Arsenic Levels and Speciation from Ingestion Exposures to Biomarkers in Shanxi, China: Implications for Human Health

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Supporting Information

ABSTRACT: Chronic exposure to arsenic (As) threatens human health. To systematically understand the health risks induced by As ingestion, we explored water and diet contributions to As exposure, and compared As in biomarkers and the arsenicosis status in a geogenic As area in China. In this study, high percentages of water (77% of n = 131 total samples), vegetables (92%, n = 120), cereals (32%, n = 25), urine (70%, n = 99), nails (76%, n = 176), and hair (62%, n = 61) contained As higher than the acceptable levels. Dietary As contributed 92% of the average daily dose (ADD) when the water As concentration was less than 10 μg/L, for which 5 out of 30 examined participants were diagnosed with arsenicosis symptoms. The distinct positive correlation between ADD and As concentrations in urine, nails, and hair suggests different applicability for these biomarkers. Methylated As as the predominant urinary As species confirms that the ingested inorganic As is methylated and is excreted through urine. In situ microdistribution and speciation analysis indicates that As is mainly associated with sulfur in nails and hair. Nails, rather than hair and urine, could be used as a proper biomarker for arsenicosis. High ADD from the environment and low excretion could result in As toxicity to humans.

INTRODUCTION

Arsenic (As) poses a severe threat to human health through exposure pathways mainly including water and diet. Drinking water with elevated As levels is the primary source of As ingestion. Moreover, As-laden groundwater, if used for irrigation, can lead to appreciable amounts of As accumulation in vegetables and cereals. Extensive research demonstrates that rice and rice products could be another As source. However, not much attention has been paid to other cereals like wheat, which are important staple foods worldwide, that could lead to significant As ingestion. The intake of As results in its detection in noninvasive human biomarkers such as urine, nails, and hair. Urine has frequently been used as a biomarker for short-term exposure because most As metabolites are excreted within 3–4 days through urine. Nails and hair indicate long-term exposure for 2–12 months, as As can be accumulated in these structures rich in keratin. Thus, the association of As levels in biomarkers and groundwater has been recently studied to evaluate its health risks.

The speciation of As in biomarkers is crucial to understand the transformation, metabolism, and toxicity of ingested As. As species in hair and nails are traditionally determined by chemical extraction, where inorganic and methylated As are detected. No As–S species have been reported using indirect chemical extraction methods, even though As readily bonds to sulphhydryl groups. In contrast, As–S species have been detected in nail and hair samples using in situ X-ray absorption near-edge structure (XANES) spectroscopy. Unfortunately, methylated As species are not distinguished from the As–S form in nails, which deserves further exploration to understand the metabolism of As.

The purposes of this study were (1) to explore the contributions of water and diet to As exposure, and (2) to compare the As in biomarkers and the arsenicosis status in a geogenic As area in China. This comprehensive study should provide insights into As exposure and its health effects.

MATERIALS AND METHODS

Sample Collection. The study area lies in Shanyin, China, as described in our previous report. Groundwater from tube wells (n = 131), garden soils (n = 19), garden vegetables (n = 120), cereals (n = 25), urine (n = 99), nails (n = 176), and hair (n = 159) samples were collected, and the details of sample collection are given in the Supporting Information (SI). A total 222 of participants (male: 128; female: 94; age ≥17) of approximately 7500 residents in 8 villages were asked to complete a self-administered questionnaire including age, sex, smoking habits, and duration of drinking groundwater. More...
participants than water samples were surveyed because some participants share one tube well. The questionnaire results indicated that the residents in the study area mostly drink untreated groundwater. Dietary habits in the area were investigated in nine families as detailed in SI Table S10. Diagnosis for skin keratosis, depigmentation, and hyperpigmentation were based on the Chinese national standards for diagnosis of arsenicosis by trained interviewers.

**Sample Preparation and Analysis.** Vegetables were washed with tap water to remove soil and dust particles, rinsed with deionized (DI) water, crushed, and dried at 60 °C for microwave digestion analysis. The cereal grains were crushed and dried for digestion. No As speciation was considered in vegetables and cereals, as inorganic As predominates in these samples.

Nail and hair samples were treated following a reported method. Briefly, nail and hair samples were washed in three steps, DI water—methanol—DI water, and then dried at 60 °C. Urine samples were centrifuged at 8000 rpm for 10 min, and the supernatant was filtered through a 0.22 μm membrane for speciation analysis.

Solid samples were digested with a microwave digestion system (MARS, CEM Corporation, U.S.) as shown in the SI. The total As concentration was determined using an atomic fluorescence spectrometer (AFS, Haiguang, P.R. China) with a detection limit of 0.6 μg/L. Groundwater samples were analyzed using a furnace atomic absorption spectrometer (FAAS, Perkin-Elmer AAS-800) with a detection limit of 0.7 μg/L.

Arsenite, arsenate, monomethylarsonic acid (MMA), and dimethylarsinic acid (DMA) in urine samples were determined using high-performance liquid chromatography (HPLC) coupled with AFS. The details of As speciation analysis are shown in the SI.

**Quality Control.** Standard reference materials (hair, GBW09101b; soil, GBW (E) 070011) were used to validate the microwave digestion analysis. A urine standard (GBW09115) was employed to justify the As speciation analysis by HPLC-AFS. Selected samples were digested and analyzed in duplicate or triplicate for quality control purposes. The relative standard derivation was calculated as an indication of method precision and was in the range of 85.4–117.5% (SI Table S3).

**μ-XRF and μ-XANES Analysis.** One nail and three hair samples were analyzed using μ-XRF and μ-XANES at beamline 15U at the Shanghai Synchrotron Radiation Facility (SSRF), China. The details of nail and hair preparation and analysis are shown in the SI. Standards including Na₂HAsO₄·7H₂O, Na₃AsO₃, MMA, DMA, and arsenic-glutathione (As(Glu)₃) as detailed in the SI were also analyzed and used in the linear combination fit.

**Health Risk Assessment.** A widely used model derived by the USEPA was applied to estimate the As impact on the residents in the study area. The As intake from water, vegetables, and cereals was considered using the following equation:

$$\text{ADD} = \text{As}_x \times \text{IR} \times \text{EF} \times \text{ED} / (1000 \times \text{AT} \times \text{BW})$$  \hspace{1cm} \text{(1)}$$

where ADD is the average daily dose (mg/kg/d); As is the As concentration (μg/L in groundwater; μg/g in vegetables and cereals based on dry weight); IR is the ingestion rate (drinking water: 1.8 L/d; various vegetables and cereals: g/d as shown in SI Table S10); EF is the exposure frequency (drinking water: 365 d/y; food: values (d/y) calculated from questionnaire, SI Table S10); ED is the exposure duration (y) from the questionnaire; AT is the average life expectancy (25 S50 y); and BW is the body weight (male: 61.0 kg; female: 53.2 kg).

Toxic and cancer risk assessment was estimated using eqs 2 and 3, respectively:

$$\text{HQ} = \text{ADD} / \text{RfD}$$  \hspace{1cm} \text{(2)}$$

where HQ is the hazard quotient (toxic risk is considered occurring if HQ > 1.00), RfD is the oral reference dose (3 × 10⁻⁴ mg/kg/d).²¹

$$R = 1 - \exp(-\text{SF} \times \text{ADD})$$  \hspace{1cm} \text{(3)}$$

where R is the cancer risk; SF is the slope factor (1.5 mg/kg/d).

### RESULTS AND DISCUSSION

**Arsenic in Groundwater, Vegetables and Cereals.** An average As concentration of 174 μg/L (ND-1160 μg/L) was found in groundwater (n = 131), with 77% exceeding the WHO’s guideline of 10 μg/L and 52% above 50 μg/L (Figure 1-I). Moreover, the average As(III) concentration (114 μg/L) was higher than As(V) (60.1 μg/L), in line with the reducing conditions in the aquifer.²⁵

![Figure 1. Boxplot of As contents in vegetables and groundwater samples (I) and cereals (II) with the sample numbers listed above the x-axis. The box represents the data between the 25th and 75th percentiles. The small box and horizontal line inside the box indicates the mean and median data. The whiskers (error bars) above and below the box indicate the 95th and 5th percentiles and dots above and below them represent outliers.](https://dx.doi.org/10.1021/es400129s)
addition to the irrigation water, other factors including genetic difference in vegetables could contribute to the variation in As accumulation.4

The As concentrations in all cereals ranged from ND to 0.42 μg/g (n = 25, Figure 1-II), with 32% of samples exceeding the permissible limit in China (0.15 μg/g in rice, 0.1 μg/g in flour, and 0.2 μg/g in other cereals).26 The wheat flour, as the local staple food, was purchased from the market, and the average As concentration (0.08 μg/g) was a little higher than the flour samples (0.021−0.054 μg/g) collected from As-contaminated areas in France.9 Meanwhile, rice, as another staple food, was not locally planted and showed a mean concentration of 0.20 μg/g, within the upper end of the global background range (0.082−0.202 μg/g).9 Low As concentrations were detected in nonstaple foods such as yellow rice (0.10 μg/g), corn (0.06 μg/g), and millet (0.06 μg/g) (Figure 1-II), which are comparable to the reported data (SI Table S11).

Averaged Daily Dose for As Exposure. By summing the ADD from drinking water, vegetables, and cereals, the residents consumed a mean total ADD of 2.6 × 10^{-3} mg/kg/day (range from 0.2 to 14.1 mg/kg/day, SI Table S7). Approximately 98.5% of ADD exceeded the RfD (3 × 10^{-4} mg/kg/day),21 indicating a potentially hazardous risk of As ingestion. However, the ADD value was lower than that in Kandal,

Table 1. Correlation Coefficientsa for Age, Sex, Arsenicosis b, EDc, ADDd and from Water Drinking (w), Vegetables (v), Cereals (c), Ln (Urine As), Hair As, and Nail As

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<th>sex</th>
<th>arsenicosis</th>
<th>ED (w and v)</th>
<th>water As</th>
<th>w-ADD</th>
<th>v-ADD</th>
<th>c-ADD</th>
<th>ADD</th>
<th>ADD</th>
<th>ln (urine As)</th>
<th>hair As</th>
<th>nail As</th>
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<td>0.299**</td>
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aCorrelation coefficient denoted as p < 0.05 (*), p < 0.01 (**) by Spearman analysis. bParticipants were divided into two populations with or without symptoms of arsenicosis. cED, exposure duration. dADD, average daily dose summing from groundwater (w), vegetables (v), and cereals (c). The number of participants or samples is 131 otherwise noted in the bracket.
In addition to water, vegetables and cereals contributed a mean ADD value of 2.3 × 10^{-7} μg/kg/day (2% exceeding the RfD) and 5.0 × 10^{-4} mg/kg/day (86% exceeding the RfD), respectively. Notably, the contribution of ADD from vegetables and cereals significantly increased with the decrease of water As concentrations (Figure 2-I, -II, and -III). For example, vegetables and cereals contributed 35% and 57% to ADD, respectively, when As concentration was below 10 μg/L (Figure 2). In line with our results, the diet may contribute approximately 75% to the total ADD when the water As concentration is below 10 μg/L.2

A significant positive correlation was found between ADD and water As concentration ($r = 0.888$, $p < 0.001$, Table 1) and ADD indeed increased with water As levels (Figure 2-IV). Moreover, a positive correlation was observed between ADD and relevant parameters including age and exposure duration. Furthermore, chronic exposure through water and diet may lead to As accumulation in the human body, which could be confirmed by the analysis of noninvasive human biomarkers.

**As Concentration and Speciation in Urine.** The mean total urinary As concentration, calculated by summing As(III), As(V), DMA, and MMA, was 56.0 μg/L (ND-551 μg/L, $n = 99$, Figure 3-I). The background As levels in urine are 10 μg/L and 100 μg/L in European countries and the United States24 and 9.1 μg/L in the control adult group in Inner Mongolia, China.13 About 70% of the analyzed urine samples exceeded 10 μg/L. Furthermore, Ln-transformed total urinary arsenic concentration increased with the increase in water As concentration ($p = 0.029$, Table 1). For example, a mean urinary As concentration of 16.2 μg/L ($n = 13$) was detected in groups with <10 μg/L As in drinking water, 45.4 μg/L ($n = 17$) in groups with 10–50 μg/L, and 110 μg/L ($n = 34$) in groups with >50 μg/L. A positive correlation was found between Ln-transformed total urinary arsenic and water ADD ($p = 0.006$), vegetables ADD ($p = 0.027$), and total ADD ($p = 0.01$) (Table 1). The outliers in Figure 3-I were due to five urine samples with high As concentrations. Higher As concentrations than 200 μg/L were observed in their drinking water, which contributed to such high urinary As concentrations.

DMA was the major As species in urine with a mean concentration of 42.6 μg/L, followed by As(III) (6.4 μg/L), MMA (5.9 μg/L), and As(V) (4.8 μg/L) (Figure 3-I). The proportion of each species in urine (SI Table S4) was in agreement with a previous study25 and showed that methylated As species metabolized in the liver are preferentially excreted through urine.

![Figure 3](image3.png)

**Figure 3.** Speciation of As in urine ($I$, $n = 99$) and total As concentration in nails ($II$, $n = 176$) and hair ($II$, $n = 159$).

As Concentration in Nails and Hair. The average As levels were 7.8 μg/g in nails ($n = 176$) and 4.2 μg/g in hair ($n = 159$) (Figure 3-II). Approximately 76% of nail and 61% of hair samples exceeded the widely accepted background level of 1.5 μg/g in nail10 and 1.0 μg/g in hair.6 A comparable study in India reported 7.2 μg/g As in nails and 3.4 μg/g As in hair with 62% of water samples exceeding 50 μg/L.31 A positive correlation between concentrations in nails ($p < 0.001$) and hair ($p = 0.001$) with water As levels was found in this study (Table 1). Moreover, nails and hair, as biomarkers of long-term exposure, indicated a positive correlation with total ADD (both $p < 0.01$) and exposure duration (both $p < 0.05$).

**Microdistribution of Arsenic and Sulfur in Nails.** The As content was higher in the margins than in the intermediate layer, as shown in the μ-XRF image (Figure 4-I). As expected, sulfur was homogeneously distributed among the sections because the nail is rich in keratin with sulphydryl groups.14 The free sulphhydryl groups in the nail margins coordinate much more easily with inorganic As than the stable disulphide, which is rich in the intermediate layer.18 In accordance with our observations, discrete layers of As were reported in the nail samples collected from a gold mining area characterized by μ-XRF mapping.18

![Figure 4](image4.png)

**Figure 4.** (I): Spatial distribution of As and S in a nail section (red outline) by μ-XRF mapping (numbers 1 and 3 represent the margins and 2 represents the intermediate). (II): As K-edge μ-XANES spectra of standards, one nail (three spots marked with numbers in (I), 73.1 μg As/g) and three individual hair samples (17.2 μg/g in a, 21.6 μg/g in b, and 17.7 μg/g in c). Experimental spectra are displayed as dashed lines. The red lines are the fit results; lines in different colors represent standard references. The solid and dashed vertical lines represent the As(Glu)3 and As(V) absorption edge, respectively. The inset percentages represent the As(Glu)3 proportion.

Arsenic Speciation in Nails and Hair Using μ-XANES. The dominant feature of nail and hair μ-XANES spectra had an absorbance edge near As(Glu)3 (Figure 4-II). The sulfur bonded As species (As-Glu) contributed 69–76% in nail and 54–64% in hair by linear combination fit (SI Table S6). The large content of As-Glu in these samples indicated that As bonds easily to the sulphhydryl groups in keratin.4,17 Relatively more As-Glu was found in nails than in hair, corresponding to the higher keratin content in nails.17

DMA (18%) was observed in the intermediate layer of nails (SI Table S6). The ingested As is primarily metabolized to a methylated form in the liver, which finally partially accumulates in the nails and hair.17,29 The nail is formed predominantly by the proximal nail matrix, where As is associated with free...
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...sulphydryl groups and is considered isolated from the metabolism during nail growth.32

**Arsenic Risk Assessment and Health Implications.** Significant positive correlations between ADD and Ln-transformed As concentrations in urine \( (n = 57, r_s = 0.341, p = 0.01) \), nails \( (n = 96, r_s = 0.412, p < 0.001) \), and hair \( (n = 95, r_s = 0.328, p = 0.001) \) were obtained (Table 1). Thus, these biomarkers can reflect the As exposure with different retention times. Notably, severe patients were identified as deviations from the general association of As concentration in their biomarkers and ADD (Figure 5). A lung cancer patient marked with a red circle ingested substantially high ADD \( (1.5 \times 10^{-2} \text{ mg As/kg/day}) \) but excreted small quantities of As through urine \( (11.9 \mu \text{g/L}) \), nails \( (3.6 \mu \text{g/g}) \), and hair \( (5.2 \mu \text{g/g}) \). Meanwhile, two more patients labeled by arrows in Figure 5 are suffering from severe skin lesions. High ADD from the environment and low excretion due to metabolism might result in As toxicity to humans, although many factors including demographic characteristics, malnutrition, and individual health status may play a role.

The average HQ was 8.6 (0.6 to 47.1) with 98% of the residents exceeding the typical risk index of 1.00 \( (n = 131, \text{SI Tables S7 and S8}) \). In addition, the mean cancer risk was about 4 in 1000 exposure \( (n = 131) \) and ranged from 4 in 10 000 to 2 in 100 (SI Table S7). Meanwhile, the cancer risk was found to be 6% for >1 in 100 exposure, 88% for >1 in 1000 exposure, and 100% for highest safe standard of 1 in 10 000 exposure (SI Table S8), which is in agreement with a previous study in Bangladesh.11 Thus, exposure to As resulted in a severe health risk.

The arsenicosis rate increased with the increase in water As concentrations (Figure 2-IV). A similar result was obtained in Inner Mongolia, China, where a significant dose–response relationship was found between skin hyperkeratosis and water As concentrations.35 Moreover, 17% of examined participants with water As concentration below 10 \( \mu \text{g/L} \) \( (n = 30) \) were diagnosed with arsenicosis symptoms. This observation was in agreement with previous reports.33,35 This extremely high rate can be attributed to the As intake from diet.

A significant correlation was observed between arsenicosis and nail As concentration \( (r_s = 0.269, p < 0.01, \text{Table 1}) \). No correlation was observed between arsenicosis and Ln (urinary As) \( (r_s = 0.143, p = 0.295) \) and hair As concentrations \( (r_s = 0.199, p = 0.051) \). Thus, arsenicosis might be predicted by nail As concentration, even though an association was found between the As concentrations in these three biomarkers (Table 1). The observed arsenicosis status was not as severe as the estimated HQ results from the health risk assessment (SI Table S7–S8). However, we cannot wait for the development of arsenicosis to make any mitigation efforts because the As chronic toxicity symptoms are indeed time-lag effects.13,36

### ASSOCIATED CONTENT

**Supporting Information**

Details of sample collection; microwave digestion of soils, vegetables, cereals, nails, and hair samples; As speciation using HPLC-AFS; statistical methods; \( \mu \text{-XRF} \) and \( \mu \text{-XANES} \) sample preparation and analysis; and additional figures and tables. This material is available free of charge via the Internet at http://pubs.acs.org.

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**Notes**

The authors declare no competing financial interest.

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### REFERENCES


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**Figure 5.** Correlation of ADD with arsenic concentrations in urine \( (p = 0.01 \text{ using Ln-transformed urinary As concentration}) \), nails \( (p < 0.001) \), and hair \( (p = 0.001) \).

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**Table S8**, which is in agreement with a previous study in Bangladesh.11 The cancer risk was found to be 6% for >1 in 100 exposure, 88% for >1 in 1000 exposure, and 100% for highest safe standard of 1 in 10 000 exposure (SI Table S8), which is in agreement with a previous study in Bangladesh.11 Thus, exposure to As resulted in a severe health risk.