

International Headquarters & Laboratory Phone 630 505 0160

WWW.WQA.ORG

A not-for-profit organization

ARSENIC FACT SHEET

Contaminant	In Water As	Maximum Contaminant Level
Arsenic (As)	When pH = 6-10	US EPA:
	As(III), arsenite as H_3AsO_3 and $H_2AsO_3^{1-}$	MCL* = 0.010 mg/L (or ppm
	As(V), arsenate as $H_2AsO_4^{1-}$ and $HAsO_4^{2-}$	MCLG** = zero
		WHO [†] Guideline = 0.010 mg/L
	Leaching from natural deposits	
Sources of Contaminant	Wood preservatives, pesticides, industrial deposits	
	Petroleum production	
	Semiconductor manufacture	
	Coal power plants	
Potential Health Effects	Serious skin problems, endocrine disruptor	
	Causes Cancer – skin, bladder, lung, kidney, liver, prostate	
	Harms cardiovascular &nervous systems	
	Iron oxide/ hydroxides	
	Activated alumina	
Treatment Methods	Iron based specialty media impregnated or coated with iron oxide/hydroxide	
Point-of-Entry Point-of-Use	Distillation	
	Titanium oxy/hydroxide	
	Anion Exchange (strong base anion exchange resins)	
	Manganese greensand	
	Reverse osmosis (RO)	
*Maximum Contaminant Level (MC	CL) - The highest level of a contaminant that is allowed in drin	king water. MCLs are set as close to
•	lable treatment technology and taking cost into consideration al (MCLG) - The level of a contaminant in drinking water belo	

risk to health. MCLGs allow for a margin of safety and are non-enforceable public health goals. WHO⁺ - World Health Organization

The presence of arsenic (As) in nature is due mainly to natural deposits of metalloids in the earth's crust and usually in ancient rock formations. Arsenic enters ground water through erosion or from manmade sources such as wood preservative, petroleum production, semi-conductor manufacture or due to misuse of animal feed additives and arsenic-containing pesticides (e.g. Paris green). Since soluble arsenic is tasteless and colorless, a chemical water analysis is necessary to detect its presence.

Higher levels of arsenic tend to be found more in ground water sources than in surface water sources (lakes and rivers) of drinking water. Compared to the rest of the United States, the western states have more water systems with arsenic levels greater than 10 micrograms per liter (µg/L, ppb).

Parts of Midwest, New England, and Texas have some systems whose current arsenic levels are greater than 10 μ g/L (or ppb).

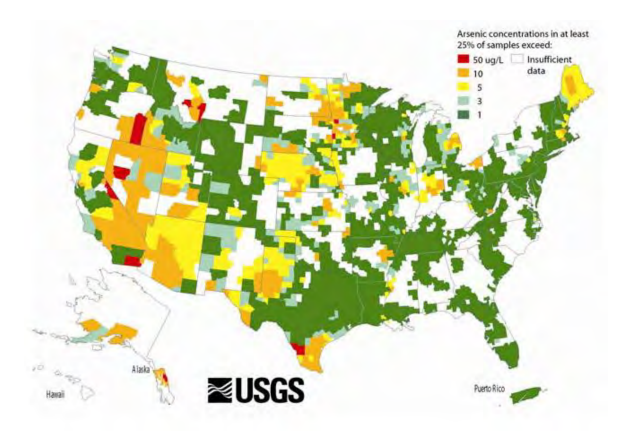


Figure 1: US Geological Survey Map of Arsenic in Groundwater (USGS, 2005).

In ground water, arsenic can combine with other elements to form inorganic as well as organic compounds; the inorganic derivatives are considered more toxic than the organic forms. The inorganic forms usually exist in potable water in two chemical valence states: as arsenite (As III) or arsenate (As V). The arsenite species exists in anaerobic/anoxic (reduced or low oxygen) waters as H_3AsO_3 or $H_2AsO_3^{1-}$. The specific form depends on the pH of the water. At pH 9.2, arsenite exists as a 50-50 mixture of H_3AsO_3 and $H_2AsO_3^{1-}$. At less than pH 8.5, arsenite exists primarily as the neutral (non-ionized) species, H_3AsO_3 (can be written as $HAsO_2$). The arsenate species exists in aerobic (oxidized) waters up to pH 9 as a mixture of $H_2AsO_4^{1-}$ and $HAsO_4^{2-}$, and as a 50-50 mixture at pH 7.0.

In order to find out whether your drinking water contains arsenic, please refer to the Natural Resources Defense Council (NRDC) website. NRDC has analyzed data compiled by the US Environmental Protection Agency (US EPA) in drinking water in 25 states over a period of 18 years. You can also get information on the presence of arsenic in your drinking water from your local Utility or state EPA. You can also call USEPA's drinking water hotline (1-800-426-4791) for more information.

As mentioned above, two conditions that dominate the behavior of arsenic in water are its state of oxidation (valence) and the pH of the water. Generally, aerated surface waters contain arsenate (As V) while the reductive well waters contain arsenite (As III). Municipally treated waters containing free available chlorine (FAC) will generally oxidize arsenite (As III) to the arsenate (As V) form. **NOTE: Chloraminated water utilizing only monochloramine (NH₂CI) will not completely oxidize As III to As V.** Generally, negatively charged (ionized) As V (as H₂AsO₄¹⁻ and HAsO₄²⁻) is much easier to remove than uncharged As III (as H₃AsO₃ or HAsO₂). Tests to determine the concentration of each form, also known as speciation, must be performed in order to choose the proper removal technique. Current technology suggests that several techniques may be used to remove the arsenate, arsenite and organic forms of arsenic from drinking water. These include iron oxide/hydroxides and activated alumina media filtration, manganese greensand filtration, strong base anion exchange resins, distillation, and reverse osmosis. Some specialty media include iron oxide/hydroxide-impregnated or coated activated alumina and anion exchange resins, as well as titanium oxy/hydroxide.

HEALTH EFFECTS

Toxicity of arsenic to humans is well known, and ingestion of as little as 100 milligrams (mg) can result in severe poisoning. Amounts in water are normally much lower, but low concentrations still can lead to chronic symptoms. The effect of arsenic, when ingested in small amounts, appears very slowly. In fact it may take several years for the poisoning to become apparent. Chronic arsenosis can in its most extreme form, cause death. Inorganic arsenic is absorbed readily from the gastrointestinal track and becomes distributed throughout the body tissues and fluids. Ingestion of inorganic arsenic leads to a number of health effects as follows:

Cancerous effects: skin, bladder, lung, kidney, nasal passages, liver and prostate cancer Non-cancerous effects: cardiovascular, pulmonary, immunological, neurological and endocrine disruption effects

The US EPA's final arsenic rule was issued in 2001. It revises the Maximum Contaminant Level (MCL) from 50 μ g/L to 10 μ g/L and sets the Maximum Contaminant Level Goal (MCLG) to zero in the drinking water. Both community water systems (CWS) and non-transient, non-community water systems (NTNCWS) are required under this rule to reduce the arsenic concentration in their drinking water to10 μ g/L or lower.

For details of toxicological and epidemiological effects of arsenic in drinking water, please refer to the National Research Council report published by National Academic Press given in the references below.

TREATMENT METHODS

	Iron oxide/hydroxides	
Residential	Activated alumina	
Point-of-Entry	Anion exchange in a fixed bed (requires regeneration)	
	Manganese greensand (requires regeneration)	
	Titanium oxy/hydroxide	
	Iron-doped anion resin and activated alumina	
	Iron oxide/hydroxides	
Residential	Activated alumina with or without iron oxide coating	
Point-of-Use	Anion exchange	
	Distillation	
	Titanium oxy/hydroxide	
	Reverse osmosis (RO)	
	Iron oxide/hydroxides	
Municipal	Activated alumina	
	Anion exchange in a fixed bed (requires regeneration) Coagulation	
	with iron hydroxide (initially added as FeCl ₃) followed by filtration	
	though a bed of adsorptive granular media	
Visit WQA org or NSE org to searc	h for products certified to WQAS-200. NSF 53, 58, 62 for arsenic reduction.	

visit wQA.org or NSF.org to search for products certified to WQAS-200, NSF 53, 56, 62 for arsenic reduction.

Iron oxide, iron hydroxide and iron coated activated alumina filtration media have shown effectiveness in removing both arsenite (As III) and arsenate (As V) from levels of over 50 parts per billion (ppb) or μ g/L to effluent levels below 5 ppb (μ g/L) for greater than 10,000 bed volumes before exhaustion. These removal systems operate best at pH less than 8. Iron-based technologies can be susceptible to competitive adsorbates, such as silicates, vanadates (e.g., VO₄³⁻) and phosphate. It has be demonstrated that each 0.5 mg/L increase in phosphate above 0.2 mg/L will reduce adsorption capacity by roughly 30% (US EPA, 2014). Therefore, a water analysis should be conducted prior to the selection of a removal technology. For reductive water from wells, where As III is expected to predominate, an oxidation step such as chlorination or ozonation is necessary if a technique other than iron based systems are used (e.g., RO and anion exchange). Potassium permanganate (KMnO₄) is also very effective in oxidizing As III to As V. Passing aqueous solutions of As III through a fixed-bed column of manganese dioxide (MnO₂) also has been shown to oxidize As III to As V and, at the same time, provide some removal capability.

Iron based specialty media are also easy to operate and have a higher capacity than AA media. Some media can remove both arsenite and arsenate even though capacities may differ. The pH can be as high as 8.5 for many of these media, however the lower the pH the better the removal performance. Activated alumina (AA) has easy maintenance and potential for non-hazardous waste disposal after exhaustion. It is highly selective for As V, so arsenites (As III) must be oxidized to arsenate (As V) and the pH lowered, preferably to <6.5 or even lower if silica is present.

Distillation is useful when only small quantities of drinking water are involved. It has been shown to reduce arsenic to less than 2 ppb (μ g/L).

Strong base anion (SBA) type I & type II resins in Cl⁻ form can effectively remove As V (arsenate, as $H_2AsO_4^{4-}$ or $HAsO_4^{2-}$). However sulfate, selenium, fluoride and nitrate ions present in water will compete with As V ions ($H_2AsO_4^{1-}$ and $HAsO_4^{2-}$) for exchange sites and may produce earlier exhaustion. Anion exchange is not effective to remove As III (arsenite, as H_3AsO_3 or HAsO2), since they are not charged (ionized), so it is necessary to oxidize it to As V by an acceptable technique (chlorination, permanganate or ozonation). Optimum pH for anion exchange removal is approximately >7. Regeneration can be done with brine (NaCl) solution.

Manganese greensand is best used for only As V reduction, so if As III is present, it should first be oxidized to As V.

NOTE: Prior to installing and utilizing regenerative treatment technologies or bulk disposable adsorptive media, it is important to contact the local, regional or state regulatory authorities to determine proper disposal requirements.

Thin film composite (TFC) Reverse Osmosis membranes are best used for only As V reduction, so, again, As III should be oxidized by chlorination or ozonation, or other oxidation technique acceptable for the specific application that will not harm the membrane.

The treatment methods listed herein are generally recognized to be effective in reducing the listed contaminants sufficiently to meet or exceed the relevant MCL. However this list does not reflect the fact that point-of-use/point-of-entry (POU/POE) devices and systems currently on the market may differ widely in their effectiveness in treating specific contaminants and performance may vary from application to application. Also the effectiveness of many of these treatment technologies is reduced by interfering reductants and competitive adorbates. Always conduct a complete water analysis prior to any treatment application. Generally it is recommended that devices and systems that are independently certified to appropriate NSF/ANSI standards should be preferably used. Whenever possible, assistance from a water professional or expert should be sought in the selection, installation and operation of a chosen technique.

Visit WQA.org to locate water professionals in your area. Note that Certified Water Specialists have passed the water treatment education program with the Water Quality Association and continue their education with recertification every 3 years.

REGULATIONS

In the United States the EPA, under the authority of the Safe Drinking Water Act (SDWA), has set the Maximum Contaminant Level Goal (MCLG) for arsenic at zero. This is the health-based goal at

which no known or anticipated adverse effects on human health occur and for which an adequate margin of safety exists. The US EPA has set this level of protection based on the best available science to prevent potential health problems. Based on the MCLG, the US EPA has set an enforceable regulation for arsenic, the Maximum Contaminant Level (MCL), at 0.010 mg/L (10 ppb). MCLs are set as close to the MCLG as possible, considering cost, benefits and the ability of public water systems to detect and remove contaminants using suitable treatment technologies. In Canada, the regulated Maximum Acceptable Concentration (MAC) for arsenic in drinking water is also set at 0.010 mg/L.

REFERENCES/SOURCES

- Federal Register (2001). "National Primary Drinking Water Regulations; Arsenic and Clarifications to Compliance and New Source Contaminants Monitoring." Retried from <u>https://www.federalregister.gov/articles/2001/01/22/01-1668/national-primary-drinking-water-regulations-</u> arsenic-and-clarifications-to-compliance-and-new-source
- Health Canada (May 2006). "Guidelines for Canadian Drinking Water Quality: Guideline Technical Document." http://www.hc-sc.gc.ca/ewh-semt/alt_formats/hecs-sesc/pdf/pubs/water-eau/arsenic/arsenic-eng.pdf
- National Resources Defense Council (February, 2000). "Arsenic in Drinking Water". Retrieved from http://www.nrdc.org/water/drinking/garsenic.asp
- US EPA (May 21, 2012). "Basic information about arsenic in drinking water". Retrieved from http://water.epa.gov/drink/contaminants/basicinformation/arsenic.cfm
- US EPA (June 3, 2013). "Drinking Water Contaminants". Retrieved from http://water.epa.gov/drink/contaminants/index.cfm
- US EPA Office of Research and Development (March 2001). "Laboratory Study on the Oxidation of Arsenic III to Arsenic V", EPA/600/R-01/02. Retrieved from http://www.epa.gov
- US EPA (October 4, 2012). "Water: Regulatory Information". Retrieved from http://water.epa.gov/lawsregs/rulesregs/

US EPA (January 21, 2014). "Arsenic Virtual Trade Show". Retrieved from http://cfpub.epa.gov/safewater/arsenic/arsenictradeshow/arsenic.cfm?action=Adsorption

ACKNOWLEDGEMENT

WQA wishes to express sincere appreciation for the unselfish contributions of the members of WQA who contributed their time and expertise toward the completion of this bulletin.

Arvind Patil, Ph.D., CWS-I Gary Hatch, Ph.D. Charles Michaud, CWS-VI Mark Brotman, CWS-VI P. Regunathan, Ph.D. Rebecca Tallon, P.E. Richard Andrew Shannon Murphy Steve VerStrat Pauli Undesser, M.S., CWS-VI Kimberly Redden, CWS-VI The Water Quality Association publishes this Technical Application Bulletin as a service to its members and the interested public. Information contained herein is based upon the most recent public data known as of the publication date and cannot take into account relevant data published thereafter. The Water Quality Association makes no recommendations for the selection of a treatment system, and expressly disclaims any responsibility for the results of the use of any treatment method or device to reduce or remove a particular contaminant.

This reference document is published by:



National Headquarters & Laboratory

4151 Naperville Road • Lisle, Illinois 60532 Tel: 630 505 0160 • Fax: 630 505 9637

Copyright © 2013 by Water Quality Association. All rights reserved. No part of this publication may be reproduced, stored in a retrieval system, or transmitted in any form or by any means, electric, mechanical, photocopying, recording, or otherwise, without the prior written permission of the publisher.