Stabilization of iron-arsenic deposits by encapsulation with montmorillonite inorganic gels

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Abstract
Encapsulation is recognized as an effective technique for enhancing the stability of hazardous waste by coating it with an inert material. In this work, an eco-friendly montmorillonite (Mt) inorganic gel with the characteristics of high viscosity, adsorption and easy preparation was developed and coated on the surface of iron-arsenic deposits (IAD) to restrain the release of arsenic (As). The encapsulation system investigated involves Mt/IAD mass ratio and aging temperature of the coated iron-arsenic deposits (C-IAD). The C-IAD was characterized by XRF, XRD, SEM-EDS, TEM, FTIR and BET. The results revealed that the IAD was completely encapsulated by the montmorillonite inorganic gel. From the experiment of stability, when IAD was coated with montmorillonite inorganic gel (Mt/IAD mass ratio 0.75) and aged at 25°C, As release decreased from 3.15 mg/L to 0.64 mg/L at pH 5 after 24 h, and then dropped to 0.11 mg/L after 7 d. Furthermore, the results indicated that the IAD encapsulated with montmorillonite inorganic gel was effective in suppressing the release of As under both weakly acidic and alkaline conditions.

Keywords: Arsenic, Encapsulation, Gel, Ion-exchange, Montmorillonite, Stabilization

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Received December 02, 2020 Accepted January 18, 2021

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1. Introduction

Arsenic (As), a nonmetallic element, is widely distributed in nature, and trace amounts of arsenic can be detected in soil, water, minerals and plants, as well as in normal human tissues [1]. Last century, As and its compounds are mainly used in alloy smelting, pesticide medicine, pigment and other industries [2]. At present, As and its compounds are restricted to some areas such as cosmetics industry because of their highly toxic and potent human carcinogen [1, 3]. Therefore, arsenic-containing residue and effluent must be treated and stored in appropriate ways [4]. As is difficult to control in solution, hence it is usually stored in the crystal structures of secondary arsenic minerals such as iron arsenate (FeAsO$_4$·12H$_2$O), iron (Fe) sulphoarsenates and sulphoarsenites, Ca, Mg and Ca-Mg arsenates, and arsenic sulfide (As$_2$S$_3$) [4]. Among them, crystalline ferric arsenate (scorodite) has been proposed as an effective storage material due to its low As release (typically less than 1 ppm As at pH 5) and its high As content (25–30 wt%) [5]. However, the synthesis of scorodite requires high temperatures and high-pressure conditions which would be an economic burden to most factories. Normally, an alternative amorphous ferric arsenate was produced under ambient-pressure conditions which reduce the release of As by increasing the molar ratio of Fe/As [6, 7]. Both scorodite and amorphous ferric arsenate are stable only at pH 4 to 7 under aerobic conditions [8, 9], and the carbonation of containing ferric may occur over a long period of time [10]. Therefore, it is beneficial to explore effective methods for enhancing the stability of iron-arsenic compounds.

Encapsulation is the process of coating hazardous waste containing heavy metals/metalloids (such as As) with inert substances to prevent their dissolution [11]. Many studies [12-14] have used encapsulation to improve the stability of solid wastes containing As.
Jing et al. [12] utilized the cement as encapsulants for high As content sludge, and the results confirmed that the formation of calcium arsenate can increase the stability of cement-treated sludge. However, calcium arsenate will react with carbon dioxide in the air to consume calcium and reduce the stabilizing effect of cement on sludge. Randall [13] also showed that Portland cement with ferrous sulfate and lime could reduce the release of As from arsenic-containing materials, and found that the ligand substitution of OH⁻ and As species will prevent the resorption of As. But both Terra-Bond™ and PFL treated samples released significantly more mass of As at higher pH (e.g. pH = 12). Ke et al. [14] founded that the crystalline polyferric sulfate coating layer can effectively inhibit the release of As from scorodite particles in both weakly acidic and alkaline solutions under both oxic and anoxic conditions. The stabilizing effects of crystalline polyferric sulfate on scorodite are mainly due to the suppression of ion exchange between PO₄³⁻ and AsO₄³⁻ and the adsorption of As by crystalline polyferric sulfate itself. However, encapsulation conditions of this method are relatively harsh, which must be carried out at 90°C and 1.5 pH. These studies indicate that encapsulation materials present as porous structure consisting of flakes may restrain the release of As from arsenic-containing waste. The purpose of this paper is to develop a material and method that can control the coating thickness without reducing the stability of arsenic-containing waste, so as to improve the fixation rate of As and reduce the total amount of waste residue. Montmorillonite inorganic gel was selected as encapsulating material because of its three-dimensional network structure and excellent cation exchange performance, which may be effective to encapsulate arsenic-containing wastes and restrain the release of anions such as AsO₄³⁻ and AsO₂⁻. Specifically this
paper will investigate the process of montmorillonite inorganic gel production, coating and aging, as well as the stability of encapsulated iron-arsenic deposits under different pH conditions.

2. Material and Methods

2.1. Materials

In this study, the water containing As was obtained from the leaching solution of pyrite cinder using 18% sulphuric acid. The acid leaching was carried out in a 3 L glass beaker at 85°C. The pulp was filtered to separation of solid-liquid after 4 h. The Fe and As contents in the filtrate were determined by inductively coupled plasma-atomic emission spectrometry (ICP-OES, Optima 8000). According to the testing results of ICP-OES, the Fe/As molar ratio was approximately 4.5:1 and the As containing in the solution was 2,260 mg/L, and the solution pH was 0.12.

Montmorillonite was provided by Beijing Innochem (A77948-500G; CAS:1318-93-0). In addition, iron power, hydrogen peroxide, sodium hydroxide, anhydrous sodium carbonate and calcium oxide were taken from Sinopharm Chemical Reagent Company. All the chemical reagents were analytical grade. Deionized water produced by a water system (Barnsted, Nanopure Ultrapure Water System, specific conductance < 0.057 μS/cm) was used for all experiments.

2.2. Preparation of Iron-Arsenic Deposits

The pH of the iron-arsenic effluent was adjusted to 0.8-1.0 by adding 20 wt% NaOH. Then added excessive reducing iron powder (1.5 times of the theoretical amount) to reduce Fe$^{III}$ to Fe$^{II}$. 
The treated effluent was stirred for 6 h at 50°C and filtered to remove the excess iron powder. Then the filtrate was added 20 wt% NaOH solution to maintain pH in the range of 5.0-5.5 when a certain amount of 30 wt% hydrogen peroxide (H$_2$O$_2$/As molar ratio of 2.0) was dripped into the solution. After stirring for 2 h at room temperature (25°C), the solution was separated and the residue was washed with deionized water (10 times of the solution) to remove impurities such as Na$^+$ and Ca$^{2+}$. The residue was dried at 100°C and subsequently used for encapsulation with montmorillonite inorganic gel.

2.3. Production of Montmorillonite Inorganic Gel

Montmorillonite inorganic gel was prepared by the sodium modification and gelatinization of calcium-based montmorillonite. Taken certain quality montmorillonite and added deionized water to prepare 5% pulp, ultrasonic crush in an ultrasound washer at 25°C for 20 min. Then sodium carbonate with 8% of the mass of montmorillonite was added as the sodium reagent and the pulp was magnetically stirred at 25°C for about 20 min. Subsequently, 4% calcium oxide was added as the gel agent and a viscous gel formed after 15 min [15]. During the production of montmorillonite inorganic gel, the stirring rate was kept at 100 r/min.

2.4. Ageing of Iron-Arsenic Deposits with Montmorillonite Inorganic Gel

The synthetic IAD and montmorillonite inorganic gel were thoroughly blended together and aged under various conditions, including Mt/IAD mass ratio (0, 0.1, 0.25, 0.5, 0.75, 1, 1.25 and 1.5) in the blended product and aging temperatures (25, 50, 75, 100, 125, 150 and 175°C). The aged gel-coated products were ground using a pestle and mortar before sample testing.
2.5. Stability Testing

The aged gel-coated products were placed in a tube with a cock and used acetate buffering solution to evaluate their long-term stability. Stability testing was done under different conditions (pH at 3, 5, 7, 9 and 11) in order to better assess the efficacy of the encapsulation capability of the montmorillonite inorganic gel. Acetate buffering solution of pH 3 was obtained by dissolving 5.7 mL of glacial CH₃CH₂OOH with reagent water to a volume of 1 L. Other buffers of different pH (pH at 5, 7, 9 and 11) were obtained as follows: 1 M NaOH solution was added dropwise to a buffer of pH 3 until the predetermined pH value was reached. In order to sample multiple times during the long-term stability test, the liquid to solid ratio (L/S) was kept at 40 for all tests [11]. The tube was placed and fixed on a horizontal vibrating device with vibrating frequency at 110 times/min under room temperature (22°C). Samples were taken on a regular basis with 5 mL syringes and then filtered using 0.2 μm membranes, and the filtered liquor was diluted with a 4% nitric acid solution to prepare for analysis of Fe and As concentrations using ICP.

2.6. Analysis and Characterization

The contents of Fe and As in the solid were obtained by dissolving the solid with HCl and HNO₃ and then using ICP-OES to measure the content of elements in the solution as described in our previous paper [16]. The pH value was detected by pH meter (pHs-3E, Leici, China). The chemical analysis for starting materials was carried out with WD XRF fluorescence spectrometer Axios mAX 4 kW by PANalitycal company, equipped with Rh source. XRD measurements were
carried on a Philips X’Pert Pro diffractometer equipped with Cu Kα radiation (λ = 0.15418 nm).

The total metal ions concentrations in the solution were measured by inductively coupled plasma-atomic emission spectrometry (ICP-OES, Optima 8000). The surface morphology and microstructure of the samples were characterized by the high-resolution field emission transmission electron microscope (TEM, JEM-2100F) and the scanning electron microscope (SEM, model 6700F, Tokyo, Japan) with energy dispersive spectrometer (EDS). Elemental mapping of particle cross-sections was achieved by mounting particles in cold-setting epoxy resin, polishing the samples and coating with a gold/palladium layer before obtaining X-ray maps with a Hitachi S-3000N variable pressure scanning electron microscope (VP-SEM). Teller (BET) specific surface area and pore structures were measured by N₂ adsorption-desorption isotherms at 77 K with ASAP 2020 apparatus (Micromertics, USA). The infrared absorption test was conducted to indicate the structures of the precipitates using Fourier transform infrared spectrometer (FTIR, Excalibur 3100).

3. Results and Discussion

3.1. The Chemical Composition Analysis of Raw Materials

The chemical composition of the montmorillonite, montmorillonite inorganic gel, IAD and C-IAD (Mt/IAD 0.75; aged at 25°C) is presented in Table S1. The chemical analysis showed that montmorillonite and montmorillonite inorganic gel has a high content of SiO₂ (over 66 wt%), the increase of Na₂O and CaO in gel is due to the addition of sodium reagent and gel agent in the preparation process of gel. IAD showed the significant presence of Fe₂O₃ (34.573 wt%), the presence of SO₃ (28.625 wt%) and As₂O₃ (11.182 wt%) proved that IAD was prepared from
acidic arsenic-containing effluent. Moreover, C-IAD exhibited high content of SiO$_2$ (42.145 wt%) and Fe$_2$O$_3$ (21.483 wt%), which was the mixture of gel and IAD.

3.2. The Theoretical Basis of The Formation of Montmorillonite Inorganic Gel

Montmorillonite is a layered mineral composed of very fine water-containing aluminosilicate with substantial isomorphic substitution, belonging to the monoclinic crystal system. Its crystal structure is a 2:1 layered silicate with each layer composed of one aluminum (magnesium) oxygen octahedral sheet sandwiched between two tetrahedral silicate sheets [17, 18]. Montmorillonite has strong cation exchange performance and water swelling performance, so the high-valent cations in the crystal structure are easily replaced by low-valent large-radius cations [19]. However, As usually exists in the form of two anions (AsO$_4^{3-}$ and AsO$_2^{-}$), and will not be affected by the cation exchange performance of montmorillonite. The montmorillonite used in this study was Ca-montmorillonite, which was first modified by adding sodium carbonate into the slurry during the preparation of the gel. When sodium modification, one Ca$^{2+}$ is usually replaced by two Na$^+$. Na-montmorillonite has a higher Zeta potential, the particles are not easy to agglomerate, and the montmorillonite sheets have good dispersibility [20]. After the addition of the gelling agent CaO, the end faces of montmorillonite sheets were connected with each other by Ca$^{2+}$ under the action of static electricity to form an inorganic gel with a three-dimensional grid structure [21].

Fig. 1 shows the XRD patterns of the montmorillonite, IAD, montmorillonite inorganic gel (aged at 25°C and 175°C) and C-IAD (Mt/IAD 0.75; aged at 25°C). And the standard peaks of the corresponding reference materials are shown in Fig. S1. XRD pattern of montmorillonite
exhibits intense diffraction peaks at angles \((2\theta)\) of 5.9, 19.8, 20.8, 26.6, 29.4 and 31.5°, indicating that the main mineral phases of montmorillonite are calcium montmorillonite (Ca-Mt), silica \((\text{SiO}_2)\) and calcium carbonate \((\text{CaCO}_3)\). After the preparation of the gel with montmorillonite, the peaks of the Ca-Mt decrease significantly and shifted to the right, the peak type became flat and broad. Meanwhile, according to Bragg’s equation: \(\lambda = 2d\sin\theta\), the \(d\) decreased from 1.49nm to 1.28nm, indicating that there is residual Na-Mt in the gel.

### 3.3. Characterization of Gel Coating

Fig. 2(a) presents the contents of Fe and As in the IAD and C-IAD with different Mt/IAD mass ratios aged at 100°C. The results demonstrate that Fe content is decreased from 22.18 to 10.77% and As content is decreased from 7.95 to 3.46% as the Mt/IAD mass ratio increased from 0 to 1.5. X-ray diffraction patterns of the C-IAD (aged at 100°C) with different Mt/IAD mass ratios are shown in Fig. 2(b). There are no characteristic peaks in IAD (Mt/IAD mass ratio is 0), indicating that they are poorly-crystalline [7]. Only one broad peak is observed in the range of \(2\theta = 10-30°\). This broad peak disappeared obviously even at the very low Mt/IAD mass ratio of 0.1 indicating that the montmorillonite inorganic gel successfully covered the surface of the precipitate. The diffraction peaks of the montmorillonite are gradually discernible with the increase of montmorillonite dosage, and we conclude that the peaks in the coated products correspond to the two mineral phases of IAD and montmorillonite inorganic gel.

SEM images of the IAD and C-IAD (aged at 25°C) under different magnification and the TEM image of the C-IAD (Mt/IAD 0.75; aged at 100°C) are illustrated in Fig. 3. It can be seen that the surface of the uncoated IAD is relatively smooth. As can be seen in Fig. 3(a)-(c), the
surface of the coated product is getting rougher as the Mt/IAD mass ratios increase from 0 to 0.75. As shown in Fig. 3(b), the surface of the coated product (Mt/IAD 0.1) is coated with fine flakes, and the coating is relatively uniform. Fig. 3(c) shows that the flakes accumulate randomly on the solid surface and more irregular porous structures appear when the Mt/IAD mass ratio is 0.75. The TEM image of Fig. 3(d) presents that the montmorillonite inorganic gel coating is closely bound to IAD and its thickness is uneven. The special morphology indicates that the increase of montmorillonite dosage may produce a certain adsorption capacity [22, 23].

Fig. 4 exhibits SEM and backscattered electron (BSE) images of the cross section of the coated products (Mt/IAD 0.75; aged at 25°C) along with the associated EDS mapping. Fe, Si and As elemental X-ray maps of Fig. 4(b) confirm that the core material of the coated products is IAD and the montmorillonite inorganic gel has formed a mineralized matrix around the IAD. This coating layer is not uniform. As we can see in Fig. 4(c), BSE image shows that region B is dark in color and obviously different from other regions, indicating that the region B is an independent phase different from regions A and C. And SEM-EDX radial compositional analysis results at line L in Fig. 4(d) show that the Si/Fe ratio is obviously different in the A, B, and C regions, which is related to their phase. Combining Fig. 4(c) and Fig. 4(d), we speculate that through the coating of montmorillonite inorganic gel, it has a high probability of forming a tightly bound intermediate layer between the gel and IAD. Obviously, it can be seen that region A is montmorillonite inorganic gel, region C is IAD, and region B may be the ferric silicate intermediate layer between IAD and gel [11].

Fourier transform infrared (FTIR) spectrum of montmorillonite, IAD, montmorillonite inorganic gel (aged at 25°C and 175°C) and C-IAD (Mt/IAD 0.75; aged at 25°C) are shown in
Fig. 5. At 400-800 cm\(^{-1}\), the spectra of IAD shows the bending and stretching vibration peaks of O-As-O and As-O at 432 and 610 cm\(^{-1}\), which are the main differences between IAD and C-IAD [12, 24, 25]. And the bands of O-As-O and As-O are not shown in the spectra of C-IAD because they are masked by the bending stretching vibration peak of Si-O-Si band as well as weak stretching vibration peak of Si-O band, which usually occurs in the range of 400-500 cm\(^{-1}\) and ~660 cm\(^{-1}\) [26, 27]. All samples have a complex group of bands at \(\sim 1,060 \text{ cm}^{-1}\) that is attributed to the asymmetric and symmetric stretching vibrations of Si-O bonds [27, 28]. The increase in the strength of this peak indicates the polymerization of silicate. These differences reflect its more extensive polymerization of the silicate due to the encapsulation of montmorillonite inorganic gel. The weak shoulder at 839 cm\(^{-1}\) is attributed to the out-of-plane bending of CO\(_3^{2-}\), and the band at 1,456 cm\(^{-1}\) correspond to the asymmetric stretching of CO\(_3^{2-}\) [27]. All samples produced essentially identical spectra at 1,638 and 3,429 cm\(^{-1}\), where the peaks are assigned to OH\(^-\) and H\(_2\)O groups [11, 24, 27]. The 3,626 cm\(^{-1}\) band in these samples probably corresponding to CaOH groups [27].

The specific surface and the pore structures of IAD and C-IAD (Mt/IAD 0.75, aged at 25\(^\circ\)C) are obtained by the N\(_2\) adsorption-desorption isotherm (see Fig. 6). A hysteresis loop is present in the adsorption isotherm of IAD in Fig. 6(a) indicating the presence of abundant mesopores [29]. Pore size distribution analysis (Fig. 6(b)) shows that both IAD and C-IAD have narrow pore distribution in the range of 10-20 Å, and the average pore radius of the IAD and the C-IAD are 51.4 Å and 38.4 Å, respectively. Correspondingly, the specific surface area of C-IAD (86.902 m\(^2\)g\(^{-1}\)) is larger than that of IAD (30.989 m\(^2\)g\(^{-1}\)). The noticeable increase in surface area of C-IAD is attributed to the encapsulation of montmorillonite inorganic gel, which has a porous
network structure [21, 30, 31]. And C-IAD has a smaller pore radius, which helps to restrain the
release of As.

3.4. Ageing of Montmorillonite Inorganic Gel

The aging process of C-IAD is essentially the hydration process of the gel. Hydration of the gel
system is a complicated process and mainly affected by temperature. The XRD patterns of the
montmorillonite inorganic gel aged at 25℃ and 175℃ are illustrated in Fig. 1. As can be seen,
compared with the 25℃ gel, the intensity of the XRD spectra of Na-Mt and CaCO₃ in 175℃ gel
is reduced, while that of SiO₂ is enhanced. The XRD results show that the increase of aging
temperature will promote the hydration process and increase the crystallinity of SiO₂ [27]. The
FTIR spectral bands of the montmorillonite inorganic gel aged at 25℃ and 175℃ can be seen in
Fig. 5. When the temperature rises from 25 to 175℃, there is an increase in the intensity of Si-O,
Si-O-Si, CO₃²⁻ and CaOH peaks at 1,060, 446, 519, 839, 1,456 and 3,626 cm⁻¹, and a decrease in
the intensity of H₂O peaks at 1,638 and 3,429 cm⁻¹. These observations indicate that the increase
of aging temperature will lose the OH⁻ group and increase the degree of polymerization of the
silicate chain [26, 32]. The conclusion is consistent with the results of XRD analysis, where the
characteristic peaks strength of SiO₂ also increased when the aging temperature increased from
25℃ to 175℃ in Fig. 1.

3.5. Stability Experiments

3.5.1. Effect of the Mt/IAD mass ratio
Fig. 7(a) displays the results of the leaching concentrations of Fe and As in the filtrate from the leaching toxicity tests at pH 5.0 for 24 h. The pure IAD and the C-IAD with different Mt/IAD mass ratios (0.1, 0.25, 0.5, 0.75, 1, 1.25 and 1.5) are all aged at 25°C. It can be seen that the leaching concentrations of Fe decrease from 42.9 to 1.611 mg/L when the ratio of Mt/IAD is 0.1 and the Fe release is not detectable when the Mt/IAD mass ratio increase to 0.75. The release of As from uncoated IAD is 3.152 mg/L at pH 5.0 after 24 h. For IAD coated with montmorillonite inorganic gel, the As in the leaching solution decreases to 1.723 mg/L when the Mt/IAD mass ratio is 0.1. However, the restrain of As release hardly increases with the further increase of montmorillonite dosage. In Fig. 7(a), it can be clearly seen that there are two inflection points on the As concentration curve, respectively at the Mt/IAD of 0.1 and 0.75. Linear fitting was carried out for line segment ab, bc and cd, and it can be known that the slope was -14.29, -0.49 and -0.16 respectively. The change in slope indicates that the effect of montmorillonite inorganic gel on As release hardly increases with the increase of Mt/IAD mass ratio after reaching 0.75. Combined with Fig. 2(a), after the Mt/IAD mass ratio reaches 0.75, the reduction of As leaching toxicity may only be due to the decrease of As content in the coated products. The results showed that montmorillonite inorganic gel could effectively inhibit the leaching of Fe and As, even at the very low Mt/IAD mass ratio of 0.1.

3.5.2. Effect of the leaching pH

The C-IAD prepared with 0.75 times the mass ratio of Mt/IAD, which after aging for 24 h in the oven at 25°C is subjected to stability testing under different pH conditions (pH at 3, 5, 7, 9 and 11) over 7 d. Fig. 7(b) indicates the relationship of As concentration with time under different pH
conditions. As can be seen in Fig. 7(b), the concentration of As decreases with the increase of time under different pH conditions, indicating that montmorillonite inorganic gel had stable adsorption characteristics. And it can be seen that the C-IAD is more stable in weakly acidic and neutral conditions (pH at 5 and 7) and less stable in acidic or alkaline conditions (pH at 3, 9 and 11). Fig. 7(b) also shows that the As concentration showed an increasing trend in the first three days at pH 9 and 11. The reason is that the adsorption rate of montmorillonite to arsenic under alkaline condition is slow [33, 34], which is lower than the leaching rate of arsenic in the early stage. The poor stability of C-IAD under acidic conditions (pH = 3) probably owe to the fact that the montmorillonite gel itself is an alkaline material and acid conditions will destroy its network structure. Although the stability of C-IAD is lower in acidic environment (As release 2.47 mg/L after 24 h), it is still higher than that of uncoated IAD in weakly acidic condition (As release 3.15 mg/L after 24h in Fig. 7(a)). Therefore, it can be concluded that the montmorillonite inorganic gel coating could restrain the release of As from IAD in both weakly acidic and alkaline conditions.

3.5.3. Effect of the aging temperature

Fig. 8(a) illustrates the results of As release from IAD and C-IAD (Mt/IAD mass ratio 0.75) under the weakly acidic conditions for 7 d using an acetate buffering solution of pH 3. It can be seen that the As release is indeed reduced when IAD are coated with montmorillonite inorganic gel. After 7 d of leaching test, the As release from C-IAD (aged at 25°C) decrease from 0.64 to 0.11 mg/L, but the As release from uncoated IAD increase from 3.15 to 3.49 mg/L. The results show that montmorillonite inorganic gel could not only restrain the release of As in IAD, but
also adsorb As in solution. Fig. 8(b) shows the relationship between As concentration in the leaching solution (pH = 5.0) and aging temperature after 24 h. As can be seen in Fig. 8(b), the As concentration first increases and then decreases as the temperature increases. And aged sample has the highest initial As release (1.403 mg/L) at 100°C. Combined with the analysis of the aging of montmorillonite inorganic gel, we can know that the release of As increases as the aging temperature increases from 25°C to 100°C. It may be caused by the reduction of water molecules and OH− groups in the gel, and further caused the collapse of the silicate layer. After the temperature exceeds 100°C, the layer structure collapses to a point and the gel structure becomes denser, increasing its inhibition on As release. The results are consistent with Xiandong Cong, who said that oven dry (110°C) of gel not only has the collapse of the layer structure, but also changed in the polymerization of silicate chains [35].

In summary, when IAD was coated with montmorillonite inorganic gel (Mt/IAD mass ratio 0.75) and aged at 25°C, As release decreased from 3.15 mg/L to 0.11 mg/L at pH 5 after 7 d. The inhibition rate of inorganic montmorillonite gel to arsenic release was 96.51%, which was higher than that of aluminum phosphate gel (90%) [36] and aluminum chloride gel (31.82%) [24]. The excellent encapsulation ability of montmorillonite inorganic gel may be attributed to the combination of the ferric silicate intermediate layer formed by the reaction of the montmorillonite gel with IAD and the adsorption capacity of the montmorillonite itself.

3.6. Mechanism Analysis

Through the stability test of C-IAD prepared and aged under different conditions, it can be seen that the use of montmorillonite inorganic gel has the ability to inhibit the release of As and
adsorb As in solution. The possible inhibitory mechanism of montmorillonite inorganic gel on the As release is shown in Fig. S2. The silicate flakes in the montmorillonite inorganic gel first adhered to the IAD uniformly as a single layer and then piled up in a grid to form a porous structure as the amount of montmorillonite increased. An intermediate layer of iron silicate may have formed between IAD and the montmorillonite inorganic gel. The intermediate layer may have been formed by the silicate reacting with Fe$^{III}$ on the surface of IAD, since the stability constant of the iron-silicate complex is much higher than the complexes that form between arsenate and Fe$^{III}$ (at least 7 orders of magnitude difference) [11]. This behavior can be explained by an “ion exchange”-type process between SiO$_3^{2-}$ and AsO$_4^{3-}$ according to reaction (1):

$$2\text{FeAsO}_4 \cdot x\text{H}_2\text{O} + 3\text{SiO}_3^{2-} \rightarrow \text{Fe}_2(\text{SiO}_3)_3 \cdot x\text{H}_2\text{O} + 2\text{AsO}_4^{3-}$$ (1)

Montmorillonite itself has a certain adsorption capacity [37, 38], so As released during the ion exchange reaction and As in the aqueous solution may be adsorbed in the gel network. This can be demonstrated by our results in Fig. 4(c)-(d), where the gels in region A also contain small amounts of As. And the adsorption of arsenic on montmorillonite is mainly through the attraction of arsenic by the metal hydroxyl groups (such as Al-OH, Mg-OH and Fe-OH) on the surface and inside of montmorillonite to form outer sphere complex and inner sphere complex [39, 40].

4. Conclusions

Montmorillonite inorganic gels were used to stabilize the hazardous arsenical material. The major findings in the current study include: (1) The characterization results showed that the IAD was encapsulated completely by montmorillonite inorganic gel on its surface. Optimum encapsulating conditions are the Mt/IAD mass ratio of 0.75 and the aging temperature of 25°C.
(2) The stability tests presented that the coated of IAD with montmorillonite inorganic gel appears to be effective in restraining the release of As under both weakly acidic and alkaline solutions (pH 3 to 11). Under optimal encapsulating conditions, after equilibrating in a buffer solution at pH 5 for 24 h, the As released from uncoated IAD and C-IAD was 3.15 mg/L and 0.64 mg/L, respectively. What's more, the As released from uncoated IAD increased to 3.50 mg/L and the As released from C-IAD decreased to 0.11 mg/L after 7 d. (3) Arsenic is immobilized in the montmorillonite inorganic gel through a combination of encapsulation and adsorption. Through encapsulation, a resistant iron-silicate layer may have formed between the IAD and the montmorillonite inorganic gel by an “ion exchange” reaction between $\text{SiO}_3^{2-}$ and $\text{AsO}_4^{3-}$. Through adsorption, As released by ion exchange is adsorbed in the porous network structure of montmorillonite inorganic gel.

Acknowledgement

This work was financially supported by the Innovation Academy for Green Manufacture, Chinese Academy of Sciences (No. IAGM-2019-A05) and the National Key Research and Development Program (No. 2019YFC1904204).

Author Contributions

J.Y. (Master student) conducted all the experiments and wrote the articles. Y.W. (Associate professor) got funding and modified the article. Y.W. (Master student) conducted supplementary experiments and reviewed the data. X.L. (Ph.D. student) assisted in some of the experiments. S.Y. (Professor) conducted experimental supervision, paper review and financial support.
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Fig. 1. XRD patterns of the montmorillonite, IAD, montmorillonite inorganic gel (aged at 25℃ and 175℃) and C-IAD (Mt/IAD 0.75; aged at 25℃).
Fig. 2. Element content and XRD patterns of C-IAD (aged at 100°C) with different Mt/IAD mass ratios: (a) Element content of Fe and As; (b) XRD patterns.

Fig. 3. SEM and TEM images of the IAD and C-IAD: (a) surface structure of IAD; (b) surface structure of C-IAD (Mt/IAD mass ratio is 0.10); (c) surface structure of C-IAD (Mt/IAD mass ratio is 0.75); (d) TEM image of C-IAD (Mt/IAD mass ratio is 0.75).
**Fig. 4.** SEM image, backscattered electron (BSE) image and elemental X-ray maps of the cross section of C-IAD (Mt/IAD 0.75; aged at 25°C): (a) SEM image; (b) SEM image of M area in Fig. 4(a); (c) Backscattered electron (BSE) image of Fig. 4(b); (d) SEM-EDX radial compositional analysis at line L in Fig. 4(c).

**Fig. 5.** FT-IR spectra of montmorillonite, IAD, montmorillonite inorganic gel (aged at 25°C and 175°C) and C-IAD (Mt/IAD 0.75; aged at 25°C).
Fig. 6. Nitrogen adsorption–desorption isotherms (a) of IAD and C-IAD (Mt/IAD 0.75; aged at 25℃) and (b) the corresponding pore size distribution curve calculated from the desorption branch of the nitrogen isotherm by the BJH method.

Fig. 7. Stability experiments: (a) Fe and As release from IAD and C-IAD (aged at 25℃) after 24h of equilibration at pH 5.0; (b) As release from C-IAD (Mt/IAD = 0.75, aged at 25℃) under different pH conditions.
**Fig. 8.** As release from IAD and C-IAD (M/IAD = 0.75) under different aging temperatures: (a) after 7 d of equilibration at pH 5.0 and (b) after 24 h of equilibration at pH 5.0.