

1 **Inorganic arsenic exposure and neuropsychological development of children of 4-5**
2 **years of age living in Spain**

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34

35 **Abstract**

36 Early-life exposure to inorganic arsenic (iAs) may adversely impact health later in life. To
37 date, evidence of iAs adverse effects on children’s neurodevelopment comes mainly from
38 populations highly exposed to contaminated water with conflicting results. Little is known
39 about those effects among populations with low iAs exposure from food intake. We
40 investigated the cross-sectional association between exposure to iAs and
41 neurodevelopment scores among children living in Spain whose main route of exposure
42 was diet. Arsenic species concentrations in urine from 400 children was determined, and
43 the sum of urinary iAs, dimethylarsinic acid, and monomethylarsonic acid was used to
44 estimate iAs exposure. The McCarthy Scales of Children’s Abilities was used to assess
45 children’s neuropsychological development at about 4-5 years of age. The median
46 (interquartile range) of children’s sum of urinary iAs, MMA, and DMA was 4.85 (2.74 -
47 7.54) µg/L, and in adjusted linear regression analyses the natural logarithm transformed
48 concentrations showed an inverse association with children’s motor functions (β , [95%
49 confidence interval]; global scores (-2.29, [-3.95, -0.63]), gross scores (-1.92, [-3.52, -
50 0.31]) and fine scores (-1.54, [-3.06, -0.03]). In stratified analyses by sex, negative
51 associations were observed with the scores in the quantitative index (-2.59, [-5.36, 0.17])
52 and working memory function (-2.56, [-5.36, 0.24]) only in boys. Our study suggests that
53 relatively low iAs exposure may impair children’s neuropsychological development and
54 that sex-related differences may be present in susceptibility to iAs related effects; however,
55 our findings should be interpreted with caution given the possibility of residual
56 confounding.

57

58 **Keywords:** urinary arsenic species, inorganic arsenic, children, neuropsychological
59 development, dietary arsenic, developmental toxicology, neurodevelopment, environment,
60 and McCarthy Scales of Children’s Abilities.

61

62 **1. Introduction**

63 Arsenic is a ubiquitous element in the environment that occurs in different oxidation states
64 (-3, 0, +3, +5) in both organic and inorganic forms that constitute total arsenic (referred to
65 as “arsenic” in this study) (WHO, 2001). Intake of inorganic arsenic (iAs), including
66 arsenite (As^{III}) and arsenate (As^{V}), is an established cause of cancer of the lung, skin, and
67 bladder and a possible cause of others, with accumulating evidence of effects on non-cancer
68 health outcomes such as neurological, cardiovascular, respiratory and metabolic diseases
69 (IARC, 2012; Nachman et al., 2017; Sanchez et al., 2016; Tsuji et al., 2015). The
70 metabolism of iAs involves a series of reduction and oxidative methylation processes
71 catalyzed by the enzyme arsenic-methyltransferase with S-adenosylmethionine as the
72 methyl group donor that results in the formation of the pentavalent monomethylarsonic
73 acid (MMA) and dimethylarsinic acid (DMA) that are primarily excreted in the urine
74 (Antonelli et al., 2014; Jansen et al., 2016; Tseng, 2009). The trivalent forms of iAs, MMA,

75 and DMA are considered to be more toxic forms with MMA^{III} having the highest toxicity
76 followed by iAs^{III} (Tseng, 2009). Direct ingestion of DMA and MMA in the pentavalent
77 form may be excreted in the urine unchanged potentially posing less toxic effects (Buchet
78 et al., 1981; Cohen et al., 2006; Meharg et al., 2014; Molin et al., 2015; Tseng, 2009). The
79 sum of urinary iAs and methylated arsenic species concentrations (i.e. MMA and DMA) is
80 considered a reliable biomarker of short-term exposure to iAs from all sources, and it also
81 appears to be a reliable source of long-term exposure among individuals with consistent
82 patterns of exposure such as child populations whose diet is generally of lower food
83 diversity (EFSA, 2009; Kile et al., 2009; Marchiset-Ferlay et al., 2012; Navas-Acien et al.,
84 2009; Signes-Pastor et al., 2017b). Oxidative stress is considered to be a potential
85 mechanism of iAs toxicity, and increasing evidence suggests that this mechanism may be
86 responsible for iAs related neurotoxicity and impaired neurodevelopment (Grandjean and
87 Landrigan, 2014; Tolins et al., 2014).

88
89 A growing number of epidemiologic studies suggest that children's iAs exposure adversely
90 impacts health later in life, including neurodevelopment (EFSA, 2009; Freire et al., 2018;
91 Grandjean and Landrigan, 2014; Nachman et al., 2017; Tolins et al., 2014; Tsuji et al.,
92 2015; Wasserman et al., 2014); however, the consistency and generalizability of these
93 findings has not been established yet, especially among populations whose main exposure
94 source is diet. This includes the Spanish population for whom ingested iAs and organic
95 arsenic is likely to be associated with rice and marine product consumption, respectively
96 (Cubadda et al., 2016; EFSA, 2009; Kurzius-Spencer et al., 2014, 2013; Navas-Acien et
97 al., 2009; Signes-Pastor et al., 2017b). Among populations whose main exposure route to
98 iAs is from food intake, consumption of fish/seafood products needs to be carefully taken
99 into account. These contain high concentrations of arsenobetaine (AsB), a putative non-
100 toxic organic form excreted in urine unchanged, which may cause exposure
101 misclassification of iAs if total urinary arsenic is used as a biomarker of exposure (Forns
102 et al., 2014; Molin et al., 2015; Navas-Acien et al., 2011). Biotransformation of other
103 fish/seafood organosensicals excreted in urine as DMA or direct ingestion of DMA or MMA
104 similarly can be problematic in the assessment of iAs intake (Jones et al., 2016; Meharg et
105 al., 2014; Molin et al., 2015). Currently there is a lack of information regarding the
106 association between early-life neuropsychological development and iAs exposure based on
107 urinary arsenic speciation among populations with access to arsenic drinking water lower
108 than the WHO guideline value of 10 µg/L (Forns et al., 2014; WHO, 2011). In water,
109 arsenic is mostly present as iAs, and relatively low levels of arsenic drinking water have
110 been negatively associated with school-age children's full intelligence quotient (IQ) in the
111 U.S. (Wasserman et al., 2014; WHO, 2011). In water arsenic-contaminated areas of
112 Bangladesh, India and Mexico inverse associations were reported between iAs exposure,
113 assessed using arsenic concentrations in water, urine and blood, and children's cognitive
114 function (Hamadani et al., 2011; Mst. Nasrin Nahar et al., 2014; Parvez et al., 2011; Rosado

115 et al., 2007; Wasserman et al., 2011; WHO, 2011). However, other studies in Bangladesh
116 focused on areas with arsenic-contaminated water have not found evidence of child
117 neuropsychological development in relation to urinary arsenic (Hamadani et al., 2010;
118 Tofail et al., 2009). Further, although a few studies have suggested sex-related differences
119 in iAs-associated neurodevelopmental outcomes, this has not always been observed, and
120 thus further investigations are needed (Hamadani et al., 2011; Llop et al., 2013; Rosado et
121 al., 2007; Sanchez et al., 2016).

122

123 In populations with access to low arsenic drinking water, i.e. $< 10 \mu\text{g/L}$, food is considered
124 to be the major source of iAs exposure (Cubadda et al., 2016; EFSA, 2009; Kurzius-
125 Spencer et al., 2014, 2013), and yet little is known regarding the potential association
126 between dietary iAs exposure and childhood neuropsychological development. In this
127 study, we investigated whether early-life exposure to dietary iAs levels adversely affects
128 children's neuropsychological development. We focused on a population of children of
129 approximately 4-5 years of age living in Spain for whom diet is expected to be the major
130 iAs exposure source (Signes-Pastor et al., 2017b, 2017a). We further explored the
131 possibility of sex-related differences in susceptibility to iAs related neuropsychological
132 outcomes.

133

134 **2. Material and methods**

135

136 **2.1. Study population.** The study population was derived from the mother-child pair
137 participants in the INMA – *Infancia y Medio Ambiente* - Environment and Childhood
138 project, a prospective population-based birth cohort study conducted in multiple regions
139 around Spain (www.proyectoinma.org). The general design of INMA has been previously
140 described in detail (Guxens et al., 2012). Briefly, women participants of the INMA project
141 were recruited at the beginning of their pregnancy (2003 - 2006) at their reference primary
142 health care centers or public hospitals and were followed-up until delivery ($n = 2,625$). All
143 women met the inclusion criteria of ≥ 16 years old, singleton pregnancy, non-assisted
144 conception and delivery scheduled at the reference hospital. Their children were enrolled
145 at birth and were followed-up during infancy and childhood. Informed consent was
146 obtained from all participants in each phase, and the hospitals ethics committees in the
147 participating regions approved the study. For the present study, 100 children, evenly
148 distributed between boys and girls, were randomly selected to provide a urine sample from
149 each sub-cohort located in the Spanish regions of Asturias, Gipuzkoa, Sabadell, and
150 Valencia (overall $n = 400$) (Signes-Pastor et al., 2017b, 2017a). To date a total of 400
151 INMA 4-year-old children have had urinary arsenic species concentrations analyzed.

152

153 **2.2. Neuropsychological assessment.** Overall, children's neuropsychological
154 development was assessed at the median age of 4.5 years (standard deviation of 0.6 years)

155 with a standardized version of the McCarthy Scales of Children's Abilities (MSCA)
156 adapted to the Spanish population (McCarthy, 2009). The MSCA was selected because of
157 its reliability and validity, and wide use in research related to environmental health and
158 neurodevelopment including prior studies by INMA (Andiarena et al., 2017; Forns et al.,
159 2012; Nagle, 1979). For children from the sub-cohorts of Asturias, Gipuzkoa and Sabadell
160 ($n = 300$) the MSCA test was performed at the same time urine samples were collected,
161 along with children's weight and height measured and a food frequency questionnaire
162 (FFQ) at a median age of 4.4 years (standard deviation of 0.2 years); for the Valencian
163 children ($n = 100$) the neuropsychological assessment was carried out at the median age of
164 5.8 years (standard deviation of 0.1 years). The urine samples, children's weight and height,
165 and the FFQ for the Valencian children were collected at a median age of 4.4 years
166 (standard deviation of 0.1 years). Trained psychologists administered the MSCA test. The
167 MSCA test included a battery of 18 subtests (i.e. construction with cubes, puzzle, pictorial
168 memory, vocabulary, calculation, beating sequence, verbal memory, right-left orientation,
169 leg coordination, arm coordination, imitative action, copying of drawings, drawing of a
170 child, numerical memory, verbal fluency, counting and distribution, opposites, and concept
171 formation). The MSCA subtests were grouped into the original function scales of general
172 cognitive, verbal, perceptive-performance, quantitative index, memory, and motor
173 function. With further classification of the MSCA subtests, we obtained the new function
174 scales of executive, working memory, visual and verbal span, verbal memory, gross motor,
175 fine motor, and cognitive function of the posterior cortex as described in detail previously
176 (Julvez et al., 2011). We previously calculated and reported high intraclass coefficients for
177 the original function scales (> 0.78), and reasonably high Cronbach's alpha coefficients (\geq
178 0.70) with the new function scales. Further details appear in the prior INMA publication
179 (Valera-Gran et al., 2017).

180

181 **2.3. Sample preparation and chemical analyses.** Arsenic speciation analyses were
182 carried out in spot urine samples (Signes-Pastor et al., 2017a). Urine samples were
183 collected in 100 mL polypropylene containers and immediately stored at or below -20°C ,
184 then a 5 mL aliquot from each child in the study were shipped on dry ice to the Institute
185 for Global Food Security at Queen's University Belfast (QUB), Northern Ireland, for
186 arsenic speciation analyses including AsB, DMA, MMA, and iAs. Before speciation, urine
187 samples were centrifuged, and analytical grade hydrogen peroxide was added to convert
188 any arsenite to arsenate to facilitate subsequent chromatographic detection by ion
189 chromatography (IC) with inductively coupled plasma mass spectrometry (ICP-MS). In
190 each analytic batch, blank and replicate samples of the urine lyophilized material
191 ClinChek[®] - Control level I (Recipe Chemicals + Instruments GmbH in Munich, Germany)
192 were included for quality control. Urine samples were normalized for urine dilution using
193 specific gravity measured with a clinical refractometer. The average recovery percentages
194 and standard deviations of the arsenic species based on several replicate samples of the

195 urine lyophilized material ClinChek[®]- Control level I ($n = 33$) were $115 \pm 2\%$ for i-As, 97
196 $\pm 2\%$ for MMA, $94 \pm 2\%$ for DMA, and $90 \pm 2\%$ for AsB. The mean and range
197 concentrations of the arsenic species reference values in the urine lyophilized material
198 ClinChek[®] - Control level I are as follows: 4.55 (2.73 - 6.37) $\mu\text{g/L}$ for i-As, 2.50 (1.50 -
199 3.50) $\mu\text{g/L}$ for MMA, 9.8 (5.88 - 13.7) $\mu\text{g/L}$ for DMA, and 16.8 (12.6 - 21.0) $\mu\text{g/L}$ for AsB.
200 The limit of detection (LOD) for arsenic speciation, calculated from DMA calibration, was
201 0.011 $\mu\text{g/L}$ (Signes-Pastor et al., 2017a).

202

203 **2.4. Questionnaire.** In the 1st trimester of pregnancy a maternal questionnaire was
204 administered to gather information regarding parental sociodemographic and
205 socioeconomic characteristics such as the number of previous live births (i.e. 0, 1, 2, or 3),
206 maternal age at conception (years), maternal highest attained level of education (i.e.
207 primary, secondary, or university), and social class according to the International Standard
208 Classification of Occupants (ISCO88) (i.e. upper - I+II, middle - III, or lower - IV+V)
209 (International Labor Office (ILO), 2012). Trained staff measured children's weight (kg)
210 and height (m) at the same time the urine samples were collected following standard
211 protocols to calculate the body mass index (BMI) in kg/m^2 . At the same time, parents
212 reported children's diet including consumption of rice and fish/seafood with a validated
213 FFQ (Signes-Pastor et al., 2017b; Vioque et al., 2016). All the aforementioned covariates
214 were among those considered while identifying potential confounders (see Supplemental
215 Material, **Figure S1**, for further details)

216

217 **2.5. Statistical analyses.** For all statistical analyses, observations with missing data for at
218 least one covariate were excluded in addition to children who did not complete the
219 neuropsychological development test. Summary statistics were calculated for each
220 variable: median (range and interquartile range) for continuous variables and n (%) of each
221 level for categorical variables. We calculated the sum of iAs (i.e. arsenite and arsenate),
222 DMA and MMA (referred to as "sum of urinary arsenic" in this study) as a biomarker of
223 iAs exposure. The distribution of children's urinary arsenic species concentrations and sum
224 of urinary arsenic were right skewed, so they were natural logarithm transformed (ln-
225 transformed) before statistical analysis. All scores from the neuropsychological MSCA
226 function scales were standardized to a mean of 100 points with a standard deviation of 15.

227

228 The association between children's sum of urinary arsenic concentrations ln-transformed
229 (continuous) and neuropsychological function scores was firstly assessed using univariate
230 linear regression models (Model 0 in Supplemental Material, **Table S1**). Then, multiple
231 linear regression models adjusted for potential confounders were computed (Model 1 in
232 **Table 3** and in Supplemental Material, **Table S1**). The potential confounders were
233 identified using the directed acyclic graph (Textor et al., 2017), and the selected minimally
234 sufficient adjustment set contained: maternal highest attained level of education (i.e.

235 primary, secondary, or university), child's sex (i.e. girls or boys), BMI (continuous), age
236 at MSCA testing (continuous) and calorie adjusted consumption of rice and fish/seafood
237 (continuous) (Supplemental Material, **Figure S1**). The adjusted models were also used to
238 explore the association between children's sum of urinary arsenic concentrations and the
239 neuropsychological scores according to sex in stratified analysis and by including the main
240 effects along with the interaction term (i.e. ln-transformed sum of urinary arsenic
241 concentrations * sex). We carried out multiple sensitivity analyses in the models: i)
242 children's sum of urinary arsenic concentrations were calibrated for fish/seafood
243 consumption using a mathematical method previously described that uses AsB as a
244 biomarker of fish/seafood intake (Model 2 in Supplemental Material, **Table S1**) (Jones et
245 al., 2016), ii) influential points identified with the Bonferroni outlier test of the "car"
246 package were excluded (Fox and Weisberg, 2011), iii) children's hair mercury
247 concentrations analyzed at 4 years were added in the core models as potential confounder
248 (Model 3 in Supplemental Material, **Table S1**), iv) analysis restricted to children with low
249 urinary AsB (i.e. < 1 µg/L) as an indicator of exclusion of fish/seafood consumption (Model
250 4 in Supplemental Material, **Table S1**) (Jones et al., 2016), vi) and finally, we explored the
251 association between children's ln-transformed sum of urinary arsenic concentrations and
252 the neuropsychological scores adjusting for sub-cohort location (i.e. Asturias, Gipuzkoa,
253 or Sabadell) in addition to the potential confounders described in the core models
254 (Supplemental Material, **Table S2**). Children from Valencia were excluded in the sub-
255 cohort adjusted models to circumvent collinearity between sub-cohort location and age at
256 MSCA test. All analyses were carried out with the R software for statistical computing
257 version 3.5.1 (R Core Team, 2014). A threshold of p -value < 0.05 was used to define
258 associations as statistically significant.

259

260 **3. Results**

261

262 Of the 400 children evaluated, 361 (90%) were ultimately included in the analyses because
263 they did not contain missing values in neither neuropsychological development test nor
264 other covariates. Our study sample contained 185 (51%) girls and 176 (49%) boys.
265 Children's median (interquartile range) sum of urinary arsenic concentrations was 4.85
266 (2.74 - 7.54) µg/L overall, and 4.76 (2.36 - 7.48) µg/L and 4.96 (3.09 - 7.60) µg/L for the
267 girls and boys, respectively. Almost all children reported school attendance at 4 years
268 across all sub-cohort locations. Refer to **Table 1** for further details.

269

270 We also assessed characteristics of the study population stratified by the median
271 concentration of 4.85 µg/L of the sum of urinary arsenic. Children with ≥ 4.85 µg/L also
272 had higher concentrations of urinary AsB with a median of 15.95 µg/L versus 5.41 µg/L (p
273 < 0.001). We did not observe statistically significant differences with other characteristics
274 of the study population (**Table 2**).

275

276 We observed a negative linear association between ln-transformed sum of urinary arsenic
277 concentrations and the scores from the original global motor function ($\beta = -2.29$, 95%
278 confidence interval (CI) = [-3.95, -0.63], $p = 0.007$), the derived gross motor function ($\beta =$
279 -1.92 , 95% CI = [-3.52, -0.31], $p = 0.020$) and fine motor function ($\beta = -1.54$, 95% CI = [-
280 3.06, -0.03], $p = 0.046$) after adjustment for maternal highest attained level of education,
281 child's sex, BMI, age at MSCA testing, and calorie adjusted consumption of rice and
282 fish/seafood (**Table 3**). We did not observe any clear association with the remaining MSCA
283 function scores and children's ln-transformed sum of urinary arsenic concentrations (**Table**
284 **3**).

285

286 In the stratified analyses by sex, we found negative trends between boy's ln-transformed
287 sum of urinary arsenic concentrations and the scores of quantitative index and with the
288 derived working memory function ($\beta = -2.59$, 95% CI = [-5.36, 0.17], $p = 0.066$, and $\beta = -$
289 2.56 , 95% CI = [-5.36, 0.24], $p = 0.073$, respectively), which were supported by low p -
290 values in the interaction term ln-transformed sum of urinary arsenic concentrations and sex
291 in the core models ($p = 0.065$ and $p = 0.052$, respectively). Further, we observed a stronger
292 negative trend with an average of 5-fold higher regression coefficient between ln-
293 transformed sum of urinary arsenic concentrations and the remaining neuropsychological
294 function scores in boys compared to girls, but they did not achieve statistical significance
295 (**Table 3**).

296

297 We did not observe any major changes in the regression coefficients between models (i.e.
298 unadjusted (Model 0), adjusted for confounders (Model 1), with calibrated children's sum
299 of urinary arsenic for consumption of fish/seafood (Model 2), and adjusted for children's
300 hair mercury concentrations (Model 3 in Supplemental Material, **Table S1**). The restrictive
301 analysis including only children who did not consume fish/seafood also followed similar
302 trends; however, wider confidence intervals were observed owing to the small dataset ($n =$
303 49) (Model 4 in Supplemental Material, **Table S1**). The results from the adjusted sub-
304 cohort location models, excluding children from Valencia, followed the trend of our
305 primary findings; however, the regression coefficients were attenuated (Supplemental
306 Material, **Table S2**). The mathematically calibrated urinary arsenic species concentrations
307 (i.e. iAs, DMA and MMA) and their sum removed any association with urinary AsB
308 concentrations and had Pearson's correlation coefficients ($r < 0.017$, $p > 0.745$). Calibrated
309 children's ln-transformed sum of urinary arsenic concentrations did not appreciably alter
310 the association with the scores in the original scale of global motor function ($\beta = -2.11$,
311 95% CI = [-3.86, -0.36], $p = 0.018$) and strengthened the negative association between ln-
312 transformed sum of urinary arsenic concentrations and children's scores on fine motor
313 function ($\beta = -1.82$, 95% CI = [-3.41, -0.22], $p = 0.026$) (Model 2 in Supplemental Material,
314 **Table S1**). In contrast, the regression coefficient between calibrated children's ln-

315 transformed sum of urinary arsenic concentrations and the scores in gross motor was
316 modestly attenuated and lost statistical significance ($\beta = -1.38$, 95% CI = [-3.08, 0.32], $p =$
317 0.112) (Model 2 in Supplemental Material, **Table S1**). Similar results were obtained when
318 adjusting for children's hair mercury concentrations (Model 3 in Supplemental Material,
319 **Table S1**). We did not observe any major change in the sensitivity statistical analyses when
320 excluding the identified outliers ($n = 10$) (data not shown).

321

322 **4. Discussion**

323

324 In this study, sum of urinary arsenic concentrations including iAs, MMA, and DMA were
325 used as a biomarker of iAs exposure. We observed that the sum of urinary arsenic
326 concentrations was negatively associated with the scores in the neuropsychological
327 assessment of global, gross and fine motor function among children of approximately 4-5
328 years of age living in Spain after adjusting for potential confounding factors. Our findings
329 also suggest that boys may be more susceptible to iAs neurotoxicity. In particular, we found
330 a stronger negative trend between ln-transformed sum of urinary arsenic concentrations
331 and children's scores in the neuropsychological quantitative and working memory function
332 scales for boys compared to girls.

333

334 In Spain, drinking water usually complies with the EU drinking water iAs regulation, set
335 at 10 $\mu\text{g/L}$ (The Council of the European Union, 1998) with a reported median level < 1
336 $\mu\text{g/L}$ (Espejo-Herrera et al., 2013; Palau Miguel and Guevara Alemany, 2011). Thus, diet
337 is expected to be the main source of iAs exposure for our study population (Davis et al.,
338 2017; Signes-Pastor et al., 2017b). Spain is the second largest producer of rice in the EU
339 and rice consumption is strongly rooted in the Spanish gastronomic culture (Comission,
340 2015; Signes-Pastor et al., 2017b). Rice contains about 10-fold higher iAs compared to
341 other cereals and the concentrations vary geographically (Meharg et al., 2009; Meharg and
342 Zhao, 2012). We have previously reported that rice consumption in our study population
343 was correlated with an increase of urinary iAs, and more weakly with the sum of urinary
344 arsenic concentrations (Signes-Pastor et al., 2017b). Using the median cut point as in **Table**
345 **2**, the difference was not statistically significant, which may be in part because the
346 concentrations of arsenic in rice vary widely and in our previous work in Spain ranges from
347 37 to 407 $\mu\text{g/kg}$ (Signes-Pastor et al., 2016). Also, lack of associations or strong
348 correlations may be related to misclassification of reporting of rice intake using a FFQ that
349 asks about intake over the past year, and not the time period reflective of urinary excretion
350 of arsenic (e.g., the past few days). Fish/seafood consumption is also an important part of
351 the Spanish diet and it contributes to the ingestion of AsB, and tends to dominate exposure
352 to organic arsenic from food intake in the Spanish and other populations with similar
353 gastronomic cultures (Navarro Serrano et al., 2016; Taylor et al., 2016). In this study, the
354 AsB concentrations contributed to over half of the sum of all urinary arsenic species

355 analyzed (i.e. median (interquartile range) of $[\text{AsB} / (\text{iAs} + \text{MMA} + \text{DMA} + \text{AsB}) * 100]$
356 equals 67.0% (41.4% - 86.8%)) and was correlated with children's fish/seafood
357 consumption (Signes-Pastor et al., 2017b), and thus, was critical to remove from our
358 analysis of iAs exposure.

359

360 Numerous studies have reported detrimental effects on neuropsychological development
361 of children living in areas with arsenic-contaminated drinking water with urinary arsenic
362 concentrations 1-2 orders of magnitude higher compared to the levels found in this study
363 (Mst Nasrin Nahar et al., 2014; Mst. Nasrin Nahar et al., 2014; Parvez et al., 2011; von
364 Ehrenstein et al., 2007; Wasserman et al., 2011; WHO, 2011). Although iAs exposure in
365 our study population was low, we observed negative associations between iAs exposure
366 and children's scores in the neuropsychological motor function scales that involve skills
367 such as playing with a ball and drawing. For each interquartile range increase in exposure
368 we found a decrease of over 2 points in the scores for global motor and gross motor scores,
369 and 1.5 points in the scores for the fine motor function.

370

371 Only a few studies have been conducted in populations with low drinking water arsenic
372 concentrations (Forns et al., 2014; Freire et al., 2018; Wasserman et al., 2014). In a cross-
373 sectional study from Maine, among ~10-year-old children, home tap water with arsenic \geq
374 5 $\mu\text{g/L}$ was associated with reductions in full-scale IQ, and with all index scores, i.e.
375 working memory, perceptual reasoning, and verbal comprehension (Wasserman et al.,
376 2014). A recent study from INMA has reported that arsenic levels in placenta were
377 associated with decrements in global and verbal executive function and quantitative
378 abilities, and could also be a risk factor for motor impairment in children of 4-5 years of
379 age (Freire et al., 2018). Another prior study from INMA carried out in the sub-cohort of
380 Sabadell did not find associations between maternal urinary arsenic concentrations during
381 pregnancy and children's neuropsychological development at the age of 4 years (Forns et
382 al., 2014). However, total urinary arsenic concentrations including AsB was used leaving
383 open the likelihood of exposure misclassification (Feldmann and Krupp, 2011; Jones et al.,
384 2016; Molin et al., 2015, 2014, 2012; Signes-Pastor et al., 2017b). In this study, iAs
385 exposure was estimated with sum of urinary iAs, MMA, and DMA. We have previously
386 reported lack of correlation between fish/seafood consumption and urinary iAs, MMA, and
387 DMA concentrations (Signes-Pastor et al., 2017b). However, urinary DMA from
388 biotransformation of organoselenicals from marine product consumption (i.e. arsenosugars
389 and arsenolipids) may still overestimate iAs. Thus, we adjusted for fish and seafood
390 consumption and performed several sensitivity analyses (Jones et al., 2016; Molin et al.,
391 2015, 2014, 2012; Signes-Pastor et al., 2017b). Indeed, to address the potential for
392 overestimation of exposure from fish/seafood consumption (Signes-Pastor et al., 2017b),
393 we calibrated children's urinary arsenic species concentrations using a residual-based
394 method (Jones et al., 2016). Nevertheless, our analyses using adjusted or calibrated sum of

395 urinary arsenic concentrations for fish/seafood consumption generally did not result in
396 appreciable changes in our findings. Similar results were observed when adjusting for
397 children's hair mercury concentrations as a biomarker of fish/seafood intake (Elhamri et
398 al., 2007). Also, similar findings were obtained when we restricted our analysis to only
399 children without fish/seafood consumption (i.e. urinary AsB < 1 µg/L), which despite the
400 small sample size ($n = 49$) produced an inverse trend between exposure to iAs and
401 children's scores in global and fine motor function. Rice contains iAs but also DMA and
402 potentially traces of MMA (Meharg and Zhao, 2012) that may be excreted in the urine
403 unchanged raising concerns of potential iAs exposure misclassification, and therefore we
404 adjusted the regression models for rice intake. Cadmium exposure has been associated with
405 impaired child development (Forns et al., 2014; Freire et al., 2018; Kippler et al., 2012),
406 and thus we analyzed cadmium concentrations in rice from Spain as a potential exposure
407 source; however, we found levels almost undetectable owing to its cultivation under
408 flooded conditions (Arao et al., 2009; Signes-Pastor et al., 2016). Information on children's
409 cadmium level of exposure in our study population is not available yet; however, we would
410 expect levels to be lower than those of children from an industrial and mining region in
411 southwestern Spain and possibly more similar to that reported in children of 6-8 years in
412 Germany or 6-11 years in the U.S. (Rodríguez-Barranco et al., 2014). A preliminary
413 analysis of 5-year-old children from the New Hampshire Birth Cohort Study do not suggest
414 a strong correlation between the children's urinary iAs and cadmium concentrations ($n =$
415 389 ; Spearman $r = 0.2$) (personal communication). In order to address residual confounding
416 from mercury exposure as a risk factor (Freire et al., 2018), we adjusted our core models
417 for children's hair mercury concentrations. Children's diet differed by sub-cohort location
418 (Supplemental Material, **Table S3**) along with their urinary AsB, MMA, and iAs
419 concentrations, but not DMA (Signes-Pastor et al., 2017a). However, they did not differ in
420 their sum of urinary arsenic concentrations (Supplemental Material, **Table S3**). In order to
421 account for geographical differences in metal exposure (Freire et al., 2018), we adjusted
422 for sub-cohort location excluding children from Valencia because of collinearity between
423 sub-cohort location and age at MSCA test and the results followed the trend of our main
424 findings, but the strength of the associations were attenuated. We did not consider exposure
425 to lead and manganese as risk factors (Freire et al., 2018), and that is a limitation of our
426 study since they could result in residual confounding if they were strongly associated with
427 iAs exposures; however, we do not expect that to be the case.

428

429 Our sex-stratified analyses are based on relatively small sample sizes, and therefore caution
430 must be taken in the interpretation of the results. Our findings suggest that boys may be
431 more susceptible to iAs neurotoxicity compared to girls particularly for cognitive tasks
432 related to numerical function, and temporarily storing and managing information. For each
433 interquartile range increase in exposure we found a decrease of 2.6 points in the scores for
434 the quantitative index and working memory among boys. In contrast, in a study from

435 Bangladesh, pre- and post-natal exposure to iAs was inversely associated with verbal and
436 full scale IQ in girls of 5 years of age (Hamadani et al., 2011). In an industrial polluted area
437 in Mexico, an inverse association was identified between urinary arsenic concentrations
438 and problem solving, vocabulary and attention scores among boys, and with memory
439 among girls at the age ranging from 6 to 8 years (Rosado et al., 2007). Sex-related
440 differences in susceptibility to metals toxicity have been associated with differences in
441 patterns of exposure, gastrointestinal absorption, metabolism and detoxification (Llop et
442 al., 2013; Tseng, 2009); however, information regarding early-life gender differences in
443 susceptibility to iAs neurotoxicity is scarce and will require further investigation (Llop et
444 al., 2013).

445

446 This study is among the first to assess the association between iAs exposure, mainly from
447 diet, and neuropsychological development of children taking part in a well-designed cohort
448 (Gascon et al., 2017), and despite the relatively small size of the study population and
449 relatively low level of iAs exposure, we observed associations between children's iAs
450 exposure and the scores in various neuropsychological function scales. Our results should
451 be interpreted cautiously given the cross-sectional design of the study that precludes us
452 from determining temporality and thus limits any inferences about causality. We adjusted
453 for several potential confounding factors, but the effect of unknown factors such as other
454 environmental/dietary factors or residual confounding remains a possibility. A particularly
455 small sample size was used in the sex-stratified analyses with limited statistical power.
456 Children's daily rice and fish/seafood consumption were measured in personal interviews
457 with parents using a validated FFQ (Vioque et al., 2016). The FFQ is considered a reliable
458 method to assess usual diet in epidemiologic studies (Willett, 2012). In this study, the
459 validity of the FFQ was examined by comparing the nutrient values from FFQ with the
460 average nutrient values of three 24 hour dietary recalls, and with the concentrations in
461 blood specimens for several vitamins (i.e. carotenoids, folate, vitamin B12, vitamin C and
462 α -tocopherol) (Vioque et al., 2016). A mathematical method independent to the data
463 recorded on the FFQ was applied to calibrate children's sum of urinary arsenic
464 concentrations for fish/seafood intake. Further, we carried out analysis adjusting for
465 children's hair mercury concentrations, sub-cohort location, and analysis including only
466 children without fish/seafood consumption. In general, sensitivity analyses supported our
467 primary findings, with some attenuation with adjustment for sub-cohort location possibly
468 due the reduced statistical power. Further, multiple testing could have led to false positive
469 results, and therefore our finding should be interpreted with caution and be explored if they
470 persist in further follow-up assessments (Blakesley et al., 2009; Rothman, 1990).

471

472 In conclusion, our study focused on a population with low arsenic in drinking water but
473 who consume iAs in their diet, exposure to iAs was related to certain domains of
474 neuropsychological function scores, in particular motor development. Our findings, along

475 with others, support the reduction of iAs exposure particularly during critical
476 developmental windows early in life.

477

478 **Competing financial interests**

479 All authors declare they have no actual or potential competing financial interests.

480

481 **Conflict of interest**

482 The authors do not have conflicts of interest to declare.

483

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511

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Table 1: Selected characteristics of the study population for the entire dataset and stratified by sex (minimum; interquartile range; maximum) for continuous and *n* (%) for categorical variables.

Selected characteristics of the study population	All (<i>n</i> = 361)	Girls (<i>n</i> = 185)	Boys (<i>n</i> = 176)	<i>p</i> -value
<i>Children:</i>				
Sum of urinary arsenic concentrations (µg/L) ¹	4.85 (0.12; 2.74 - 7.54; 84.46)	4.76 (0.21; 2.36 - 7.48; 84.46)	4.96 (0.12; 3.09 - 7.60; 47.65)	0.393
Urinary AsB (%) ²	67.0 (3.4; 41.4 - 86.8; 100)	67.8 (5.9; 44.4 - 86.8; 100)	66.5 (3.4; 37.4 - 86.8; 100)	0.493
Rice consumption (g/day)	27.2 (0.9; 27.2 - 39.9; 155.2)	26.1 (5.2; 16.2 - 37.8; 142.2)	28.5 (0.9; 18.9 - 40.9; 155.2)	0.373
Fish/Seafood consumption (g/day)	39.9 (10.5; 31.7 - 48.8; 102.3)	40.6 (10.5; 32.5 - 49.3; 102.3)	38.4 (11.2; 29.9 - 48.6; 91.1)	0.078
Sub-cohort (<i>n</i>)	Asturias	96 (27)	48 (26)	0.932
	Gipuzkoa	90 (25)	47 (25)	
	Sabadell	76 (21)	41 (22)	
	Valencia	99 (27)	49 (26)	
BMI (kg/m ²)	16.0 (11.5; 15.2 - 17.2; 25.0)	16.0 (12.9; 15.2 - 17.1; 23.5)	15.9 (11.5; 15.3 - 17.3; 25.0)	0.578
<i>Maternal:</i>				
Age at enrollment (years)	31 (21; 29 - 34; 43)	31 (21; 29 - 34; 43)	31 (21; 29 - 34; 42)	0.277
Social class	Upper - I+II	83 (23)	43 (23)	0.483
	Middle - III	106 (29)	59 (32)	
	Lower - IV+V	172 (48)	83 (45)	
Highest attained level of education	Primary	70 (19)	35 (18)	0.929
	Secondary	148 (41)	75 (41)	
	University	143 (40)	75 (41)	
	0	198 (55)	103 (56)	
Number of previous live births	1	141 (39)	68 (37)	0.278
	2	21 (6)	14 (7)	
	3	1 (0)	0 (0)	
			1 (1)	

For test of differences by sex, we used Welch's t-test or Wilcoxon's rank test for continuous variables, and Chi-square or Fisher's exact test for categorical variables. BMI = Body mass index.

¹DMA + MMA + iAs.

²AsB (%) = (AsB/(iAs + MMA + DMA + AsB)) * 100.

Table 2: Selected characteristics of the study population stratified by the median of the sum of urinary arsenic species concentration (4.85 µg/L) (minimum; interquartile range; maximum) for continuous and *n* (%) for categorical variables.

Selected characteristics of the study population		< 4.85 µg/L (<i>n</i> = 180)	≥ 4.85 µg/L (<i>n</i> = 181)	<i>p</i> -value
<i>Children:</i>				
Sex	Girls	96 (53)	89 (49)	0.493
	Boys	84 (47)	92 (51)	
Rice consumption (g/day)		26.7 (0.1; 18.2 – 36.4; 155.2)	27.9 (0.9; 18.7 – 42.3; 96.8)	0.587
Fish/Seafood consumption (g/day)		39 (10.5; 29.6 - 48.1; 88.5)	40.1 (14.9; 33.5 - 50.0; 102.3)	0.090
Urinary arsenobetaine (µg/L)		5.41 (0.05; 1.24 - 17.47; 3,569)	15.95 (0.29; 5.90 - 59.00; 1,098)	< 0.001
	Sub-cohort			
	Asturias (<i>n</i>)	49 (27)	47 (26)	0.863
	Gipuzkoa (<i>n</i>)	45 (25)	45 (25)	
	Sabadell (<i>n</i>)	40 (22)	36 (20)	
	Valencia (<i>n</i>)	46 (26)	53 (29)	
BMI (kg/m ²)		15.9 (11.5; 15.2 – 16.9; 25.0)	16.1 (12.9; 15.2 – 17.5; 21.0)	0.546
<i>Maternal:</i>				
Enrollment	Age (years)	31.0 (21.0; 29.0 - 34.2; 43.0)	31.0 (24.0; 29.0 - 34.0; 42.0)	0.624
	Upper - I+II	42 (23)	41 (23)	
Social class	Middle - III	50 (28)	56 (31)	0.807
	Lower - IV+V	88 (49)	84 (46)	
Highest attained level of education	Primary	37 (21)	33 (18)	0.583
	Secondary	69 (38)	79 (44)	
	University	74 (41)	69 (38)	
Number of previous live births	0	91 (51)	107 (59)	0.150
	1	74 (41)	67 (37)	
	2	14 (8)	7 (4)	
	3	1 (1)	0 (0)	

For test of differences by sex, we used Welch's t-test or Wilcoxon's rank test for continuous variables, and Chi-square or Fisher's exact test for categorical variables. BMI = Body mass index.

Table 3: Association between children’s sum of urinary arsenic concentrations (ln-transformed) and the McCarthy Scales of Children’s Ability scores standardized to a mean of 100 points with a standard deviation of 15 according to child sex.

McCarthy Scales of Children's Abilities	Model 1 (n = 361) ^a				Girls (n = 185) ^c				Boys (n = 176) ^c				Interaction (n = 361) ^{b,d}	
	β	95% CI		<i>p-value</i>	β	95% CI		<i>p-value</i>	β	95% CI		<i>p-value</i>	<i>p-value</i>	
Original functions	General cognition	-0.86	-2.43	0.71	0.281	-0.08	-2.00	1.84	0.937	-1.87	-4.58	0.84	0.176	0.213
	Verbal	-0.20	-1.88	1.49	0.819	0.71	-1.37	2.79	0.502	-1.54	-4.43	1.34	0.293	0.208
	Perceptual-performance	-1.30	-2.79	0.20	0.090	-0.94	-2.78	0.90	0.313	-1.56	-4.14	1.03	0.236	0.539
	Quantitative index	-0.91	-2.58	0.77	0.288	0.28	-1.84	2.39	0.796	-2.59	-5.36	0.17	0.066	0.065
	Memory	-0.75	-2.39	0.88	0.367	0.00	-2.16	2.17	0.997	-1.63	-4.20	0.94	0.212	0.224
	Global motor	-2.29	-3.95	-0.63	0.007	-1.85	-3.84	0.15	0.069	-3.00	-5.93	-0.07	0.045	0.533
New functions	Executive	-0.28	-1.86	1.30	0.727	0.54	-1.33	2.41	0.570	-1.56	-4.35	1.23	0.270	0.188
	Visual executive	-0.53	-2.10	1.04	0.508	-0.56	-2.49	1.38	0.571	-0.43	-3.08	2.22	0.751	0.971
	Verbal executive	-0.16	-1.82	1.50	0.850	1.00	-0.92	2.92	0.307	-2.00	-5.00	0.99	0.189	0.085
	Visual and verbal span	-0.50	-2.16	1.16	0.557	-0.36	-2.63	1.92	0.757	-0.64	-3.11	1.84	0.611	0.754
	Working memory	-0.67	-2.37	1.04	0.442	0.61	-1.57	2.79	0.581	-2.56	-5.36	0.24	0.073	0.052
	Verbal memory	-0.58	-2.26	1.11	0.501	0.00	-2.11	2.12	0.999	-1.03	-3.85	1.79	0.471	0.446
	Gross motor	-1.92	-3.52	-0.31	0.020	-1.86	-3.67	-0.04	0.045	-2.27	-5.24	0.69	0.132	0.931
	Fine motor	-1.54	-3.06	-0.03	0.046	-0.98	-2.95	0.98	0.326	-2.18	-4.66	0.30	0.085	0.394
Cognitive function of posterior cortex	-1.18	-2.80	0.45	0.156	-0.24	-2.28	1.79	0.813	-2.23	-4.97	0.52	0.111	0.177	

^{a,b}Multiple linear regression models adjusted for maternal highest attained level of education (i.e. primary, secondary, or university), and children’s sex (i.e. girls or boys), BMI (kg/m²), age at MSCA (years) and calorie adjusted consumption of rice and fish/seafood (g/day).

^cMultiple linear regression models adjusted for maternal highest attained level of education (i.e. primary, secondary, or university), and children’s BMI (kg/m²), age at MSCA (years) and calorie adjusted consumption of rice and fish/seafood (g/day).

^dInteraction between children’s sum of urinary arsenic species concentrations (ln-transformed) and sex.

Supplemental Material

Table S1: Association between children's sum of urinary arsenic concentrations (ln-transformed) and the McCarthy Scales of Children's Ability scores

McCarthy Scales of Children's Abilities	Model 0 (n = 361)				Model 1 (n = 361)				Model 2* (n = 361)				Model 3# (n = 234)				Model 4 (n = 49)				
	β	95% CI		p-value	β	95% CI		p-value	β	95% CI		p-value	β	95% CI		p-value	β	95% CI		p-value	
Original functions	General cognitive	-0.74	-2.37	0.90	0.376	-0.86	-2.43	0.71	0.281	-0.80	-2.46	0.85	0.340	0.48	-1.46	2.41	0.629	-1.87	-7.41	3.67	0.499
	Verbal	-0.07	-1.79	1.64	0.933	-0.20	-1.88	1.49	0.819	-0.01	-1.79	1.77	0.991	1.45	-0.68	3.57	0.182	-0.66	-6.02	4.71	0.806
	Perceptual performance	-1.25	-2.80	0.30	0.113	-1.30	-2.79	0.20	0.090	-1.31	-2.88	0.27	0.104	-1.01	-2.87	0.85	0.285	-3.32	-8.15	1.51	0.173
	Quantitative index	-0.79	-2.48	0.91	0.362	-0.91	-2.58	0.77	0.288	-0.99	-2.75	0.77	0.270	0.63	-1.33	2.59	0.527	-0.28	-6.12	5.56	0.922
	Memory	-0.54	-2.21	1.13	0.523	-0.75	-2.39	0.88	0.367	-0.88	-2.61	0.85	0.316	0.48	-1.53	2.49	0.640	2.37	-3.07	7.81	0.383
	Global motor	-2.16	-3.82	-0.50	0.011	-2.29	-3.95	-0.63	0.007	-2.11	-3.86	-0.36	0.018	-2.12	-4.22	-0.02	0.048	-4.77	-10.03	0.49	0.074
New functions	Executive	-0.12	-1.74	1.49	0.882	-0.28	-1.86	1.30	0.727	-0.10	-1.76	1.56	0.903	1.06	-0.90	3.02	0.286	-2.13	-7.77	3.50	0.449
	Visual executive	-0.42	-2.01	1.17	0.601	-0.53	-2.10	1.04	0.508	-0.39	-2.04	1.26	0.641	0.06	-1.92	2.05	0.950	-0.51	-6.06	5.03	0.852
	Verbal executive	-0.01	-1.69	1.67	0.991	-0.16	-1.82	1.50	0.850	0.00	-1.74	1.75	0.997	1.29	-0.81	3.39	0.227	-2.39	-7.98	3.20	0.393
	Visual and verbal span	-0.35	-2.02	1.33	0.685	-0.50	-2.16	1.16	0.557	-0.49	-2.24	1.26	0.585	0.52	-1.47	2.51	0.609	3.75	-2.02	9.51	0.197
	Working memory	-0.48	-2.19	1.23	0.582	-0.67	-2.37	1.04	0.442	-0.69	-2.48	1.11	0.454	0.43	-1.63	2.48	0.683	-0.21	-6.79	6.36	0.948
	Verbal memory	-0.42	-2.12	1.28	0.625	-0.58	-2.26	1.11	0.501	-0.75	-2.53	1.03	0.410	0.42	-1.64	2.49	0.686	0.43	-4.85	5.70	0.871
	Gross motor	-1.68	-3.32	-0.04	0.045	-1.92	-3.52	-0.31	0.020	-1.38	-3.08	0.32	0.112	-1.31	-3.33	0.70	0.201	-2.52	-7.71	2.67	0.332
	Fine motor	-1.59	-3.16	-0.02	0.047	-1.54	-3.06	-0.03	0.046	-1.82	-3.41	-0.22	0.026	-1.79	-3.67	0.09	0.062	-4.77	-10.20	0.67	0.084
Cognitive function of posterior cortex	-1.14	-2.82	0.55	0.186	-1.18	-2.80	0.45	0.156	-1.18	-2.90	0.53	0.176	0.09	-1.93	2.11	0.928	-1.40	-6.68	3.89	0.596	

Model 0: Univariate models. **Model 1:** Models adjusted for maternal highest attained level of education (i.e. primary, secondary, or university), children's sex (i.e. girls or boys), BMI (kg/m²), age at MSCA test (years) and calorie adjusted consumption of rice and fish/seafood (g/day). **Model 2:** The sum of urinary arsenic species concentrations were calibrated following a mathematical method previously described (Jones *et al.* 2016) and the confounders were those included in Model 1 excluding consumption of fish/seafood. **Model 3:** In addition to the potential confounding factors included in Model 1 we added children's hair mercury concentrations analyzed at 4 years. **Model 4:** Analyses were restricted to children with low urinary AsB (i.e. < 1 µg/L) as an indicator of exclusion of fish/seafood consumption and the models were adjusted for the potential confounding factors included in Model 1.

*We calibrated children's urinary arsenic species concentrations following a methodology previously described (Jones et al. 2016). This methodology takes advantage of the fact that urinary arsenobetaine (AsB), a putative non-toxic form of arsenic excreted unchanged rapidly in urine, is an adequate biomarker of fish/seafood intake. To proceed with the calibration, the original sum of urinary arsenic concentrations (i.e. iAs + MMA + DMA) were regressed by the urinary AsB and the model residuals were extracted. Then, we added the mean level of the urinary arsenic species concentrations of participants with low AsB (<1 µg/L; $n = 49$) to the residuals, assuming that iAs exposure levels not derived from fish and seafood are similar in participants with low and high AsB concentrations (Jones et al. 2016). Finally, the calibrated children's urinary arsenic concentrations were included as an independent variable in the multiple linear regression core models adjusted also for potential confounding factors to assess the association with children's neuropsychological scores.

#Among children included in our models only 234 had their hair mercury concentrations analyzed.

Table S2: Association between children’s sum of urinary arsenic concentrations (ln-transformed) and the McCarthy Scales of Children’s Ability scores adjusted by sub-cohort location.

McCarthy Scales of Children's Abilities		Asturias, Gipuzkoa, and Sabadell (<i>n</i> = 262) ^a			
		β	95% CI		<i>p</i> -value
Original functions	General cognition	-0.63	-2.42	1.16	0.487
	Verbal	-0.13	-2.04	1.77	0.891
	Perceptual-performance	-0.51	-2.22	1.20	0.558
	Quantitative index	-1.35	-3.28	0.57	0.168
	Memory	-1.02	-2.89	0.85	0.285
	Global motor	-1.75	-3.61	0.10	0.064
New functions	Executive	-0.10	-1.88	1.68	0.911
	Visual executive	0.08	-1.75	1.91	0.929
	Verbal executive	-0.21	-2.04	1.62	0.818
	Visual and verbal span	-0.52	-2.43	1.40	0.595
	Working memory	-1.07	-3.00	0.86	0.275
	Verbal memory	-0.81	-2.79	1.17	0.420
	Gross motor	-1.48	-3.30	0.34	0.110
	Fine motor	-1.16	-2.86	0.55	0.182
	Cognitive function of posterior cortex	-1.06	-2.91	0.79	0.261

Models adjusted for maternal highest attained level of education (i.e. primary, secondary, or university), children’s sex (i.e. girls or boys), BMI (kg/m²), age at MSCA test (years), sub-cohort location (i.e. Asturias, Gipuzkoa, or Sabadell), and calorie adjusted consumption of rice and fish/seafood (g/day).

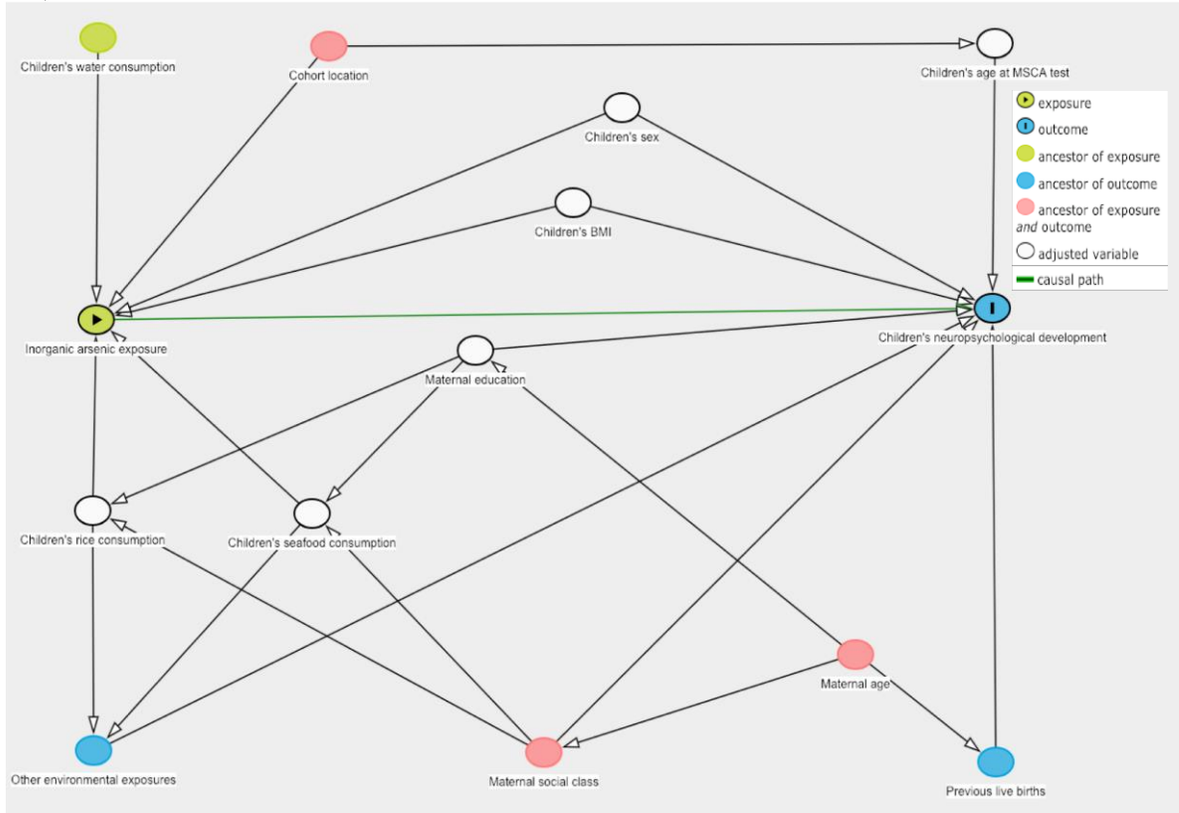
Table S3: Selected characteristics of the study population stratified by sub-cohort location (minimum; interquartile range; maximum) for continuous and *n* (%) for categorical variables.

Selected characteristics of the study population		Asturias (<i>n</i> = 96)	Gipuzkoa (<i>n</i> = 90)	Sabadell (<i>n</i> = 76)	Valencia (<i>n</i> = 99)	<i>P</i> -value
<i>Children:</i>						
Sex	Girls	48 (50)	47 (52)	41 (54)	49 (49)	0.932
	Boys	48 (50)	43 (48)	35 (46)	50 (51)	
Rice consumption (g/day)		26.5 (0.9; 11.2 – 37.5; 72.5)	22.2 (3.7; 9.9 – 34.1; 83.3)	30.9 (7.2; 21.9 – 48.0; 155.2)	28.2 (0.9; 22.2 – 39.3; 142.2)	0.000
Fish/Seafood consumption (g/day)		40.1 (14.7; 32.5 - 50.2; 93.1)	37.1 (18.2; 30.4 - 45.1; 102.3)	43.8 (21.1; 36.9 - 56.0; 82.7)	38.0 (10.5; 26.9 - 47.2; 75.0)	0.000
Sum of urinary arsenic concentrations (µg/L) ¹		4.81 (0.31; 3.04 - 7.19; 84.46)	4.85 (0.12; 2.70 - 8.49; 69.60)	4.76 (0.22; 2.14 - 6.48; 49.1)	5.23 (0.37; 2.95 - 7.80; 28.49)	0.615
BMI (kg/m ²)		16.1 (11.5; 15.3 – 17.5; 21.0)	16.1 (13.2; 15.3 – 17.2; 22.8)	15.6 (12.9; 15.0 - 17.0; 25.0)	15.9 (12.6; 15.2 - 16.9; 21.0)	0.371
<i>Maternal:</i>						
Enrollment	Age (years)	32.0 (21.0; 29.0 - 35.0; 42.0)	32.0 (25.0; 29.0 - 35.0; 43.0)	30.5 (22.0; 28.8 - 34.0; 40.0)	30.3 (21.0; 27.0 - 33.0; 42.0)	0.004
	Social class	Upper - I+II	18 (19)	33 (37)	14 (18)	
Social class	Middle - III	24 (25)	27 (30)	27 (36)	28 (28)	0.007
	Lower - IV+V	54 (56)	30 (33)	35 (46)	53 (54)	
Highest attained level of education	Primary	19 (20)	7 (8)	22 (29)	22 (22)	0.000
	Secondary	38 (40)	31 (34)	31 (41)	48 (48)	
	University	39 (41)	52 (58)	23 (30)	29 (29)	
Number of previous live births	0	59 (61)	46 (51)	40 (53)	53 (54)	0.632
	1	30 (31)	39 (43)	31 (41)	41 (41)	
	2	7 (7)	4 (4)	5 (7)	5 (5)	
	3	0 (0)	1 (1)	0 (0)	0 (0)	

For test of differences by sex, we used Kruskal-Wallis rank test for continuous variables, and Chi-square exact test for categorical variables. BMI = Body mass index.

¹DMA + MMA + iAs.

Figure S1: Directed acyclic graph showing the minimal sufficient adjustment set (Textor et al. 2017).



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