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1 Assessment of arsenic species in human hair, toenail and urine and 2 their association with water and staple food

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24 Abstract

Arsenic intake from household drinking/cooking water and food may represent a significant exposure pathway to induce cancer and non-cancer health effects. This study is based on the human biomonitoring of 395 volunteers from 223 households with private water sources located in rural Punjab, Pakistan. This work has shown the relative contribution of water and staple food to arsenic intake and accumulation by multiple biological matrix measurements of inorganic and organic arsenic species, while accounting for potential confounders such as age, gender, occupation, and exposure duration of the study population. Multi-variable linear regression showed a strong significant relationship between total arsenic (tAs) intake from water and concentrations of tAs, inorganic arsenic (iAs), monomethylarsonic acid (MMA), dimethylarsinic acid (DMA) in urine and toenail samples. tAs intake from staple food (rice and wheat) also showed a strong significant relationship with hair tAs and iAs. The sole impact of staple food intake on biomarkers was assessed and a significant correlation found with all of the urinary arsenic metabolites. Toenail was found to be the most valuable biomarker of past exposure to inorganic and organic arsenic species of dietary and metabolic origin.

Keywords: Monomethylarsonic acid, dimethylarsonic acid, toenail arsenic, dietary
 exposure, urinary arsenic metabolites.

66 **1. Introduction**

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68 Human exposure to toxic inorganic arsenic (iAs) via water is a recognized public 69 health and scientific concern (1). Recently detected arsenic concentrations in food 70 have also raised the question as to the contribution from food. Based on evidence of 71 carcinogenicity in humans, the International Agency for Research on Cancer (IARC) 72 classified arsenic and iAs compounds as 'carcinogenic to humans' (Group 1) and 73 classified dimethylarsinic acid (DMA) and monomethylarsonic acid (MMA) as 74 'possibly carcinogenic to humans' (Group 2B) (2). A sequence of reduction and 75 methylation reactions in the human body metabolises iAs into monomethylarsonic 76 acid (MMA), which is further methylated to DMA (3, 4). Following ingestion, iAs 77 compounds are well-absorbed by humans at an estimated rate of 50 and 95% (5).

78 Most of the ingested arsenic is excreted as methylated arsenic within 1-3 days 79 following exposure although a part of it is stored in sulphydryl-rich tissue such as skin, nail and hair (6). Average per day growth rates for fingernails (0.1 mm), 80 toenails (0.1 and 0.03–0.5 mm) and hair (0.2 to 1.12 mm) depict exposure during the 81 82 last 6, 12–18 and 3-12 months, respectively (7-9). This makes nail and hair effective 83 biomarkers of past exposure, however arsenic toxicokinetics depend on the forms of 84 arsenic and variations in association with various factors such as age, sex, nutritional 85 status and genetic polymorphisms (10). Types and levels of excreted methylated 86 arsenic as a useful biomarker may vary with such factors although few studies have assessed their impact (11-14). Arsenic speciation in hair toenail/nail has been 87 inadequately performed, whilst the association of arsenic intake from water and food 88 89 with inorganic and organic arsenic species in hair, toenail and urine has also been insufficiently studied. For this reason, the present study aimed to assess the impact 90 91 of arsenic exposure in a population of rural settings of the Punjab province, Pakistan 92 using urine, hair and toenail biomarkers. The specific objectives of this research 93 were set to (1) assess human exposure to As through measurement of total arsenic (tAs) and arsenic species in hair, toenail, and urine, and; (2) study the impact of 94 95 dietary exposure (including water) on the internal dose of arsenic species in relation 96 to potential modifiers.

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100 **2. Materials and Methods**

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2.1 Study area and study participants

104 The study villages were located within four districts of Pakistan (Kasur, Sahiwal, 105 Bahawalpur and Rahim Yar Khan), where at least one ground water source was found to be contaminated with arsenic above 50 µg L⁻¹. The sampling frame 106 107 consisted of 398 volunteers (223 households in villages Chak-46/12-L, Chak-48/12-I and Chak 49/12-I, Badarpur, Basti Balochan and Basti Kotla Arab) enrolled 108 109 and interviewed in our previous studies aimed to assess household ground water 110 arsenic concentrations (15) and dietary consumption patterns (16). Residents of 111 these villages were mostly dependent on the household ground water sources 112 (wells, hand pumps) installed 8 to 44 years ago and previously found to have tAs of 113 0.48 to 3090.00 μ g L⁻¹ (15). The participants were non-smoking males and females who used their household ground water for drinking and food preparation did not eat 114 seafood, use any homeopathic or herbal medicines and were not away from their 115 houses for more than a week during the sampling months of August-October, 2014 116 for collection of urine, hair and toenail samples. Pregnant women were excluded 117 118 from the study and after all exclusions, urine (n=395), toenail (n=20) and hair (n=19) 119 samples were collected.

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121 **2.2** Collection of urine, hair and toenail samples

123 Spot urine samples from 246 males and 149 females of six villages o Punjab province, Pakistan were collected in labelled sterile 2 oz polyethylene urine 124 125 collection containers and kept in an ice box at 4 °C prior to return to the laboratory. 126 All urine samples were transferred to a field freezer within 2 hours for storage at -20°C and transported to the National Water Quality Laboratory, where crea tinine was 127 determined on a 1 mL sub-sample. All samples were then shipped with dry ice to the 128 Brooks Applied Laboratory (BAL), USA by air, stored at -70 °C, and finally measured 129 130 for urinary arsenic metabolites within 4 months.

Using ethanol-rinsed stainless-steel scissors, a full strand of hair sample was obtained by the sampling team from the nape of the head as near as possible to the scalp (at a distance of 1 cm from scalp). Hair samples were stapled on cardboard, placed in sealed plastic bags and stored at room temperature until analysis.

135 Participants were asked to remove nail polish, if any, and collect their toenail clippings from all toes using the provided stainless steel clippers (7, 17). These were 136 137 placed in individual polyethylene bags, shipped to BAL and stored at ambient 138 temperature (20℃) until analysis.

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140 2.3 Urine samples processing and analysis

142 Urinary concentrations were corrected for creatinine concentrations, which were determined by the Jaffe method as described by Bonsnes and Hertha (18). This 143 144 correction was done by dividing the concentration of arsenic metabolites (µg L⁻¹) by U-Cre (g L⁻¹) to express urinary arsenical species as $\mu g g^{-1}$ creatinine. 145

146 Frozen urine samples were thawed to room temperature and centrifuged at 3000 147 rpm for 10 min and the resultant supernatants were diluted 10-fold with ultrapure water and analyzed for tAs following U.S. Environmental Protection Agency method 148 149 1638 (mod.) using inductively coupled-plasma dynamic reaction cell-mass spectrometry (Model: ELAN DRC II ICPMS, Perkin Elmer SCIEX, Concord, Ontario, 150 151 Canada). For measurement of urinary arsenic species i.e. arsenate (AsV), arsenite 152 (AsIII), MMA, DMA and arsenobetaine (AsB), aqueous samples were filtered through 153 a 0.45-µm filter. The filtered aliquot were analysed by high-performance liquid 154 chromatography system (Dionex GP-40) coupled to an inductively coupled plasma -155 mass spectrometer (ICP-MS) (Agilent 7700x ICPMS, Agilent Technologies) following the method described by Hata (2007). Urine samples after processing were rapidly 156 157 analysed to ensure appropriate preservation of organic species. Since As(III) can 158 oxidize to As(V) (5) during samples handling and laboratory processing, thus urinary 159 iAs was presented as the sum of As(III) and As(V). The limits of detection were 0.1 μ g L⁻¹ for tAs, As(III), DMA, and AsB, 0.3 μ g L⁻¹ for As(V) and 0.2 μ g L⁻¹ for MMA. 160

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162 2.4 Hair and toenail samples processing and analysis

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164 Each hair sample was cut to a length of 0.125-inch (0.3-cm), representing 165 approximately the last two months of As exposure before sampling. Past studies 166 evaluating the external contamination of hair and nail have reported that washing procedures effectively removed the exogenous As from toenail and hair samples (20, 167 21). Thus, external contamination from hair and toenail clipping samples was 168

169 removed by immersing samples three times in 5 ml of a 0.5% Triton TX-100 solution 170 and shaking thoroughly by hand for 30 seconds. Samples were rinsed three times 171 with 18.2 M Ω deionised water (DIW) and then twice with HPLC grade acetone (21). 172 Hair samples underwent the same cleaning and digestion procedure as toenail 173 samples. Polycarbonate filters (0.4 µm) and an anti-static device were used for the 174 transfer of hair samples between vessels. Following rinsing, samples were dried overnight at room temperature and weighed. Following USEPA method 3050b (22), 175 176 an aliquot of dried toenail or hair sample was prepared by adding multiple additions 177 of HNO₃ and hydrogen peroxide (H₂O₂) and heating at 95 °C \pm 5 °C. After cooling, 178 the volume was made up to 100 mL with DIW, centrifuged and stored at room 179 temperature until analysed exclusively for endogenous arsenic and its species. Total 180 arsenic was measured using the technique of inductively coupled-plasma dynamic 181 reaction cell-mass spectrometry (Model: ELAN DRC II ICPMS, Perkin Elmer, 182 Shelton, CT, USA). All sample extracts for arsenate (AsV), arsenite (AsIII), MMAs, and DMAs quantitation were also analyzed employing an Agilent 7700 CRC ICP-MS 183 184 with a Dionex GP40 HPLC (IC) Systems.

For speciation, an aliquot of filtered sample was injected using a Dionex HPLC onto an anion-exchange column and mobilized isocratically using an alkaline (pH > 7) eluent. The mass-to-charge ratio (m/z) of As at mass 75 was monitored using an Agilent 7700, whilst selenium at m/z 82 was monitored as an internal standard. Retention times for eluting species were compared to NIST traceable known standards for species identification.

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192 **2.5 Quality assurance**

Species data was provided by the analysis of NIST (National Institute of Standards 194 195 and Technology) traceable standard reference materials (SRMs-1640A, trace 196 elements in natural water). Background contamination was monitored using 197 laboratory fortified blanks for urine analysis. Duplicate measurements were made on 10% (n = 40) of urine samples for total arsenic and arsenic species. The 198 199 reliability of the arsenic species determination was evaluated by analysing samples in duplicate and spiking the samples with As(III), As(V), MMA, DMA and AsB. 200 201 Arsenic measured in SRMs-1640A was 7.59 \pm 0.36 tAs μ g kg⁻¹ (n = 6), within the certified range of 8.010 \pm 0.067 µg kg⁻¹, yielding a mean recovery of 96%. The spike 202

recoveries of tAs, AsIII, AsV, DMA, MMA and AsB in digests of matrix spikes (n=31), matrix spike duplicates, duplicates (n=40) and laboratory fortified blank (n=6) met the data quality standards in terms of relative percent difference (RPD) of <25%, percent recovery of 75 to 125% and completeness of 80%.

207 For quality control of hair and nail samples, method blanks, blank spikes, standard 208 reference materials (SRMs) and duplicates were treated in the same way as the 209 samples and incorporated into each digestion batch and analytical run. Human hair 210 SRM (NCS DC 73347 from China National Analysis Centre for Iron and Steel 211 Beijing, China) was used for both hair and nail samples. Arsenic measured in SRM NCS DC 73347 was 274 \pm 0.5 tAs µg kg⁻¹ (n = 2), within the certified range of 280 \pm 212 213 50 µg kg⁻¹, yielding a mean recovery of 98%. There is no available SRM of human 214 hair or nail containing certified concentration for arsenic species. The organic 215 species represented a minimum fraction of tAs in SRM NCS DC 73347, whilst iAs was more than 65% of the extraction indicated as the main proportion of As in hair. 216 The spike recoveries of tAs, iAs, DMA and MMA in digests of matrix spikes (n=2), 217 218 matrix spike duplicate (n=2), duplicate (n=2), blank spikes (n=2), and post spikes 219 (n=2) were 83-92% for hair and 93-123% for toenail.

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221 **2.6 Statistical analysis**

The analysed tAs represents the sum of As species as well as other unidentified 223 224 forms of As species, whilst the SumAs is defined as the sum of urinary iAs, MMA 225 and DMA. Mass balance was assessed by the difference of tAs intake and tAs 226 excreted assuming the mean 24-h urine volume of 1.5 L day⁻¹ (based on urine output of 2.0 L day⁻¹ for men and 1.6 L day⁻¹ for women given by EFSA, 2010). Urine, 227 toenail and hair As concentrations had positively skewed distributions therefore 228 229 geometric transformations applied for statistical analysis. For this analysis, concentrations below the limit of detection (LOD) of the test methods were replaced 230 231 by a value equal to half of the LOD.

ANOVA and student's t-test were used to test for differences in natural log transformed values of urine, toenail and hair arsenic concentrations between different subgroups with respect to age (≤16 and >16 years), gender, ground water tAs concentration, occupation and exposure duration of the study population. Multivariable linear regression models were constructed to assess significant predictors of

biomarkers while controlling for possible confounding factors for this study 237 population. The independent variables were log-transformed values of daily As 238 239 intake from water and staple food (rice and wheat). The dependent variables were 240 log-transformed concentrations of toenail and hair (tAs, iAs, MMA and DMA), and 241 urine (tAs, iAs, MMA, DMA and SumAs). Considered potential confounders were 242 age, gender, occupation and exposure duration. Before multi-variable analyses, 243 bivariate analyses (Pearson correlations) were conducted to assess associations 244 between potentially confounding factors and biomarkers. Factors associated with a P-value<0.1 were first selected then the factors with the weakest P-value were 245 246 inserted in the multi-variable linear regression model using forward selection. The 247 multi-variable models were checked for multicolinearity and goodness of fit. Microsoft 248 Excel, SPSS 24.0 (IBM, New York, NY, USA) and GraphPad Prism 7.0 were used 249 for statistical analyses. The statistical significance level of P≤0.05 was set for the 250 multi-variable analysis.

3. Results and Discussion

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3.1 Study population characteristics

255 Data on the estimated daily total arsenic (tAs) intake of this study population from water, rice and wheat was obtained from previously published studies (15, 16) and 256 257 further, as yet, unpublished work (Table 1). The study participants living in rural region of Punjab Province, Pakistan had an age range of 3-80 years at the time of 258 259 sampling with 37% female participants and 10% participants above 60 years of age. 260 The household's drinking/cooking water was found to have a GM tAs concentration 261 of 55.33 µg L⁻¹ and a range of 0.48-3090 µg L⁻¹, with 89% of sources above the WHO provisional guideline value (10 μ g L⁻¹) for arsenic in drinking water (23). 262

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3.2 Urinary biomarker levels in relation to population subgroups

The GMs for the concentrations of urinary tAs (234.43 μ g g⁻¹ creatinine), iAs (26.98 µg g⁻¹ creatinine), MMA (23.32 µg g⁻¹ creatinine) and DMA (142.80 µg g⁻¹ creatinine) for all study participants and for different demographic and behavioural subsets are shown in Tables 2. The DMA metabolite was the predominant form of As in urine (representing 71% of the sum of urinary arsenic metabolites), followed by iAs (13%) and MMA (12%). This conforms to the findings of Melak, Ferreccio (24) indicating As
excretion as iAs (10–20%), MMA (10–15%) and DMA (60–75%) depending on interindividual variation. AsB generated as a result of seafood ingestion, was not
detected in this study population.

The significant impact (P < 0.001) of ground water tAs concentration (<10 µg L⁻¹, 10-275 276 50 μ g L⁻¹ and >50 tAs μ g L⁻¹) on urinary arsenic metabolites (Table 2) was found in 277 concordance with the other studies on low arsenic regions (25, 26). There was a 278 significant age-dependent trend for urinary tAs concentrations (P = 0.032) whilst 279 males had significantly higher concentrations of urinary tAs, iAs, MMA, SumAs $(P \le 0.05)$ than females. The trend of higher MMA excretion in men than women 280 281 $(27.72 \text{ vs. } 17.47 \mu \text{g} \text{g}^{-1} \text{ creatinine})$ was consistent with previous investigations (27, 282 28). This difference was reported to be linked with choline synthesis under the effect 283 of estrogen in women of childbearing age (12, 13). Estrogen contributes to the synthesis of choline by regulating the phosphatidylethanolamine methyltransferase 284 (PEMT) pathway (29). Non-intensive labour occupations were associated with 285 significantly increased tAs, iAs and SumAs concentrations (P < 0.05) compared to 286 287 labour intensive occupations (Tables 2). Exposure duration (≤14 and >14 years) did 288 not have a significant impact on urinary concentrations (data not shown).

Mass balance was estimated to determine which source provided the majority of the tAs intake. Out of tAs intake (842.69 μ g day⁻¹) from total water intake water (799.47 μ g day⁻¹) and staple food (43.22 μ g day⁻¹), the mean tAs excreted in urine was 591.18 μ g day⁻¹. The remaining 251.51 μ g day⁻¹ was assumed to be internally absorbed and/or excreted in faeces. The tAs intake from the consumption of food (43.22 μ g day⁻¹) represents only 7.31 % of the excreted tAs.

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3.3 Toenail and hair biomarkers levels in relation to population subgroups

A significant increase in toenail and hair concentrations of tAs and its species ($P \le 0.001$) was found with increasing drinking/cooking water tAs concentration (<10 µg L⁻¹ to >50 tAs µg L⁻¹) except for hair DMA (Table 3). The binding of iAs, dietary and/or metabolically produced DMA and MMA with sulfhydryl nails is reported to be partly dependent on the concentration available in the blood (30). Thus, this study participants with longer exposure duration (>14 years) had significantly higher 305 concentration of toenail and hair tAs and iAs, indicative of prolonged exposure306 (Table 3).

Age and gender in this study population did not show a significant impact on toenail and hair concentrations (data not shown). Type of occupation (labour intensive and non-labour professions) showed no impact. Despite the higher outdoor activities of participants engaged in labour intensive occupations (services, farmers, wives of farmers contributing in farming), significantly higher toenail DMA in this study participants engaged in non-labour intensive occupations (general house wives, students, tailors, teachers and un-employed) was unclear (Table 4).

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315 Intercorrelations among exposure biomarkers

The concentration of urinary iAs was significantly correlated with urinary MMA (r=0.905, P \leq 0.0001) and DMA (r=0.884, P \leq 0.0001). Whilst, urinary MMA was significantly associated with DMA (r=0.912, P \leq 0.0001). Urinary iAs was significantly correlated with toenail tAs (r=0.484, P=0.036), toenail iAs (r=0.494, P=0.031), hair tAs (r=0.513, P=0.030) and hair iAs (r=0.487, P=0.040). A significantly strong association between hair tAs (r=0.779, P \leq 0.0001) and toenail (tAs) also exist.

323 Significant positive intercorrelations between urinary, toenail and hair arsenic 324 species suggest that either of these may be used as biomarkers of arsenic exposure, 325 however these biomarkers reflect the As exposure over different time periods as 326 mentioned in section 1.

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328 3.4 Multi-variable linear regression analysis of relations between tAs intake 329 and exposure biomarkers 330

331 Multi-variable linear regression analysis revealed a positive significant relationship 332 between the tAs intake from drinking/cooking water and urinary tAs, iAs and MMA 333 after adjusting for gender, occupation and exposure durations for this study 334 population (Table 4). The association between urinary arsenic metabolites and 335 drinking water arsenic concentrations in household water sources in the rural area of 336 Punjab are in line with the results of multi-variable regression models from previous studies (31, 32) indicating a positive relation between estimated intake of tAs from 337 338 drinking water and urinary As species adjusting for gender (Table 4).

A significant positive association existed between tAs intake of this study population from staple food and those of urinary arsenic metabolite concentrations when 341 adjusted for gender and occupation. The predictor variables such as 342 drinking/cooking water and food tAs intakes both showed significance with response 343 variables i.e. toenail tAs, iAs, MMA, DMA and hair tAs and iAs, indicating the mean 344 change in the response variable for one unit of change in the predictor variable while 345 holding gender, occupation and exposure duration as constant (Table 4). The influence of gender, exposure duration and occupation subgroups on urine, hair and 346 347 toenail tAs and arsenic species suggests the possible underlying reasons. These include metabolic, inter-individual, social-demographic and behavioural variability, 348 349 growth rate of skin appendages, health status, nutrition or exogenous contamination 350 from dust or soil in crop field and kinetic models for peripheral tissues (30). This study participants living in rural Punjab exposed to tAs (water) <1 μ g L⁻¹ and <10 μ g 351 L^{-1} showed a staple food tAs intake of 0.485 µg kg⁻¹ bw day⁻¹ (n=5) and 0.733 µg 352 kg⁻¹ bw day⁻¹ (n=50) respectively. No significant impact of tAs intake from food was 353 354 found on urinary arsenic metabolites below 1 µg L⁻¹. However, participants exposed to <10 μ g L⁻¹ tAs concentration of drinking/cooking water (n=50) showed significant 355 356 Pearson correlation (P < 0.05: data not shown) between tAs intake from food and 357 urinary arsenic metabolites, suggesting the sole contribution of food in human 358 exposure to arsenic.

359 The regression model coefficients (Table 4) showed that for every additional unit of 360 tAs intake from water in this study, an average increase of urinary tAs by 220.74 µg g⁻¹ creatinine (urine), 1944.96 µg kg⁻¹ (toenail) and 755 µg kg⁻¹ (hair) was expected. 361 362 Compared to this, tAs intake from food shows increased tAs concentration by an average of 456.23µg g⁻¹ creatinine (urine), 5721.58 µg kg⁻¹ (toenail) and 4272.70 µg 363 364 kg⁻¹ (hair). This increase due to food tAs intake was higher by an average factor of 365 3.6 when compared to values derived from model coefficient of water tAs intake. 366 These findings showed that water and food tAs intake were found as the strongest 367 predictors of all of the urinary and toenail biomarker concentrations. When compared to food, drinking/cooking water was a relatively stronger predictor as seen by 368 369 adjusted R-square values (Table 4). Though the sample size of toenail and hair could 370 constitute a limitation of this study, the degree of significant associations (Table 4) 371 revealed that toenail arsenic speciation is a more precise biomarker of effects, a 372 potential determinant of prolonged arsenic exposure and indicative of critical arsenic 373 related health effects. In the same context, an elevated risk of cutaneous melanoma

374 (33) and lung cancer (34) was reported in persons with higher toenail arsenic 375 concentrations.

4. Conclusions 376

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The consumption of drinking/cooking water containing range of total arsenic 378 379 concentrations in household hand pumps/wells of six rural settings of Pakistan 380 significantly increased the absorbed dose of tAs, iAs and its mono- and di-381 methylated arsenic in urine, hair and toenail of study participants under the influence 382 of certain biological and behavioural modifiers such as gender, exposure level, 383 occupation and exposure duration. Levels of these species in biological matrices of rural residents of arsenic affected region of Punjab, Pakistan can also increase 384 385 significantly due to exposure through frequent consumption of staple foods such as rice and wheat. The levels of tAs, iAs and its mono- and di-methylated arsenic in 386 387 urine, hair and toenail were also influenced by certain biological and behavioural 388 modifiers such as gender, exposure level, occupation and exposure duration. 389 Association of toenail arsenic with water and food intake of arsenic can be observed 390 as a more favourable biomarker of arsenic exposure than urine and hair.

Given the critical role of highly reactive and genotoxic intermediate trivalent forms 391 392 of MMA and DMA produced from methylation of inorganic arsenic, this study 393 underscores the need to determine these trivalent forms in association with 394 potentially modifying effects of dietary and occupational exposure along with 395 confounding factors such as smoking, nutrients, genetics, education on arsenic 396 accumulation and excretion.

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404

405 **Competing interests**

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407 The authors declare that they have no competing/conflicting interests.

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Table-1: Selected characteristics of study participants who provided urine, hair and toenail samples

Characteristics	n	GM (min-max)	Data source
Study participants	398		This study
Urine samples	395		
Hair samples	19		
Toenail samples	20		
Age			Rasheed, Slack
≤16 years	66		(16)
>16 years	332		
Gender			
Male	249		
Females	149		
Body weight (Kg)	398	52.19 (9-105)	
Exposure duration from ground water tAs (vears)		14.7 (3-44)	Rasheed, Kay (15)
8-13	212		· · /
13-15	62		
15-44	124		
tAs concentration in household ground water			
(rg -) Overall	398		
<10	50		
10-50	145		
>50	203		
Estimated daily tAs intake (ug kg ⁻¹ bw day ⁻¹)			
Drinking/cooking water	398	3.217 (0.02-236.510)	
Participants consumed rice only	4	0.176 (0.122-0.226)	Rasheed, Slack
Participants consumed wheat only	230	0.609 (0.194-2.234)	(16)
Participants consumed staple food (wheat+rice)	164	0.589 (0.275-2.0235)	
Occupation category		, , , , , , , , , , , , , , , , , , ,	
Labour non-Intensive (n=149)			
House wives (general)	45		
Students	75		
Tailors	4		
Teachers	4		
Un-employed	21		
Labour intensive (n=249)			
Farmers	186		
Wives/family member of farmers (contributing in farming)	56		
Services	7		
GM:Geometric mean	•		·

Table-2: Geometric means [GM (min-max)] for creatinine adjusted urinary 517 **arsenic metabolites (\mu g g^{-1} creatinine)**

Characteristics	n	Urine Creatinine	tAs	iAs	DMA	ММА	Sum As
Overall	verall 385		234.43 (7.78-8743.59)	26.98 (0.139-1411.11)	142.80 (0.08-2353.53)	23.32 (0.08-615.31)	201.38 (0.30-4375.76)
Age			, , , , , ,	, í			
≤16 years	62	0.92 (0.56-1.56)	302.38 (27.55-8743.59)	30.44 (0.23-1357.24)	162.99 (0.13-1704.08)	26.52 (0.14-615.31)	230.81 (0.49-3676.63)
>16 years	323	1.02 (0.35-2.55)	223.17 (7.78-3969.70)	26.36 (0.14-1411.11)	139.22 (0.08-2353.54)	22.75 (0.08-611.11)	196.17 (0.30-4375.76)
p-values (t test)		0.03	0.032	0.424	0.411	0.415	0.395
Gender							
male	241	1.03 (0.35-2.55)	267.13 (7.78-8743.59)	30.60 (0.14-1411.11)	158.71 (0.08-2353.54)	27.72 (0.08-611.11)	226.30 (0.30-4375.76)
female	144	0.96 (0.54-2.01)	188.97 (10.30-4510.20)	21.85 (0.23-1357.24)	119.67 (0.11-1955.22)	17.47 (0.14-615.31)	165.65 (0.49-3676.63)
p-values (t test)		0.02	0.002	0.013	0.052	0.001	0.020
tAs in water used for drinking and cooking (μg L ⁻¹)							
<10	50	0.98 (0.50-1.93)	113.76 (10.297-760.60)	14.53 (1.43-123.03)	87.45 (9.167-488.46)	11.81 (1.38-102.02)	116.75 (12.12-677.58)
10-50	140	0.97 (0.41-2.55)	163.46 (18.636-1233.33)	19.72 (1.29-229.62)	118.15 (9.93-967.68)	16.59 (0.76-251.28)	159.13 (17.80-1220.64)
>50	195	1.02 (0.35-2.45)	360.50 (7.778-8743.59)	39.60 (0.14-1411.11)	185.54 (0.08-2353.54)	35.47 (0.08-615.31)	274.25 (0.30-4375.76)
p-values (ANOVA)		0.350	0.0005	0.0005	0.0005	0.0005	0.0005
Occupation							
Labour intensive	242	1.024 (0.53-2.55)	213.44 (7.78-2563.64)	24.34 (0.14-381.11)	129.06 (0.08-1990.91)	21.16 (0.08-415.45)	182.22 (0.30-2767.36)
Labour non-Intensive	143	0.96 (0.50-2.03)	274.62 (12.62-8743.59)	32.11 (0.23-1411.11)	169.47 (0.13-2353.54)	27.49 (0.14- 615.31)	238.49 (0.488-4375.76)
n volues (t test)	1	0.036	0.019	0.042	0.061	0.067	0.046

Table-3: Geometric means [GM (min-max)] for arsenic and arsenic species in toenail and hair (μ g kg⁻¹)

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Characteristics	n	n	Toenail, GM(min-max) (μg kg⁻¹)				Hair, GM(min-max) (µg kg ⁻¹)			
Characteristics	(toenail)	(hair)	tAs	iAs	MMA	DMA	tAs	iAs	MMA	DMA
Overall	20	19	1942.18	1756.91	79.44	21.88	702.16	653.25	1.43	2.64
			(586-27500)	(557-22000)	(6-955)	(0.8-432)	(67.0-3100.0)	(84-10700)	(0.5-55)	(0.5-123)
tAs in water										
used for										
drinking and										
cooking (µg L ⁻¹)										
<10	5	5	593.06	568.43	32.82	0.91	73.82	90.06	2.46	3.36
			(586-599.2)	(559.2-578)	(26-39)	(0.8-1)	(67-94.1)	(84.0-95)	(2.0-3.0)	(2.0-7)
10-50	4	3	1321.94	1217.91	32.72	12.88	1006.65	830.73	0.72	10.12
			(602-4070)	(557-3840)	(6-97)	(6-57)	(352-2760)	(325-2250)	(0.6- 0.9)	(0.6-69)
>50	11	11	3830.19	3352.55	163.93	112.52	1771.87	1505.83 (438-	1.35	1.64
			(1190-27500)	(1270-22000)	(77-955)	(25-432)	(531-13100)	10700)	(0.5-55)	(0.5-123)
p-values (ANOVA)			0.0005	0.001	0.001	0.0005	0.0005	0.0005	0.0005	0.364
Occupation										
Labour intensive	13	12	1766.00	1627.03	67.12	11.81	504.73	507.16	1.39	1.69 (0.5-69)
			(586-27500)	(559-22000)	(6-955)	(0.8-432.0)	(67-13100)	(84-10700)	(0.5-17)	
Non-Labour	7	7	2317.31	2026.26	108.65	68.75	1236.57	1008.19	1.52	5.66
intensive			(605.0-4660)	(557-4070)	(48-209)	(9.1-310.0)	(352-4610)	(325-3590)	(0.5-55)	(0.5-123)
p-values (t test)			0.53	0.59	0.25	0.04	0.190	0.34	0.89	0.28
Exposure duration										
≤14 years	13	12	1277.25	1163.46	66.54	13.48	330.06	329.59 (84.0-	1.57	1.34
			(586.0-4660)	(557.0-4070)	(26.0-209)	(0.8-310.0)	(67.0-3770)	3140)	(0.5-17.0)	(0.5-7.0)
>14 years	7	7	4229.75	3777.22	110.41	53.74	2561.18	2110.63	1.23	8.50
			(2060.0-27500)	(1840-22000)	(6.0-955)	(6.0-432)	(615.0-13100)	(669.0-0700)	(0.5-55.0)	(0.5-123
p-values (t test)			0.012	0.009	0.331	0.123	0.005	0.004	0.703	0.119

Table-4. Multi-variable linear regression analysis of associations between log transformed values of estimated daily intake of tAs (μ g kg⁻¹ bw day⁻¹) and exposure

biomarkers

Independent variable	Biological Matrix	Biomarkers	β coefficient	Std. Error	p-value	Model Adjusted R ²
		tAs	0.307	0.028	0.0005	0.276 ²
		iAs	0.3	0.038	0.0005	0.168 ²
	Urine	DMA	0.229	0.042	0.0005	0.069 5
		MMA	0.284	0.04	0.0005	0.158 ²
		Sum As	0.259	0.038	0.0005	0.104 5
the intoke from		tAs	0.348	0.063	0.0005	0.612 ³
drinking water	Teeneil	iAs	0.342	0.056	0.0005	0.660 ³
uninking water	roenali	DMA	0.672	0.08	0.0005	0.606 5
		MMA	0.24	0.122	0.008	0.294 5
		tAs	0.443	0.073	0.0005	0.792 ¹
	Hair	iAs	0.386	0.07	0.0005	0.764 ¹
		DMA	-0.291	0.159	0.15	0.243 5
		MMA	0.009	0.19	0.958	-0.17 ⁵
		tAs	0.577	0.106	0.0005	0.122 ²
		iAs	0.894	0.132	0.0005	0.105 ⁵
	Urine	DMA	0.773	0.143	0.0005	0.068 5
		MMA	0.866	0.138	0.0005	0.136 ²
		Sum As	0.812	0.131	0.0005	0.088 5
the intelse from	Toenail	tAs	1.017	0.291	0.003	0.547 ¹
tAs intake from staple diet		iAs	0.995	0.265	0.002	0.587 ¹
		DMA	2.698	0.598	0.0005	0.504 4
		MMA	1.131	0.336	0.003	0.352 5
	Hair	tAs	1.725	0.357	0.0005	0.718 ¹
		iAs	1.547	0.322	0.0005	0.718 ¹
		DMA	-1.139	0.700	0.128	0.258 ¹
		MMA	0.043	0.591	0.943	-0.169 ⁵

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