INTRODUCTION

Rice (Oryza sativa L.) is one of the main foods in many countries, and this explains why annual production is high. The United States Department of Agriculture (USDA) reported that production in 2014 was 474.86 million tons (ODEPA, 2014), and the major producers were China, India, Indonesia, Bangladesh, and Vietnam. Some of these countries report soil and groundwater contamination with high arsenic (As) levels (Rahman et al., 2014a). Rice is characterized as an efficient As accumulator because of the biogeochemical characteristics of agricultural soils and its ability to absorb and transport As (Kumar et al., 2015). The International Agency for Research on Cancer has classified inorganic arsenic (iAs) as arsenite (AsIII) and arsenate (AsV) as type 1 carcinogenic agents (IARC, 2012); hence, its importance for public health. Therefore, how to decrease As exposure in rice is an important concern for the scientific community (Rahman et al., 2014a). This review explains possible mechanisms involved in As absorption that contaminate the rice plant through the soil and water, and mentions studies that have been conducted to minimize the risk for human exposure.

Arsenic in rice cultivation

It is estimated that more than 150 million people worldwide have been affected by increased As concentrations. Oral ingestion is the major human route of exposure (Welna et al., 2015). Arsenic contamination has been reported in many places in the world, such as Bangladesh, India, China, Turkey, Argentina, Chile, and Colombia (Bundschuh et al., 2012; Gan et al., 2014; Tong et al., 2014). This element is widely distributed in nature, and its accumulation in soils is caused by the use of pesticides, fertilizers, and the burning of fossil fuels; its level in soils depends on the anthropogenic activity, the distance from the source of contamination, as well as contaminant dispersion (Bastías et al., 2013a; 2013b).

Toxicity of arsenic species and their occurrence in rice cultivation

Arsenic toxicity is based on species (Jomova et al., 2011). The rice grain contains both organic and inorganic As (iAs). Arsenite (AsIII) and dimethylarsenic acid (DMA) are the typical species. Furthermore, it can contain arsenate (AsV), monomethylarsenic acid (MMA), and occasionally tetramethylarsonate. Other authors...
include monomethylarsenic acid (MMA), arsenobetaine (AsB), and arsenocholine (AsC) (Meharg and Zhao, 2012).

Inorganic As is mainly found as AsV in aerobic environments and water, while AsIII prevails in anoxic and neutral environments. However, they can interconvert when there are changes in the redox potential, pH, and the occurrence of microorganisms (Zhao et al., 2010; Nearing et al., 2014). The bacteria that reduce (AsV) and oxidize AsIII are found in the soil. Arsenate reduction by microorganisms occurs through two mechanisms: AsV reduction serves as a terminal electron acceptor during anaerobic respiration and detoxification in which AsV is reduced into AsIII and pumped out of the microbial cells (Zhao et al., 2010).

It has been observed that AsIII is 10 times more toxic than AsV; monomethylarsonate (MMAIII) and dimethylarsonate (DMAIII) are more toxic than AsV and AsIII. In turn, these are more toxic than monomethylarsonate (MMAV) and dimethylarsenate (DMAV). The least toxic is trimethylarsenic oxide (TMAO) (Rahman and Hassler, 2014).

Biomethylation of iAs species has been considered as a potential detoxification mechanism of microorganisms; this results in the production of trivalent methylarsenic species (MMAIII and DMAIII) as shown in Figure 1. Both MMAIII and DMAIII are intermediaries in the biosynthetic route of pentavalent methylarsenic species (MMAV and DMAV), and they are the most toxic forms of As (Dopp et al., 2010). Trivalent intermediaries are structurally different from their pentavalent counterparts; they are more reactive and less carcinogenic (Rahman and Hassler, 2014).

![Figure 1. Possible routes of arsenic methylation by microorganisms and its excretion outside the phytoplankton microbial cell.](image)

AsV: Arsenate, AsIII: arsenite, GSH: glutathione, MMA: monomethylarsonate, DMA: dimethylarsenate.

Table 1. Elements in the soil that interact with arsenic.

<table>
<thead>
<tr>
<th>Elements in soil</th>
<th>Importance</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Iron</td>
<td>Fe serves as a strong adsorbent for As.</td>
<td>Zhao et al., 2010; Pan et al., 2014</td>
</tr>
<tr>
<td></td>
<td>Reductive dissolution of Fe oxyhydroxides under a reducing environment releases adsorbed As, leading to enhanced As availability to plants.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Fe-reducing bacteria are linked to As mobilization.</td>
<td></td>
</tr>
<tr>
<td>Phosphorus</td>
<td>Arsenate enters plant cells via phosphate transporters and also interferes with phosphate metabolism.</td>
<td>Rahman et al., 2014b</td>
</tr>
<tr>
<td>Sulfur</td>
<td>S helps to detoxify As through the complexation of arsenite with thiol-rich peptides; these compounds can also help to maintain As in the roots and restrict its translocation to the tillers.</td>
<td>Rai et al., 2011; Rahman and Hassler, 2014</td>
</tr>
<tr>
<td></td>
<td>It is particularly important to use adequate amounts of S in soil fertilization in As-contaminated environments.</td>
<td></td>
</tr>
<tr>
<td>Silicon</td>
<td>Si fertilization can be an effective strategy to decrease As accumulation in rice grown in As-contaminated soil because it increases the levels of antioxidant enzymes, isozymes, cysteine, GSH, and NPSH; it also reduces As-induced lipid peroxidation in rice.</td>
<td>Tripathi et al., 2013</td>
</tr>
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</table>

To reduce the ingestion of As, it is essential to know and identify the mechanisms involved in the accumulation and detoxification of rice. Its molecular mechanisms of intake, metabolism, and translocation have been identified in recent years. The chemistry of As in the rhizosphere is complex and controlled by several factors. There are four important elements that take part in As intake in rice: Fe, P, S, and Si (Table 1) (Zhao et al., 2010).

Iron (Fe) plays an important role in the biogeochemical cycle of As. Iron oxyhydroxides in the soil and/or on the plant root surface function as a strong As adsorbent. The reductive dissolution of Fe oxyhydroxides in a reducing environment releases As that is adsorbed by the rice plant, which leads to greater bioavailability of the metalloid. Oxygen is transported in the roots through root aerenchyma where it is consumed by the adjacent tissue cells or diffused toward the root apex or the rhizosphere. Oxygen transfer from the aerenchyma to the rhizosphere is called radial oxygen loss (ROL). It can oxidize elements in the rhizosphere soil, for example, Fe2+ to Fe3+, and cause the precipitation of toxic metals in the rhizosphere soil and on the root surface (Pan et al., 2014).

Mobility of the metal is then altered in the rhizosphere. Aerenchymae develop in the root plants to transfer O2 from the aerial parts of the plant to the roots, which results in the oxidation of ferrous iron into ferric iron and the precipitation of Fe oxides or hydroxides on the root surface (Pan et al., 2014). The Fe coating can sequester metals in wetland plant roots and modify As translocation from the roots to the shoots (Rahman et al., 2014c). Rhizosphere interactions therefore play a key role in controlling As bioavailability in rice cultivation (Deng et al., 2010).

Chemical reactions could occur in reducing environments under anoxic conditions that dissolve Fe minerals to which As is bound (Michael, 2013). The correlation between available soil Fe and pH is negative and highly significant, while the correlation is weak between available soil Fe and As found in the rice grain. One of the main Fe sources is the Fe-Mn-As complex, which creates a more stable binary complex in the active site of the enzyme than AsIII (Rahman and Hassler, 2014).
fraction. The weak correlation between available Fe and iAs in the rice grain could be due to the Fe-Mn-As fractions or pH that influence available Fe (Jiang et al., 2014).

Another element is phosphorus (P). Because it is an analogue of AsV phosphate (Pi), this element could enter the cells through Pi transporters, which can interfere with the metabolism of the latter. Phosphate competes with AsV and also with root adsorption of oxides and Fe hydroxides. When P concentration increases, As uptake decreases. Some studies with different types of soil at different P and Fe concentrations found that there is a positive correlation between available Fe and iAs concentrations in the grain, while the correlation is negative when soil has available P (Jiang et al., 2014). It is also assumed that As mobility in organic acid-rich soils is reduced because the acid acts as a binding agent and/or in the formation of insoluble compounds that do not allow As to be absorbed by the roots (Jiang et al., 2014). Arsenate can be incorporated in the ATP (adenosine triphosphate) by replacing phosphates in the binding sites. This substitution, along with the inhibition of oxidative phosphorylation and interference in the correct ATP synthesis, increases AsV toxicity (Rahman et al., 2014b).

In the case of sulfur (S), this helps to detoxify As through the binding of AsIII with thiol-rich peptides. This formation of compounds can also help to maintain As in the roots and restrict its translocation to the tillers. Arsenite is similar to sulphhydryl groups existing in cysteine residues that have a detrimental effect on the general metabolism of proteins, which increases its toxicity (Rai et al., 2011). Arsenite inhibits pyruvate dehydrogenase by binding to dihydrodiposphoamide sulphhydryl groups by decreasing the conversion of pyruvate into acetyl-coenzyme A (CoA) in el original tenia una referencia aqui (Bergquist et al., 2009). It also inhibits glutathione production that protects cells against oxidative damage and phytochelatin (PC) production (of which glutathione is a crucial component) as a result of which phytoplankton and plants lose the ability to detoxify numerous heavy metals (Rahman and Hassler, 2014). Maintaining enough S in soil fertilization can be particularly important in environments contaminated with As.

The role of silicon (LSi1) transporters in As uptake is particularly important because similarities already exist between silicic acid and arsenious acid; this allows arsenious acid to penetrate into the root by the silicic acid transporter. The presence of Si decreases As phytotoxicity and increases the levels of antioxidant enzymes and their isozymes. It also enhances the concentrations of cysteine, GSH, and NPSH, and reduces As-induced lipid peroxidation in rice, Silicon fertilization can also be an effective strategy to decrease As accumulation in rice grown in As-contaminated soil (Tripathi et al., 2013).

Proposed mechanisms for arsenic translocation in rice plant

Arsenic is able to translocate from the roots to the tiller in rice. Some studies, conducted with different types of rice, have shown that there are differences in As concentration in plant parts with a decrease from the root to the grain (Welna et al., 2015). The highest As concentrations were in the root (248 ± 65 mg kg⁻¹) and the lowest in the grain (1.25 ± 0.23 mg kg⁻¹) (Abedin et al., 2002). Some research studies have suggested that there is low As accumulation in the rice grain although plants grow in contaminated environments with high concentrations of the metalloid. Under hydroponic conditions, As concentrations in roots was 107.5 mg kg⁻¹, while they decreased between 0.15 and 0.42 mg kg⁻¹ in the grain (Abedin et al., 2002).

Metal-accumulating plants can translocate this element rapidly and efficiently to the tillers via the xylem. This implies that the xylem is loaded with this metalloid, passes it on to the vacuoles, and this is due to the characteristics of the root cell tonoplast (Rascio and Navari-Izzo, 2011). The root absorbs As as AsV and AsIII. Arsenate penetrates because it is highly similar to the Pi transporters that belong to the PHT1 family (Ali et al., 2009). The plasma membrane of the root cells also has a higher density of Pi/AsV transporters than other plants that are not As accumulators because of constitutive gene overexpression (Rascio and Navari-Izzo, 2011). Arsenite is absorbed through the aquaglycoprotein NPs (nodulin-like intrinsic proteins) (Meharg and Jardine, 2003) and by LS1 transporters (Rascio and Navari-Izzo, 2011). The methylated forms of As, DMA, and MMA are absorbed through the aquaporins and use the same glyceral mechanism (Rahman et al., 2011).

Arsenate in the root cells is reduced by As reductase (AR) to AsIII and leads to the conversion of glutathione (GSH) in its oxidized form (GSSG). Arsenite is transformed into trimethylarsenic oxide (TMAOV) and trimethylarsine oxide (TMAOIII). The final product of the methylation route and the As volatile species are released into the environment. Another As detoxification route occurs by PC synthesis because of the condensation of three amino acids: cysteine, glutamate (Glu), and glycine (Gly). Finally, the sequestration of the AsIII-PC compound occurs within the vacuole through the activity of the ABC transporters (Rahman and Hassler, 2014). Although AsIII is more toxic than AsV, it can bind to proteins or peptides that contain thiol groups, such as glutathione, phytochelatins, and metallothioneins, and form an inactive compound (Tsai et al., 2009) that protects the cell components of this toxin (Ali et al., 2009). The AsIII compounds are sequestered by the vacuole, but this mechanism is still unknown (Kumar et al., 2015). The main As compounds are AsV and AsIII that are found in the xylem sap of rice. Certain studies report that AsIII predominates in 80% of the analyzed sap samples in different rice species, including in those grown in AsV-rich soils; this suggests that reduction occurs in the root cells before passing to the xylem and the rest of the plant parts (Su et al., 2010). Other studies demonstrate the role played by the rice plasma membrane intrinsic proteins (OsPiP) in the permeability of AsIII (Mosa et al., 2012).

Of the total As (ToAs) absorbed by the root, only a very small part is sequestered in the leaf and grain vacuoles
because the reduction and sequestration mechanisms are similar to those of the roots. The presence of different As species in the phloem is a prerequisite for it to be distributed to the rest of the plant (Yu et al., 2010). The possible As absorption mechanisms are shown in Figure 2.

**Factors influencing arsenic absorption in the rice plant**

There are studies that demonstrate a significant correlation between root porosity, tolerance, and As accumulation in rice (Wu et al., 2011). Other studies show that radial oxygen loss (ROL) rates were negatively correlated with As concentrations in rice roots among different genotypes. Rice genotypes with higher porosity can sequester less As of the Fe coating in the root, reduce its translocation to the plant, and finally to the grain (Deng et al., 2010).

Soil As accumulation over a long period of time increases its bioavailability for absorption by plant roots. Arsenic concentration is clearly dependent on rice variety (Rahman et al., 2014c), grain size and color (Meharg et al., 2008), procurement processes, and production environment (Zhao et al., 2010). Some studies report that As contamination in the rice grain is positively influenced by the amount of As in paddy irrigation water; however, no influence has been observed for As concentration in soils (Zavala and Duxbury, 2008).

The genotypes TD71 and Yinjingruanzhan contain less iAs and ToAs in their grains than genotypes IAPAR9 and Nanyangzhan. Genetic variation and As speciation are the two factors that influence plant As absorption (Wu et al., 2011; Pan et al., 2014). Regardless of genotype, all the studies show that iAs species predominate in rice grains, but some authors report that the genotype Nanyangzhan mainly contains DMA (Pan et al., 2014). Other authors have reported that different rice genotypes grown in greenhouses had As concentrations ranging between 8.3 and 14.0 mg kg$^{-1}$ in the leaf and between 0.87 and 1.52 mg kg$^{-1}$ in the grain; however, this study suggests that rice accumulates significant As concentrations under favorable growth conditions (Jomova et al., 2011).

*Figure 2. Possible mechanisms of arsenic uptake in rice plants (adapted from Ali et al., 2009).*

When grain size increases, As concentration increases (Zavala and Duxbury, 2008). Findings related to color showed that As concentration is 0.196 ± 0.111 mg kg⁻¹ in brown rice, 0.127 ± 0.087 mg kg⁻¹ in white rice, and 0.07 ± 0.05 mg kg⁻¹ for other colors (Zavala and Duxbury, 2008; Sharma et al., 2014). The highest amount of As is in brown rice because its outer layers have a higher content of the metalloid.

Some studies suggest that rice plants lack the ability to methylate As, and its speciation in the grain is mainly attributable to environmental factors such as the methylated As species found in soil (Ma et al., 2014). The main species found in rice from Europe, Bangladesh, India, and the USA are As[III], DMA[IV], and As[V] (Welna et al., 2015).

Arsenic species change in aquatic environments because of the role played by microorganisms (Figure 1), which are particularly active in terms of As methylation under anaerobic conditions of rice cultivation (Lomax et al., 2012). Multiple theories have been proposed to explain the influence of microorganisms on As biotransformation. These reactions (generally oxidation or reduction) are produced by the microorganisms to protect themselves from the toxic effects of this metalloid or generate energy and ensure their growth (Mateos et al., 2010); the biotransformation includes the oxidation of As[III] to As[V] and methylation to methylarsenic forms (Mateos et al., 2010). Among the microorganisms with the proven ability to generate these biotransformations are algae (Cyanidium) and some bacteria (Hydrogenobacter acidophilus). The oxidation mechanism of As[III] by microorganisms includes the chemolithotrophic metabolism used as a source of energy (Rahman and Hassler, 2014).

**Current legislation for arsenic level in rice grain**

The As concentration permitted by the Codex Alimentarius is 0.3 mg kg⁻¹ for ToAs and 0.2 mg kg⁻¹ for iAs in raw rice (Codex Alimentarius, 2013). Chile has established 0.5 mg kg⁻¹ for cereals and legumes (RSA, 2011) and the same limits permitted by the Codex Alimentarius for rice (Villanueva, 2014).

The accuracy for identifying As compounds depends on the analysis method that is selected (Nearing et al., 2014). Many techniques have been used and innovated to increase both the specificity level and accuracy in results. Most of the studies use the inductively coupled plasma-mass spectrometry (ICP-MS) technique coupled to high performance liquid chromatography (HPLC). This is the internationally validated method used to determine ToAs. The two most reported complementary analytical methods for As speciation are X-ray absorption spectroscopy (XAS) and electrospray mass spectrometry (ESI-MS); however, other studies propose different techniques (Table 2).

There are different methods in HPLC that have been used because they have high selectivity and a low detection limit (< 50 pg), for example, ion exchange (Raber et al., 2012) or ion-pairing reverse phase (Dufailly et al., 2011). HPLC-ICP-MS requires procedures to extract As compounds that can be in methanol-water and water extractions. An intermediate heating step with diluted nitric acid to extract iAs species is necessary to increase crop efficiency; efficiency is 90% when this methodology is used in rice samples. It has been observed in all cases that some As compounds are more easily extracted than others, and this depends on the selected extraction method and sample matrix. A disadvantage of this technique is that the chromatograms provide unstructured information about the separated compounds and the number of patterns is limited (Nearing et al., 2014).

To obtain the necessary detection limits in the XAS method, radiation based on synchrotron is used (giving typical detection limits of approximately 1 to 10 mg kg⁻¹ depending on the experimental conditions). The principles of X-ray fluorescence are generally used in the XAS method for As in environmental samples (Nearing et al., 2014). The X-ray absorption spectra are characterized by a sharp increase in absorption at specific X-ray photon energies giving rise to an absorption edge that characterizes the element. The problem is that the energy is the same for similar As compounds isolated from the same environments; this is one of the limiting factors of the technique. The most common use of XAS for As speciation is the analysis of X-ray absorption near-edge structure (XANES), which reveals the most probable chemical environment around As by matching the spectral characteristics, including white-line energy, to the standard compounds. This method has provided good results for the speciation of As compounds in rice (Maher et al., 2013). Studies comparing both methods demonstrate that compounds such as As[III]-S have not been found by the traditional method, but have been found by XAS; this has been attributed to its oxidation during extraction with nitric acid. Analysis with XAS found As[III]-S compounds in 55.6% of As in rice endosperm and 12.9% in the husk (Carey et al., 2012).

The use of ESI-MS to complement HPLC-ICP-MS in determining different As species in rice allowed the tentative identification of DMA, a species not previously found in rice. This method has provided good results in identifying As[III].

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### Table 2. Principal genetic modifications to obtain seeds with lower ability to accumulate arsenic (Kumar et al., 2015).

<table>
<thead>
<tr>
<th>Source</th>
<th>Target</th>
<th>Gene</th>
<th>Consequence</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oryza sativa</td>
<td>O. sativa</td>
<td>As[III] S-adenosyl transferase (arsM)</td>
<td>Decreases As in rice grain</td>
<td>Kumar et al., 2015</td>
</tr>
<tr>
<td>O. sativa</td>
<td>O. sativa</td>
<td>Phosphate transporter (PH1.8)</td>
<td>Increases phosphate and As[V] absorption and translocation</td>
<td>Wu et al., 2011</td>
</tr>
<tr>
<td>O. sativa</td>
<td>O. sativa</td>
<td>Response to phosphate 2 requirement (PHR2)</td>
<td>Increases tolerance to As and greater As[V] efflux capacity</td>
<td>Wu et al., 2011</td>
</tr>
<tr>
<td>Saccharomyces cerevisiae</td>
<td>O. sativa</td>
<td>Arsenate reductase (ACR3)</td>
<td>Increases As[III] efflux</td>
<td>Kumar et al., 2015</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Decreases As accumulation in rice grain</td>
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</tbody>
</table>
compounds (Cao et al., 2015). Another methodology used to
determine and quantify As is radiochemical neutron activation
analysis (RNAA). It uses the irradiation in a nuclear reactor
with $1.8 \times 10^{12} \text{ n cm}^{-2} \text{s}^{-1}$ neutron flux; its As detection limit
in the sample is in the ng g$^{-1}$ range (Halder et al., 2014). In
recent years, analytical techniques have been worked on for
As speciation and coupling of a hydride generator coupled
to ICP-MS to measure ToAs in rice. Total As is directly
measured by this method without any previous treatment of
samples. The technique determines all the main As species:
As$^{\text{III}}$, DMA, MMA, and As$^{\text{V}}$; it eliminates the difference of
sensitivity among different organic and inorganic As species
when measuring ToAs (Sengupta and Dasgupta, 2009).

While it is true that methods have been developed to
allow the detection of ToAs along with its species, reagents
introduced in the different techniques to treat samples could
transform the species; it is therefore important to develop
analytical methods that directly detect the original species
in the rice grains (Welna et al., 2015). Nondestructive
techniques have been used for this, such as X-ray fluorescence
spectrometry (XRF), X-ray absorption near edge structure
(XANES), instrumental neutron activation analysis (INNA),
and near-infrared spectroscopy (NIRS). No chemical
products are used in these instrumental methods and the
manipulation of the sample is reduced to washing and/or
drying and grinding before analysis (Carey et al., 2012).

**Risk of As from rice diet in human health**

No epidemiological studies have been published about the
health effects associated with ingesting iAs through rice
cConsumption. Bioavailability should be included in the
assessment to evaluate the risk of iAs on human health. There
are many possible pathways for human exposure to As, but
rice is the second most important in many countries (Rahman
et al., 2008). It is important to initiate epidemiological
studies about the implication of As on health because it will
take decades to understand how elevated rates of As in rice
affects human lifetime exposure (Rahman et al., 2014c).

Diet, lipids in particular, has an important effect on
As bioavailability. Not all the constituents found in the
food matrix are bioavailable (Bastías et al., 2013b). Some
studies have demonstrated that thiol groups are strongly
bound to As$^{\text{III}}$ in certain protein-rich rice varieties, which
can influence its bioavailability (Rahman et al., 2014c).
Other studies demonstrated a relationship between diet
composition and MMA$^{\text{III}}$ levels found in urine after iAs
intake; furthermore, those whose diet was relatively high in
vegetables had less probability of developing skin lesions
related to As and exhibit differences in the metabolic rate of
As consumed (Koch et al., 2013).

After the oral intake of As-contaminated rice,
gastrointestinal digestion can release a fraction of the bound
matrix and make it accessible. The As fraction that is not
absorbed through the small intestine epithelium reaches the
colon. Here microorganisms decompose the remainder of
the food and more As is released. Some studies have revealed
that microorganisms are important contributors in the As
speciation changes. Studies exist that explain the role of
bacterial biota in the gastrointestinal tract over a wide range
of biotransformations that the metalloid undergoes during
digestion, including oxidation, reduction, methylation, and
thiolation (Alava et al., 2015).

A high amount of As has been found released in the
food matrix at the gastric level that can be altered by food
composition. Therefore, the difference in diet composition
can also differ from the amount of As reaching the
colon and the amount of As available for pre-systemic
metabolism. The acidic conditions of the stomach increase
As bioaccessibility more than in the intestine (Alava et al.,
2015). Other studies have observed that iAs$^{\text{V}}$ is released
proportionally more easily than DMA$^{\text{V}}$ from the rice matrix
during gastrointestinal digestion. It has been found that iAs
species are more likely to bind to the protein matrix of the
endosperm cells, especially by the amino acid compounds
that contain thiol (Alava et al., 2015).

Different studies show that real exposure to ToAs and iAs
through foods depends on the type of food processing, time,
temperature, and medium, especially cooking medium. It is
absolutely necessary to cook rice because it is not consumed
raw (Rahman et al., 2014c). Arsenic concentration has been
observed to be lower in dry rice husking than wetting rice
with contaminated water before husking it. Another method
is to cook one part rice in six parts water; the disadvantage
of this method is the higher consumption of energy and water
(Rahman et al., 2014c).

**Remediation techniques**

Arsenic affects humans and is a public health problem. Better
agricultural irrigation practices are therefore proposed to
help reduce contamination; one of these techniques is that
crops be grown in raised organoponics instead of flooded
fields (Rahman et al., 2014c). The code of good agricultural
practices for rice cultivation in Chile also proposes using
only certified seeds. Selecting the variety to be sown should
be based on seeds that are better adapted to the production
zone, and the number of seeds per hectare will vary according
to the type used, soil preparation conditions, sowing system,
and water management.

Studies of soil and water remediation continue using
phytoremediation and mycoremediation. In the case of
As, the use of a hyperaccumulator fern, *Pteris vittata* L.,
has been proposed (Rascio and Navari-Izzo, 2011). The
limitation of this method is that the plant absorbs As without
using it and can be transferred back to the food chain (Singh
et al., 2015). Fungi have the ability to counteract As toxicity
by transforming organic forms with reduced toxicity (Tsai
et al., 2009). The behavior of *Glomus geosporum* (Gg),
*G. mosseae* (Gm), and *G. versiforme* (Gv) was studied to
minimize As absorption by the rice plant; it was found that
species, taken separately or mixed, can be used because
the amount of As decreases in all cases (Chan et al., 2013).
Redox changes mediated by rhizosphere microorganisms
can influence As absorption in the rice plant (Jia et al., 2014). However, the participation of microbes in As oxidation and reduction during the interaction with the root has not been studied in detail.

Using seeds with modified As absorption genes is another alternative. By restricting the influx of As, its toxicity is avoided (Rahman et al., 2014c). The main variations have been performed on phosphate transporter genes (Wu et al., 2011) and *arsM* (Adenosyltransferase), which decrease As in rice. The gene *PHT 1;8* and *PHR 2* increase root tolerance to As in the rice plant and also increase the efflux ability of As. The main genetic modifications to obtain seeds with lower As accumulation ability are shown in Table 2.

Soil fertilization with Si is advantageous because it decreases ToAs concentration in the straw and grain, 78% and 16%, respectively, although adding Si increases As concentration in the soil solution. Silicon decreases iAs concentration in the grain by 59% (Tripathi et al., 2013). On the other hand, Fe is an important factor in regulating bioavailability and As absorption in rice plants because it creates a zone around the roots in which it is oxidized and precipitated to form a sheath, and As has a high binding affinity with the precipitated Fe hydroxides that act as an As filter to reduce its absorption by the rice plant (Garnier et al., 2010). Sulfur improves the formation of Fe coating in the rhizosphere and reduces As accumulation, but adding nitrate decreases the formation of Fe coating on the root surface of rice; however, S reduces As absorption by the rice plant (Hu et al., 2015).

**CONCLUSIONS**

The root of the rice plant is able to absorb and accumulate large amounts of As, but only small amounts are translocated to the grain and tillers. Arsenic concentrations in rice tissues decrease from the root to the grain. Information about As translocation in rice is sparse and research is directed toward studying the molecular mechanism of absorption and accumulation in the grain because it has not yet been explained. Some rice varieties have been developed that are resistant to high soil As concentrations and are not able to translocate the metalloid toward the root.

Many studies suggest that not all ingested iAs accumulated in the gastrointestinal tract is absorbed into the bloodstream and produces toxicity. It is therefore recommended that As species found in different types of diets be studied, as well as the influence of bacterial biota of the gastrointestinal tract on the bioavailability of different As species and their release after intake. Arsenic bioavailability in imported or domestic Chilean rice should be evaluated, thus contributing to studies estimating its exposure and providing a more precise risk estimate.

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