Arsenic speciation in Chinese Herbal Medicines and human health implication for inorganic arsenic

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Abstract
Rice and drinking water are recognized as the dominant sources of arsenic (As) for human intake, while little is known about As accumulation and speciation in Chinese Herbal Medicines (CHMs), which have been available for many hundreds of years for the treatment of diseases in both eastern and western cultures. Inorganic arsenic was the predominant species in all of CHMs samples. The levels of inorganic arsenic in CHMs from fields and markets or pharmacies ranged from 63 to 550 ng/g with a mean of 208 ng/g and 94 to 8683 ng/g with a mean of 1092 ng/g, respectively. The highest concentration was found in the Chrysanthemum from pharmacies. It indicates that the risk of inorganic As in CHMs to human health is higher in medicines from markets or pharmacies than that collected directly from fields. Some CHMs may make a considerable contribution to the human intake of inorganic arsenic.

1. Introduction

Herbal or medicinal plant products in various forms have been available for many hundreds of years for treatment of diseases in both Eastern and Western cultures (Chan, 2003). A survey indicated that about 70–80% of the world populations, particularly in the developing countries, rely on non-conventional medicine mainly from herbal sources in their primary healthcare reported by WHO (Akerle, 1993; Chan, 2003). Chinese Herbal Medicines and their manufactured products recorded in the Chinese Pharmacopoeia have been approved for safe use and have been widely consumed for thousands of years as home remedies or for prevention and treatment of diseases in China (Chan, 2003; Chinese Pharmacopoeia Commission, 2005). Such increase in popularity has also brought concerns and fears over the adverse effects of herbal medicines, thus the quality, efficacy and safety of the products from herbal and natural sources available in the market should have been proven. Therefore, contamination or adulteration of CHMs with heavy metals such as lead, mercury, cadmium or arsenic has attracted much attention (Au and Reddy, 2000; Koh and Woo, 2000), especially arsenic contamination through natural and anthropogenic pathways (Koch et al., 2007; Liu et al., 2010a,b). There are concerns that CHMs can accumulate As from growth environments and conditions, such as soil/plant uptake, irrigation water and from atmospheric deposition, and conditions during dryness, storage, transport and manufacturing processes (Chan, 2003; Liang et al., 1998; Liu et al., 2010a). China has a maximum contaminant concentration (Green Standard of Medicinal Plants and Preparations for Foreign Trade and Economy in China) for arsenic in CHMs of 2.0 mg/kg. Some investigations of total arsenic in CHMs from both domestic and foreign markets have illustrated that total arsenic concentrations in some CHMs samples exceeded this legislative standard (Cooper et al., 2007; Liu et al., 2010a).

Arsenic is a ubiquitous element in the natural environment. The International Agency for Research on Cancer (IARC) regards inorganic arsenic as a carcinogen with a linear dose response for chronic exposure (IARC (International Agency for Research on Cancer), 1973; NRC (National Research Council), 2001). Because inorganic As is considered to be more toxic than methylated species – MMA and DMA (Marin et al., 1992; NRC (National Research Council), 2001; Schoof et al., 1999), the assessment of human health risk associated with As in edible plants or foodstuffs mainly depends on the concentrations of inorganic arsenic (Muñoz et al., 2002; Schoof et al., 1999; Tsuji et al., 2007). Moreover, Food and
Agriculture Organization/World Health Organization (FAO/WHO) has a provisional tolerable weekly intake (PTWI) for inorganic arsenic of 15 μg/kg bodyweight (2.1 μg/kg bodyweight (b.w.)/d) (FAO/WHO (Food and Agriculture Organization/World Health Organization), 1993). Allowable Daily Intake (ADI) of inorganic As for one person with 60 kg bodyweight would be 126 ng/g. However, the EFSA (European Food Safety Authority) Panel on Contaminants in the Food Chain (CONTAM Panel) suggested that PTWI of 15 μg/kg b.w. established by the Joint FAO/WHO Expert Committee on Food Additives (JECFA) is no longer appropriate as data had shown that inorganic arsenic causing cancer of the lung, urinary bladder and skin had been reported at exposures lower than those reviewed by the JECFA. CONTAM Panel assessed the risks of arsenic in food to human health. The inorganic arsenic exposure from food and water across 19 European countries had been estimated to range from 0.13 to 0.56 μg/kg bodyweight (b.w.) per day for average consumers and from 0.37 to 1.22 μg/kg b.w. per day for 95th percentile consumers (European Food Safety Authority (EFSA), 2009). Chinese Government, considering the adverse effects of inorganic As on human health, had issued the maximum levels of inorganic arsenic in foods in GB2762-2005, such as rice flour (150 ng/g), wheat flour (100 ng/g), non-staple cereals (200 ng/g) and vegetables (50 ng/g/60 kg body weight) in Food Standards (China, 2005).

Most recent studies have focused on the total As levels in CHMs, and little information is available on arsenic species in CHMs and the human health implications of inorganic As in CHMs. More research is therefore needed into human intakes of inorganic arsenic in CHMs and its risk to health. In addition, Chinese Government had put more emphasis on the effects of inorganic As on human health and issued the regulatory standards for inorganic arsenic in foods in GB2762-2005, but they have not well built up the related certified reference materials (CRMs) and developed the protocols for analysis of inorganic arsenic in terrestrial edible plants, especially Chinese Herbal Medicines. Therefore, the one objective of this study was to screen a suitable extraction method for As species in CHMs. In general, arsenic in CHMs comes from growth environments and conditions, or CHMs accumulate arsenic under conditions during dryness, storage, transport and manufacturing processes. For this reason, some samples were directly collected from field to represent pure/unprocessed plant materials with un-impacted As species. We also collected CHMs samples from markets or pharmacies, considering the processes could impact on the As species in the medicines. Arsenic speciation in CHMs taken directly from fields and from markets or pharmacies was extracted using that selected method and analyzed by HPLC-ICP-MS. The risk to human health of inorganic arsenic in CHMs has also been examined.

2. Materials and methods

2.1. Survey

An-guo city in Hebei province has been well known for cultivating and processing herbal medicines in China for hundreds of years. According to the local cultivated species, planting areas and locations, 12 typical herb medicine species were selected and samples were directly collected from field or purchased from markets or pharmacies (n = 197 in total). Based on a preliminary survey of total As in CHMs (Liu et al., 2010a), a part of samples (n = 65) with high levels of total As in CHMs were selected to explore As speciation in this study.

In general, the roots of some herb plants can be used as medicines only, which were collected as samples in this study, including Indigowoad Root (Isatis indigotica; Banlangen in Chinese), Large-head Atractylodes Rh (Atractylodes macrocephala; Baihu in Chinese), Danshen (Salvia miltiorrhiza), Radix Saposhnikoviae (Saposhnikovia divaricata; Fangfeng in Chinese), Radix Astragali (Astragalus membranaceus, Huangqi in Chinese), Tatarian Aster (Aster tataricus; Ziwan in Chinese), Rhizoma anemarrhenae (Anemarrhena asphodeloides; Zhiwu in Chinese), Radix Trichosanthis (Trichosanthes kirilowii; Tianhuafen in Chinese). Except for herbal roots, other organs of herbs could be medicines, such as shoots of Schizonepeta (Schizonepeta tenuifolia; Jingjie in Chinese) and Indigowoad Leaf (Isatis indigotica Fort., Daqingye in Chinese) and fruits of Trichosanthis (Trichosanthes kirilowii; Gualou in Chinese) and flowers of Chrysanthemum (Dendranthema morifolium; Juhua in Chinese and well known as tea as well as in eastern and western countries).

Both root and leaf of Isatis digotica and both root and fruit of Trichosanthes kirilowii are used as medicines in general.

2.2. Samples preparation

All samples were washed with ultrapure water (18.2 MΩ), and then were oven-dried at 65 °C until a constant weight was reached. The dry samples were powdered using a stainless-steel ball-mill. All samples were stored in desiccators at room temperature before extraction.

2.3. Chemicals

AnalR Nitric acid (HNO₃) (70%) and hydrogen peroxide (H₂O₂) (30%) were used as reagent grade and were obtained from Beijingchemical works; Trifluoroacetic acid (TFA) (CH₃CH₂O₂) (99%) were obtained from Sigma–Aldrich. Sodium arsenate (Na₅AsO₄·12H₂O) was purchased from Chemical Reagent Factory of Union in Beijing, and sodium arsenite (Na₂AsO₃) was purchased from Merck (Germany). Methylarsonic acid (MAA) and dimethylarsinic acid (DMAA) were purchased from PA (USA). Indium (In), the internal standard in ICP-MS, was a high purity stock supplied from Agilent (USA). The HPLC mobile phase was prepared using diammonium hydrophosphate ((NH₄)₂HPO₄) and ammonium nitrate (NH₄NO₃), which all were analytical reagent (AR) from Fucheng Chemicals (Tianjin) and Shantou Chemicals (Guangdong), respectively. The certified reference material (CRM) of rice flour (GBW10010) from the National Research Center for Standard Materials in China was used to validate the analyses. Ultrapure water (18.2 MΩ) in preparation for solution was produced by Milli-Q Element system from Millipore (USA).

2.4. Arsenic speciation extraction

Method screening: to obtain the suitable extraction method of arsenic speciation in CHMs, four methods to extract As species were evaluated in this study according to TFA extraction method from Heitkemper et al. (2001) and microwave extraction with 1% HNO₃ solution in Zhu et al. (2008). No certified reference material (CRM) for arsenic speciation in herbal medicine, the CRM of Chinese rice flour GBW10010, arsenic speciation published in Zhu et al. (2008), was used to validate analysis. We consider that the herbal medicine is normally roots or shoots different from rice flour, so Panax notoginseng (Sanqi in Chinese), one of herbal medicines, was run for arsenic speciation extraction with rice CRM. Arsenic speciation in rice flour CRM and herbal medicine sample Panax notoginseng were extracted by the following methods:

1. Microwave method for extracting arsenic species: Ultra-pure water, 1% HNO₃, 1% HNO₃ + 1% H₂O₂ (Table 1). The CRM and 0.2000 g of Panax notoginseng were weighed into 50 ml polypropylene centrifuge tubes and 10 ml of solutions was added, respectively. Covered centrifuge tubes were allowed to stand overnight at room temperature. All tubes were randomized, heated in a microwave accelerated reaction system (CEM Microwave Technology Ltd, USA) using three temperature steps: (1) maintained samples at 55 °C for 10 min; (2) 75 °C for 10 min; (3) 95 °C for 30 min. And each step ramped to temperature over a 5 min period.

2. Extraction with 2M TFA (Trifluoroacetic acid). TFA has been commonly used to extract As species in rice plant samples (Abedin et al., 2002; Liu et al., 2006; Williams et al., 2005). The sample (0.2000 g) was weighed into the quartz glass digestion tubes, and steeped in 2 ml of 2 mol/l TFA solution. The tubes were then left overnight at room temperature, followed by digestion heating block at 100 °C for 5 h until the solution evaporated to dryness. If not, the temperature was elevated to 160 °C to dry the content. When the tubes were cooled down, the residues and solution were vortexed by ultra-pure water to 10 ml. Samples were then centrifuged at 7000 rpm for 8 min, and the supernatant was collected and passed through a 0.45 μm × 13 mm nylon filter.

Based on our findings, microwave extraction with 1% HNO₃ solution was the most suitable method for arsenic species extraction in CHMs (Table 1). Therefore, milled subsamples of CHMs were extracted with 1% HNO₃ using the microwave oven system.

2.5. Arsenic speciation determination

Arsenic speciation analysis was performed using high performance liquid chromatography inductively coupled plasma-mass spectrometry (HPLC-ICP-MS, Agilent 1200 series and Agilent ICP-MS7500, Agilent Technologies, USA,) connected to a PXR-X1000 10 μm anion-exchange column (250 × 4 mm). The injection volume of the sample was 20 μl. The mobile phase consisted of pre-filtered (0.45 μm) 5 mM NH₄NO₃ and 5 mM (NH₄)₂HPO₄ adjusted to pH 6.2 (HNO₃) with flow rate at 1.0 ml/min.
min. Retention time of peaks were identified by mixed arsenic speciation standards (10 µg As/l of arsenite, arsenate, dimethylarsinic acid (DMA\textsuperscript{V}) and monomethylarsonic acid (MMA\textsuperscript{V})). Arsenic species were quantified by external calibration with DMA\textsuperscript{V} standard solutions (0, 5, 10, 15, and 20 µg of As/l). All data was normalized with the Internal standard.

2.6. Total digestion and determination of arsenic

For total As concentration in CHMs, milled subsamples (about 0.2 g) were weighted into a 50 ml covered polypropylene centrifuge tubes and 2 ml of concentrated HNO\textsubscript{3} was added. The tubes were left overnight at room temperature and then heated in a microwave using the same temperature program as documented for As speciation extractions and finally made up to the volume of 25 ml with ultra-pure water. Hydride generation and an atomic fluorescence spectrometry (AFS-2202E, Beijing Haiguang Analytical Instrument Co., Beijing, China) and inductively coupled plasma mass spectrometry (ICP-MS; Agilent 7500, Agilent Technologies, USA) were used in the As measurement. CRM and blanks were run during digestion for data quality controls.

2.7. Speciation quality control

To check the stability of arsenic species in 1% HNO\textsubscript{3} solution, four blank spikes with 10 µg As/l of arsenite, arsenate, DMA\textsuperscript{V} and MMA\textsuperscript{V}, respectively were taken in triplicate and then extracted in the microwave oven. Arsenic species in extract solutions were analyzed by HPLC-ICP-MS. The recoveries for the blank spikes of arsenite, arsenate, DMA\textsuperscript{V}, and MMA\textsuperscript{V} using 1% HNO\textsubscript{3} were 92 ± 2.2%, 99 ± 0.4%, 118 ± 0.5%, 123 ± 0.3%, respectively. The mean limits of detection (LOD) for arsenite, arsenate, DMA\textsuperscript{V}, and MMA\textsuperscript{V} by HPLC-ICP-MS were 0.065, 0.080, 0.079, 0.065 ng/g.

2.8. Statistical analysis

Statistical analyses were undertaken using Origin Pro8.0 for graphic analysis and using SPSS 18.0 for ANOVA data analysis.

3. Results and discussion

3.1. Recoveries of As Speciation extractions

The recoveries of arsenic species extracted from rice CRM (GBW10010) or herbal medicine Panax notoginseng were different for the four extract solutions (Table 1). In general, the recoveries from rice CRM (GBW10010) were higher than those from Panax notoginseng according to the different extraction solutions. The reasonable explanation is that rice CRM is grain flour and Panax notoginseng is powder of roots. For ultra-pure water extraction, whether rice CRM (GBW10010) or herbal medicine Panax notoginseng, recoveries were the lowest, accounting for 51% and 64%, respectively, compared to other extraction solutions. The higher recovery of As species from rice CRM extracted by 1% HNO\textsubscript{3} + 1% H\textsubscript{2}O\textsubscript{2} (128%) was probably because of a low concentration of arsenic in this CRM (Zhao et al., 2010). Better and stable recoveries from TFA extraction were observed in Table 1, 84% for rice CRM and 74% for the herbal medicine sample, but TFA can reduce arsenate to arsenite during extraction. Much higher concentrations of As\textsuperscript{III} were observed in TFA extraction compared to others. Therefore, many reports just presented the combined concentration of inorganic arsenic composed of As\textsuperscript{III} and As\textsuperscript{V} (Abedin et al., 2002; Ackerman et al., 2005; Heitkemper et al., 2001; Liu et al., 2006; Smith et al., 2008, 2006; Vela and Heitkemper, 2004; Williams et al., 2005). The extract solution 1% HNO\textsubscript{3} has been used widely to extract arsenic species from plant tissue samples, especially for rice flour (Raab et al., 2009; Zhu et al., 2008). It has also been proved successfully in extracting As species from rice shoot, bran, husk, and root, when exactly the same procedure used for the grain was applied in these plant sections, obtaining recoveries of 94 ± 11 (n = 9), 91 ± 13 (n = 3), 81 ± 13 (n = 3), and 106 ± 4% (n = 6), respectively (Zhu et al., 2008), Raab et al. (2009) also demonstrated good extraction efficiency (104%) using 1% HNO\textsubscript{3} with microwave extraction for rice flour CRM. For Panax notoginseng, recovery was the highest in 1% HNO\textsubscript{3} extract solution and the proportions of four arsenic species were stable. Moreover, a little amount of unidentified As species were detected at the same retention time, located by external calibration (species sum/total As)*100.

![Table 1](image)

Summary of arsenic speciation and extraction recoveries in certified reference material (CRM) of GBW10010 (rice flour) and the Panax notoginseng obtained from pharmacies using four extraction methods (Means ± SE, n = 3).

<table>
<thead>
<tr>
<th>CRW10010</th>
<th>As species extraction</th>
<th>Total As (ng/g)</th>
<th>As species (ng/g)</th>
<th>Recovery (%)a</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>As\textsuperscript{III}</td>
<td>As\textsuperscript{V}</td>
</tr>
<tr>
<td>Ultra-pure water</td>
<td>102</td>
<td>24 ± 1</td>
<td>17 ± 0.5</td>
<td>11 ± 0.3</td>
</tr>
<tr>
<td>1% HNO\textsubscript{3}</td>
<td>102</td>
<td>68 ± 7</td>
<td>18 ± 3</td>
<td>27 ± 1</td>
</tr>
<tr>
<td>1% HNO\textsubscript{3} + 1% H\textsubscript{2}O\textsubscript{2}</td>
<td>102</td>
<td>13 ± 0.4</td>
<td>80 ± 77</td>
<td>27 ± 2</td>
</tr>
<tr>
<td>2 mol/l TFA</td>
<td>102</td>
<td>81 ± 1</td>
<td>86 ± 2</td>
<td>84 ± 1</td>
</tr>
<tr>
<td>Panax notoginseng</td>
<td>Ultra-pure waterb</td>
<td>5985</td>
<td>374 ± 4</td>
<td>1744 ± 20</td>
</tr>
<tr>
<td>1% HNO\textsubscript{3}c</td>
<td>5985</td>
<td>685 ± 19</td>
<td>2729 ± 46</td>
<td>77 ± 0</td>
</tr>
<tr>
<td>1% HNO\textsubscript{3} + 1% H\textsubscript{2}O\textsubscript{2}</td>
<td>5985</td>
<td>129 ± 4</td>
<td>2777 ± 49</td>
<td>104 ± 12</td>
</tr>
<tr>
<td>2 mol/l TFAa</td>
<td>5985</td>
<td>2976 ± 96</td>
<td>236 ± 39</td>
<td>79 ± 1</td>
</tr>
</tbody>
</table>

a Recovery = (species sum/total As)\texttimes 100.
b Contained an unidentified As species at 20 ng/g.
c Contained an unidentified As species at 42 ng/g.
d Contained an unidentified As species at 122 ng/g.

3.2. Arsenic speciation in Chinese Herbal Medicines

The concentrations of arsenic species in CHMs samples collected directly from fields are shown in Table 2. Only As\textsuperscript{III} and As\textsuperscript{V} were detected in the extract solutions of CHMs with the roots as medicines. The concentrations ranged at 472–182 ng/g for As\textsuperscript{III} and at 1.1–264 ng/g for As\textsuperscript{V}. Arsenite was the predominant species in the extracts of 63% samples, accounting for 60–99% of total As. Arsenate was the main species in the remaining samples with the proportion at 66–74% (Table 2). There was quite a big difference in the dominant As species in the medicines from fields, which might depend on the growing conditions and their genetic properties. There were no MMA or DMA found in roots of CHMs. Furthermore, arsenic species in those CHMs with shoots or flowers as medicines also followed that inorganic arsenic is predominant. Only a low level of DMA\textsuperscript{V} (<10%) was found in the flowers of D. morifolium. Recent report showed that arsenic could not be methylated in planta and soil microorganisms are responsible for As methylation and DMA\textsuperscript{V} in plants mainly comes from soil solution (Lomax et al., 2012). DMA\textsuperscript{V} could be more slowly taken up by roots compared to inorganic As, but translocated rapidly from root to shoot (Li et al., 2009; Raab et al., 2007). This might explain that DMA\textsuperscript{V} was detected in flowers of D. morifolium and not in roots of CHMs.
Arsenic speciation in CHMs samples collected from markets or pharmacies is listed in Table 3. In general, inorganic arsenic, AsIII and AsV, was the predominant species with the proportion over 94% like that in CHMs from fields. DMA\(^\text{\textregistered}\) was only detected in roots of *A. asphodeloides* and shoots of *T. kirilowii* from markets at 7.3 ng/g and 15.8 ng/g, respectively. The peak of un-identified As speciation was detected at retention time between 500 s and 600 s, separated after As\(^\text{\textregistered}\) peak in a few samples including *L. indigotica, A. asphodeloides* and *D. morifolium* from markets (Table 3 and see Inline Supplementary Fig.S1(b)), It would be done further analysis about this un-identified arsenic species in CHMs in the future. The DMA\(^\text{\textregistered}\) percentage was less than 5% of total arsenic in these samples. Whether AsIII and AsV concentrations or total As concentrations in CHMs from markets or pharmacies (*L. indigotica* (shoots), *A. Tataricus, A. asphodeloides, T. kirilowii* (root), *T. kirilowii* (fruits), *D. morifolium*) were much higher than those in the same CHMs collected directly from fields (Tables 2 and 3), accounting for 19–30 times. This indicated that arsenic accumulation and speciation in these CHMs from markets have been impacted during dryness, storage, transport and manufacturing processes of herbs. Moreover, the highest mean concentrations of AsIII and AsV were 2443 ng/g, 1560 ng/g and 3790 ng/g in flowers of *D. morifolium* from markets and pharmacies, which is related to the process of sulfur fumigation of the flowers of *D. morifolium*. In general, sulfur fumigation is commonly used to reduce the damage of insects and fungus on herbs, especially during processes of the dryness and storage of Chrysanthemum flowers. Sulfur powder, from natural stone with sulfur, is normally combined with a little amount of arsenic and produces the gas with arsenic contaminating Chrysanthemum flowers during fumigation. Although State Food and Drug Administration in China issued the regulation to ban using sulfur power to fumigate CHMs in 2004 for harmful gases emission, human health assessments of arsenic accumulation and speciation in Chrysanthemum flowers from the markets should do further investigation to make this point clear.

The levels of inorganic As in edible plants are closely related to the toxicity of arsenic to human health (Schoof et al., 1999; Zavala et al., 2008). When data from all of field collected samples plotted together, a striking pattern can be observed that the concentration of inorganic As increased with increasing total As (*r* = 0.720, *P* < 0.01) (Fig. 1a). The same pattern also appeared in data plot between inorganic As and total As in CHMs from markets or pharmacies (*r* = 0.987, *P* < 0.01) (Fig.1b). This relationship between inorganic As and total As was also reported in rice grain samples by Xu et al. (2008) and Zavala et al. (2008).

The recoveries of arsenic species varied from 48% to 160% for different CHMs samples from fields and markets or pharmacies (Tables 2 and 3). The mean recovery of As species from shoot of *T. kirilowii* was 158% and the recovery from root of *S. miltiorrhiza* was around 50%. The variation of recoveries for arsenic species has been reported in rice samples by Heitkemper et al. (2001) from 31 to 75% and in bran and flour of wheat by Zhao et al. (2010) from 57%
to 218% using the same extract solution (1% HNO₃) as that in this study. The former article indicated that it was difficult to explain the large differences in extraction efficiency and recovery among different samples. Zhao et al. (2010) gave the explanation for anomalously high recovery of 218% in white flour of wheat that total As concentration in this sample was very low at 21 ng/g, compared with 50 e 205 ng/g in other samples. Our results also followed this explanation (Tables 2 and 3).

For As species determination in the samples from this survey, the percentages of methylated As were no more than 10% for all CHMs growing in Anguo area of Hebei Province. However, the proportion of methylated arsenic in the *Panax notoginseng* sample was 31% of total As using the same method. In addition, unidentified As speciation in *Panax notoginseng* differed from that in other CHMs samples. The diversity of As speciation and its proportion between samples may be attributed to the different herbal varieties and growth regions relevant to *Panax notoginseng* growing in Yunnan province and 12 species of CHMs growing in Hebei province. The soil properties and climate conditions are totally different between these two areas and lead to differences in methylated As levels among CHMs. Williams et al. (2006, 2005) also investigated arsenic speciation in different rice varieties from different locations around the globe to explore the contributions of rice to arsenic exposure and demonstrated the predominant species detected in rice from European, Bangladeshi, and Indian being inorganic arsenic but DMA being a main component in rice from the USA. Although Zavala et al. (2008) indicated that genetic differences lead to arsenic speciation variation finally, processing and cultivating conditions are also reasons for the diversity of arsenic speciation in edible plants (Meharg et al., 2008; Smith et al., 2008).

### 3.3. Human health implication for inorganic As

The predominant arsenic species in samples of CHMs using 1% HNO₃ extraction was inorganic As. Inorganic arsenic concentrations ranged from 62.7 to 550 ng/g (*n* = 26) with a mean of 208 ng/g in CHMs from fields (Fig. 1a) and ranged from 93.5 to 8683 ng/g (*n* = 39) with a mean of 1092 ng/g in CHMs from markets or pharmacies (Fig. 1b). The Chinese Maximum Contaminant Levels (MCLs) of inorganic As in foods was presented in 2005, such as rice (150 ng/g), wheat flour (100 ng/g), non-staple cereals (200 ng/g) and vegetables (50 ng/g/China Food Standard Agency, 2005), not including the maximum limit of inorganic As level for CHMs. If the MCL of inorganic As in non-staple cereal crops (200 ng/g) was referred in this study for the similar growing conditions to discuss the health risk assessment of inorganic arsenic in CHMs, about 65% of the CHMs from field and 25% of the CHMs from markets or pharmacies would be legal for sale in China; If compared to MCL of inorganic As in vegetables (50 ng/g), inorganic As concentrations in all of CHMs exceeded this limitation.

To estimate the daily intake of arsenic from CHMs, the dose ingested should be taken into account. In accordance with the conservative principle, the maximum dose of each CHMs prescribed by the Chinese Pharmacopoeia was selected in the risk calculation (for example 9 g/d was selected as the dose in the risk calculation when 4.5 e 9 g/d was recommended for *S. divaricata* in Chinese Pharmacopoeia) (Chinese Pharmacopoeia Commission, 2005). If the inorganic As concentration in CHMs is 200 ng/g (MCL) and consumption of 30 g CHMs by a 60 kg person daily, the proportion of the daily intake value to the ADI (126 mg) from FAO/WHO is 4.76%, which can be explained that a potential risk will exist if the proportion value of daily intake for inorganic As in CHMs is higher than 4.76%. So, the percentage of inorganic As in CHMs to ADI was calculated and shown in the Fig. 2. According to our data, daily contribution of inorganic arsenic in all samples from field

![Fig. 1. Relationship between the concentrations of inorganic As and total As in CHMs (a: samples collected directly from fields, r = 0.720, P < 0.01, n = 26; b: samples collected from markets or pharmacies, r = 0.987, P < 0.01, n = 39).](image)

![Fig. 2. Comparison of percentages of dietary exposure for inorganic arsenic inducing by CHMs collected from fields and from markets or pharmacies.](image)
ranged from 0.94 µg to 4.95 µg, and counting for 0.75—2.83% of ADI. The range of daily intake of inorganic arsenic in all samples from markets or pharmacies was from 1.29 µg to 78.2 µg with the risk values of 1.02—62.0%. The comparison of percentages of daily intake for inorganic arsenic in CHMs from fields and markets or pharmacies was shown in Fig. 2. For the CHMs of L. indigota (shoots), A. Tataricus, A. asphodeloides, T. kirilowii (root), T. kirilowii (fruits), and D. morifolium, the proportions of inorganic As daily intake to ADI in CHMs from markets or pharmacies were much higher than that from fields.

According to the highest boundary of estimated inorganic As exposure from food (1.22 µg/kg b.w. per day) published by EFSA (2009), ADI of inorganic As for one person with 60 kg body-weight will be 73.2 µg, which is only 58.1% of ADI (126 µg) from FAO/WHO. If ADI (73.2 µg) is adapted in this health risk assessment of inorganic As in CHMs, the contribution of daily intake of inorganic As from all of CHMs to health risk will increase significantly.

4. Conclusion

Inorganic arsenic was the predominant species detected in all of CHMs samples with the proportion of 94—100%. Inorganic arsenic levels in CHMs from fields and markets or pharmacies ranged from 62.7 to 550 ng/g and 94 to 8683 ng/g, respectively. The highest concentration of inorganic arsenic was found in the flowers of D. morifolium from pharmacies. The study proves useful information for a better understanding of the distribution of arsenic species in Chinese Herbal Medicines. In addition, the risk of inorganic As in CHMs from markets or pharmacies to human health should be paid more attention further.

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Appendix A. Supplementary data

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