation has been noted in the Arabidopsis *phr1*

397 mutant, which accumulates higher sulfate levels when grown under low phosphate
398 conditions indicating crosstalk between phosphate and sulfate transport (Rouached et al.,
399 2011). In fact, *PHR1* has been proposed to act both positively in the regulation of root-
400 to-shoot sulfate translocation via the sulfate transporter *SULTR1;3*, and negatively to
401 repress other sulfate transporters under phosphate deficiency (Rouached, 2011). Our
402 observation that phosphate uptake and translocation are enhanced in the *slim1* mutants is
403 consistent with the model that arsenic accumulation is a result of non-specific uptake of
404 arsenic by phosphate transporters.

405

406 In summary, we show here that the SLIM1 transcription factor plays an important role
407 in mediating arsenic resistance and in arsenic-induced gene expression. Our results
408 suggest that the arsenic sensitivity of *slim1* mutants can be explained by decreased thiol
409 production resulting in increased oxidative stress, enhanced arsenic uptake resulting in
410 increased arsenic accumulation. Interestingly, we found that the *slim1* mutant alleles do
411 not show a strong cadmium sensitivity, consistent with a recent study (Yamaguchi et al.,
412 2020) indicating a difference in the rate-limiting functions of the thiol synthesis pathway
413 in processing arsenic and cadmium. We also identify a number of genes regulated by
414 SLIM1 in an arsenic-dependent manner. Taken together, our data support a model in
415 which SLIM1 is both a positive and negative regulator of gene expression in response to
416 arsenic.

417

418 **Experimental Procedures**

419 *Arabidopsis* accessions

420 The WT *Arabidopsis thaliana* ecotype used in this study is Columbia (Col-0).
421 The *slim1-1* and *slim1-2* mutants were generated in the Col-0 genetic background and
422 were kindly provided by Dr. Akiko Maruyama (Maruyama-Nakashita et al., 2006).

423

424 ***Plant Growth Media & Conditions***

425 Seeds were surface sterilized by briefly soaking in 70% ethanol before allowing them to
426 dry in a sterile hood. For root growth experiments and enzymatic assay experiments,
427 surface sterilized seeds were plated on minimal media containing 1/10-strength
428 Hoagland solution, 1% phytoagar (Duchefa, <http://www.duchefa.com>), pH 5.6. For the
429 microarray experiments, seeds were plated on 1/2-strength MS standard medium
430 (M5519; Sigma-Aldrich, <http://www.sigmaaldrich.com>) buffered with 1 mM 2-(*N*-
431 morpholine)-ethanesulphonic acid (MES), 1% phytoagar
432 (Duchefa, <http://www.duchefa.com>) and the pH was adjusted to 5.6 with 1.0 M KOH.
433 Seeds were then stratified with cold treatment at 4°C for 48 h, and grown under
434 controlled conditions (150 $\mu\text{mol m}^{-2} \text{s}^{-1}$, 70% humidity, 16-h light at 21°C/8-h dark at
435 18°C) for the specified time. For toxic metal(loid) treatments, the specified amounts of
436 either cadmium or arsenic were added to the autoclaved base media in a sterile hood
437 prior to pouring the plates. Concentrated stock solutions of cadmium and arsenic were
438 filter-sterilized prior to use.

439

440 ***Statistical Analyses***

441 The root growth, thiol, peroxidase, superoxide, and anion data were all analyzed using
442 one-way ANOVA followed by a Tukey post-hoc test to determine significance.

443 Significance groups are indicated in the figures and key p-values are stated in the text.

444

445

446 ***Root Length Measurements***

447 For root growth experiments, surface sterilized seeds of WT, *slim1-1* and *slim1-2* were
448 plated on minimal media (2.5 mM H₃PO₄, 5 mM KNO₃, 2 mM MgSO₄, 1 mM (CaNO₃)₂,
449 1 mM MES, 1% phytoagar pH 5.7) supplemented with 30 µM Cd or 10 µM As (III)
450 (Lee et al., 2003). Plates were placed in the dark two days at 4°C for vernalization and
451 then transferred to a growth chamber. After 7 days of growth, seedlings were
452 photographed and root length was measured using ImageJ.

453

454 ***Antioxidant Enzyme Assays***

455 Seedling samples were weighed and pulverized in liquid nitrogen after treatment. The
456 powder was dissolved in pre-cooled 50 mM phosphate buffer (pH 7.8) to extract the
457 superoxide dismutase (SOD). The extract was then centrifuged at 12 000g for 10 min,
458 resulting in a crude enzyme supernatant solution. In a separate 10 ml tube, 1.9 ml
459 reaction buffer (50 mM phosphate buffer, pH 7.8, 9.9 mM L-methionine, 57 µM NBT
460 solution, 1 M EDTA-Na₂ solution, 0.0044% (w/v) riboflavin) and 0.1 ml enzyme
461 solution were mixed and placed into 250 µmol m⁻²s⁻¹ light for 20 min. Additionally,
462 another separate 10 ml tube was procured, where the enzyme solution was replaced with
463 water as a control. The reagent was added according to the above steps, where one tube

464 was placed in the light together with the sample, and the other was placed in the dark
465 where the reaction was allowed to complete. The control tube that was placed in the
466 dark was blanked, and the absorbance of each tube was measured at 560 nm. Peroxidase
467 (POD) was extracted in 50 mM phosphate buffer (pH 7.0). 30 µl of enzyme solution was
468 mixed with reaction buffer containing 1.77 mL of 50 mM sodium phosphate buffer (pH
469 7.0), 0.1 mL of 4% guaiacol and 0.1 mL of 1% (v/v) H₂O₂. Increased absorbance was
470 recorded at 470 nm for 1 min. All reported enzyme activities are means of 3-5
471 biologically independent samples, and error bars indicate the standard error of the mean
472 (SEM).

473

474 ***Arsenic Determination by ICP-MS***

475 Plant material was harvested, dried at 70°C for at least 48 hours before being aliquoted
476 and weighed. Approximately 10 mg of dried plant material was mixed with 1 ml of
477 concentrated nitric acid and digested by heating at 100°C for approximately 30 minutes
478 or until the solution became transparent and particle-free. These digests were diluted
479 with deionized water and measured by ICP-MS for total arsenic concentrations at the
480 University of Cologne Biocenter Mass Spectrometry Platform. All reported ion
481 quantities are means of 3-5 biologically independent samples, and error bars indicate the
482 standard error of the mean (SEM).

483

484 ***Anion Extraction and Measurement by Ion Chromatography***

485 To quantify the water soluble anion concentrations (phosphate, and sulfate) in plant
486 tissues, 10-30 mg of fresh tissue was harvested and flash frozen in liquid nitrogen.

487 Frozen tissue was then pulverized using a bead mill (make & model) and anions were
488 extracted by addition of 1000 μ L of sterile milliQ-water and incubating for 60 minutes
489 at 4°C while shaking at 1500 rpm. The extraction process was stopped by incubating at
490 95°C for 15 minutes. Cell debris was removed by centrifugation at 4°C for 15 minutes
491 and 100-200 μ L of supernatant was used for anion exchange chromatography. An
492 automatic ion analyzer (DX 120, Dionex Corporation, Sunnyvale, CA, United States)
493 equipped with an IonPac™ column (AS9-SC, 4 \times 250 mm; Dionex, Thermo Fisher
494 Scientific GmbH; Waltham, MA, United States) was used to separate and quantify the
495 anions. Anions were eluted with an elution buffer of 2.0 mM Na₂CO₃ and 0.75 mM
496 NaHCO₃. Ion concentrations were detected using a conductivity detector module (CDM,
497 Dionex Corporation, CA, United States). All reported anion quantities are means of 3-5
498 biologically independent samples, and error bars indicate the standard error of the mean
499 (SEM).

500

501 ***Thiol Detection By Fluorescence HPLC***

502 The thiol-containing compounds cysteine and GSH were analyzed using fluorescence
503 detection HPLC as described by (Fahey and Newton, 1987). To analyze the levels of
504 these thiol compounds, plants were grown on minimal growth media plates for 12 days
505 then transferred to fresh media plates containing either 20 μ M cadmium, 100 μ M
506 arsenate, or control minimal media. To minimize the oxidation of thiol compounds
507 during the extraction, plant seedlings were flash-frozen in liquid nitrogen immediately
508 after harvesting, and then pulverized using a bead mill and extracted as described by
509 (Krueger et al., 2009). Thiols were extracted from homogenized plant material with 1

510 mL 0.1 M HCl for 40 min at 25°C. After centrifugation for 5 min at 14,000 g and 4°C,
511 thiols in the supernatant were reduced by mixing 60 µL of the supernatant with 100 µL
512 2-(cyclohexylamino)ethanesulfonic acid (0.25 M, pH 9.4) and 35 µL DTT (10 mM,
513 freshly prepared). The mixture was incubated at 25°C for 40 min. Thiols were
514 derivatized by adding 5 µL (25 mM) monobromobimane (SigmaAldrich, Cat#B4380).
515 Derivatization was stopped by adding 110 µL methane sulfonic acid (100 mM) and
516 clarified by centrifugation for 15 min at 14,000 g and 4°C. Forty microliters of the
517 derivatization mix were used for HPLC analysis using the Dionex Ultimate 3000 HPLC
518 System. Derivatized thiols were separated in a Eurosphere 100-3 C18, 150×4 mm
519 column (Knauer), and were detected by fluorescence detection with an excitation of 380
520 nm and emission detection at 480 nm. The peaks of thiol compounds were identified and
521 quantified by comparison with cysteine and glutathione standards purchased from
522 Sigma-Aldrich. All reported thiol quantities are means of 4-5 biologically independent
523 samples, and error bars indicate the standard error of the mean (SEM).

524

525 ***Microarray Analyses***

526 To evaluate transcriptional differences in the *slim1* mutants under cadmium and arsenic
527 stress we performed microarray analyses. To obtain tissue for the microarray analysis,
528 plants were grown on ¼ MS plates for 12 days then transferred to fresh media plates
529 containing either 100 µM cadmium or 20 µM arsenite. Whole seedlings were then
530 harvested in 2mL Eppendorf tubes, flash-frozen in liquid nitrogen, and stored at -80°C
531 until further processing. The tissue was subsequently pulverized using a bead mill by
532 adding three 2.5mm glass beads to each tube and grinding for 15 seconds. RNA was

533 extracted using the Qiagen RNEasy mini kit (Cat#74104) per the manufacturer's
534 instructions (www.qiagen.com). RNA quality was assessed by spectrophotometer and
535 gel electrophoresis before submission to the University of California, San Diego Gene
536 Expression Core facility for processing. Results were analyzed using R and the
537 Bioconductor suite of microarray analytical packages as indicated in the text.

538

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542 Institutes of Health under Award Number P42ES010337 (JIS). The microarray data are
543 available through the NCBI GEO database (series record GSE138943).

544

545

546

547 **Conflicts of Interests**

548 The authors have no conflicts of interest to declare. All co-authors have seen and agree
549 with the contents of the manuscript and there is no financial interest to report. We certify
550 that the submission is original work and is not under review at any other publication.

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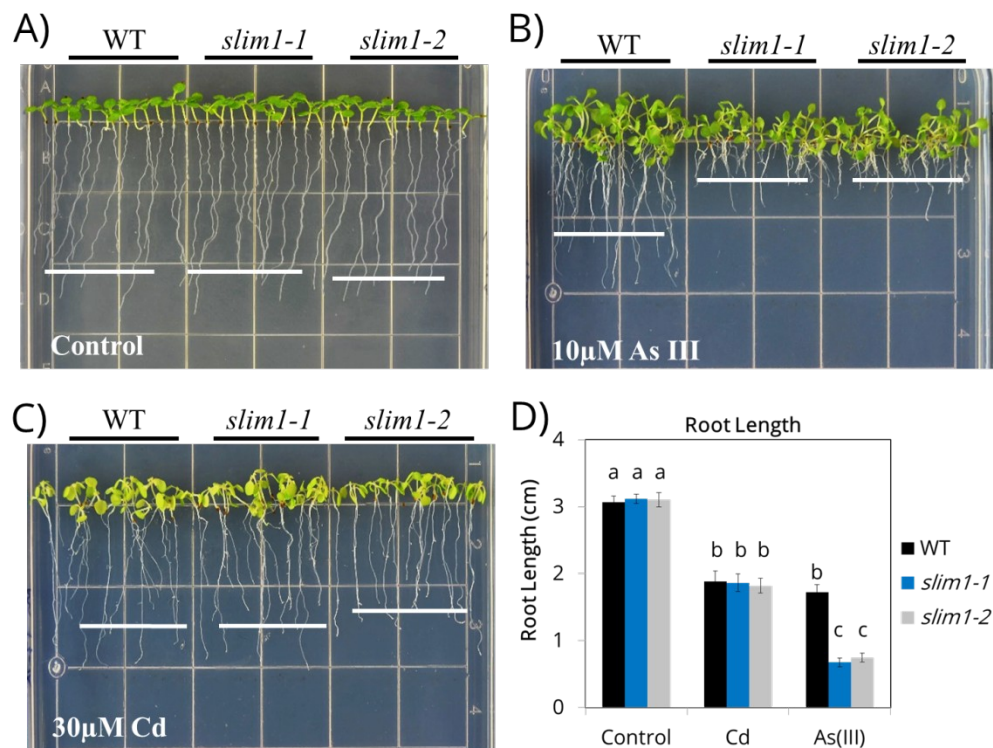
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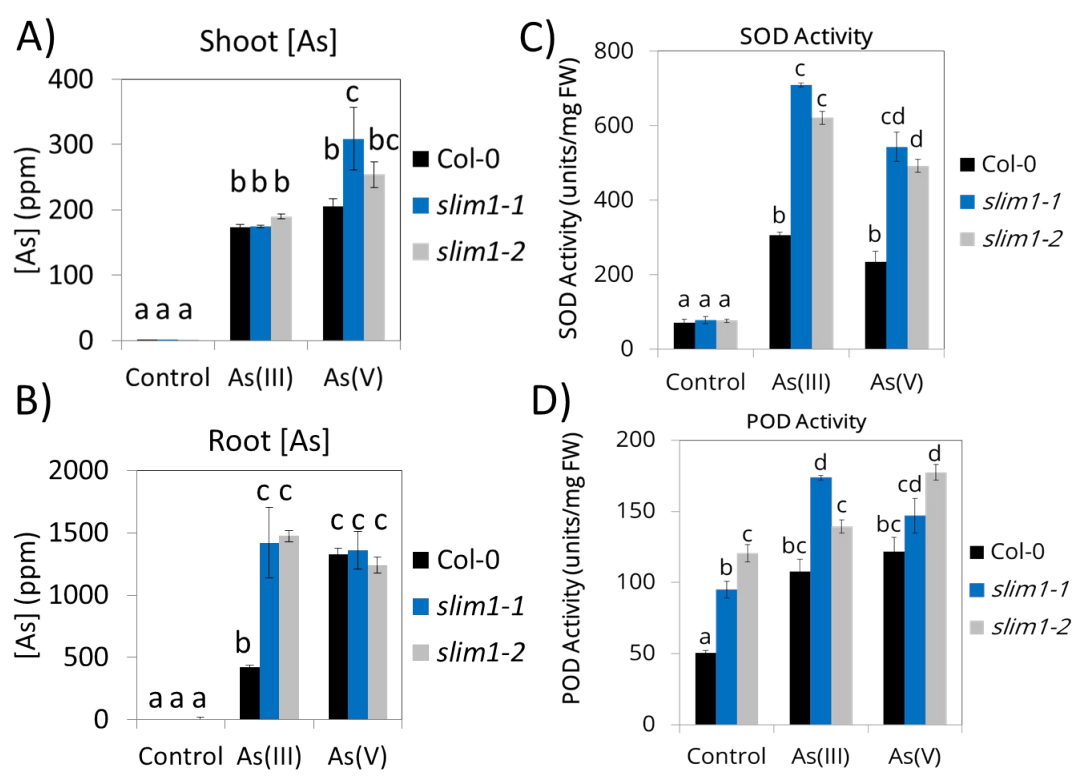
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674 **Figures & Figure Legends**

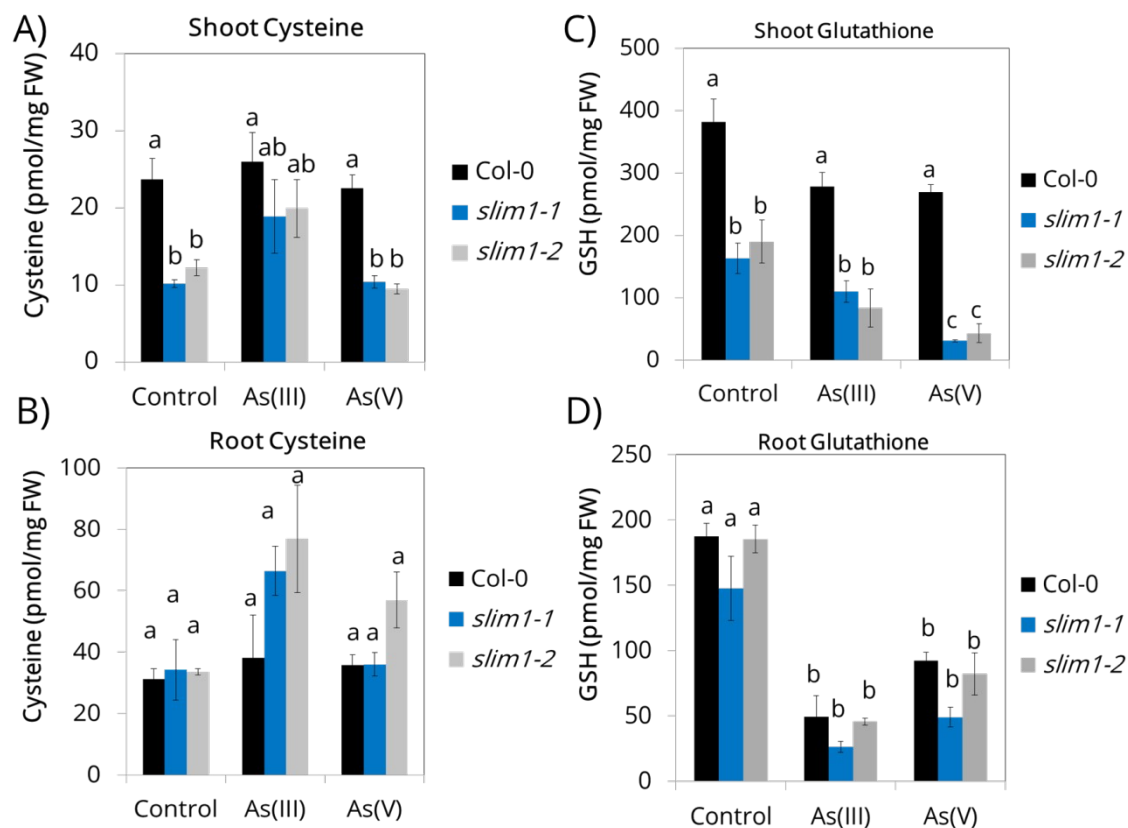


676 **Figure 1. Root growth inhibition of *slim1* mutants grown on cadmium or arsenic**
 677 **containing media.** The *slim1-1* and *slim1-2* mutant alleles were compared to wildtype
 678 controls (WT) grown on control minimal media and media containing 30 μM Cd or 10

679 μM As(III) for 7 days (A - D). Root growth was quantified using ImageJ (one-way
 680 ANOVA, Tukey HSD).

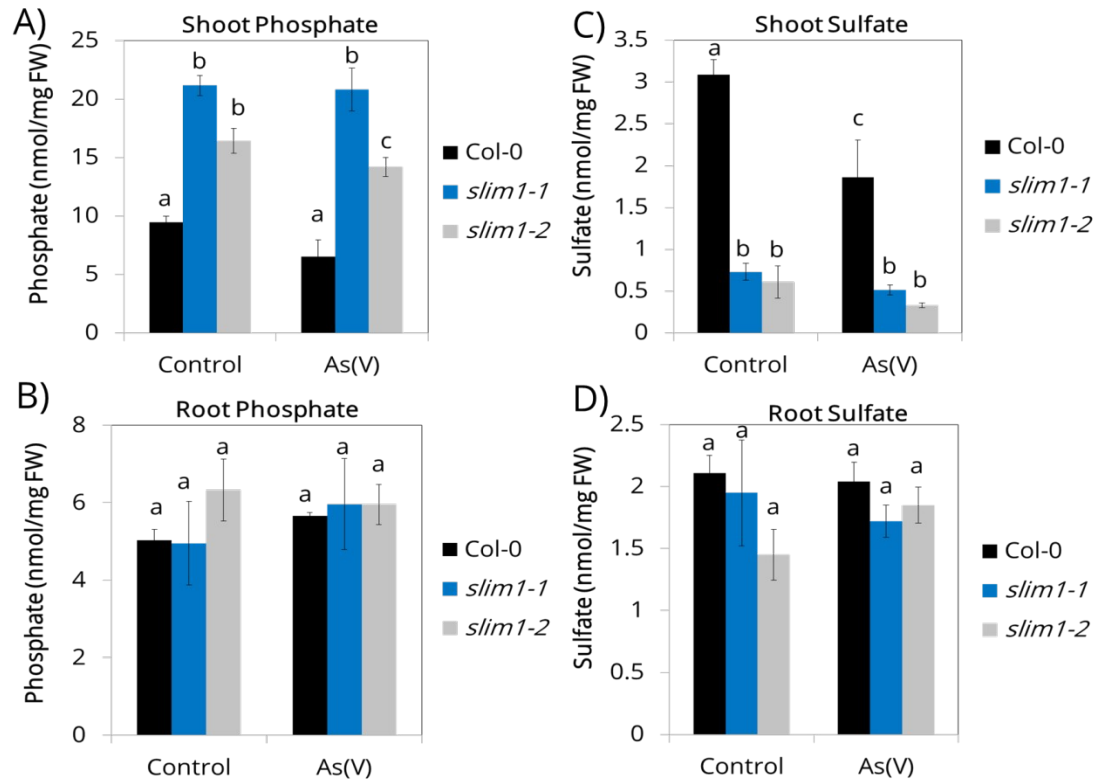


681
 682 **Figure 2. *slim1* mutants accumulate arsenic in roots and have high antioxidant**
 683 **activity when exposed to arsenic.** *slim1* mutants grown on arsenic-containing media
 684 accumulate arsenic in the shoots when grown on As(V) (A) but accumulate arsenic in
 685 the roots when grown on As(III) (B). Growth on arsenic-containing media caused an
 686 increase in superoxide dismutase (C) enzyme and peroxidase dismutase enzyme (D)
 687 activities in both the *slim1-1* and *slim1-2* mutants compared to WT controls.



688

689 **Figure 3. Thiol accumulation of *slim1* mutants grown on arsenic.** Total shoot
 690 cysteine levels in *slim1-1* and *slim1-2* compared to WT (A). Total root cysteine levels in
 691 *slim1-1* and *slim1-2* compared to WT (B). Total shoot glutathione levels for *slim1-1* and
 692 *slim1-2* compared to WT (C). Total root glutathione levels for WT, *slim1-1*, and *slim1-2*
 693 (D).



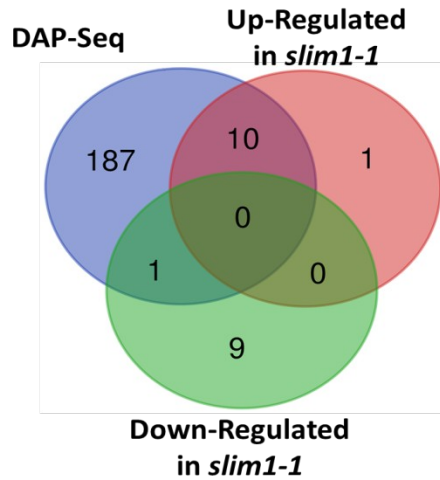
694

695 **Figure 4. Anion accumulation in *slim1* mutants grown on arsenic.** Total shoot
 696 phosphate levels in *slim1-1* and *slim1-2* compared to WT (A). Total root phosphate
 697 levels in *slim1-1* and *slim1-2* compared to WT (B). Total shoot sulfate levels for *slim1-1*
 698 and *slim1-2* compared to WT (C). Total root sulfate levels for WT, *slim1-1*, and *slim1-2*
 699 (D).

700

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702



703

704 **Figure 5. Number of genes regulated by arsenic treatment compared with SLIM1**
 705 **transcription factor DAP-Seq target genes.** Venn diagram showing the overlap of
 706 significantly upregulated genes in WT vs. *slim1-1* (pink) and down-regulated genes in
 707 WT vs. *slim1-1* (green) compared with previously published DAP-Seq targets of SLIM1
 708 (blue). Gene lists can be found in Supplementary Table S14.