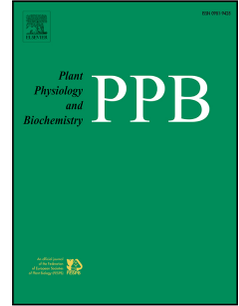


# Accepted Manuscript

Impact of double inoculation with *Bradyrhizobium japonicum* E109 and *Azospirillum brasilense* Az39 on soybean plants grown under arsenic stress

Ana L. Armendariz, Melina A. Talano, María Florencia Olmos Nicotra, Leticia Escudero, María Laura Breser, Carina Porporatto, Elizabeth Agostini



PII: S0981-9428(19)30067-1

DOI: <https://doi.org/10.1016/j.plaphy.2019.02.018>

Reference: PLAPHY 5599

To appear in: *Plant Physiology and Biochemistry*

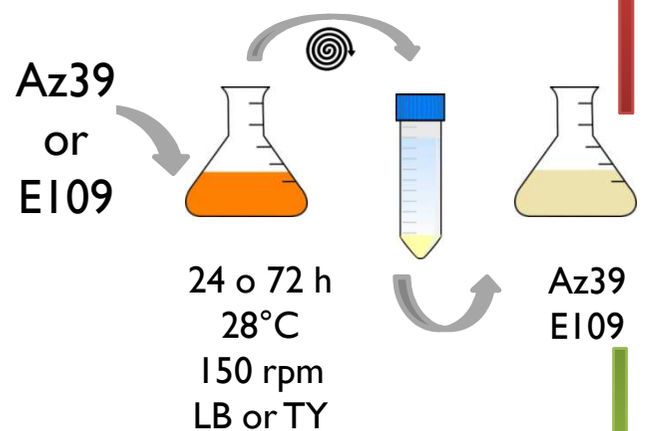
Received Date: 4 January 2019

Revised Date: 14 February 2019

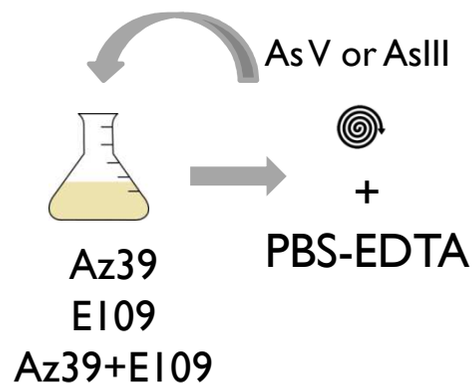
Accepted Date: 19 February 2019

Please cite this article as: A.L. Armendariz, M.A. Talano, Mari.Florencia. Olmos Nicotra, L. Escudero, Mari.Laura. Breser, C. Porporatto, E. Agostini, Impact of double inoculation with *Bradyrhizobium japonicum* E109 and *Azospirillum brasilense* Az39 on soybean plants grown under arsenic stress, *Plant Physiology et Biochemistry* (2019), doi: <https://doi.org/10.1016/j.plaphy.2019.02.018>.

This is a PDF file of an unedited manuscript that has been accepted for publication. As a service to our customers we are providing this early version of the manuscript. The manuscript will undergo copyediting, typesetting, and review of the resulting proof before it is published in its final form. Please note that during the production process errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.



### Flow cytometry analysis



Flow cytometer

**Mortality**

### In vivo analysis



Control plants

Inoculated plants

E109

Az39

E109+Az39

-Germination



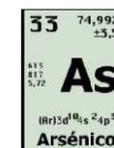
-Plant growth

-Nodule number



-Nitrogen content

-As accumulation



1 *Impact of double inoculation with Bradyrhizobium japonicum E109 and Azospirillum brasilense*  
2 *Az39 on soybean plants grown under arsenic stress*

3

4 Ana L. Armendariz<sup>(1)(+)</sup>, Melina A. Talano<sup>(1)(+)\*</sup>, María Florencia Olmos Nicotra<sup>(1)</sup>, Leticia  
5 Escudero<sup>(2)</sup>, María Laura Breser<sup>(3)</sup>, Carina Porporatto<sup>(3)</sup>, Elizabeth Agostini<sup>(1)</sup>

6

7 (1) Molecular Biology Department, Faculty of Exact, Physical, Chemical and Natural  
8 Sciences, National University of Río Cuarto, Ruta Nacional 36 Km 601, (CP 5800)  
9 Río Cuarto, Córdoba, Argentina

10 (2) Laboratory of Analytical Chemistry for Research and Development (QUIANID),  
11 Interdisciplinary Institute of Basic Sciences (ICB), UNCUYO–CONICET, Faculty of  
12 Natural and Exact Sciences, National University of Cuyo. Padre J. Contreras 1300,  
13 (CP 5500) Mendoza, Argentina.

14 (3) Research and Transference Center of Villa María (CITVM-CONICET), National  
15 University of Villa María, Arturo Jauretche 1555, (CP 5900), Villa María, Córdoba,  
16 Argentina.

17 (+) Should be considered as first authors since they have equally contributed to this work.

18 (\*) Corresponding author.

19

20 Tel.: +54 358 4676537; fax: +54 358 4676232.

21

22 e-mail: analauraarmendariz@gmail.com, mtalano@exa.unrc.edu.ar;

23 folmos@exa.unrc.edu.ar; letibelescudero@gmail.com; laurabreser45@hotmail.com;

24 cporporatto@unvm.edu.ar; eagostini@exa.unrc.edu.ar;

25 **Abstract**

26 Inoculation practice with plant growth-promoting bacteria (PGPB) has been proposed as a good  
27 biotechnological tool to enhance plant performance and alleviate heavy metal/metalloid stress. Soybean  
28 is often cultivated in soil with high arsenic (As) content or irrigated with As-contaminated groundwater,  
29 which causes deleterious effects on its growth and yield, even when it was inoculated with rhizobium.  
30 Thus, the effect of double inoculation with known PGPB strains, *Bradyrhizobium japonicum* E109 and  
31 *Azospirillum brasilense* Az39 was evaluated in plants grown in pots under controlled conditions and  
32 treated with As. First, the viability of these co-cultivated bacteria was assayed using a flow cytometry  
33 analysis using SYTO9 and propidium iodide (PI) dyes. This was performed *in vitro* to evaluate the  
34 bacterial population dynamic under 25  $\mu$ M AsV and AsIII treatment. A synergistic effect was observed  
35 when bacteria were co-cultured, since mortality diminished, compared to each growing alone. Indole  
36 acetic acid (IAA) produced by *A. brasilense* Az39 would be one of the main components involved in *B.*  
37 *japonicum* E109 mortality reduction, mainly under AsIII treatment. Regarding *in vivo* assays, under As  
38 stress, plant growth improvement, nodule number and N content increase were observed in double  
39 inoculated plants. Furthermore, double inoculation strategy reduced As translocation to aerial parts thus  
40 improving As phytostabilization potential of soybean plants. These results suggest that double  
41 inoculation with *B. japonicum* E109 and *A. brasilense* Az39 could be a safe and advantageous practice  
42 to improve growth and yield of soybean exposed to As, accompanied by an important metalloid  
43 phytostabilization.

44

45 **Keywords:** ARSENIC, PGPB, INOCULATION, PHYTOSTABILIZATION, GLYCINE MAX

46

## 47 1. Introduction

48 Arsenic (As) is a highly toxic metalloid present in the environment, being arsenate (AsV) and  
49 arsenite (AsIII) the predominant inorganic species in soil and water (Farooq et al., 2016). In plants, As  
50 interferes with critical metabolic processes such as photosynthesis and can induce water stress by  
51 reducing transpiration rate, stomatal conductance, and leaf relative water content along with reduction  
52 of xylem vessel size. In addition, this metalloid induces oxidative stress, cellular membrane damage and  
53 electrolyte leakage (Stoeva et al., 2004; Gusmán et al., 2013a,b). As consequence, a severe plant growth  
54 and reproductive capacity inhibition is often seen (Garg and Singla, 2011; Finnegan and Chen, 2012;  
55 Reichman, 2014; Armendariz et al., 2016; Bustingorri and Lavado, 2014).

56 The use of plants for contaminant removal is named phytoremediation and based on the type of  
57 biological mechanism adopted this phytotechnology is classified as phytoextraction, phytostabilization,  
58 phytotransformation, phytovolatilization, rhizofiltration or phytostimulation (Abhilash et al., 2009).  
59 Generally, plants use a variety of processes that collectively contribute to the overall effectiveness of  
60 remediation (Kumar Yavad et col., 2018). For heavy metals, several reviews have been published in the  
61 last years, mainly considering phytoextraction, phytostabilization, phytoevaporation and  
62 phytotransformation (Gomes et al., 2017; Mahar et al., 2016; Sarwar et al., 2017). Initially, these  
63 phytotechnologies focused on heavy metals/metalloids phytoextraction, while phytostabilization  
64 received less attention. Recently, phytostabilization has been revalued as a metal immobilization  
65 strategy for polluted soils (Sarwar et al., 2017). Even more, high metal/metalloid retention ability in  
66 roots takes relevance for edible plants and those which have fruits or grains for food, since the risk of  
67 contaminant introduction into the food chain is minimized (Robinson et al., 2009, Sarwar et al., 2017).  
68 The use of plant growth promoting bacteria (PGPB) can improve growth of plants exposed to  
69 metal/metalloids and even promote phytostabilization through their ability to decrease metal  
70 bioavailability. This strategy is named as assisted phytoremediation. Although many PGPB have been  
71 isolated and used for metal phytoremediation improvement (Nie et al., 2002; Ullah et al., 2015; Ma et

72 al., 2016; Titah et al., 2013; Ojuederie and Babalola, 2017; Sarwar et al., 2017), few studies have  
73 evaluated PGPB potential for As phytostabilization.

74 Soybean (*Glycine max* L.) is a legume with worldwide economic importance because of its high  
75 protein content in grains and other valuable food sub-products. For optimum yields, this crop is  
76 inoculated with symbiotic rhizobia, mainly *Bradyrhizobium japonicum* strains. Soybean-rhizobia  
77 symbiosis is an important ecological and agronomical association, since plants receive enough Nitrogen  
78 (N) supply through biological N-fixation, hence, the use of N fertilizers can be reduced (Sytnikov 2013).  
79 The association between soybean roots and *B. japonicum* bacteria results in the formation of specific  
80 organs, called nodules, where N-fixation takes place. The main products of N-fixation on soybean  
81 nodules, such as ureides (allantoin and allantoic acid), are exported to the rest of the plant where they  
82 are incorporated into aminoacids and proteins. Thus, the number of effective nodules (regularly  
83 evaluated through its red-pink colour indicative of leghemoglobine presence) is key in those crops in  
84 which N content depends mainly on biological N-fixation (Wang and Martinez-Romero 2000;  
85 Masciarelli et al., 2014; Pommeresche and Hansen, 2017).

86 Argentina presents a cultivated area of 20.3 million hectares of soybean, with a production of 58  
87 million tons (2016-2017) (Integrated Agricultural Information System Argentina, 2016). This crop is  
88 often cultivated in areas with high As concentration and/or irrigated with groundwater containing this  
89 metalloid because of crop expansion to arid and semiarid regions with low rainfall regime (Smedley and  
90 Kinniburgh, 2002; Bundschuh et al., 2010). This is of great concern because As toxicity may produce  
91 not only animal and human health problems but also negatively affect sustainable crop production. In  
92 Argentina, *B. japonicum* E109 is used for soybean inoculation since it is the commercially available  
93 strain (Cassán et al., 2009). In a previous work, we showed that under As exposure this bacterium was  
94 sensitive, mainly when exposed to AsIII since its growth was reduced a 50% for 10  $\mu$ M and almost  
95 totally reduced for 25  $\mu$ M AsIII, while for AsV from 25  $\mu$ M only a minimal reduction in growth was  
96 seen (Armendariz et al., 2015). When soybean plants were treated with As, the plant growth was  
97 significantly reduced when exposed to 25  $\mu$ M AsV and AsIII even when they were inoculated with *B.*

98 *japonicum* E109 since nodule number was reduced under these conditions (Talano et al., 2013). Other  
99 reports have also shown that soybean inoculated with other *Bradyrhizobium* strains was negatively  
100 affected by As exposure leading to significant ecological, economic and nutritional losses (Reichman,  
101 2014; Bustingorri and Lavado, 2014). Therefore, in As impacted environments the application of PGPB  
102 could not only improve As phytostabilization process, but also alleviate metal toxicity and stimulate  
103 plant growth. Hence, it could constitute an economic and effective approach for reducing metalloid  
104 impact (Ojuederie and Babalola, 2017). Considering that *B. japonicum* E109 is the commercially  
105 available strain and the only one adopted for soybean inoculation schemes in Argentina and taking into  
106 account the negative performance when inoculated in As-treated soybean plants, a strategy of  
107 combining this with other PGPB could be considered. In this sense, *Azospirillum brasilense* Az39 is a  
108 free-living bacterium that when inoculated alone or in combination with *B. japonicum* E109 has shown  
109 capacity to promote seed germination, nodule formation, and early development of soybean seedlings in  
110 As-free soils (Cassán et al., 2009). *A. brasilense* Az39 is able to produce indole acetic acid (IAA),  
111 gibberellins (GA3) and zeatin (Z), which produce morphological and physiological changes in maize  
112 and soybean young seed tissues (Cassán et al., 2009; García et al., 2017).

113 Based on this background, the aims of this work were to evaluate the *in vitro* viability of two  
114 rhizospheric strains (*B. japonicum* E109 and *A. brasilense* Az39) under AsV and AsIII exposure in  
115 single and co-cultured suspensions and to test *in vivo* the effects of double inoculation (DI) on soybean  
116 plants exposed to As. The advantages of DI, in particular on soybean germination parameters, plant  
117 growth, nitrogen content, nodule number and As accumulation were evaluated, in order to assess the  
118 feasibility of DI strategy for an efficient symbiosis and growth improvement in soybean plants under As  
119 stress.

120

121

## 122 2. Materials and Methods

### 123 2.1 Bacterial strain and growth conditions

124 Two collection strains, *B. japonicum* E109 and *A. brasilense* Az39, were used in the present  
125 work. These bacteria belong to a strain collection from the Agriculture Collection Laboratory of the  
126 Instituto de Microbiología y Zoología Agrícola (IMYZA) and Instituto Nacional de Tecnología  
127 Agropecuaria (INTA), Castelar, Argentina. The complete genome sequence of *B. japonicum* E109 is  
128 available at NCBI GenBank under the following accession number CP010313 (Torres et al., 2015)  
129 while that of *A. brasilense* Az39 is registered as CP007793 for the chromosome and CP007794 to  
130 CP007798 for the other replicons (Rivera et al., 2014). Bacterial inocula were obtained by growing *B.*  
131 *japonicum* E109 for 96 h in liquid TY medium containing vancomycin ( $4 \mu\text{g mL}^{-1}$ ) and *A. brasilense*  
132 Az39 for 24 h in LB medium. Both cultures were incubated under agitation at 200 rpm and 28 °C. ~~When~~  
133 ~~necessary, the CFU mL<sup>-1</sup> of bacterial suspension was calculated by drop count plate method~~  
134 ~~(Somasegaran and Hoben, 1994).~~

135

### 136 2.2. Bacterial in vitro studies

#### 137 2.2.1. Viability analysis of *B. japonicum* E109, *A. brasilense* Az39 and co-cultured strains 138 under arsenic stress using flow cytometry analysis

139 In order to evaluate rhizospheric strains viability under As stress, a flow cytometry analysis was  
140 performed in single or co-cultured bacterial suspensions. For that, bacterial cultures were centrifuged at  
141 10,000 rpm for 20 min at 15 °C, and the pellets were suspended in physiological saline solution (NaCl  
142 0.9%) to reach an OD<sub>620nm</sub> of 1. Finally, the bacterial suspensions were incubated separately or co-  
143 cultured in absence or presence of 25  $\mu\text{M}$  AsV or AsIII for 72 h. After that, bacterial suspensions were  
144 harvested by centrifugation and pellets were washed twice with saline phosphate buffer containing 1  
145 mM EDTA, pH 7.4 (Mandal et al., 2008). Viability evaluation was performed using the LIVE/DEAD  
146 BacLight Bacterial Viability Kit staining (Invitrogen, ThermoFisher Scientific, CA, USA), according to  
147 the manufacturer's instructions. Bacterial viability was carried out by SYTO9 and propidium iodide (PI)

148 dyes which determine cell membrane integrity. SYTO9 dye can be incorporated to live and dead  
149 bacterial cells and can be useful to determine the total cells population, while PI dye is commonly used  
150 for identify dead cells which present disrupted membranes. Bacterial suspensions were acquired on an  
151 ACCURI C6 (BD Biosciences, San Diego, CA, USA) flow cytometer and the data were analyzed using  
152 FlowJo software (Tree Star, OR, USA). To evaluate mortality of the strains treated with As, bacteria  
153 were detected by forward scatter (FSC), side scatter (SSC), and fluorescence.

154

155 **2.2.2. IAA produced by *A. brasilense* Az39 under As stress and its effect on *B. japonicum***  
156 ***E109* survival**

157 *A. brasilense* Az39 cultures grown for 24 h in LB medium were harvested by centrifugation and  
158 the pellet was suspended in physiological saline solution and adjusted to an OD<sub>620nm</sub> of 0.5. Those  
159 bacterial suspensions were supplemented with stock sodium arsenate (AsHNa<sub>2</sub>O<sub>4</sub>7H<sub>2</sub>O) (SIGMA)  
160 (AsV) and sodium arsenite (NaAsO<sub>2</sub>) (SIGMA) (AsIII) solutions to reach a final concentration of 25  
161 µM. For the control suspensions the same volume of As stocks was added as distilled water. These  
162 suspensions were incubated at 28° C and 180 rpm for 72 h. Then, IAA produced by *A. brasilense* Az39  
163 was tested as described by Glickman and Dessaux (1995) using the Salkowski reagent (H<sub>2</sub>SO<sub>4</sub>: 37.5  
164 mL; FeCl<sub>3</sub> 0.5M: 1.88 mL; H<sub>2</sub>O: 62.5 mL for 100 mL). For that, a calibration curve using commercial  
165 IAA solutions from 2 to 20 µg mL<sup>-1</sup> was used and the OD (at 530 nm) values were registered. Finally,  
166 the IAA concentration produced by *A. brasilense* Az39 was expressed as µM considering its molecular  
167 weight (175.18 g mol<sup>-1</sup>). As positive control of IAA production, *Azospirillum* sp. Cd strain was included  
168 (Kaushik et al., 2000), while non-inoculated physiological saline solution was used as negative control.

169 For the evaluation of IAA effect on *B. japonicum* E109 viability flow cytometry analysis was  
170 performed. For that, *B. japonicum* E109 culture previously grown in TY medium was centrifuged and  
171 the pellet was suspended in physiological saline solution to reach an OD<sub>620nm</sub> of 1. Then, 5 mL-fractions  
172 of that suspension were diluted 1/2 to reach a final OD<sub>620nm</sub> of 0.5 with: a) physiological saline solution  
173 with the addition of commercial IAA (final concentration 4 µM), b) cell-free supernatant from *A.*

174 *brasilense* Az39 and c) *A. brasilense* Az39 viable cells previously suspended in physiological saline  
175 suspension with an  $OD_{620nm}$  of 1. As control, *B. japonicum* E109 suspension in physiological saline  
176 solution was incubated under the same conditions. The final  $OD_{620nm}$  of *B. japonicum* E109 suspensions  
177 at all the conditions reached a value of 0.5. All these treatments were exposed to AsV or AsIII (25  $\mu$ M)  
178 adding the proper volume of concentrated stock solutions while those without As were used as control.  
179 After incubation for 72 h, bacteria were centrifuged and washed with phosphate saline buffer (PBS)  
180 with 1 mM EDTA. Then, cell mortality was evaluated by flow cytometry analysis as previously  
181 described in order to discuss the IAA effect on *B. japonicum* E109.

182

### 183 **2.3. Inoculation studies in As-treated plants**

#### 184 **2.3.1. Plant material, growth and treatment conditions**

185 Seeds of *Glycine max* cv. DM 4670 were used. They were sterilized using 70% (v/v) ethanol for  
186 1 min and then 30% (v/v) sodium hypochlorite for 10 min. They were washed thoroughly with sterile  
187 distilled water, submerged in distilled water and incubated at  $28 \pm 2$  °C with agitation for 24 h. Then,  
188 they were used for *in vitro* studies (germination test) and *in vivo* inoculation assay in pots as detailed in  
189 2.3.2. and 2.3.3. sections, respectively.

190

#### 191 **2.3.2. In vitro studies: Effect of inoculation on soybean germination under As stress**

192 To evaluate whether inoculation contributes at the initial development stage of soybean,  
193 germination index (*GI*), germination rate index (*S*), root length (cm) and relative radical elongation (*E*)  
194 were determined in seeds with single or double inoculation. For that, sterilized seeds (n=10) were placed  
195 on Petri dishes containing sterile filter paper. It was impregnated with 6 mL of: sterile water (control  
196 condition), *B. japonicum* E109 or *A. brasilense* Az39 suspension made with physiological saline  
197 solution ( $OD_{620nm}$  0.5) and equal amount of mixed bacterial suspension for DI condition. For As  
198 treatment, water or bacteria suspensions were supplemented with AsV or AsIII solutions to reach 25  $\mu$ M  
199 final concentration. The experiment was repeated three times and each condition was analyzed by

200 duplicate in each independent experiment ( $n= 60$ ).  $GI$ ,  $S$ , root length and  $E$  were determined after  
201 incubating the plates for 7 d at  $28 \pm 2$  °C in darkness.

202  $E$  and  $GI$  were calculated according to Barrena et al. (2009): ( $E = [Xf/Xc] \times 100$ ) and ( $GI =$   
203  $[(Gf/Gc) \times 100] \times E/100$ ), where:  $Xf$ = root length average of AsV or AsIII treated seeds,  $Xc$ = root  
204 length average of control seeds,  $Gf$ = germinated seeds in the presence of AsV or AsIII and  $Gc$ = seeds  
205 germinated under control conditions.  $S$  was calculated as described by Ahmed and Wardle (1994): ( $S =$   
206  $[N1/1 + N2/2 + N3/3 + \dots + Nn/n] \times 100$ ), where:  $N_1, N_2, N_3 \dots N_n$  is the proportion of seeds germinating  
207 on days 1, 2, 3 ...  $n$  throughout the experiment. In this way,  $S$  varies from 100 (if all seeds germinate on  
208 the first day) to 0 (if the seeds did not germinate at the end of the experiment).

209

### 210 **2.3.3. *In vivo* inoculation assays and responses of soybean plants under As stress**

211 Previously disinfected seeds were placed in sterile flasks and soaked with a necessary volume  
212 (28 seeds/4.3 mL) either of physiological saline solution (non-inoculated), or bacterial suspensions  
213 obtained as previously described (section 2.2.1) from *B. japonicum* E109 and *A. brasilense* Az39 and  
214 both (inoculated and double inoculated (DI)). When soybean seeds were DI, the suspension was  
215 prepared from a mixture of both microorganisms in equal parts. Then, seeds were incubated in an orbital  
216 shaker (200 rpm) for 2 h at 28 °C to allow the impregnation with bacteria. After draining the seeds from  
217 the bacterial suspensions they were kept in a laminar flow hood by 2 h to allow them to dry.  
218 Subsequently, 10 seeds (non-inoculated (NI), inoculated with *B. japonicum* E109 or *A. brasilense* Az39  
219 and those DI) were placed in pots containing 50 g of sterile perlite humidified by capillarity with 125  
220 mL distilled water (control) or 25  $\mu$ M AsV and AsIII solution. Plants were supplemented alternatively  
221 with water or free nitrogen  $\frac{1}{2}$  Hoagland solution as needed. At 14 and 21 days, plants were repeatedly  
222 treated with As, so the treatments were designated as T0, T14 and T21.

223 The experiments were carried out in a growth chamber set with controlled temperature ( $28 \pm 2$   
224 °C) under photoperiod regime [16 h light ( $200 \mu\text{mol m}^{-2} \text{s}^{-1}$ )/8 h dark] and relative humidity of 80%.  
225 After 30 d, harvested plants were divided in root, shoot and nodules. First, the nodule number was

226 counted. Dry weight of root and shoot (obtained after drying in an electric heating oven at 70 °C for 5 d)  
227 was registered. Root and shoot were frozen, homogenized with liquid N<sub>2</sub> and kept at -80 °C until their  
228 use for analytical determinations.

229

### 230 ***2.3.3.1 Total nitrogen content in soybean plants***

231 Total nitrogen content was determined in shoots by Kjeldahl Method (Reference Method) based  
232 on titration of protein and non-protein nitrogen through a digestion with concentrated sulfuric acid  
233 (AOAC, 1990).

234

### 235 ***2.3.3.2 Total As accumulation analysis***

236 Root and shoot of inoculated and NI plants were used for As quantification. Dried tissues were  
237 acid digested and total As was determined by atomic fluorescence spectrometry (AFS). For digestion,  
238 0.3 g of sample were weighed and mixed with 10 mL of concentrated HNO<sub>3</sub> (Ultrex® II Mallinckrodt  
239 Baker, Phillipsburg, NJ, USA) (30 min at 50°C and 60 min under boiling). After cooling, 2 mL of H<sub>2</sub>O<sub>2</sub>  
240 30% (Merck, Darmstadt, Germany) were added and the digestion was continued at constant boiling  
241 during 60 min. Each digested sample was left to cool, and then it was filtered and transferred to a 50 mL  
242 flask. Subsequently, 5 mL of HCl 37% (v/v) (Merck) and 2 mL of IK 25% (w/v) (JT Baker, USA) were  
243 added to the flask. Finally, ultrapure water (18 MΩ cm) (Bedford, MA, USA) was added to reach a  
244 volume of 50 mL. Arsenic was detected using a Rayleigh AF-640A atomic fluorescence spectrometer  
245 (Beijing Rayleigh analytical Instrument Corp., Beijing, China). Instrumental and experimental  
246 conditions were: lamp and wavelength: As High intensity hollow cathode lamp, 197.3 nm; main current:  
247 40mA; auxiliary current: 0 mA; reductant: 0.7% (w/v) NaBH<sub>4</sub> (Merck), carrier: 5% (v/v) HCl (Merck);  
248 reductant and carrier flow rates: 12 mL min<sup>-1</sup>, argon flow rate: 800 mL min<sup>-1</sup> and atomizer temperature:  
249 300°C. Calibration was performed against aqueous standards and blank solutions. For validation, a  
250 Perkin Elmer (Uberlingen, Germany) Model 5100ZL atomic absorption spectrometry equipped with a

251 transversely heated graphite atomizer, an As Electrodeless Discharge Lamp (EDL) and a Zeeman  
252 correction system, was used.

253

#### 254 **2.4. Statistical analysis**

255 Results are the average of at least 3 independent replicates, performed by triplicate. Mean and  
256 standard errors of the evaluated parameters were calculated and plotted using the Microsoft Excel 2007  
257 program. To determine the statistical difference between at least one pair of means, analysis of variance  
258 test (ANOVA) was used. When the assumptions of homogeneity of variance (*Levene* test) and normality  
259 (*Shapiro-Wilk* test) were not checked, corresponding transformations were performed using the  
260 appropriate functions. To determine significant differences between treatments, *Tukey* test was applied,  
261 with a significance level of 0.05 ( $p < 0.05$ ). For some parameters nonparametric analysis was performed  
262 by *Kruskal Wallis* test (Software InfoStat versión 2015; from National University of Córdoba,  
263 Argentina).

264

### 265 **3. Results and Discussion**

#### 266 **3.1. Bacterial in vitro studies**

##### 267 **3.1.1. Viability analysis of *B. japonicum* E109, *A. brasilense* Az39 and co-cultured strains under 268 *AsV* and *AsIII* treatment**

269 In order to understand how 25  $\mu\text{M}$  *AsV* and *AsIII* affects *B. japonicum* E109 and *A. brasilense*  
270 Az39 viability, single or mixed cultures were stained with SYTO9 and PI dyes and analyzed by flow  
271 cytometry. As shown in Figure 1A (representative dot plots) and Figure 1B, the metalloids increased *B.*  
272 *japonicum* E109 and *A. brasilense* Az39 mortality in single and DI cultures. In this sense, mortality  
273 increase was statistically significant only for *AsIII* treatment and *B. japonicum* E109 was more affected  
274 than *A. brasilense* Az39, since mortality values were 45% and 38%, respectively (Fig 1B). These data  
275 are in agreement with previous results obtained using conventional methodology [growth curves  
276 ( $\text{OD}_{620\text{nm}}$ ) and plate count ( $\log_{10}$  CFU  $\text{mL}^{-1}$ )] (Armendariz et al., 2015). As it is shown, As is an

277 important stress factor especially for *B. japonicum* E109, severely affecting its viability. However, co-  
278 culture of *B. japonicum* E109 and *A. brasilense* Az39 improved bacteria survival under As treatment  
279 compared with single cultures. Furthermore, this effect was more significant under AsIII treatment since  
280 co-cultured mortality decreased 21% for AsV and 13-27% AsIII treatment, compared with the mortality  
281 of single bacteria suspensions. Hence, flow cytometry was useful for identifying and quantifying viable  
282 and dead rhizobacteria in an easy, fast and efficient way as a complement to standard methods (Mandal  
283 et al., 2008; Tejerizo et al., 2015; Valdameri et al., 2015). Moreover, flow cytometry assay allowed us  
284 analyzing in an accurate and exact manner the behavior of this mixed bacterial population under As  
285 exposure. These results suggest that there may be a synergistic/cooperative effect between bacteria,  
286 which encourage us to evaluate their effectiveness under *in vivo* conditions for the improvement of  
287 soybean plants exposed to As.

288

### 289 ***3.1.2 IAA produced by A. brasilense Az39 under As stress and its effect on B. japonicum*** 290 ***E109 survival***

291 With the purpose of exploring whether IAA produced by *A. brasilense* Az39 is responsible of the  
292 increased viability of *B. japonicum* E109 in co-culture under As stress, it was incubated with  
293 commercial IAA, *A. brasilense* Az39 cell-free supernatant and *A. brasilense* Az39 bacterial suspension.  
294 *B. japonicum* E109 alone was also included as control and incubated under the same conditions. The  
295 cell-free supernatant was included to consider the presence of another potential soluble compound in the  
296 culture medium responsible of *B. japonicum* E109 survival.

297 First, IAA produced by *A. brasilense* Az39 was determined under AsV and AsIII exposure. As  
298 shown in Table 1, *A. brasilense* Az39 produced around 4-5  $\mu\text{M}$  of IAA, similar to *A. brasilense* Cd,  
299 used as a positive control, with no significant effect of 25  $\mu\text{M}$  AsV or AsIII on IAA production.  
300 Considering this, 4  $\mu\text{M}$  was chosen as the concentration of commercial IAA added to *B. japonicum*  
301 E109.

302 As it can be seen in Figure 2, when *B. japonicum* E109 was incubated with commercial IAA (4  
303  $\mu\text{M}$ ), *A. brasilense* Az39 cell-free supernatant and *A. brasilense* Az39 cells, its mortality percentage was  
304 reduced. Although IAA induced a mortality reduction effect in all conditions, the main effect was  
305 observed under AsIII treatment (Figure 2). These results indicate that IAA produced by *A. brasilense*  
306 Az39 would represent an important component associated to *B. japonicum* E109 viability under As  
307 stress. In fact, it has been reported that *B. japonicum* strains can use this compound as a carbon source  
308 (Egebo et al., 1991; Jensen et al., 1995). In addition, there is some evidence that IAA might be a signal  
309 able to coordinate bacterial behavior to enhance protection under adverse conditions (Spaepen et al.,  
310 2007 and references there in). Using *E. coli*, Bianco et al. (2006a) and (2006b) showed that IAA induces  
311 the expression of genes related to survival under stress conditions and others involved in the central  
312 metabolic pathways such as the tricarboxylic acid cycle (TCA), glyoxylate shunt and amino acid  
313 biosynthesis (leucine, isoleucine, valine and proline). These findings showing IAA as a signaling  
314 molecule shed new light on the role of IAA in bacteria-plant interactions, but can also explain bacteria-  
315 bacteria interactions in the rhizosphere. Accordingly, in the present work, this phytohormone can play a  
316 key role in the protection of the more As-sensitive bacterial partner in the *B. japonicum* E109 and *A.*  
317 *brasilense* Az39 interaction in an As-contaminated environment. In order to evaluate the advantages of  
318 double inoculating soybean plants using *B. japonicum* E109 and *A. brasilense* Az39 in an As-  
319 contaminated soil, *in vitro* studies of germination parameters as well as *in vivo* studies with plants were  
320 performed.

321

### 322 **3.2. Inoculation studies in plants treated with As**

#### 323 **3.2.1. *In vitro* studies: Effect of inoculation on soybean germination under As stress**

324 Some parameters related to germination and young stages of soybean growth such as *GI*, *S*, root  
325 length and *E* were determined in NI and inoculated seedlings treated with As.

326 In NI seeds, *GI* was significantly reduced (around 64%) under both As treatments compared to  
327 control (Table 2). Similar reduction in germination percentage was shown in our previous work by

328 concentrations from 25  $\mu\text{M}$  AsV or AsIII (Talano et al., 2013). Considering that germination percentage  
329 is sometimes a relatively low-sensitive parameter to study the toxicity of a xenobiotic and not enough to  
330 predict subsequent effect on tested plant growth (Gong et al., 1999) here we present results from other  
331 related parameters such as *S*, root length and *E*. *S* was significantly reduced (23.8%) when seeds were  
332 treated with 25  $\mu\text{M}$  AsIII, whereas root length and *E* were significantly affected by both As treatments,  
333 with a decrease of 50%. Similar results were found by Kaur et al. (2012) whom reported that As  
334 exposure (10  $\mu\text{M}$ ) caused a reduction of around 50% of radicle emergence and elongation in *Phaseolus*  
335 *aureus*. The negative effect of As on germination and early development of seedlings has been  
336 attributed to the marked decline in amylolytic enzyme activities in rice and wheat endosperms, which  
337 produce a delay in mobilization of starch (Jha and Dubey 2005; Liu et al., 2005). Also, As produced a  
338 reduction of N-assimilatory enzyme activities (nitrate reductase, nitrite reductase and glutamine  
339 synthetase) in germinating rice seeds and seedlings, with the consequent reduced vigor and impaired  
340 growth (Jha and Dubey, 2004a; Jha and Dubey, 2004b). Inhibition of proteases has been also reported in  
341 As-treated plants, thus it can explain the reduced germination of soybean since proteins are the main  
342 reserve material in the grains. Thus, the disturbance of As on sugars, N and protein metabolism of  
343 germinating seeds could explain the reduced *GI*, *S*, root length and *E* observed for As-treated soybean  
344 seeds.

345         Regarding inoculation, in the present work, no improvement was observed in seeds inoculated  
346 with *A. brasilense* Az39, which was surprising since this strain presented high tolerance to the metalloid  
347 as it was previously demonstrated (Armendariz et al., 2015). Contrarily, when As-treated seeds were  
348 inoculated with *B. japonicum* E109, all the analyzed parameters significantly increased compared to NI  
349 seeds. Similarly, a positive effect has also been reported by Dary et al. (2010) since germination of  
350 *Lupinus luteus* seeds was improved when they were inoculated with *Bradyrhizobium* sp. 750 and  
351 exposed to contaminated soils with moderated heavy metal concentration (including around 65-70 mg  
352  $\text{Kg}^{-1}$  of As).

353

### 354 3.2.2. *In vivo* inoculation assays and responses of soybean plants under As stress

#### 355 3.2.2.1. *Effect on growth and nodulation*

356 Under control condition (without As), inoculation with *B. japonicum* E109, *A. brasilense* Az39  
357 or DI produced a significant increase in shoot and root biomass compared to NI plants (Fig. 3). Plants  
358 inoculated with *B. japonicum* E109 showed an increase in root and shoot biomass of 27% and 47%,  
359 respectively, while in plants inoculated with *A. brasilense* Az39 the increase was lower (22 and 17%,  
360 respectively). However, when soybean seeds were DI no significant differences in plant biomass were  
361 found compared to single inoculations. These results agree with pre-existing data, since numerous field  
362 studies and laboratory tests have shown that *B. japonicum* E109 significantly increases soybean  
363 production (Cassán et al., 2009; Benintende et al., 2010). However, it seems that the beneficial effects of  
364 each individual strain would not be additive when they were DI. This could be explained by alteration in  
365 microbial ecology of the rhizosphere, probably by natural competition. Some evidences indicate that the  
366 production of secondary metabolites and other physiological processes in bacteria depend on population  
367 density. Therefore, the benefits that microorganisms produce in plants could not be significant if they do  
368 not reach an appropriate number or density (Barnard et al., 2007).

369 Under As stress, inoculation was an effective strategy to improve plant growth, although with  
370 less efficiency. Although there was a negative effect of As on soybean, reflected as biomass reduction,  
371 the damage was more severe in NI plants (Fig. 3). Inoculation with *A. brasilense* Az39 or *B. japonicum*  
372 E109 separately produced an increase in root and shoot biomass in As-treated plants, but this effect was  
373 higher for those inoculated with *B. japonicum* E109. Considering DI, there was a significant growth  
374 improvement of As-treated plants, although it was statistically significant only for AsV treatment.  
375 Similarly, Reichman (2007; 2014) observed that inoculation with *B. japonicum* CB1809 promoted  
376 soybean, wheat and sunflower growth when exposed to AsV compared to those NI plants. In addition,  
377 other authors have reported better results in canola and rice growth when inoculated with  
378 *Brevundimonas diminuta* and *Enterobacter cloacae* CAL2, respectively, under As stress (Nie et al.,  
379 2002; Singh et al., 2016). On the other hand, there are few reports on *Azospirillum* strains inoculated in

380 As-treated plants. This is not surprisingly because our previous results indicated that *A. brasilense* Az39  
381 did not promote germination parameters in the presence of As. Similarly, Lyubun et al. (2006) neither  
382 found significant differences in biomass of wheat plants inoculated with *A. brasilense* Sp245 growing in  
383 presence of As compared to NI ones.

384         Regarding nodulation under control conditions, the number of effective nodules was not  
385 modified in DI plants compared to those inoculated with *B. japonicum* E109 (Fig. 4). Under As  
386 treatment, the nodule number was significantly reduced compared with control, mainly by 25  $\mu\text{M}$  AsIII.  
387 However, in DI plants the nodule number significantly increased compared *B. japonicum* E109  
388 inoculated plants, under AsIII stress. Several authors have described that nodulation of legumes is  
389 generally reduced or inhibited in As-contaminated soils (Carrasco et al., 2005; Mench et al., 2006;  
390 Talano et al., 2013). For instance, Reichman (2007) reported that the nodule number in soybean plants  
391 inoculated with *B. japonicum* CB1809 was reduced by 90% in the presence of 5  $\mu\text{M}$  AsV. In addition,  
392 in As-treated plants of *Vigna mungo* and *Medicago* sp. inoculated with highly As-resistant bacterial  
393 strains this parameter was also reduced (Pajuelo et al., 2008; Mandal et al., 2011). This decrease would  
394 be related to the toxic effect of As on roots, mainly with reduction or damage of radical hairs which  
395 would affect the sensitivity, or the low expression level of several nodulin genes, which have a  
396 fundamental role in the infection thread formation (Pajuelo et al., 2008; Lafuente et al., 2010). More  
397 recently, La Fuente et al., (2015) using the model legume *Medicago truncatula* and *Ensifer* (syn.  
398 *Sinorhizobium*) *medicae* MA11, a highly As-resistant bacterium, found a strong reduction of nodule  
399 number under AsIII treatment with a median inhibitory concentration ( $\text{ID}_{50}$ ) of 20  $\mu\text{M}$ . The author  
400 emphasized that nodulation was the most sensitive process comparing the AsIII- $\text{ID}_{50}$  for plant growth,  
401 seed germination, shoot and root length, nodulation and other physiological parameters.

402         In the present work, the lower nodulation in As-treated plants inoculated with *B. japonicum*  
403 E109 would be a consequence of root biomass reduction and minor number of root hairs as available  
404 infection points. In addition, since *B. japonicum* E109 is highly sensitive to As, mainly AsIII  
405 (Armendariz et al., 2015), a smaller number of bacteria are alive for colonization and symbiosis is

406 reduced. Other explanation about As deleterious effect would be related with metalloids injuries on root  
407 structure. The toxicity of As would also be related with delicate regulatory events through gene  
408 modulation during rhizobia-legume interaction. Recently, La Fuente et al., (2015) studied the effect of  
409 As on *M. truncatula*-*E. medicae* MA11 symbiosis through transcriptomic meta-analysis. In this  
410 experimental model, the enhancement of chalcone synthase transcripts (involved in the first step of  
411 legume-rhizobia cross-talk) and the repression of 13 subsequent nodulation genes codifying for Nod  
412 factors (involved in perception, infection, thread initiation and progression, and nodule morphogenesis)  
413 suggests that plants are impaired to establish symbiotic interactions under AsIII stress. This focus  
414 involving transcriptomic analysis of As-treated plants inoculated with rhizobia would complement the  
415 advances made with 'arsenomic' approach which includes the study of non-legume plants or legume-  
416 rhizobia interaction but without stress. Certainly, more studies in this line but under As exposure would  
417 allow elucidating the effect of the metalloid on symbiotic interactions from a global perspective.

418

#### 419 **3.2.2.2 Total N content**

420 In control condition, the total N content in shoots (Fig. 5) was higher when soybean plants were  
421 inoculated with *B. japonicum* E109, and also when they were DI. Contrarily, plants inoculated with *A.*  
422 *brasilense* Az39 did not show considerable increase in N content compared to control NI plants. In the  
423 presence of As, N content of NI plants did not change while As-treated plants inoculated with *A.*  
424 *brasilense* Az39 showed higher N content although without significant difference. Contrarily, in plants  
425 inoculated with *B. japonicum* E109 As treatment produced reduction in N content (around 20-25%),  
426 which could be explained by the considerable reduction in nodule number, as it was previously shown  
427 (Fig 4), and the reduction of nitrogenase activity in nodules of As-treated plants (data not shown). In  
428 addition, soybean root nodules derived from plants treated with both AsV and AsIII showed a pale pink  
429 or whitish inner coloration as compared to the intense red color of control plant nodules. This result  
430 indicates a lower concentration of leghemoglobin thus, higher O<sub>2</sub> concentration diffuses inside the  
431 nodule and nitrogenase activity decreases (Kundu et al., 2003). It is important to remark that, in DI

432 plants, the N content increased under As treatment. These results suggest that *A. brasilense* Az39, a  
433 highly As-tolerant strain, would be efficient in N-fixing under As stress, slightly improving N content in  
434 As-treated plants when compared with those inoculated only with *B. japonicum* E109. It shows that  
435 addition of *Azospirillum* strain to inoculation programs would give better results in plant growth

436

### 437 **3.3.3 Effect of inoculation on As accumulation in soybean plants**

438 As shown in Fig 6, the pattern of As accumulation changed depending on the bacterium used. In  
439 general, inoculation produced a reduction in As concentration in roots independently of As chemical  
440 species, except for DI plants treated with AsIII. In this case, the root accumulated higher As content,  
441 constituting a good strategy for an efficient phytostabilization of As, even more when these plants had  
442 low As accumulation in shoots. In fact, inoculated plants mainly those with *A. brasilense* Az39 or DI  
443 showed reduced As concentration in aerial parts and consequently lower As translocation compared  
444 with NI plants.

445 In a similar way, different plant species inoculated with plant growth promoting bacteria from  
446 *Staphylococcus*, *Bacillus*, *Acinetobacter* genera and others, have shown reduced As uptake and minor  
447 accumulation in aerial parts, grains and/or other edible parts of plants as a result of the bacterial  
448 inoculation (Srivastava et al., 2013; Das et al., 2016; Das and Sarkar, 2018). Therefore, those bacteria  
449 can be accounted for an efficient As phytostabilization. This finding emphasizes the important role of  
450 inoculation strategies to avoid high translocation and As accumulation in aerial parts of plants, mainly  
451 those which produce seeds/grains, fruits or are themselves vegetable foods for human and/or animal  
452 consumption. In this sense, inoculation could be helpful to avoid transference of As to food chain.  
453 However, it is important to consider that depending of bacterial strain and As chemical species, results  
454 can differ.

455 The presence of microorganisms affects the bioavailability of As in soybean rhizosphere. In this  
456 sense, it is known that bacteria are able to promote the mobility of metals and metalloids either by  
457 acidification and changes in the redox state of the medium, production of chelating agents or

458 siderophores and accumulation and/or adsorption in the biomass or exopolysaccharides (EPS) (Zubair et  
459 al., 2016 and references cited therein). Therefore, the different results obtained in the present work can  
460 be related with bacteria abilities for As metabolism in the rhizosphere as well as with tolerance  
461 mechanisms such as EPS and biofilm production (Armendariz et al., 2015). Joshi and Juwarkar (2009)  
462 reported that the ability of *Azotobacter* spp. to chelate Cd and Cr in EPS explained the low adsorption of  
463 metals by *Triticum aestivum*. In the present work, the lower content of metalloid in roots of AsIII-  
464 treated plants inoculated with individual strains (*B. japonicum* E109 or *A. brasilense* Az39) could be  
465 explained by their increased biofilm production under 25  $\mu$ M AsIII treatment, as shown in Armendariz  
466 et al. (2015). Biofilm would retain As and/or adsorbed it on the polymeric matrix frequently formed by  
467 EPS, thus leaving lower As concentration available for root (Rajkumar et al., 2012). In addition, other  
468 explanation for the lower As concentrations in roots inoculated with single bacterium would be the high  
469 As content translocated to aerial parts, which would depend on the metabolism of AsIII in the  
470 rhizosphere, uptake transporters and movility in plant tissues.

471 In the present work, it is important to remark that double inoculation of soybean plants, in  
472 particular under AsIII treatment, improved As-phytostabilization, hence reducing not only As lixiviation  
473 in soils but also As translocation to aerial parts and consequently, the potential risk of introducing this  
474 contaminant into the food chain. In addition, soybean plants treated with AsIII and DI showed a better  
475 growth and higher N content compared with NI plants. Also, it seems that the presence of both bacteria  
476 in soybean rhizosphere would contribute positively with nodule formation, probably as a result of the  
477 protective role of *A. brasilense* Az39 on *B. japonicum* E109 survival through IAA production.

478 As shown for soybean, legumes often accumulate As (and metals) mainly in root (Pajuelo et al.,  
479 2007, 2011; Reichman, 2007; El Aafi et al., 2012), and this fact is adequate for metal phytostabilization  
480 (Dary et al., 2010; El Aafi et al., 2012), as it reduces metal/loids mobilization in the plant rhizosphere  
481 with a scarce translocation to shoot (Mendez and Maier, 2008). In this sense, autochthonous legumes  
482 and resistant rhizobia are the most effective partnerships for many cases of metal-polluted soil  
483 restoration (Maynaud et al., 2013). However, when rhizobia are highly sensible to As, its combination

484 with resistant bacteria could be a synergistic way to improve plant and inoculation performance under  
485 stressful condition.

486

#### 487 **4. Conclusion**

488 Combining complementary properties of strains used for inoculation such as N-fixing ability  
489 from a poorly As-tolerant symbiotic strain (*B. japonicum* E109) with a highly As-tolerant free-living  
490 bacterium (*A. brasilense* Az39) is a good strategy to attenuate the As deleterious effect on soybean  
491 plants. A synergistic effect when both bacteria were co-cultured was observed through flow cytometry  
492 assays under As exposure. Despite there could be many factors involved in that protection, IAA  
493 produced by *A. brasilense* Az39 could be one beneficial metabolic relation that would reduce *B.*  
494 *japonicum* E109 mortality, mainly under AsIII treatment. Independently of the inoculation scheme used,  
495 single or combined, it produced positive effects on growth of As-treated plants. It is important to remark  
496 that DI plants significantly promoted plant growth, total nodule number and N content under As  
497 treatment. Regarding As accumulation, DI inoculation caused a reduction in As content in shoot and  
498 root of plants treated with AsV, while those exposed to AsIII showed higher retention of As in roots  
499 with low translocation to aerial parts. This would constitute an improvement of plant phytostabilization  
500 potential when exposed to AsIII, helping with As immobilization and consequently reducing As entry  
501 into the food chain. These results would allow considering DI strategy using *B. japonicum* E109 and *A.*  
502 *brasilense* Az39 as a safe and advantageous practice for the improvement of growth, yield of soybean  
503 crops and safe grain consumption for foods.

504

#### 505 **5. Acknowledgments**

506 ALA is a CONICET scholarship. MFON is a postdoctoral fellow. MAT, MLB, LE, CP and EA  
507 are members of the research career from Consejo Nacional de Investigaciones Científicas y Técnicas  
508 (CONICET) (Argentina). This work was supported by FONCYT (PICT 828/13), SeCyT-UNRC (PPI

509 2016-2018) and CONICET (PIP). We really appreciate and acknowledge to Dr. Sabrina Ibañez and Ana  
510 Laura Wevar Oller for language revision.  
511

ACCEPTED MANUSCRIPT

512 **6. Figures Legends**

513 Figure 1. Bacterial viability after incubation with or without 25  $\mu\text{M}$  of AsV or AsIII in saline solution  
514 for 72 h at 28°C. A) Representative dot plots [SYTO9 green fluorescence intensity (FL1-A) vs. PI red  
515 fluorescence intensity (FL3-A)] of the bacterial suspensions analyzed by flow cytometry. *B. japonicum*  
516 E109, *A. brasilense* Az39 and co-incubated strains (E109+Az39) in saline solution for 72 h (control),  
517 positive death control (Heat-killed), 25  $\mu\text{M}$  of AsV or AsIII. B) Bar graphs show the percentages of cell  
518 mortality obtained by flow cytometry and represent the mean  $\pm$  SE (n = 6). Different letters indicates  
519 significant differences (Tukey's test,  $p < 0.05$ ).

520 Figure 2. Effect of IAA on *B. japonicum* E109. Percentages of cell mortality incubated alone (E109),  
521 with commercial IAA (E109+IAA), with supernatant produced by *A. brasilense* Az39 (E109+SNT) or  
522 co-incubated with *A. brasilense* Az39 (E109+Az39) analyzed by flow cytometry. All samples were  
523 incubated with or without 25  $\mu\text{M}$  of AsV or AsIII in saline solution for 72 h at 28°C. Results represent  
524 the mean  $\pm$  SE (n = 5). Different letters indicates significant differences (Tukey's test,  $p < 0.05$ ).

525 Figure 3. Effect of As on root and shoot fresh weight of soybean plants non-inoculated (NI), inoculated  
526 with *B. japonicum* E109, *A. brasilense* Az39 or double inoculated (DI). The results represent the mean  $\pm$   
527 SE (n = 40). Different letters indicates significant differences (Test de Kruskal Wallis,  $p \leq 0,05$ ).

528 Figure 4. Effect of As on nodule number formed after inoculation with *B. japonicum* E109 or with *B.*  
529 *japonicum* E109 and *A. brasilense* Az39 (DI). The results represent the mean  $\pm$  SE (n = 40). Different  
530 letters indicate significant differences (Test de Tukey,  $p \leq 0,05$ ).

531 Figure 5. Nitrogen content in aerial parts of soybean plants non-inoculated (NI), inoculated with *B.*  
532 *japonicum* E109, *A. brasilense* Az39 or both strains (DI) treated with AsV and AsIII (25  $\mu\text{M}$ ). The  
533 results represent the mean  $\pm$  SE (n = 2). Asterisks represent significant difference with the  
534 corresponding NI plants under AsV and AsIII treatment.

535 Figure 6. Arsenic accumulation in roots or aerial parts of soybean plants non-inoculated (NI) or  
536 inoculated with *B. japonicum* E109, *A. brasilense* Az39 or with both strains (DI). The results represent  
537 the mean  $\pm$  SE (n = 3).

538

539 **References**

- 540 Abhilash, P.C., Jamil, S., Singh, N .(2009) Transgenic plants for enhanced biodegradation and  
541 phytoremediation of organic xenobiotics. *Biotechnol Adv* 27: 474-488.
- 542 Ahmed, M., Wardle, D.A. (1994) Allelopathic potential of vegetative and flowering ragwort (*Senecio*  
543 *jacobaeal* L.). *Plants against associated pasture species*. *Plant Soil* 64: 61-68.
- 544 AOAC (1990) *Official Methods of Analysis of Association of Official Analytical Chemists*.15th  
545 Edition. Arlington, Virginia (USA). Method 960.52.
- 546 Armendariz, A.L., Talano, M.A., Travaglia, C., Reinoso, H., Wevar Oller, A.L., Agostini, E. (2016)  
547 Arsenic toxicity in soybean seedlings and their attenuation mechanisms. *Plant Physiol Biochem*. 98:  
548 119-127.
- 549 Armendariz, A.L., Talano, M.A., Wevar Oller, A.L., Medina, M.I., Agostini, E. (2015) Effect of arsenic  
550 on tolerance mechanisms of two plant growth-promoting bacteria used as biological inoculants. *J.*  
551 *Environ. Sci. (China)*. 33: 203-210.
- 552 Barnard, A.M., Bowden, S.D., Burr, T., Coulthurst, S.J., Monson, R.E., Salmond, G.P. (2007) Quorum  
553 sensing, virulence and secondary metabolite production in plant soft-rotting bacteria. *Philos. Trans.*  
554 *R. Soc. Lond. B. Biol. Sci.* 362: 1165-1183.
- 555 Barrena, R., Casals, E., Colón, J. Font, X., Sánchez, A., Puentes, V. (2009) Evaluation of the ecotoxicity  
556 of model nanoparticles. *Chemosphere* 75(7): 850-857.
- 557 Benintende, S., Urich, W., Herrera, M., Gangge, F., Sterren, M., Benintende, M. (2010) Comparación  
558 entre coinoculación con *Bradyrhizobium japonicum* y *Azospirillum brasilense* e inoculación simple  
559 con *Bradyrhizobium japonicum* en la nodulación, crecimiento y acumulación de N en el cultivo de  
560 soja. *Agriscientia*. 2: 71-77.
- 561 Bianco, C., Imperlini, E., Calogero, R., Senatore, B., Amoresano, A., Carpentieri, A., Pucci, P., Defez,  
562 R. (2006a) Indole-3-acetic acid improves *Escherichia coli*'s defences to stress. *Arch. Microbiol.* 185:  
563 373-382.
- 564 Bianco, C., Imperlini, E., Calogero, R., Senatore, B., Pucci, P., Defez, R. (2006b) Indole-3-acetic acid  
565 regulates the central metabolic pathways in *Escherichia coli*. *Microbiol. Sgm* 152: 2421-2431.
- 566 Bundschuh, J., Litter, M.I., Nicoli, H.B., Hoinkis, J., Bhattacharya, P. (2010) Identifying occurrences of  
567 groundwater arsenic in Latin America: A continent wide problem and challenge. *Arsenic in*  
568 *Geosphere and Human Disease; Page Proceedings of Arsenic in the Environment Conference in*  
569 *Taiwan; London.UK, Tyler and Francis Group*. pp. 512-516.
- 570 Bustingorri, C., Lavado, R.S. (2014) Soybean as affected by high concentrations of arsenic and fluoride  
571 in irrigation water in controlled conditions. *Agric. Water Manage.* 144: 134-139.

- 572 Carrasco, J.A., Armario, P., Pajuelo, E., Burgos, A., Caviedes, M.A., López, R., Chamber, M.A.,  
573 Palomares, A.J. (2005) Isolation and characterization of symbiotically effective Rhizobium resistant  
574 to arsenic and heavy metals after the toxic spill at the Aznalcollar pyrite mine. *Soil Biol. Biochem.*  
575 37: 1131-1140.
- 576 Cassán, F., Perrig, D., Sgroy, V., Masciarelli, O., Penna, C., Luna, V. (2009) *Azospirillum brasilense*  
577 *Az39* and *Bradyrhizobium japonicum* E109, inoculated singly or in combination, promote seed  
578 germination and early seedling growth in corn (*Zea mays* L.) and soybean (*Glycine max* L.).  
579 *European J. Soil Biol.* 45: 28-35.
- 580 Dary, M., Chamber-Pérez, M.A., Palomares, A.J., Pajuelo, E. (2010) 'In situ' phytostabilisation of  
581 heavy metal polluted soils using *Lupinus luteus* inoculated with metal resistant plant-growth  
582 promoting rhizobacteria. *J. Hazard. Mat.* 177: 323-330.
- 583 Das, J., Sarkar, P. (2018) Remediation of arsenic in mungbean (*Vigna radiata*) with growth  
584 enhancement by unique arsenic-resistant bacterium *Acinetobacter lwoffii*. *Sci. Total Environ.* 624:  
585 1106-1118.
- 586 Das, S., Jean, J.S., Chou, M.L., Rathod, J., Liu, C.C. (2016) Arsenite-oxidizing bacteria exhibiting plant  
587 growth promoting traits isolated from the rhizosphere of *Oryza sativa* L.: Implications for mitigation  
588 of arsenic contamination in paddies. *J. Hazard. Mat.* 302: 10-18.
- 589 Egebo, L.A., Nielsen, S.V., Jochimsen, B.U. (1991) Oxygen-dependent catabolism of indole-3-acetic  
590 acid in *Bradyrhizobium japonicum*. *J. Bacteriol.* 173: 4897-4901.
- 591 El Aafi, N., Brhada, F., Dary, M., Filali Maltouf, A., Pajuelo, E. (2012) Rhizostabilization of metals in  
592 soils using *Lupinus luteus* inoculated with the metal resistant rhizobacterium *Serratia* sp. MSMC541,  
593 *Internat. J. Phytoremediation*, 14(3): 261-274.
- 594 Farooq, M.A., Islam, F., Ali, B., Najeeb, U., Mao, B., Gill, R. A., Zhou, W. (2016) Arsenic toxicity in  
595 plants: Cellular and molecular mechanisms of its transport and metabolism. *Environ. Exp. Bot.* 132:  
596 42-52.
- 597 Finnegan, P.M., Chen, W. (2012) Arsenic toxicity: the effects on plant metabolism. *Frontiers Physiol.* 3:  
598 1-18.
- 599 García, J.E., Maroniche, G., Creus, C., Suárez-Rodríguez, R., Ramirez-Trujillo, J.A., Groppa M.D.  
600 (2017) *In vitro* PGPR properties and osmotic tolerance of different *Azospirillum* native strains and  
601 their effects on growth of maize under drought stress. *Microbiol. Res.* 202: 21-29.
- 602 Garg N., Singla P. (2011). Arsenic toxicity in crop plants: physiological effects and tolerance  
603 mechanisms. *Environ. Chem. Lett.* 9, 303-321.
- 604 Glickman, E., Dessaux, Y. (1995) A critical examination of the specificity of the Salkowski reagent for  
605 indolic compounds produced by phytopathogenic bacteria. *Appl. Environ. Microbiol.* 61: 793-796.

- 606 Gomes M.A., Hauser-Davis R.A., de Souza A.N., Vitória A.P. (2017) Metal phytoremediation: General  
607 strategies, genetically modified plants and applications in metal nanoparticle contamination.  
608 Ecotoxicol. Environ. Saf. 134: 133-147.
- 609 Gong, P., Wilke, B.M., Fleischmann, S. (1999) Soil-based phytotoxicity of 2,4,6-trinitrotoluene to  
610 higher plants. Arch. Environ. Contam. Toxicol. 36: 152-157.
- 611 Gusmán, G.S., Oliveira, J.A., Farnese, F.S., Cambraia, J. (2013a) Arsenate and arsenite: the toxic  
612 effects on photosynthesis and growth of lettuce plants. Acta Physiol. Plant. 35: 1201-1209.
- 613 Gusman, G.S., Oliveira, J.A., Farnese, F.S., Cambraia, J. (2013b) Mineral nutrition and enzymatic  
614 adaptation induced by arsenate and arsenite exposure in lettuce plants. Plant Physiol. Biochem. 71:  
615 307-314.
- 616 Jensen, J.B., Egsgaard, H., Van Onckelen, H., Jochimsen, B.U. (1995) Catabolism of indole-3-acetic  
617 acid and 4- and 5-chloroindole-3-acetic acid in *Bradyrhizobium japonicum*. J. Bacteriol. 177: 5762-  
618 5766.
- 619 Jha, A.B., Dubey, R.S. (2004a) Carbohydrate metabolism in growing rice seedlings under arsenic  
620 toxicity. J. Plant Physiol., 161: 867-872.
- 621 Jha, A.B., Dubey, R.S. (2004b) Arsenic exposure alters activity behaviour of key nitrogen assimilatory  
622 enzymes in growing rice plants. Plant Growth Regul. 43: 259-268.
- 623 Jha, A.B., Dubey, R.S. (2005) Effect of arsenic on behaviour of enzymes of sugar metabolism in  
624 germinating rice seeds. Acta Physiologiae Plant. 27(3): 341-347.
- 625 Joshi, P.M., Juwarkar, A.A. (2009) *In vivo* studies to elucidate the role of extracellular polymeric  
626 substances from *Azotobacter* in immobilization of heavy metals. Environ. Sci. Technol. 43: 5884-  
627 5889.
- 628 Kaur, S., Singh, H.P., Batish, D.R., Negi, A., Mahajan, P., Rana, S., Kohli, R.K. (2012) Arsenic (As)  
629 inhibits radicle emergence and elongation in *Phaseolus aureus* by altering starch metabolizing  
630 enzymes vis-à-vis disruption of oxidative metabolism. Biol. Trace Elem. Res. 146: 360-368.
- 631 Kaushik, R., Saxena, A.K., Tilak, K.V.B.R. (2000) Selection of Tn5:lacZ mutants isogenic to wild type  
632 *Azospirillum brasilense* strains capable of growing at sub-optimal temperature. World J. Microbiol.  
633 Biotechnol. 16: 567-570.
- 634 Kumar Yadava, K., Gupta, N., Kumar, A., Reece, L.M., Singh, N., Rezaia, S., Ahmad Khan, S. (2018)  
635 Mechanistic understanding and holistic approach of phytoremediation: A review on application and  
636 future prospects. Ecological Engineering 120: 274-298.
- 637 Kundu, S., Trent, J.T., Hargrove, M.S. (2003) Plants, humans and hemoglobins. Trends Plant. Sci. 8:  
638 387-393.

- 639 Lafuente, A., Pajuelo, E., Caviedes, M.A., Rodríguez-Llorente, I.D. (2010) Reduced nodulation in  
640 alfalfa induced by arsenic correlates with altered expression of early nodulins. *J. Plant Physiol.* 167:  
641 286-291.
- 642 Lafuente, A., Pérez-Palacios, P., Doukkali, B., Molina-Sánchez, M.D., Jiménez-Zurdo, J.I., Caviedes,  
643 M.A., Rodríguez-Llorente, I.D., Pajuelo, E. (2015) Unraveling the effect of arsenic on the model  
644 *Medicago-Ensifer* interaction: a transcriptomic meta-analysis. *New Phytol.* 205(1): 255-272.
- 645 Liu, X., Zhang, S., Shan, X., Zhu, Y.G. (2005) Toxicity of arsenate and arsenite on germination,  
646 seedling growth and amylolytic activity of wheat. *Chemosphere* 61: 293-301.
- 647 Lyubun, Y.V., Fritzsche, A., Chernyshova, M.P., GertDudel, E., Fedorov, E.E. (2006) Arsenic  
648 transformation by *Azospirillum brasilense* Sp245 in association with wheat (*Triticum aestivum* L.)  
649 roots. *Plant Soil* 286: 219-227.
- 650 Ma, Y., Rajkumar, M., Zhang, C., Freitas, H. (2016) Beneficial role of bacterial endophytes in heavy  
651 metal phytoremediation. *J. Environ. Manag.* 174: 14-25.
- 652 Mahar A., Wang P., Ali A., Awasthi M.K., Lahori A.H., Wang Q., Li R., Zhang Z. (2016) Challenges  
653 and opportunities in the phytoremediation of heavy metals contaminated soils: a review *Ecotoxicol.*  
654 *Environ. Saf.* 126: 111-121.
- 655 Mandal, S.M., Gouri, S.S., De, D., Das, D.K., Mondal, K.C., Pati, B.R. (2011) Effect of arsenic on  
656 nodulation and nitrogen fixation of blackgram (*Vigna mungo*). *Indian J. Microbiol.* 51: 44-47.
- 657 Mandal, S.M., Pati, B.R., Das, A.K., Ghosh, A.K. (2008) Characterization of a symbiotically effective  
658 *Rhizobium* resistant to arsenic: Isolated from the root nodules of *Vigna mungo* (L.) Hepper grown in  
659 an arsenic-contaminated field. *J. Gen. Appl. Microbiol.* 54(2): 93-99.
- 660 Masciarelli, O., Llanes, A., Luna, V. (2014) A new PGPR co-inoculated with *Bradyrhizobium*  
661 *japonicum* enhances soybean nodulation. *Microbiol. Res.* 169(7-8): 609-615.
- 662 Maynaud, G., Brunel, B., Mornico, D., Durot, M., Severac, D., Dubois, E., Navarro, E., Cleyet-Marel,  
663 J.C., Le Quéré, A. (2013) Genome-wide transcriptional responses of two metal-tolerant symbiotic  
664 *Mesorhizobium* isolates to Zinc and Cadmium exposure. *BMC Genomics* 14: 292.
- 665 Mench, M., Vangronsveld, J., Beckx, C., Ruttens, A. (2006) Progress in assisted natural remediation of  
666 an arsenic contaminated agricultural soil. *Environ. Pollut.* 144: 51-61.
- 667 Méndez, M.O., Maier, R.M. (2008) Phytostabilisation of mine tailings in arid and semiarid  
668 environments: an emerging remediation technology. *Environ. Health Perspectives* 116: 278-283.
- 669 Nie, L., Shah, S., Burd, G.I., Dixon, D.G., Glick, B.R. (2002) Phytoremediation of arsenate  
670 contaminated soil by transgenic canola and the plant growth promoting bacterium *Enterobacter*  
671 *cloacae* CAL2. *Plant Physiol. Biochem.* 40: 355-361.

- 672 Ojuederie, O.B., Babalola, O.O. (2017) Microbial and plant-assisted bioremediation of heavy metal  
673 polluted environments: A Review Int. J. Environ. Res. Public Health 14: 1504.
- 674 Pajuelo, E., Carrasco, J.A., Romero, L.C., Chamber, M.A., Gotor, C. (2007) Evaluation of the metal  
675 phytoextraction potential of crop legumes. Regulation of the expression of o-acetylserine (thiol)  
676 lyase under metal stress. Plant Biol. 9: 672-681.
- 677 Pajuelo, E., Rodriguez-Llorente, I.D., Dary, M., Palomares, A.J. (2008) Toxic effects of arsenic on  
678 Sinorhizobium-Medicago sativa symbiotic interaction. Environ. Pollut. 154: 2003-2011.
- 679 Pajuelo, E., Rodríguez-Llorente, I.D., Lafuente, A., Caviedes, M.A. (2011) Legume-rhizobium  
680 symbiosis as a tool for bioremediation of heavy metal polluted soils. In: Khan, M.S., Zaidi, A., Goel,  
681 R., Musarrat, J. (eds.) Biomangement of metal contaminated soils. Environmental Pollution,  
682 Heidelberg, Germany: Springer, 20: 95-123.
- 683 Pommeresche and Hansen, (2017) Examining root nodule activity on legumes. FertilCrop Technical  
684 Note 1-4.
- 685 Rajkumar, M., Sandhya, S., Prasad, M.N.V., Freitas, H. (2012) Perspectives of plant-associated  
686 microbes in heavy metal phytoremediation. Biotechnol. Adv. 30: 1562-1574.
- 687 Reichman, S.M. (2007) The potential use of the legume-rhizobium symbiosis for the remediation of  
688 arsenic contaminated sites. Soil Biol. Biochem. 39: 2587-2593.
- 689 Reichman, S.M. (2014) Probing the plant growth-promoting and heavy metal tolerance characteristics  
690 of *Bradyrhizobium japonicum* CB1809. Eur. J. Soil Biol. 63: 7-13.
- 691 Rivera, D., Revale, S., Molina, R., Gualpa, J., Puente, M., Maroniche, G., Paris, G., Baker, D., Clavijo,  
692 B., McLay, K., Spaepen, S., Peticari, A., Vazquez, M., Wisniewski-Dyé, F., Watkins, C., Martínez-  
693 Abarca, F., Vanderleyden, J., Cassán, F. (2014) Complete genome sequence of the model  
694 rhizosphere strain *Azospirillum brasilense* Az39, successfully applied in agriculture. Genome  
695 Announc. 2(4):e00683-14.
- 696 Robinson, B.H., Banuelos, G., Conesa, H.M., Evangelou, M.W., Schulin, R. (2009) The  
697 phytomanagement of trace elements in soil. Crit. Rev. Plant Sci. 28: 240-266.
- 698 Sarwar, N., Imran, M., Shaheen, M.R., Ishaque, W., Kamran, M.A., Matloob, A., Rehim, A., Hussain,  
699 S. (2017) Phytoremediation strategies for soils contaminated with heavy metals: Modifications and  
700 future perspectives. Chemosphere. 171: 710-721.
- 701 Singh, N., Marwaa, N., Mishra, S.K., Mishra, J., Verma, P.C., Rathaur, S., Singh, N. (2016)  
702 *Brevundimonas diminuta* mediated alleviation of arsenic toxicity and plant growth promotion in  
703 *Oryza sativa* L. Ecotoxicol. Environ. Saf. 125: 25-34.
- 704 Smedley, P.L., Kinniburgh, D.G. (2002) A review of the source, behaviour and distribution of arsenic in  
705 natural waters. Appl. Geochem. 17: 517-568.

- 706 Spaepen, S., Vanderleyden, J., Remans, R. (2007) Indole-3-acetic acid in microbial and microorganism-  
707 plant signaling. *FEMS Microbiol. Rev.* 31: 425-448.
- 708 Srivastava, S., Verma, P.C., Chaudhry, V., Singh, N., Abhilash, P.C., Kumar, K.V., Sharma, N., Singh,  
709 N. (2013) Influence of inoculation of arsenic-resistant *Staphylococcus arlettae* on growth and arsenic  
710 uptake in *Brassica juncea* (L.) Czern. Var. R-46
- 711 Stoeva, N., Berova, M., Zlatev, Z. (2004) Physiological response of maize to arsenic contamination.  
712 *Biol. Plant.* 47: 449-452.
- 713 Sytnikov DM (2013) How to increase the productivity of the soybean rhizobial symbiosis. In: A  
714 comprehensive survey of international soybean research-Genetics, Physiology, Agronomy and  
715 Nitrogen Relationships, Board J. (Ed) Chapter 4: pp 61-82.
- 716 Talano, M.A., Cejas, R.B., González, P.S., Agostini, E. (2013) Effect of sodium arsenate and arsenite on  
717 soybean development and in the symbiotic interaction with *Bradyrhizobium japonicum*. *Plant*  
718 *Physiol. Biochem.* 63: 8-14.
- 719 Tejerizo, G.T., Bañuelos, L.A., Cervantes, L., Gaytán, P., Pistorio, M., Romero, D., Brom, S. (2015)  
720 Development of molecular tools to monitor conjugative transfer in rhizobia. *J. Microbiol. Methods.*  
721 117: 155-163.
- 722 Titaha, H.S., Abdullaha, S.R.S., Mushrifah, I., Anuar, N., Basri, H., Mukhlisin, M. (2013) Effect of  
723 applying rhizobacteria and fertilizer on the growth of *Ludwigia octovalvis* for arsenic uptake and  
724 accumulation in phytoremediation. *Ecological Engineering* 58: 303-313.
- 725 Torres, D., Revale, S., Obando, M., Maroniche, G., Paris, G., Peticari, A., Vazquez, M., Wisniewski-  
726 Dyé, F., Martínez-Abarca, F., Cassán, F. (2015) Genome sequence of *Bradyrhizobium japonicum*  
727 E109, one of the most agronomically used nitrogen-fixing rhizobacteria in Argentina. *Genome*  
728 *Announc* 3(1):e01566-14.
- 729 Ullah, A., Heng, Sun, HussainMunis, M.F., Fahad, S., Yang, X. (2015) Phytoremediation of heavy  
730 metals assisted by plant growth promoting (PGP) bacteria: A review. *Environ. Exp. Bot.* 117: 28-40.
- 731 Valdameri, G., Kokot, T.B., Pedrosa F. de O., de Souza, E.M. (2015) Rapid quantification of rice root-  
732 associated bacteria by flow cytometry. *Lett. Appl. Microbiol.* 60(3):237-241.
- 733 Wang, E., Martínez-Romero, E. (2000) *Sesbania herbacea*-*Rhizobium huautlense* nodulation in flooded  
734 soils and comparative characterization of *S. herbacea*-nodulating rhizobia in different environments.  
735 *Microb Ecol* 40: 25-32.
- 736 Zubair, M., Shakir, M., Ali, Q., Rani, N., Fatima, N., Farooq, S., Nasir, I.A. (2016) Rhizobacteria and  
737 phytoremediation of heavy metals. *Environ. Tech. Rev.* 5: 112-119.

**Table 1.** IAA production by *A. brasilense* Az39 incubated in saline solution for 72 h at 28°C under As treatment. Positive control: *Azospirillum brasilense* Cd. Results represent the mean  $\pm$  SE (n = 8).

	IAA production ( $\mu$ M)	
	Az39	AzCd
Control	3.9 $\pm$ 1.1	4.5 $\pm$ 0.3
AsV	4.1 $\pm$ 0.5	4.5 $\pm$ 0.7
AsIII	5.6 $\pm$ 1.1	5.7 $\pm$ 1.1

**Table 2.** Germination parameters of soybean seedlings treated with 25  $\mu$ M AsV and AsIII. Effects of inoculation with *B. japonicum* E109, *A. brasilense* Az39 and double inoculation.

	<i>Treatment</i>	<i>Germination index (IG)</i>	<i>Speed of germination index (S)</i>	<i>Radical length (cm)</i>	<i>Radical relative elongation (E)</i>
NI	Control	100.0 $\pm$ 0.0 <sup>b</sup>	93.8 $\pm$ 2.4 <sup>a</sup>	8.3 $\pm$ 0.4 <sup>b</sup>	100.0
E109	Control	110.9 $\pm$ 1.9 <sup>ab</sup>	94.0 $\pm$ 1.7 <sup>a</sup>	9.4 $\pm$ 0.5 <sup>ab</sup>	113.0
Az39	Control	109.3 $\pm$ 5.7 <sup>ab</sup>	93.0 $\pm$ 2.3 <sup>a</sup>	9.5 $\pm$ 0.4 <sup>ab</sup>	115.0
E109+Az39	Control	119.1 $\pm$ 2.6 <sup>a</sup>	91.0 $\pm$ 3.3 <sup>ab</sup>	10.2 $\pm$ 0.5 <sup>a</sup>	123.0
NI	AsV	46.2 $\pm$ 1.7 <sup>de</sup>	89.7 $\pm$ 4.8 <sup>ab</sup>	4.1 $\pm$ 0.2 <sup>de</sup>	49.5
E109	AsV	60.7 $\pm$ 1.4 <sup>c</sup>	89.6 $\pm$ 3.3 <sup>ab</sup>	5.9 $\pm$ 0.3 <sup>c</sup>	62.9
Az39	AsV	53.3 $\pm$ 5.9 <sup>cde</sup>	90.6 $\pm$ 5.1 <sup>ab</sup>	5.1 $\pm$ 0.3 <sup>cde</sup>	53.9
E109+Az39	AsV	55.2 $\pm$ 2.7 <sup>c</sup>	86.3 $\pm$ 6.0 <sup>ab</sup>	5.9 $\pm$ 0.4 <sup>c</sup>	58.0
NI	AsIII	45.8 $\pm$ 1.0 <sup>de</sup>	71.5 $\pm$ 1.7 <sup>bcd</sup>	3.9 $\pm$ 0.2 <sup>e</sup>	47.4
E109	AsIII	60.9 $\pm$ 1.7 <sup>c</sup>	77.6 $\pm$ 2.6 <sup>abc</sup>	5.7 $\pm$ 0.3 <sup>cd</sup>	60.8
Az39	AsIII	39.7 $\pm$ 2.5 <sup>e</sup>	55.7 $\pm$ 3.2 <sup>cd</sup>	3.7 $\pm$ 0.3 <sup>e</sup>	39.1
E109+Az39	AsIII	50.6 $\pm$ 1.4 <sup>cde</sup>	64.0 $\pm$ 3.1 <sup>d</sup>	5.2 $\pm$ 0.4 <sup>c</sup>	50.5

NI: non-inoculated seeds.

Fig 1

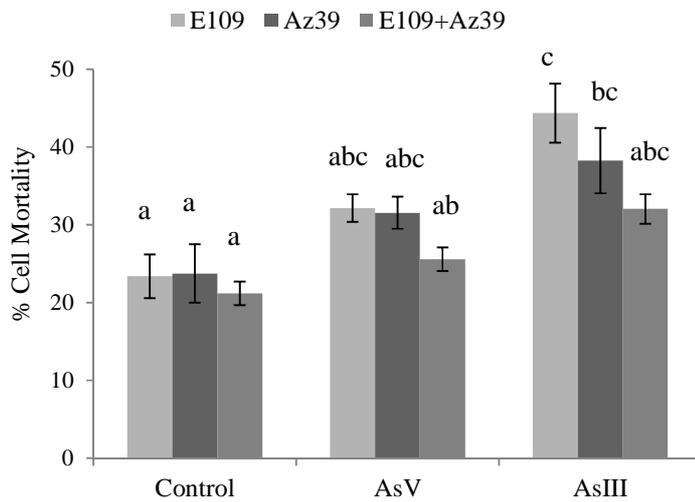
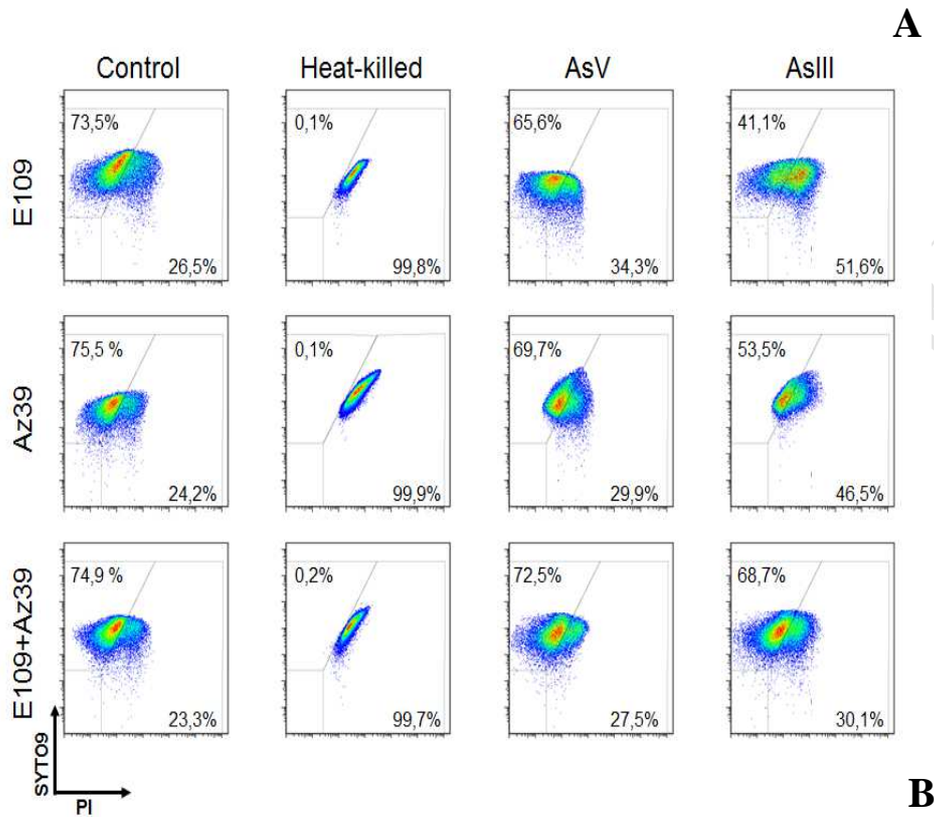


Fig 2

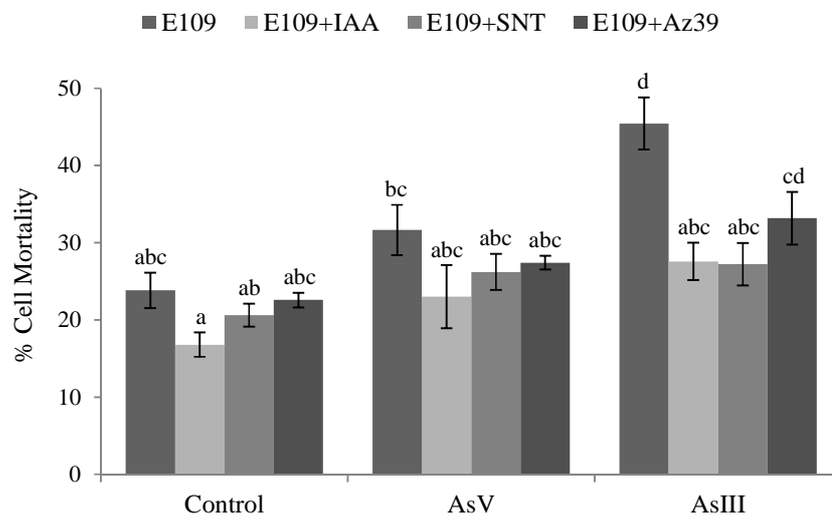
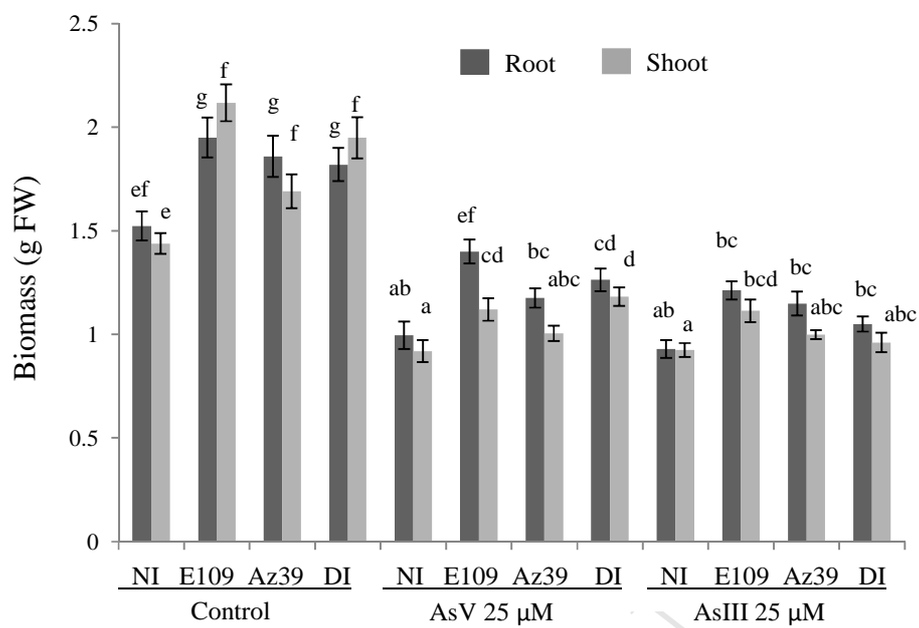


Fig 3



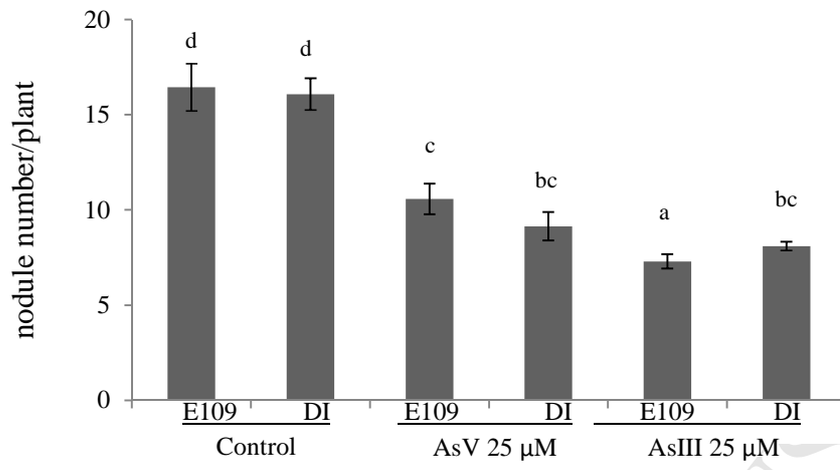
**Fig 4**

Fig 5

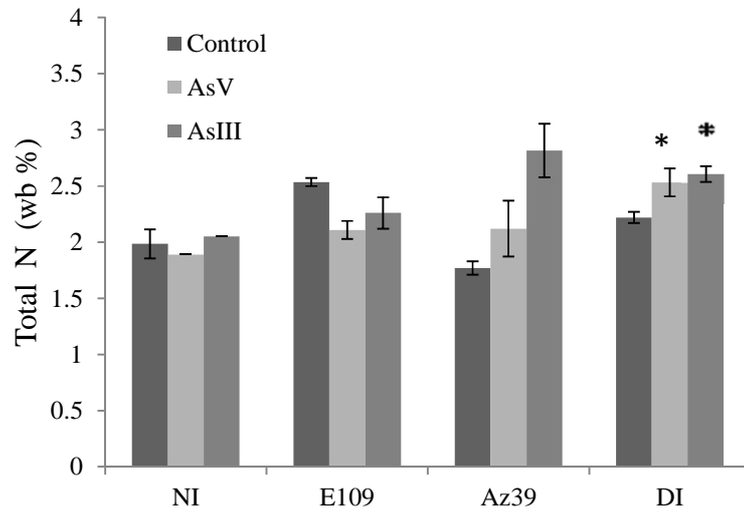
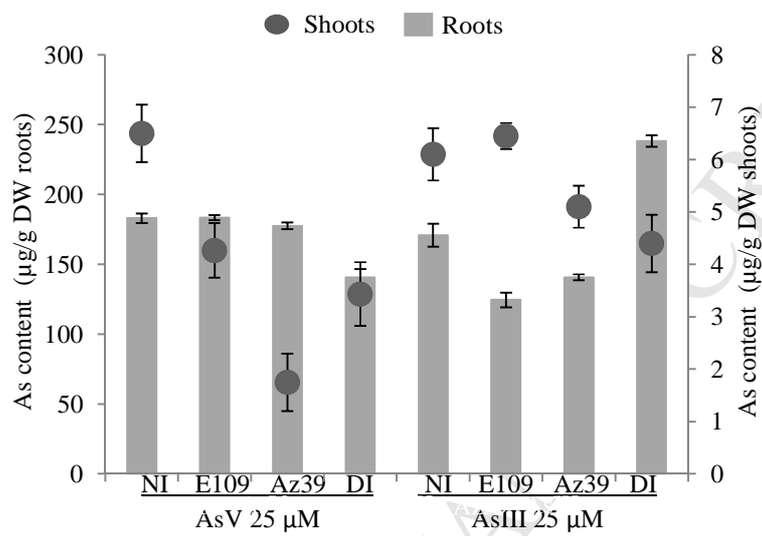


Fig 6



**Highlights**

-Flow cytometry revealed synergism between two rhizospheric bacteria when exposed to As.

-Indole acetic acid produced by *A. brasilense* Az39 would protect *B. japonicum* E109 when exposed to As.

-Plant growth improvement, increase of nodule number and N content was observed in double inoculated plants treated with As.

-Double inoculation strategy promoted As phytostabilization potential of soybean plants.

**Contributions**

MAT and EA conceived and planned the experiments. ALA y MAT carried out the inoculation experiments, MFON and MLB carried out cytometry assays and LE made the arsenic quantification. ALA, MAT, MFON, MLB, CP and EA contributed to the interpretation of the results. MAT wrote the manuscript with input from all authors. EA, LE and CP provided critical feedback and helped shape the research and manuscript analysis.