

EFFICIENT METHODS FOR ARSENIC REMOVAL FROM GROUNDWATER

G. BADALIANS GHOLIKANDI¹, H.R. ORUMIEH² & R. RIAHI³

¹Power and Water University of Technology (PWUT), Water Research Institute (WRI), Tehran, Iran.

²Pars Arianab Consulting Engineers, Esfahan, Iran.

³Water Research Institute (WRI), Water and Wastewater Research Center (WWRC), Tehran, Iran.

ABSTRACT

Arsenic may be found in water that has flowed through arsenic-rich rocks. Arsenic is a toxic, trace element that is ubiquitous in nature. It can easily be transported from the sediment to the surrounding pore-water. Severe health effects have been observed in populations drinking arsenic-rich water over long periods in countries worldwide. A 2007 study found that over 137 million people in more than 70 countries are probably affected by arsenic poisoning of drinking water. In groundwater, arsenic combines with oxygen to form inorganic pentavalent arsenate and trivalent arsenite. Most arsenic treatments fall into four process categories: ion exchange, membrane process, adsorption, or chemical precipitation. This study investigates the potential of removing arsenic from groundwater by using two process categories – activated alumina and lime softening. Arsenic adsorption by commercially available activated alumina is surveyed and its efficiency investigated. We have incorporated some of the valuable literature on arsenic remediation by adsorption. According to results of three activated alumina pilot studies, considering influence of adsorption time, temperature, pH, alumina quantity, arsenic concentration, and different alumina production resources, it has high efficiency for arsenic removal. Adsorption isotherm for both species of arsenic (III and V) is compatible with both Freundlich and Langmuir models (correlation coefficient >0.93). The prevalent pH range for arsenate was between 6 and 8. Modified activated alumina can remove arsenate at the influent pH of 8.1 ± 0.4 to below the maximum concentration level (MCL). The exhausted media passed the Toxicity Characteristic Leaching Procedure (TCLP) test with respect to arsenic. Lime softening operated within the optimum pH range of more than 10.5 is likely to provide a high percentage of arsenic removal (90%) for influent concentrations of up to 0.05 mg/L. It may be difficult to reduce consistently to 0.01 mg/L by lime softening alone. Systems using lime softening may require secondary treatment to meet that goal.

Keywords: activated alumina, arsenic contamination, groundwater, removal efficiency, softening.

1 INTRODUCTION

Groundwater is a remarkable source for supplying drinking water as well as in sustainable irrigated crop production in Iran [1]. Extent and severity of arsenic contamination of groundwater is a crucial issue and a threat to human health and food safety. Alarming information has emerged in recent decades about the widespread presence of arsenic (As) in groundwater used to supply drinking water in many countries on all continents (Fig. 1) [9]. A 2007 study found that over 137 million people in more than 70 countries are probably affected by arsenic poisoning of drinking water. Hundreds of millions of people, mostly in developing countries, daily use drinking water with arsenic concentrations several times higher than the World Health Organization (WHO) recommendation of 0.01 mg/L of water [3, 4]. Dangerous arsenic concentrations in natural waters are now a worldwide problem and often referred to as a 20th–21st century calamity. High arsenic concentrations have been reported recently from the USA, China, Chile, Bangladesh, Taiwan, Mexico, Argentina, Poland, Canada, Hungary, Japan, and India [5]. In 2001, US–EPA published a new 0.01 mg/L standard for arsenic in drinking water [6].

Arsenic is a crystal-shape metalloid which is brittle in nature and grayish white in color. Arsenic rarely occurs in a free state and is found largely in combination with sulfur, oxygen,

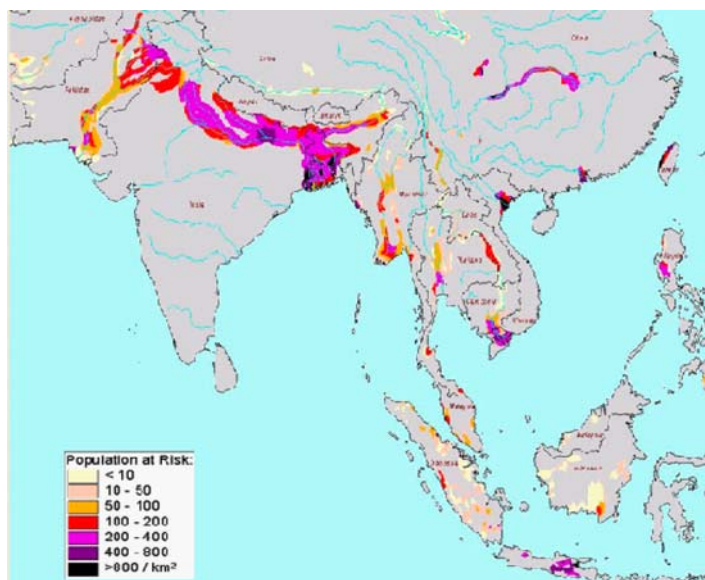


Figure 1: Population at arsenic contaminated groundwater risk [14].

and iron. Unlike other heavy metalloids and oxyanion-forming elements, arsenic can be mobilized under a wide range of oxidizing and reducing conditions at the pH values typically found in groundwater (pH 6.5–8.5). Whereas all other oxyanion-forming elements are found within the $\mu\text{g/L}$ range, arsenic can be found within the mg/L range. Arsenic has four main chemical forms having oxidation states, -3 , 0 , $+3$, and $+5$, but in groundwater its predominant forms are inorganic oxyanions of trivalent arsenite (As III) or pentavalent arsenate (As V) [7]. The toxicity of different arsenic species varies in the order arsenite>arsenate>monomethylarsonate>dimethylarsinate. Trivalent arsenic is about 60 times more toxic than arsenic in the oxidized pentavalent state, and inorganic arsenic compounds are about 100 times more toxic than organic arsenic compounds [8]. The organic forms of arsenic are quantitatively insignificant and are found mostly in surface waters or in areas severely affected by industrial pollution [9]. The relative concentrations of As III to As V vary widely, depending on the redox conditions in the geological environment [10]. The two most important factors controlling the speciation of arsenic and, to some extent, solubility are pH and redox potential. Under oxidizing conditions at pH less than 6.9, H_2AsO_4^- is the dominant species, whereas HAsO_4^{2-} predominates at higher pH. Under reducing conditions at a pH value less than 9.2, the uncharged arsenite species H_3AsO_3 is dominant. In contrast to the pH dependency of As V , As III was found virtually independent of pH in the absence of other specifically adsorbed anions. Most often, more trivalent arsenic than pentavalent arsenic is found in reducing groundwater conditions, whereas the converse is true in oxidizing groundwater conditions. Unlike other toxic trace metals whose solubility tends to decrease as pH increases, most oxyanions, including arsenate, tend to become more soluble as pH increases. When most other metals become insoluble within the neutral pH range, arsenic is soluble at even near-neutral pH in relatively high concentrations. That is why groundwater is easily contaminated with arsenic and other oxyanions [11, 12]. The severe arsenic intoxication occurs when the element affects the nervous system which may cause coma and, in intoxications

with 70–80 mg/L, may be even mortal. Digestive system, nervous system, respiratory system, and skin all are highly sensitive to the element and arsenic may cause skin cancer, keratinization, and hyperpigmentation.

If a completely satisfactory, arsenic-free water source cannot be established, the short-term goal should be to reduce arsenic levels in drinking water as much as possible, as quickly as possible, even if regulatory standards cannot be immediately met. It should be recalled that health effects of arsenic are dose-dependent, and a partial solution is better than no solution. The implementation of a temporary solution should not be used as a reason to delay design and implementation of a long-term plan [13]. Most arsenic treatments fall into four process categories: ion exchange, membrane process, adsorption, or chemical precipitation. Ion-exchange treatments are very limited in their ability to remove arsenic because of exchange competition from other anions found in groundwater. Membrane processes are very effective at removing arsenic from groundwater, but the cost is high. Accordingly, adsorption and chemical precipitation processes are being explored for low-cost, effective treatments [6, 12]. Historically, the most common technologies for arsenic removal have been coagulation with metal salts, lime softening, and iron/manganese removal. Coagulation processes are sometimes unable to efficiently remove arsenic to the recommended low levels. As a result, various alternate technologies have been developed or adapted that are capable of removing arsenic to trace levels. One of these advanced treatment options is activated alumina [13]. Present review aimed to study the efficiency of two arsenic removal methods; adsorption using activated alumina and lime softening, to investigate a safe and feasible method for removing arsenic from drinking water in simple household treatment systems especially for the contaminated groundwater resources in West Iran.

2 MATERIAL AND METHODS

Two process categories are considered: (1) activated alumina; (2) lime softening. Three pilot studies are investigated:

(a) A field program to evaluate the performance of ALCAN AAFS-50 arsenic removal media [15]. This field trial program was carried out by 2002 to evaluate the performance of this arsenic removal media at Well # 15, Kirtland Air Force Base, New Mexico. Well # 15 historically has water with arsenic concentration more than 0.01 mg/L. One of the media tested was ALCAN AAFS-50; a ferric coated activated alumina, with Density-dry 1.12 g/cm³, BET (Brauner, Emmett, and Teller) surface area 139.8 cm²/g, size of media 28 × 48 mesh, volume of media loaded 5.0 L, and empty bed contact time 5.0 minutes. The scanning electron microscope (SEM) analysis of the virgin AAFS-50 and the exhausted media grains after the completion of the column tests with a JEOL JSM-6300V scanning electron microscope equipped with an energy-dispersive X-ray emission analyzer, and the toxicity characteristic procedure (TCLP) test for exhausted media, according to the EPA-Method 1311 (1992) were also carried out. To ensure that the arsenic in the well water was in the pentavalent form before entering the media columns, the arsenic feed water was treated with 5% commercial sodium hypochlorite solution to a level of approximately 1 mg/L of residual chlorine in a holding tank. Source groundwater was obtained downstream from a pump station. This water entered a high-density polyethylene tank via a flow control valve. The water from the feed tank passed through individual flow totalizers before entering the bottom of each individual media column. Each media column was divided into two sections. The bottom contained graded gravel, which served as a containment base for the media and as a uniform water flow distributor. On top of the gravel bed laid the arsenic sorption media bed

of uniform size composition. The water flowed up through each packed column at the flow rate of approximately 1.0 L/min at an empty bed contact time (EBCT) of 5 minutes. At the EBCT of 5 minutes, the bed behaved as a packed column. Both influent and effluent samples (250 mL) were collected for laboratory analyses on a daily basis. The more common operational control parameters and the water quality parameters were measured on a daily basis. The field setup was operated until breakthrough, defined by an effluent water arsenic concentration of 0.005 mg/L. After breakthrough, each individual column was emptied, and samples of the saturated media were sent for SEM analysis and TCLP test.

(b) Batch experiments carried out by 2004 to evaluate the performance of ALCAN AAFS-50 arsenic removal media at some villages in West Iran [16]. The efficiency of modified activated alumina with iron compounds with trade name ALCAN AAFS-50 was studied. Equilibrium batch experiments were carried out using shaker incubator and arsenic was analyzed with Silver Diethyl Dithiocarbamate Spectrophotometric (SDDC) method. The media tested was a ferric coated activated alumina, with Density-dry 0.91 g/cm³, BET surface area 200–250 cm²/g, and size of media 28 × 48 mesh. Effects of initial concentration of arsenic, adsorbent dose, oxidation state of arsenic, reaction time, pH, and oxidation with chlorine on adsorption were studied.

(c) An activated alumina pilot plant operated from 1997 to 1999 to evaluate the performance of Alcoa medium for arsenic removal from drinking water [17]. An activated alumina pilot plant was installed and operated to evaluate the performance of Alcoa medium for arsenic removal from drinking water. Source water was supplied by 700-ft-deep bedrock well. The capacity of this well was 190–220 L/sec. The total soluble arsenic concentration was in average 0.062 mg/L. About 30% of the soluble arsenic existed as As(III) and the rest was As(V). Particulate arsenic was not detected in the raw water. After passing through the cartridge filters, the filtered water splits into the two activated alumina trains (two parallel sets of two tanks in series). Each fiberglass mineral tank (132.1 cm tall by 40.6 cm in diameter) contains about 1.8 m³ of Alcoa DD-2 activated alumina. The depth of the activated alumina bed is about 84 cm. The influent flows downward through the activated alumina bed and the treated water returns to the top of the tank through a 1-inch polyvinyl chloride riser tube. The differential pressure across the medium was about 3 psi. The hydraulic loading rate to each tank is 0.085 L/cm².sec and the EBCT is 4.3 minutes. Physical characteristics of Alcoa DD-2 activated alumina (14 × 28 mesh) are: white granule, surface area 250 m²/g, total pore volume 0.395 cc/g, total porosity 56.2%, bulk density 620–830 kg/m³, and abrasion loss 1.6 wt%. Its chemical composition (wt %) is: Al₂O₃ (92.2), Na₂O (0.90), Fe₂O₃ (0.08), SiO₂ (0.09), loss on ignition (water, 6.5), Alumina XRD phase amorphous, boehmite, and gamma. The first set of the tanks (TA1 and TA2) were used as roughing filters and the second set (TB1 and TB2) were used as polishing filters. During the preliminary sampling phase of this study, water samples were collected at four locations: (1) at the inlet; (2) after the roughing tank of train 1 (TA1); (3) after the polishing tank of train 1 (TB1); and (4) at the combined effluent of trains 1 and 2. Preliminary sampling consisted of biweekly sample collection and analysis. Biweekly long-term sampling and analysis was performed for 36 weeks (18 events).

The pilot plant operated by 2006 to evaluate the performance of the softening method by lime for arsenic removal from drinking water [18]. The Plant includes following units: (1) chemical pollutant stock tank: this unit comprises of a polyethylene tank with 80 L volume and 50 L effective volume, an electric stirrer including an electromotor with 1450 rpm, axe and stirring blade, and a dosing pump. The pollutant solutions including arsenic solutions are prepared in this tank and injected to the fixed-level tank; (2) fixed-level tank: a polyethylene tank

Table 1: Samples' preparation [18].

Pilot stage	Concerned content of arsenic in raw water (mg/L)	System rate (L/min)	Pollutant stock injected (mg/L)	Arsenic concentration in the injected pollutant stock (mg/L)	1000 mg/L arsenic added to pollutant stock (mg/L)
A	0.05	3	5	1.875	18.6
B	0.1	3	5	3.714	37.2
C	0.15	3	5	5.571	55.9
D	0.2	3	5	7.428	74.4

with 500 L volumes equipped with float and electromechanical stirrer with 1450 rpm velocity. The tank is connected to the water supply line from one side and its level remains fixed. The other side of the tank is connected to the dosing pumps, which injects the chemical pollutant solution to the tank, and the electromechanical stirrer makes a homogenous mixture with the raw water; (3) rapid mixing unit: the unit comprises of three parts: rapid mixture tank, stirrer with variable rpm, and stirrer blades. The unit is made of Plexiglas and all the calculations have been done based on Camp's theory presented in 1965. The section of the tank is quadratic with 20 cm sides and its effective depth is 20 cm and total depth of the tank is 30 cm. For prevention from formation of eddy current, a baffle on the walls is installed; (4) coagulation unit: the tank of the unit is made of polyethylene and has an effective volume of 200 L. An electromotor with zero to 60 rpm and paddle blades is used for stirring the tank; (5) clarification unit: the cross-section of this unit is rectangular, made of Plexiglas, and the duration of current inertia is 4 hours; the effective volume of the pool is 628 L and its depth is 70 cm. The sludge resulting from sedimentation is moved through the sloped bottom of the pool to the collecting hopper which is located in the center of the pool; (6) chlorine and lime water mixture preparation and injection unit: the mixture was prepared separately in polyethylene tanks in rapid mixture unit (Table 1).

Samples were collected at three locations: (1) after fixed-level tank for raw water; (2) after flocculation tank; and (3) after clarification tank. Table (2) shows the analytical results. In all stages of the test the pH level and hardness of the raw water were continuously controlled. Considering the importance of coagulation and flocculation process, sampling was considered before and after the unit to determine the process efficiency. To ensure the hardness (TH = 236 mg/L as CaCO₃), the feed water was treated with 93.64 mg calcium chloride (CaCl₂, 77% purity). The Plant was operated at four stages (A, B, C, & D) with initial concentrations of 50, 100, 150, and 200 mg/L respectively. Different As (V) concentrations in feed water were prepared using arsenic pentoxide (As₂O₅). The techniques for analyses followed the APHA-Standard Methods [19].

3 RESULTS AND DISCUSSION

3.1 Activated alumina

For 25 years, activated alumina has been the preferred media for arsenic removal from water. The ALCAN specialty aluminas have developed a new, improved grade ActiGuard AAFS50

with up to three times the capacity of standard activated alumina [8]. Historically, regeneration was the accepted method of making activated alumina systems cost-effective. Regeneration, however, has some drawbacks, including the handling and storage of chemicals, additional processing steps, and the disposal of regeneration effluent. According to the product information documents, the material is a granule and doesn't break down or deteriorate with use as iron-based media. The low pressure drop provides a system requiring less maintenance. Pre-filtered water will allow the adsorptive system to operate virtually without back washing. ALCAN AAFS-50's capacity for As (III) adsorption is about 40% its capacity for As (V). Oxidation is recommended to assist with As (III) removal, when possible. Simple chlorination will oxidize As (III) to As (V) allowing complete removal [8].

(a) The field trial program was carried out at Well # 15, Kirtland Air Force Base, New Mexico showed that this media was able to remove arsenate and meet the total arsenic maximum contaminant level (MCL) of 0.01 mg/L in drinking water [15]. The arsenate removal capacity was defined at a breakthrough effluent concentration of 0.005 mg/L arsenic. At an influent pH of 8.1 ± 0.4 (Temperature 24–26°C, average arsenic concentration 0.013 ± 0.0009 mg/L), the arsenate removal capacity of AAFS-50 was 33.5 mg As (V)/L of dry media (0.03 mg As(V)/g of media on a dry basis). Silicate, fluoride, and bicarbonate ions were removed by ALCAN AAFS-50; chloride, nitrate, and sulphate ions were not removed by the media. The average conductivity values of the influent and treated water were 421630 and 437647 $\mu\text{S}/\text{cm}$, respectively. The turbidity of both the influent and treated water were below detection, indicating that both the influent and treated water were not turbid and well below the turbidity maximum contaminant level of 1 normal turbidity unit (NTU). At operating pH more than 7.0, the structural integrity of the AAFS-50 media was good, as indicated by the constant flow rate observed for the duration of the test. The exhausted media passed the toxicity characteristic procedure (TCLP) test with respect to arsenic (0.0001 mg/L) and therefore could be disposed as non-hazardous waste. This study was limited to the adsorption of As (V) because the feed water was pre-chlorinated. Groundwater contaminated with arsenic may contain As (III) species, and therefore, pre-oxidation of As (III) to As (V) is required unless the media performance is evaluated for the removal of As (III). The scanning electron microscope (SEM) analysis of the virgin AAFS-50 and the exhausted media grains after the completion of the column tests did not show any changes in the structural morphology of the media. This indicates that the media integrity was maintained and that the bed characteristics did not change, an observation further substantiated by the stable water flow rate (1.0 L/min) observed throughout the test (volume of water treated until breakthrough: 12.986 L) (Fig. 2) [13].

(b) Considering contamination of drinking water to arsenic in some villages in West Iran, and according to results of this pilot study, modified activated alumina has high efficiency for removal of arsenic and can reduce arsenic concentration under WHO guidelines [16, 20]. Adsorption isotherm for both species of arsenic (III and V) is compatible with both Freundlich and Langmuir models (correlation coefficient > 0.93). Removal efficiency increases as a result of increasing dose and reaches to 98% for As (V) during 2 hours. Adsorption is first order reaction and removal rate for concentration of 0.421 and 1.15 mg/L was 91% and 66% respectively. Removal efficiency increases 1.54 times from 61% to 94% with increasing of adsorption time from 15 min to 60 min for primary concentration of 0.25 mg/L. For As (III) the highest removal was 30% with adsorbent dose = 2000 mg/L. Our study showed that absorption of arsenic has direct relation with increasing of pH up to 8, then decreases with increasing of pH up to 14. For arsenate the highest removal was observed at pH between 6 and 8 and removal efficiency was



Figure 2: Analytical results [15].

higher than 90%. The highest removal for primary concentration of 0.5 mg/L with adsorbent dose 2000 mg/L, was 95% at pH 8 that was achieved during 2 hours. Totally for activated alumina used at research pH can not affect removal efficiency. The prevalent pH range at monitoring stations was 6 to 8. Chloride ion could decrease adsorption of activated alumina 2.3%. According to sulfate, the rate was 11%. Oxidation with 4 mg/L chlorine could convert 99% of As (III) to As (V) (initial concentration of As = 0.55 mg/L. For chlorine this amount was 61%. Removal of As (III) using chlorine oxidation increased from 19.6% to 94% at pH 6. Column study showed that treatment system was able to remove arsenic of water containing 1 mg/L As(V) up to 19000 BVs with EBCT = 10 min that equalled to 133 operation days. With half EBCT (5 min) the treated BVs decreased to 17000(70 operation days) (Figs. 3 to 9) [3, 16].

(c) An activated alumina pilot plant was installed and operated to evaluate the performance of Alcoa medium for arsenic removal from drinking water [17]. Total arsenic concentration at the inlet ranged from 0.034 to 0.076 mg/L with an average of 0.054 mg/L. Samples collected after the roughing tank contained total arsenic of 0.014 to 0.051 mg/L with an average of 0.039 mg/L. Samples collected after the polishing tank contained total arsenic of 0.0004 to 0.027 mg/L with an average of 0.007 mg/L. The combined treated water (i.e. outlet) had total arsenic concentrations ranging from 0.001 to 0.022 mg/L with an average of 0.007 mg/L. The average removal percentages were 26.5% and 87.3% by the roughing and polishing tanks, respectively. The average overall arsenic removal efficiency was 86.5% during the long-term sampling. As (III) concentrations averaged 0.008 mg/L at the inlet, 0.003 mg/L after the roughing tank, 0.0006 mg/L after the polishing tank, and 0.0007 mg/L at the outlet location. Because no oxidative treatments were performed ahead of the activated alumina columns, conversion of As (III) to As (V) was rather unlikely. The removal of As (III) would occur either through a direct sorption of As (III) or via some unexplained conversions of As (III) to As (V) prior to adsorption (Fig. 10). Clifford and Lin [5] had observed some unplanned oxidation of As (III) to As (V) within alumina columns, which resulted in better-than-expected performance for arsenic removal. Nonetheless, the adsorption of As (III) onto activated

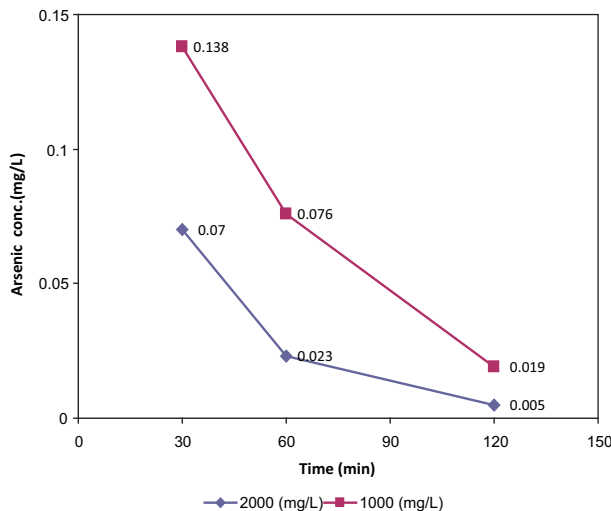


Figure 3: As (V) removal using different quantities of activated alumina, arsenic's initial concentration is 0.25 mg/L [16].

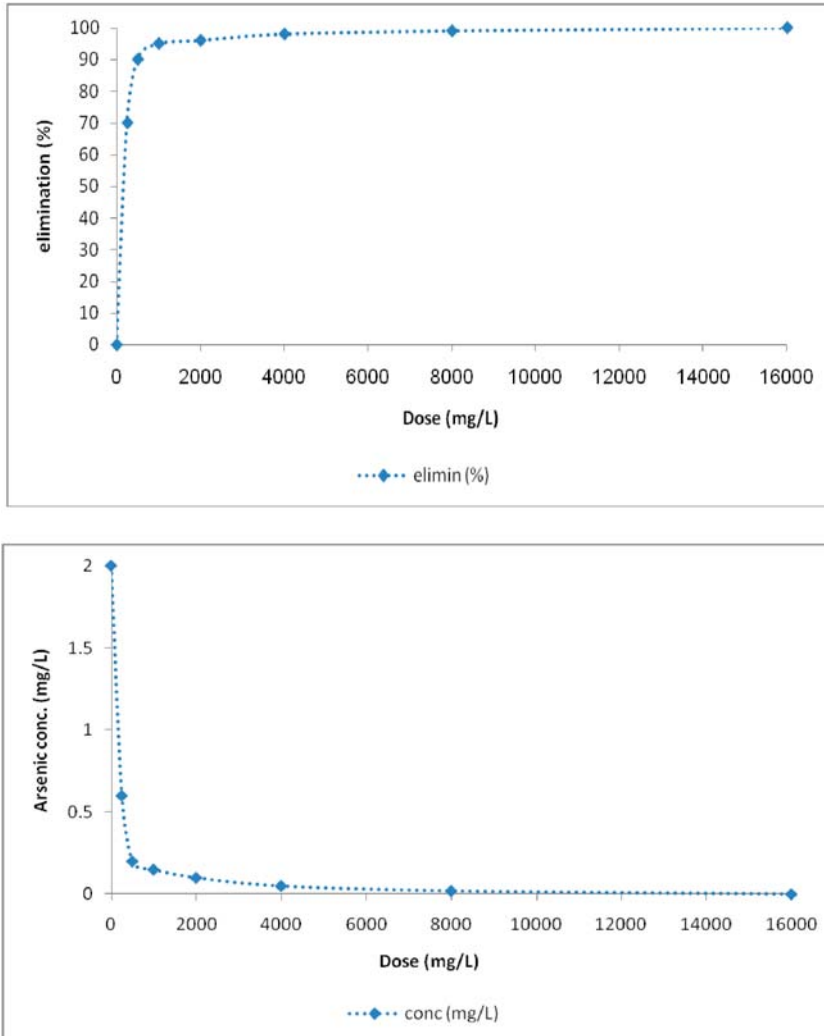


Figure 4: As (V) adsorption using activated alumina; arsenic's initial concentration is 2 mg/L; experience time 24 hours [20].

alumina has been reportedly far less than that of As (V) [22]. Therefore, arsenite should be oxidized to arsenate before activated alumina treatment [7, 8].

The average As (V) concentrations were 0.0452 mg/L at the inlet, 0.0399 mg/L after the roughing tank, 0.007 mg/L after the polishing tank, and 0.006 mg/L at the outlet location. The arsenic in the finished water consists almost entirely of As (V).

The activated alumina capacity of TA1 was nearly exhausted after treating 10.050 BV of water. The arsenic breakthrough occurred in TA2 after treating 9.156 BV of water. The estimated activated alumina capacity was 0.30 g/kg (219 g/m³). This value is comparable to the capacity of 0.26 g/kg reported by Fox [24] in his study with a source water of pH 8.3 and containing 0.05 to 0.35 mg/L As (V). Clifford [23] estimated the practically achievable

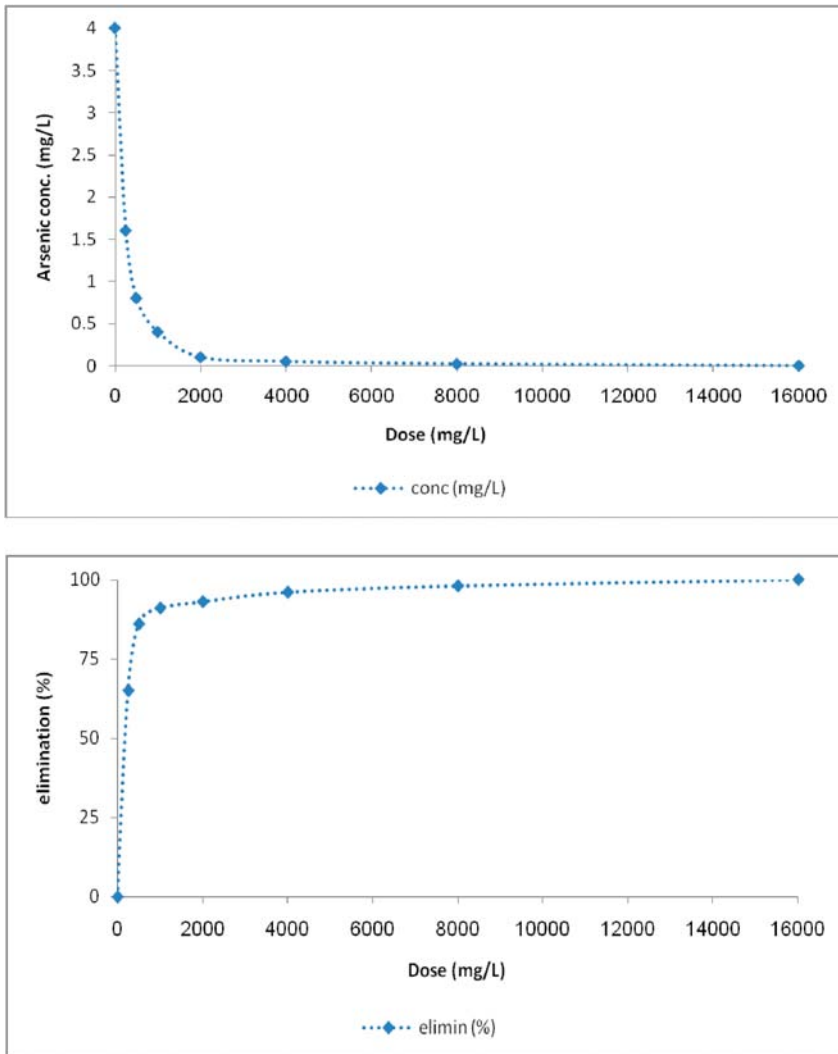


Figure 5: As (V) adsorption using activated alumina; arsenic's initial concentration is 4 mg/L; experience time 24 hours [16].

column capacity based on the pH 6.0 operation with a source water containing 0.1 mg/L As (V) to be 1400 g As (V)/m³ of alumina.

None of the results from analysis of spent activated alumina from this Plant indicate exceedances of TCLP limits.

3.2 Lime softening

The softening by lime method is normally used for decreasing the water hardness. The water hardness is the result of calcium and magnesium ions content of water, which are removed through the relevant process in the form of CaCO₃ and Mg (OH)₂ precipitates.

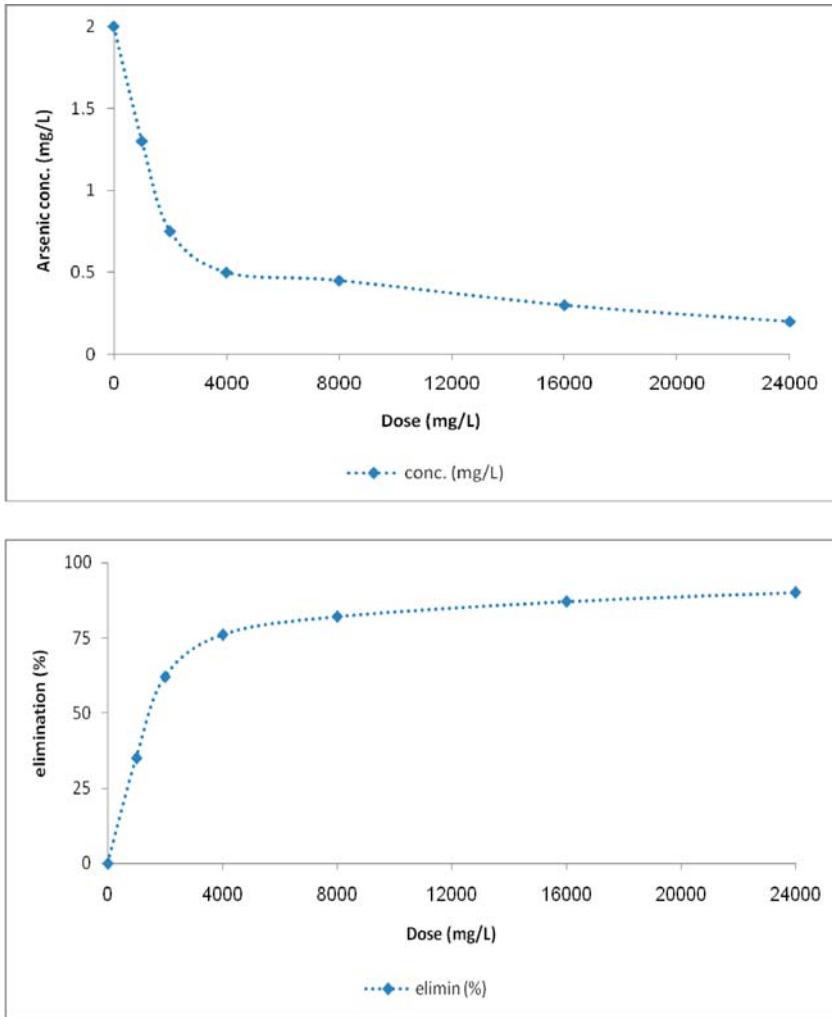


Figure 6: As (III) adsorption using activated alumina; arsenic's initial concentration is 2 mg/L; experience time 24 hours [20].

For softening process, the lime must be used separately or combined with soda; it depends on the type of hardness. The lime should be used by low hardness or carbonate hardness; in case that the hardness is caused by a non-carbonate material, the lime or lime with soda must be used. Sometimes, the caustic soda is also used instead of lime or lime and soda to decrease produced sludge [9, 18]. Table 2 shows the analytical results of pilot operation.

Lime softening operated within the optimum pH range of more than 10.5 is likely to provide a high percentage of arsenic removal (about 90%) for influent concentrations of up to 0.05 mg/L. It may be difficult to reduce consistently to 0.01 mg/L by lime softening.

Table 2: Analytical results [18].

Row	Sample code	Total hardness (mg/L as CaCO ₃)	pH	Arsenic concentration (mg/L)	Arsenic removal (%)
1	A-1-1	247	4.47	0.041	—
2	A-3-1	145	10.31	0.017	58.8
3	A-3-2	151	10.55	0.016	61.1
4	A-3-3	137	11.12	0.006	87.0
5	B-1-1	230	7.36	0.097	—
6	B-3-1	154	8.97	0.073	25.6
7	B-3-2	143	9.62	0.060	38.8
8	B-3-3	121	10.94	0.020	79.7
9	C-1-1	251	7.63	0.146	—
10	C-3-1	170	9.59	0.095	35.4
11	C-3-2	153	10.02	0.089	39.6
12	C-3-3	124	10.87	0.017	88.5
13	D-1-1	244	7.19	0.210	—
14	D-3-1	167	9.33	0.140	33.4
15	D-3-2	152	9.86	0.136	35.1
16	D-3-3	134	11.07	0.037	82.3

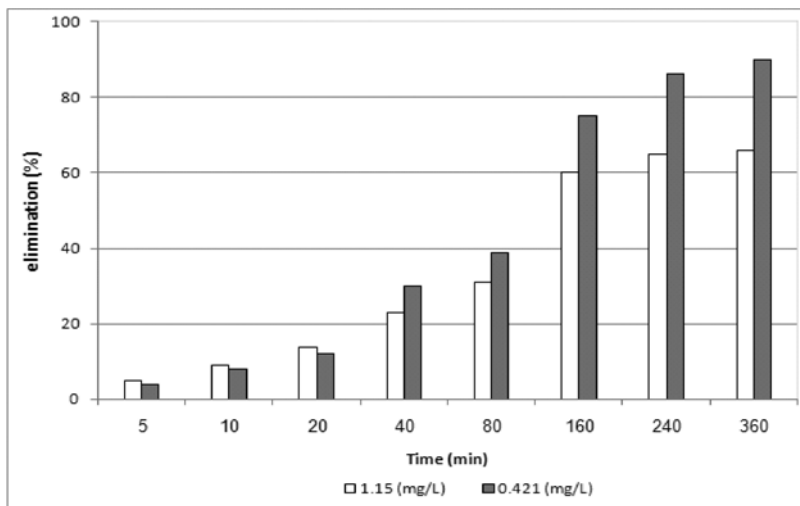


Figure 7: Higher concentration arsenic removal using 500 mg/L activated alumina; pH 5.8 [16].

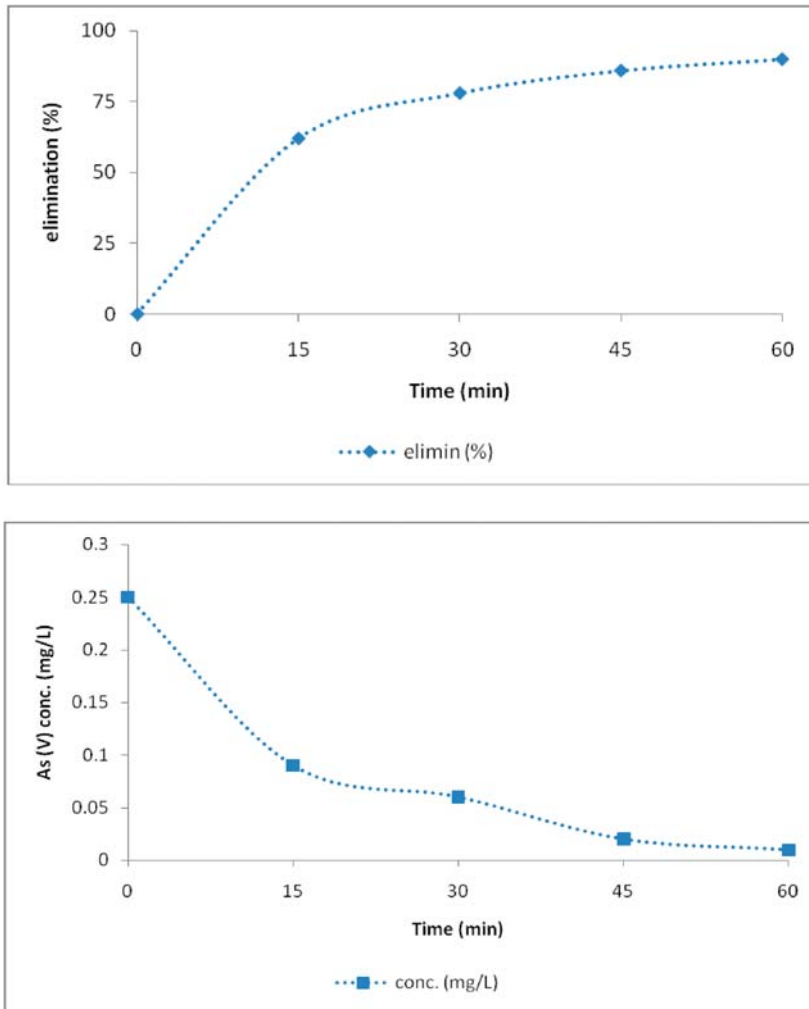


Figure 8: Influence of adsorption time on As (V) removal using activated alumina; arsenic's initial concentration is 0.25 mg/L; alumina concentration 2000 mg/L [16].

tening alone. Systems using lime softening may require secondary treatment to meet that goal.

4 CONCLUSIONS

Exposure to arsenic contaminated drinking water is a major threat to human health. Millions of people across the world are exposed to arsenic contaminated drinking water with concentrations far in excess of the 0.01 mg/L maximum permissible level established by the World Health Organization. Two different arsenic removal process categories were investigated in this paper. According to the results of three operated activated alumina pilot plants, we can conclude that this media could have application as a safe adsorbent for removal of arsenic

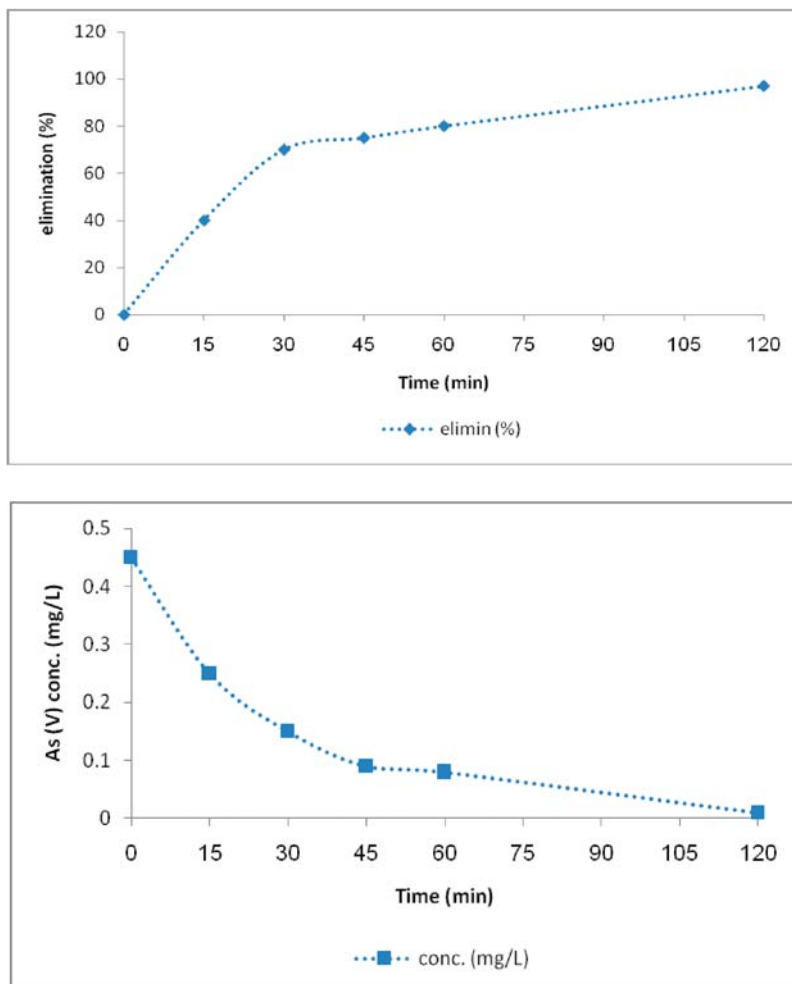


Figure 9: Influence of adsorption time on As (V) removal using activated alumina; arsenic's initial concentration is 0.445 mg/L; alumina concentration 2000 mg/L; pH 5.8 [16].

from drinking water in treatment systems in form of adsorptive column. Experiments using different concentrations of arsenic showed that adsorption of arsenic on activated alumina are a first order reaction; rate of reaction is dependent on initial concentration of arsenic. Arsenite should be oxidized to arsenate before activated alumina treatment. None of the results from analysis of spent activated alumina from operated plants indicate exceedances of TCLP limits. The SEM analysis of the virgin modified activated alumina and the exhausted media grains didn't show any change in the structural morphology of the media. Arsenite removal capacity is much less than for arsenate (more than 90%). Activated alumina is relatively well known and commercially available but needs replacement after four or five regenerations.

Lime softening is proven effective in laboratories and at pilot scale and its efficiency should be largely independent of scale. Disadvantages include extreme pH more than 10.5 and large volume of waste generated. It is inexpensive, but more expensive than coagulation

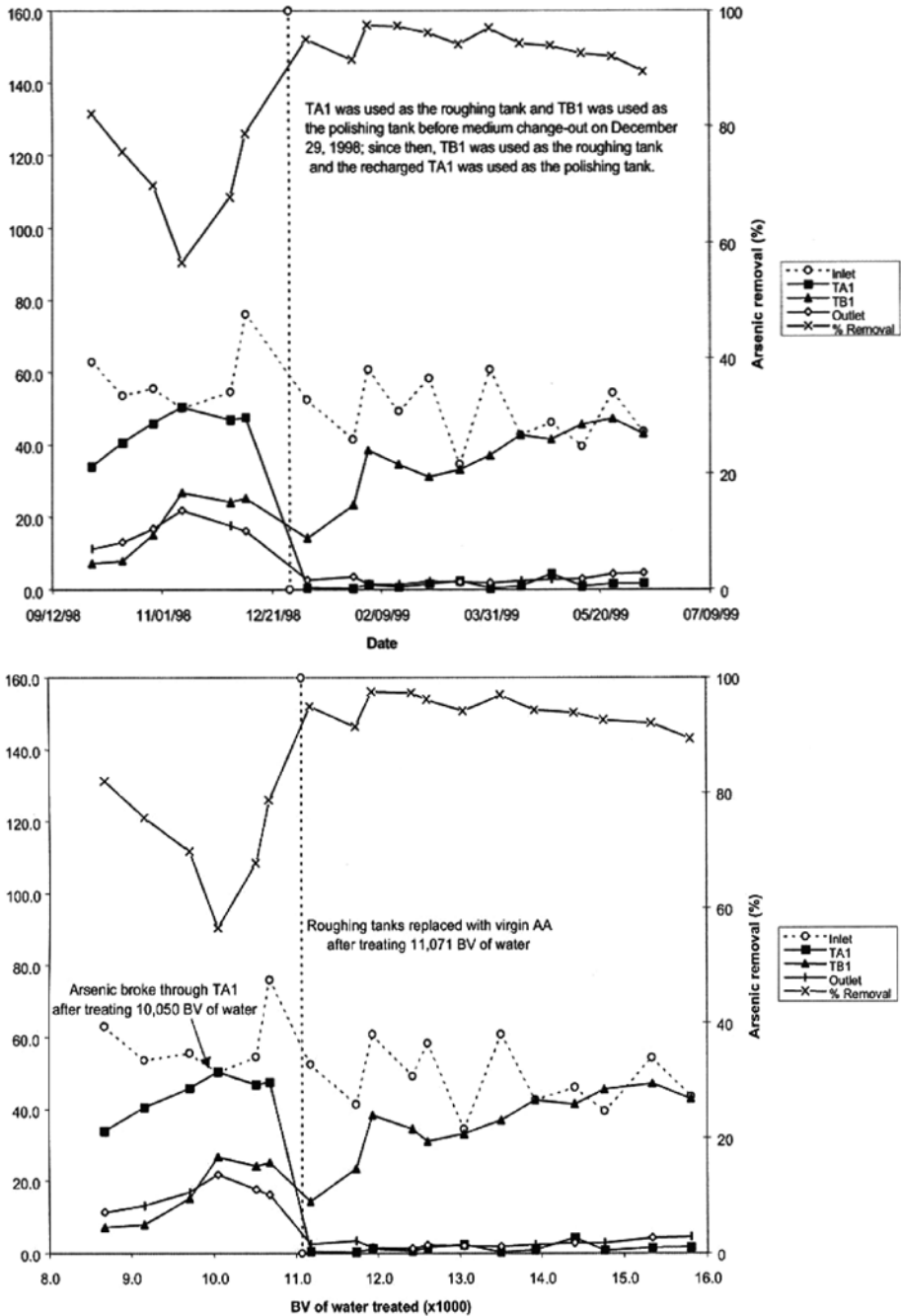


Figure 10: Total arsenic analytical results during long-term sampling [17].

with iron salts or alum because of larger doses required, and wastes handling. The chemicals are available commercially but readjustment of pH is required. Removal efficiency for arsenic (III) is 30–60% and for arsenic (V) it is more than 90%.

ACKNOWLEDGMENTS

Financial support was provided by the National Water and Wastewater Company (NWWC) and Iranian National Management and Planning Organization. Special thanks to water and wastewater authority of province Kurdistan and Faculty of Public Health Tehran University of Medical Sciences.

REFERENCES

- [1] *Kurdistan Province Rural Water and Wastewater*, Report, 2005.
- [2] Haque, A.A.M., Thwe, H.M., Jayasuriya, H.P.W., Hossain, H.Z., Rahman, M., Harunur-Rashid, M. & Matsumura, K., Groundwater arsenic contamination: food safety and human health hazards in Bangladesh. *Chiang Mai University Journal of Natural Science*, **6(2)**, pp. 321–339. 2007.
- [3] World Health Organization (WHO), *Arsenic in drinking water*, www.who.int/media-center/factsheets/fs210/en/index.html. 2010.
- [4] World Health Organization, (1996, 1999, 2001), *Guidelines for Drinking Water Quality, Arsenic and Arsenic Compounds, Arsenic in Drinking Water*, Geneva.
- [5] Mohan, D., Pittman Jr., C.U., *Arsenic removal from water/wastewater using adsorbents*, a critical review, Elsevier, 1–42. 2007.
- [6] US Environmental Protection Agency (EPA), (1998, 1999, 2000, 2001), *Research Plan for Arsenic in Drinking Water - Treatment, Technologies and Costs for Removal of Arsenic from Drinking Water*, National Primary Drinking Water Regulation, Office of Ground Water & Drinking Water.
- [7] Smedley, P.L., Kinniburgh, D.G., A review of the source, behavior and distribution of arsenic in natural waters, *Applied Geochemistry* **17**, pp. 517–568. 2002.
- [8] Idswater, *ActiGuard AAFS50 Activated Alumina for the Removal of Arsenic from Potable Water*, www.idswater.com/water/us/activated_alumina/457/products.html. 2010.
- [9] Reynolds, T.D., Richards, P.E., *Operational and Process Units in Environmental Engineering*. PWS Publishing Company, Boston, London, 2000.
- [10] Jain, C.K, Ali, I., *Arsenic: Occurrence, Toxicity and Speciation Techniques*. ater Research **34 (17)**, pp. 4304–4312. 2000.
- [11] Badalians Gholikandi, G., *Water Chemistry*, 2nd edition, Publ. Nopardazan, Tehran, Iran, 2006.
- [12] Vu, K.B., Kaminski, M.D., Nunez, L., *Review of Arsenic Removal Technologies for Contaminated Groundwaters*, ANL-CMT-03/2, Argonne National Laboratory, Illinois 60439, University of Chicago, USA. 2003.
- [13] Johnston, R., Heijnen, H., *Safe Water Technology for Arsenic Removal*, citeseerx.ist.psu.edu/.../arsenic/.../Mohan_As_removal_adsorbents.pdf. 2002.
- [14] Ravenscroft, P, Homepage, Documents. www.rgs.org/.../Arsenic+the+geography+of+a+global+problem.htm, 2010.
- [15] Reynolds, T.D., Richards, P.E. (2000), *Operational and Process Units in Environmental Engineering*, Boston, London: PWS Pub. Co.
- [16] Mosaféri, M., Mesdaghinia, A.R., *Removal of Arsenic from Drinking Water Using Modified Activated Alumina*, *Iranian Water and Wastewater Journal*, No. **55**, Esfahan, Iran. 2005.

- [17] Wang, L., Chen, A., Fields, K., Arsenic Removal from Drinking Water by Ion Exchange and Activated Alumina Plants, USEPA/600/R-00/088, Cincinnati, USA. 2000.
- [18] Badalians Gholikandi, G., Abbaspour, R., *Technical and Financial Analysis of Groundwater Remediation System, Chemical Contaminants Treatment*, national research program, PWUT, Tehran, 2006.
- [19] APHA., *Standard Methods for the Examination of water and wastewater*. American Public Health Association: Washington, D. C. 2005.
- [20] Badalians Gholikandi, G., Sadrzadeh Ardebili, M., Riahi, R., Orumieh, HR., *Arsenic Polluted Groundwater: Epidemiological Study and Efficient Removal Method*, Environmental Health Risk V, Proceeding book, WIT, UK, pp. 133–142, 2009.
- [21] Clifford, D., Lin, C.C. (1991), *Arsenic III and Arsenic V Removal from Drinking Water in San Ysidro, New Mexico*, USEPA/600/2-91/011, Cincinnati, USA.
- [22] Clifford, D., *Ion Exchange and Inorganic Adsorption*. In American Water Works Association (AWWA)(Eds.), *Water Quality and Treatment: A Handbook of Community Water Supplies*. 5th ed., NY, McGraw-Hill. 1999.
- [23] Clifford, D., Rosenblum, E., *The Equilibrium Arsenic Capacity of Activated Alumina*, USEPA CR-807939-02, Cincinnati, USA. 1982.
- [24] Fox, K., Field Experience with Point-of-Use Treatment Systems for Arsenic Removal, *J. AWWA*, pp.94–101. 1989.
- [25] UStoday *Arsenic in drinking water seen as threat*, http://www.ustoday.com/news/world/2007-08-30-553404631_x.htm, 2007.