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Adsorption Study using Optimised 3D Organised Mesoporous Silica Coated with Fe and Al Oxides for Specific As(III) and As(V) Removal from Contaminated Synthetic Groundwater

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Abstract:

This work presents the possibility of optimising 3D Organised Mesoporous Silica (OMS) coated with both iron and aluminium oxides for the optimal removal of As(III) and As(V) from synthetic contaminated water. The materials developed were fully characterised and were tested for removing arsenic in batch experiments. The effect of total Al to Fe oxides coating on the selective removal of As(III) and As(V) was studied. It was shown that 8% metal coating was the optimal configuration for the coated OMS materials in removing arsenic. The effect of arsenic initial concentration and pH, kinetics and diffusion mechanisms was studied, modelled and discussed. It was shown that the advantage of an organised material over an un-structured sorbent was very limited in terms of kinetic and diffusion under the experimental conditions. It was shown that physisorption was the main adsorption process involved in As removal by the coated OMS. Maximum adsorption capacity of 55 mg As(V).g⁻¹ was noticed at pH 5 for material coated with 8% Al oxides while 35 mg As(V).g⁻¹ was removed at pH 4 for equivalent material coated with Fe oxides.
Highlights:

- 3D and 2D Organised Mesoporous Silica were efficiently coated with different Al to Fe oxides ratio
- 8 % metal oxide loading was estimated to be the most appropriate content for efficient As removal
- Successful tailoring of adsorbent toward specific As(III) to As(V) ratio by varying Al to Fe oxide content was confirmed
- Tailoring of the adsorbents produced is highly dependent on the pH of the water treated as well as the level of contaminant concentration
- No superior diffusion of pollutants into organised materials over unorganised materials was demonstrated

Keywords: Organised Mesoporous Silica, iron oxide, aluminium oxide, arsenic removal, diffusion
1 Introduction

Arsenic contamination is a problem affecting all countries throughout the world. However the countries in south and south-east Asia are considered the most vulnerable places facing this crisis which might impact many lives in the coming years [1]. The drinking water guideline from the World Health Organisation for arsenic levels was lowered to 10 ppb in 1993 [2]. Since then, many governments have modified their legislation to adopt this guideline value as a national standard. Nevertheless, a few countries, which are facing severe arsenic contamination are still using the previous value of 50 ppb, Bangladesh being one of them [1,3]. The adsorption technology has been extensively used to remove toxics from water because of its low running costs, low maintenance demand and the effectiveness of the process to guarantee a good drinking water quality [4,5]. The different types of materials used for arsenic removal can be divided into three major categories: organic or natural ore, industrial by-product and man-made sorbents mainly relying on metal oxides. The optimisation of material removal capacity relies very often on the increase of adsorbent surface area and pore volume, the doping of materials with oxide with specific affinity toward As, or, the addition of specific chemical surface groups for either chemisorption or ion-exchange of dissolved ions [6,7]. The use of nano-sized metal oxides such as Organised Mesoporous Silica has been regarded as a major technique is enhancing pollutant removal [8]. OMS materials are very versatile materials, easy to tailor and adapt to any specific pollutant. The development of OMS was first reported by researchers at the Mobil Oil Corporation [9]. Since then, many researchers have worked on the application of organised materials mainly for catalysis, air treatment and water treatment applications [10]. Organised mesoporous silicas are interesting materials due to their high surface area, narrow pore size distribution range and high porosity [11]. The most interesting feature of OMS is the
possibility to control, to a certain extent, most of its physical characteristics: mean pore size, porosity and surface area. OMS are useful materials as they can be a support matrix for many doping of interest. The addition of metal oxides in OMS is largely reported, mainly for its use in catalytic reactions [12] and [13] and in the removal of specific pollutant [14]. Production of mesoporous carbon can also be prepared using OMS as a template [14] and these materials have been successfully applied as adsorbents for water pollution control [15]. The same idea can be applied in order to create mesoporous metal oxides [16].

Previously, the potential for optimising the As adsorption capacity of coated OMS has been briefly presented [17]. In this current work a comprehensive and full detailed study is presented, covering both the production of the coated OMS sorbents and the impact on the As adsorption characteristics of these materials.

Li reported a coating process for iron oxides loading onto OMS for enhanced As(V) adsorption. Li showed that a layer by layer coating process was an efficient technique for iron loading into OMS in regards to As(V) adsorption [18].

Other studies reported innovative technics to load iron oxides into Organised Mesoporous Carbon (OMC), aluminium oxides into OMS or the direct production of Organised Mesoporous Iron oxides and their used in removing arsenic or fluoride [19–22]. The removal capacities reported were in the range of 0.16 to 4.1 mg.g\(^{-1}\) for As(V), 4.0 to 7.3 mg.g\(^{-1}\) for As(III) and 75.1 mg.g\(^{-1}\) for F\(^{-}\).

The originality of the present work relies onto the possibility of tailoring an adsorbent to match a specific groundwater contaminated by a pollutant present under two different oxidation stages, namely As(V) and As(III). Different ratio of iron and aluminium oxides were coated onto the 2D and 3D OMS materials and used to control the selectivity of the adsorbents for potential arsenic removal. The chemical and physical properties of the coated Al and Fe oxides OMS are fully characterised and discussed. Moreover adsorption of both
As(III) and As(V) are fully investigated, the effect of pH, the kinetic and diffusion of the As species into these materials are compared. Finally, the benefit of using organised materials over unorganised materials both in terms of adsorption removal capacity and diffusion or kinetic is discussed. The unorganised iron and aluminium oxides used as references were produced under the same synthesis conditions.

2 Materials and methods

2.1 Synthesis of 2D and 3D-Organised mesoporous silica coated with Fe and Al oxides

Organised Mesoporous Silica, KIT-6 (3D organised) and SBA-15 (2D organised) were produced using a modified method presented by Rumplecker and previously reported [17,23]. Briefly, for the production of SBA-15, a solution of 252 g of deionised water and 7.7 g of HCl at 37 % was used to dissolve 13.9 g of Pluronic P123. After dissolution of the surfactant, 25.0 g of TEOS was added. The solution was stirred in Duclan glass bottles for 24 h at 35 °C using a silicon oil bath. The bottles were then placed in an autoclave to undergo a hydrothermal treatment (at 40 °C, 60 °C, 80 °C or 100 °C) for 24 h under static conditions. The solid product was then filtered without any washing and dried for 48 h at 95 °C in an oven. The surfactant was removed by calcination in flow of air at 550 °C for 6 h with a temperature ramp of 1 °C.min⁻¹ and a flow of air set at 100 mL.min⁻¹. Similar steps were followed to produce the KIT-6 materials with some alterations: 9.0 g of P123, 325 g of deionised water and 17.4 g HCl were used. After complete dissolution of the surfactant, 9.0 g of butanol was added. Further steps were the same as for SBA-15 synthesis except the drying time which was reduced to 24 h. Mesoporous silica were named KIT-XX °C and SBA-XX °C where XX is the hydrothermal temperature used in the synthesis. The 3D-organised mesoporous silica KIT-100 °C was selected for aluminium and iron doping because of its higher mesoporosity. This process was carried out using a solvent evaporation method; 0.8 M
aluminium and iron nitrate in absolute ethanol solutions were used to coat the mesoporous silica. Different loading percentage were studied: 1 % to 8 % in one impregnation step and 15 % to 30 % in three impregnation steps. The percentage loading is based on the internal pores filling by $\text{Al}_2\text{O}_3$ or $\text{Fe}_2\text{O}_3$. The selected amount of iron or aluminium solution was added at once onto the mesoporous silica and was left under stirring for 1 h. After homogeneous dispersion of the solution, the HDPE bottles were placed in an oven at 60 °C to undergo solvent evaporation. The resulting powder was then calcined at 210 °C or 250 °C into a chamber furnace to eliminate the nitrate salts following the reactions presented in Equation (1) and (2). Nitrate is usually decomposed into $\text{NO}_2$, $\text{NO}$, $\text{NO}_2$ and $\text{NO}_3^-$ for clarity reason the nitrogen species were reported as $\text{N}_2$ [24]. Coated mesoporous silica were named KIT-YY ZZ % where ZZ is the percentage of coating and YY the corresponding metal salt (Al or Fe).

\[
\text{Fe(NO}_3)_3.9\text{H}_2\text{O} \xrightarrow{210 \degree \text{C}} 0.5\text{Fe}_2\text{O}_3 + 1.5\text{N}_2 + 3.75\text{O}_2 + 9\text{H}_2\text{O} \quad (1)
\]

\[
\text{Al(NO}_3)_3.9\text{H}_2\text{O} \xrightarrow{250 \degree \text{C}} 0.5\text{Al}_2\text{O}_3 + 1.5\text{N}_2 + 3.75\text{O}_2 + 9\text{H}_2\text{O} \quad (2)
\]

A set of mixed iron and aluminium coated mesoporous KIT-100 °C was produced by varying the relative percentage of iron and aluminium while keeping constant the total metal loading percentage at 8 %. The same coating procedure as for single metal oxide coating was followed. Mixed metal coated mesoporous silicas were named KIT-8 %-Fe/Al-AA/BB where AA and BB are the iron and aluminium relative content. In order to compare the impact of the 2D and 3D organisation on the diffusion of As(III) and As(V), raw unorganised iron and aluminium oxides were produced. The synthesis procedure followed was similar to the one used for metal oxide coating.

2.2 Adsorbents characterization

The BET (Brunauer, Emmett and Teller) surface area, porosity and average pore size of the different materials produced were measured using a Nova 4200e Surface area and pore size
analysed from Quantachrome Instruments. XRD (X-Ray Diffraction) analysis was carried out to first ascertain the organised structure of the OMS materials produced. The equipment used was a PANalytical X’pert model from Philips. Radiation was carried out by a Cu K-alpha 1 lamp at wavelength 0.154056 nm. A FEI Tecnai F20 Field emission high-resolution TEM (Transmission Electron Microscopy) was used to verify the 2D and 3D organised structure of OMS and coated OMS. The TEM used had a 200kV accelerating voltage giving it sufficient resolution for level imaging at the nm scale. The OMS and coated OMS samples fine powder were dispersed in acetone to ensure good dispersion of the particle and limit their agglomeration. The samples were then spread onto a CuO grid for analysis. An EOL JSM-6500F field emission SEM equipment was used for SEM-EDX (Scanning Electronic Microscope with Energy Dispersion X-Ray) analysis. Samples were analysed by FTIR (Fourier Transformation Infra-Red) using the KBr tableting method. A PerkinElmer Spectrum One FT-IR spectrometer instrument was used. The immersion technique was used to measure the PZC of the raw KIT material or KIT-100 °C as well as the PZC of KIT-Fe 8 % and KIT-Al 8 % materials. The procedure presented by Bourikas et al. was followed, [25]. A solid to liquid ratio of 1 % w/w was used. The corresponding materials were weighted in 60 mL plastic jars and solutions of different initial pH having an ionic strength of 0.1 M NaCl were used. Time before equilibrium was reached was selected as 24 h.

2.3 As(III) and As(V) adsorption onto coated OMS experiments

2.3.1 Chemicals and equipment used

The chemicals used during this work were all received from Sigma Aldrich and were all of reagent grade quality or higher. Deionised water with a resistivity of 18.2 MΩ.cm was used in the different solutions. Appropriate amounts of Na₂HAsO₄·7H₂O and NaAsO₂ were dissolved in deionised water to produce As(V) and As(III) stock solutions of 1000 ppm
respectively. 100 mg.L\(^{-1}\) of NaHCO\(_3\) was added in solutions used in adsorption experiments as a pH buffer. The pHs of all solution were adjusted using dilute solutions of HCl and NaOH.

2.3.2 Optimisation experiments

The first set of experiments explored the effect of iron and aluminium oxides percentage loading onto the OMS. KIT materials loaded with increasing Fe or Al oxides loading percentage were used to remove As(III) or As(V). 1.0 g.L\(^{-1}\) of material was used in 50 mL solution of 50 ppm of arsenic at pH 7 [17]. Solutions were stirred on a horizontal shaker for 48 h. The second set of experiments explored the effect of co-coating Fe and Al oxides onto the OMS at different ratios and studied the potential of tuning the adsorption capacity toward the arsenic species. Arsenic adsorption experiments using As(III) and As(V) in single or binary systems were carried out. The same experimental conditions were used: 1 g.L\(^{-1}\) of adsorbent, 50 mL of 50 ppm of As solution at pH 7 in 60 mL glass flasks and 48 h experiment time on a horizontal shaker working at 100 rpm.

2.3.3 pH effect on optimised sorbents and isotherm of adsorption

The effect of pH onto As(III) and As(V) adsorption onto some selected coated OMS, KIT-Al 8 % and KIT-Fe 8 %, were carried out in batch systems. The materials exhibited very high As removal capacity, so, initial adsorption concentration of 55 ppm of As(III) and As(V) was used alongside a dosage ratio of 1 g.L\(^{-1}\). Initial pHs ranging from 2 to 12 and equilibrium pHs were recorded by a Thermo Scientific Orion 3 Star pH meter equipped with a Camlab pH probe accurately calibrated over the study pH range. The isotherms of adsorption of arsenic species onto selected adsorbents were performed with initial As concentrations ranging from 1 to 50 ppm at pH 7.
2.3.4 Arsenic kinetics of adsorption onto 2D and 3D organised mesoporous silica coated with Fe and Al oxides and raw Fe and Al oxides

The influence of organised or unorganised structured materials onto the kinetic removal of As species was tested in 1 L beaker containing 1 L solution. The adsorbent dosage was 1 g L\(^{-1}\), initial pH 7 and the magnetic stirrers used working at 300 rpm. The kinetics data points come from regularly taken samples of 5 mL which were diluted to 10 mL using deionised water. Experiments were carried out over 48 h with initial concentration of 50 ppm approximately.

2.4 Concentration Analysis

The only dissolved element monitored in this study was arsenic and its analysis was carried out by ICP-OES. A Thermo Scientific IRIS Intrepid model was used at a working wavelength of 189.0 nm. Before being analysed, the samples were filtered through 13µm cut off cellulose acetate filters and acidified at 2 % HNO\(_3\). Data points obtained are the average of three consecutive analyses of the same sample.

2.5 Mathematical modelling

2.5.1 Isotherms models

The equilibrium data was modelled using Langmuir and Freundlich adsorption models, under Sigma plot 11.0 software. These models are usually represented by Equations (3) and (4) [26].

Langmuir model:

\[
q_e = \frac{q_b b_e C_e}{1 + b_e C_e}
\]  

(3)

Freundlich model:

\[
q_e = K_f C_e^{n_e}
\]  

(4)
where $q_e$ is the equilibrated adsorption capacity for a given initial concentration in mg.g$^{-1}$, $q_m$ is the maximum pollutant adsorption capacity expressed in mg.g$^{-1}$ and $C_e$ is the concentration in bulk solution at equilibrium in mg.L$^{-1}$. $b_L$ is the Langmuir constant in L.mg$^{-1}$ and $K_F$ is the Freundlich coefficient in mg$^{1-n}$.g$^{-1}$.L$^n$. $n$ is the Freundlich equation constant and is dimensionless. The separation factor $R_L$ presented by Weber and Chakravorti is known as the separation factor and defined by equation (5) [27].

$$R_L = \frac{1}{1 + b_L C_o}$$

(5)

where $C_o$ is the initial pollutant concentration in mg.L$^{-1}$. This separation factor is used to characterise the nature of the adsorption with unfavourable adsorption having a $R_L > 1$, $0 < R_L < 1$ for favourable reaction and $R_L = 0$ for irreversible reaction.

The Dubinin-Radushkevich (DR) isotherm is very often used to model adsorption isotherm data and determine the mean free energy of adsorption [28]. By using equation (6) and (7) the adsorption data can be modelled, while using equation (8), the mean free energy of adsorption $E$, expressed in kJ.mol$^{-1}$ can be calculated [29].

$$q_e = q_s \times e^{-\frac{k_{ad} \varepsilon_{RD}}{2}}$$

(6)

$$\varepsilon_{RD} = RT \ln \left(1 + \frac{1}{C_e}\right)$$

(7)

$$E = \frac{1}{\sqrt{2k_{ad}}}$$

(8)

where $q_s$ is the theoretical isotherm saturation capacity calculated in mg.g$^{-1}$, $k_{ad}$ the DR isotherm constant in mol$^2$.J$^{-2}$ and $\varepsilon_{DR}$ is the DR isotherm variable expressed in J.mol$^{-1}$.

2.5.2 Non-linear kinetics models

The pseudo first and pseudo second order models were applied to the kinetic experiments data obtained using Equations (9) and (10) [30,31].

11
Pseudo first order model:

\[
\ln \left( \frac{q_t - q_e}{q_e} \right) = -k_1 t
\]  \hspace{1cm} (9)

Pseudo second order model:

\[
\frac{1}{(q_e - q_t)} = \frac{1}{q_e} + k_2 t
\]  \hspace{1cm} (10)

where \( q_t \) is the amount of pollutant adsorbed at a specific time \( t \) in mg.g\(^{-1}\); \( k_1 \) and \( k_2 \) are the pseudo-first order kinetic constant and the pseudo second order kinetic constant expressed in in min\(^{-1}\) and in g.mg\(^{-1}\).min\(^{-1}\), respectively.

2.5.3 Diffusion modelling

The diffusion models were developed to study the diffusion of adsorbate inside the adsorbents. One of the first hypotheses used in these models is that the bulk liquid diffusion is not a limiting step, which implies that stirring energy is sufficient. In this study, two different diffusion models were applied: the intraparticle diffusion model and the Boyd diffusion model. The intraparticle model links the adsorbed quantity of pollutant at a given time with the time \( t \) following Equation (11), [32,33].

\[
q_t = k_{ad} t^{0.5}
\]  \hspace{1cm} (11)

where \( k_{ad} \), the intraparticle diffusion rate is a constant expressed in mg.g\(^{-1}\).h\(^{0.5}\) and \( i \) represents the diffusion stage number. The plot of \( q_t \) versus \( t \) is linear when the intraparticle diffusion is the main resistance step in the experimental conditions.

The Boyd diffusion model is usually expressed by the Equation (12), [34].

\[
\frac{q_t}{q_e} = 1 - \left( \frac{6}{\pi^2} \right) \sum_{n=1}^\infty \left( \frac{1}{n^2} \right) \exp(-n^2B \times t)
\]  \hspace{1cm} (12)
where $B$ is a coefficient expressed in h$^{-1}$. Resolving equation (12) can be done using Equations (13) and (14) based on the work of Reichenberg, [35].

For $q/t/q_e \geq 0.85$

$$B \times t = 0.4977 - \ln\left(1 - \frac{q}{q_e}\right)$$

(13)

For $q/t/q_e < 0.85$

$$B \times t = \left(\sqrt{\pi} - \left[\frac{\pi \cdot \frac{q}{q_e}}{3}\right]\right)^2$$

(14)

Using $B$ an average diffusion coefficient can be approximated using equation (15) as explained by Reichenberg, [35].

$$B = \frac{\pi^2 \times D_{\text{eff}}}{r}$$

(15)

where $r$ is the radius of the material particle. If the resulting plot of $B\times t$ versus $t$ is a straight line passing through the origin then the main resistance is based on intraparticle diffusion and $D_{\text{eff}} = D_{\text{intra}}$ [34]. Using an optimisation tool, the different diffusion constant were calculated for each diffusion step identified [34].

### 3 Results and discussion

#### 3.1 Characteristics of adsorbents samples

##### 3.1.1 Surface area, pore size distribution and porosity analysis

The pore size distributions of the uncoated KIT materials produced at different temperatures are presented in Figure 1 (a). It can be seen that the average pore diameter increases for KIT materials with an increase in the hydrothermal curing temperature. The wider average diameter obtained was evaluated at 6.4 nm for the KIT-100 °C material. In Figure 1 (b) it can
be noted that the KIT-100 °C displays a higher mesopores volume than the SBA-100 °C. The increase in the N\textsubscript{2} adsorbed volume is higher in the region of 0.6 and 0.8 relative pressure. Moreover, the steady higher increase of N\textsubscript{2} sorbed in the region of 0.1 and 0.5 relative pressure proves that the micropore volume of KIT-100 °C is also higher than the SBA-100 °C. Finally, the macropores volume of both materials is very limited as shown by the plateau achieved for both N\textsubscript{2} adsorption isotherms at relative pressure between 0.8 and 1.0. It can also be noted that both isotherms follow the type IV isotherm with a sharp increase and rather wide hysteresis loop type H1 as defined by IUPAC [36]. These features are characteristic of organized mesoporous materials with a narrow pore size distribution. Due to the superior physical characteristics displayed by KIT-100 °C mesoporous silica, this material was then selected for iron and aluminium oxides coating.

Table 1 presents the BET analysis results of the two series of KIT-100 °C materials coated by increasing iron or aluminium oxide content. It was found that, after 8 % of the pores were filled by iron oxides, the coated OMS presented a significant loss in surface area and porosity. This same difference is noted after 3 % of coating in the case of aluminium oxide coating. For samples coated by 15 % of metal oxides and higher, the coating procedure was achieved in two steps which might have increased the pore blockage and might have increased the amount of remaining unoxidized nitrate groups in the pores. In the case of aluminium oxides, the high presence of remaining unoxidized nitrate groups is clearly responsible for the early pore filling. The presence of remaining nitrate groups was identified by FTIR analysis and is discussed below.

Table 2 summarises the physical characteristics of the materials used in different adsorption experiments. The BET analysis measured a very low surface area and porosity for the Al oxide sample. The very low measurement obtained can be accounted for by the release of species from the remaining nitrate or an obstruction of the pores by nitrate molecules. Al
oxides produced using conventional sol-gel method usually display surface area in the region of 200 m$^2$.g$^{-1}$ and porosity around 0.4 cc.g$^{-1}$ [37].

3.1.2 XRD analysis: organised structure of material and coating identification

The amorphous nature of SBA-15 and KIT-6 materials produced at 100 °C was confirmed as no silica oxides crystal was identified, as no major peaks were identified at high diffraction angle, see Figure 2 (a). The XRD spectrum of both materials show the organised nature of their respective structure as indicated by the low angle peaks noticed at 0.7 ° and 1.1 ° respectively (Figure 2 (b)). These peaks are characteristic of organised mesoporous materials and are always reported in this region [38] and [39]. The XRD analysis on the coated OMS did not allow a clear identification of the iron and aluminium oxides coated on the surface, probably due to the fact that most of oxides were present inside the pores. Iron oxides were identified as Hematite, while aluminium oxides could not be specifically identified as seen in Figure 3, even if Boehmite is the most probable crystal formed.

3.1.3 TEM imaging analysis

Figure 4 (a) and (d) presents the 3D organised KIT-6 structure and Figure 4 (b) and (e) the 2D organised structure of SBA-15. It is difficult to clearly identify the 3D organised pattern but the 2D pattern is clearly visible in both directions in Figure 4 (b). Coating SBA-15 internal walls with iron oxide appears to lead to particles growth in-homogeneously distributed into the pores, as seen in Figure 4 (e) and (f). The coating process is in reality closer to particle growth inside the pores rather than an homogeneous coating, as some gaps can be spotted inside the pores, Figure 4 (f). There was no oxide growth noticed outside the pores for SBA-Fe 8 %, KIT-Fe 8 %, SBA-Al 8 % or KIT-Al 8 % materials.

3.1.4 FTIR analysis: chemical surface groups identification

SBA-100 °C and KIT-100 °C were analysed by FTIR prior to coating by iron and aluminium oxides. KIT-Fe 8 % and KIT-Al 8 % were studied by FTIR before and after adsorption by 15
As(III) and As(V) of 50 ppm at pH 7. The FTIR analysis of KIT-100 °C and SBA-100 °C is presented in Figure 5 (a) while KIT-100 °C, KIT-Fe 8 % and KIT-Al 8 % are shown in Figure 5 (b). The peak at 1635 cm\(^{-1}\) indicates the presence of free water molecules and bonded water molecules onto the two silica samples. The broad peak at 3430 cm\(^{-1}\) is attributed to the O-H stretching bonds. In the present case, this peak comes from the water molecules bonded onto the silica since all hydroxyl (OH\(^{-}\)) groups from silanol (Si-O-H) groups have been removed during the calcination process at 800 °C [40] and [41]. The double peak at 1160 and 1070 cm\(^{-1}\) is characteristic of Si-O bonds as well as the peaks noticeable at 965, 800 cm\(^{-1}\) and the one starting at 500 cm\(^{-1}\) [42]. There is no major difference between the two sample spectra as they are both amorphous silica. The KIT-Fe 8 % spectrum shows a wider and higher peak at 3430 cm\(^{-1}\) revealing a higher content of O-H bonds. These bonds come from a higher presence of bonded water, as seen with the higher peak at 1635 cm\(^{-1}\), and the presence of Fe-OH groups. The clear higher peak at 1170 cm\(^{-1}\) compared to the peak at 1070 cm\(^{-1}\) is characteristic of Fe-O bonds as presented by Martinez et al. [42]. The KIT-Al 8 % spectrum presents an even wider and higher peak in the region of 3500 to 3000 cm\(^{-1}\) revealing a very high increase in O-H bonds. These bonds can be attributed to Al-OH surface groups. The presence of a sharp peak at 1386 cm\(^{-1}\) proves that -NO\(_3\) groups from the aluminium nitrate salt used during the coating process were not completely removed during the calcination step at 250 °C. This peak is related to -N bonding and can relate to any nitrogen containing species (N\(_2\), NO, N\(_2\)O). The relative increase of the peak at 1200 cm\(^{-1}\) reveals the presence of Al-O bonds. The spectrum thus obtained is very similar to the one produced by Shin et al. for the adsorption of phosphate from water [43].

3.1.5 SEM-EDX

An EDX analysis was carried out only on KIT-Fe 8 % sample in order to verify the iron coating process. The identification spectrum and the quantitative analysis of a small portion
of the sample are reported in supporting information, Figure S1 and Table S1. The total iron content of the sample is quite high (~ 11 %) on the sample area analysed. The real sample iron content is even higher when the carbon counted is removed as that is due to the carbon coating (~ 14.5 %) in sample preparation. This value can be compared with the 8 % Fe$_2$O$_3$ content coated into the pores. This value means that 8 % of the pore volume is occupied by Fe$_2$O$_3$. Using the KIT-6 BET porosity of 1.356 cc.g$^{-1}$ and assuming Fe$_2$O$_3$ density being 5.24 g.cc$^{-1}$, the 8 % Fe$_2$O$_3$ pores loading results in nearly 20 % Fe w/w in KIT-Fe 8 % as shown in equation (16).

$$Fe \% \ w/w = BET_{porosity} \times \% \ Fe_2O_3 \times d_{Fe_2O_3} \times \frac{M_{Fe}}{M_{Fe_2O_3}} \approx 20 \%$$ (16)

The SEM-EDX analysis gives then a good approximation of the metals content (14.5 % instead of 20 %) in the coated OMS samples. The average iron content measured is slightly lower than the theoretically coated iron weight attributable to the iron oxide being coated inside the pores.

3.2 Optimised sorbents analysis

3.2.1 Effect of the percentage of metal oxides coating onto As(III) and As(V) adsorption

It was shown that, after 8 % loading of iron or aluminium oxides into KIT materials, very little improvement of As(III) or As(V) was observed as shown in Figure 6. The percentage removal evolution generally followed the surface area available (Table 1). The KIT-Al material showed a very high As(V) removal capacity, while also exhibiting a very low removal capacity for As(III). In comparison, KIT-Fe materials possess a higher As(III) adsorption capability than As(V) and these two removal capacities are very close. Hence, 8 % metal oxide coatings were used for subsequent optimisation experiments. This percentage coating is a good optimum balance between surface area, pore availability and arsenic removal efficiency.
3.2.2 Fe and Al oxide relative loading effects onto As(III) and As(V) removal in single and binary solution systems

The removal of As(III) and As(V) by 8% oxides coated OMS materials having different Al to Fe oxides ratio is presented in Figure 7. Adsorption capacity achieved for As(III) and As(V) in single and binary systems are presented as a function of the Fe oxides percentage coating onto the 8% loaded OMS, the Al oxides percentage being the remaining percentage. It was found that the arsenate adsorption capacity decreases when the Fe oxides coating content decreases, or when the Al oxides coating percentage increases. This was expected as the Al oxides displayed the higher As(V) adsorption capacity. The As(III) adsorption capacity was nearly constant, independently of the Al to Fe oxides coating configuration tested. The results thus show that the best coating configuration, in terms of overall adsorption capacity, i.e. As(III) + As(V), is 10% Fe oxide and 90% Al oxide coating. Similar results have been reported for removal of As(III) and As(V) using iron oxide coating activated alumina [44]. The simultaneous removal of As(III) and As(V) did not show major differences, as seen in Figure 7 (b). The arsenate and total arsenic removal capacity of the coated OMS generally follow the same trend in single or binary configuration, and the As(III) and As(V) adsorption capacity are very similar in both cases. However, the materials with Fe oxides coating percentages between 27 to 75 display a total As removal capacity higher than the As(V) adsorption capacity in single configuration. This result shows that parts of the adsorption sites for As(III) and As(V) are different, indicating that the total removal capacity of the materials increases in binary configuration. Also, in the single As species configuration, As(III) adsorption capacity is around 6.67 mg.g\(^{-1}\) and for As(V) it reaches 10.60 mg.g\(^{-1}\). In the binary system, the As(V) adsorption capacity remains similar at around 10.20 mg.g\(^{-1}\) and the As(III) adsorption capacity drops down to 1.37 mg.g\(^{-1}\). In this case, the maximum number of As adsorption sites increases slightly, from 10.60 to 11.57 mg.g\(^{-1}\) with 18
As(III) adsorption being predominantly favoured. In the case of low Fe oxide coatings percentage i.e. < 50 %, the As(V) adsorption is generally favoured over As(III) adsorption in terms of total arsenic removal. It was easier to tailor the coated OMS materials toward As(V) than As(III) due to the superior As(V) adsorption capacity of Al oxide coatings. The results demonstrated the ability to tailor one material capacity toward the characteristics of a specific groundwater. Nevertheless, the possibility of optimising the coated OMS materials is limited by the maximum adsorption capacity of the pure single metal oxides and cannot cover the full range of groundwater As(III) to As(V) ratio range, i.e. As(III) : As(V) from 0 : 1 to 1 : 0. The range covered by the optimisation technique presented in this paper for As(III) : As(V) is 0.16 : 0.84 to 0.97 : 0.13 and is presented in detail in Table 3.

3.3 pH effect analysis

The pH study of As(III) and As(V) adsorption onto KIT-Al 8 % and KIT-Fe 8 % is presented in Figure 8. It was found that only KIT-Al-8 % was able to remove completely As(V) at around pH 5 and 55 ppm. The profiles of As(V) adsorption capacity in function of pH of KIT-Al 8 % and KIT-Fe 8 % have similar trends but seem to be shifted by 1 pH value. The PZC of KIT-Al 8 % and KIT-Fe 8 % were evaluated at pH 5.4 and 4.5 respectively. The higher PZC of aluminium coated KIT material is partly responsible for the better performance of KIT-Al 8 % in removing As(V) at neutral pH when compared to KIT-Fe 8 % material. As(III) removal is not influenced by pH change until pH 9, when arsenite ions start to be negatively charged (pKₐ ~ 9.2) and both materials become negatively charged as well (pH > PZC). It is noticed that the As(III) removal capacity of KIT-Fe 8 % is better than its As(V) removal capacity only at high pH. At pH between 7 and 9, KIT-Al 8 % has better As(V) than As(III) removal capacity while, at the same pH range, KIT-Fe 8 % As removal
capacity are inversed. This pH study thus reveals that it is only possible to tailor the As(III) to As(V) removal ratio of the coated OMS for As removal at pH between 7 and 9.

3.4 Concentration effect analysis

Figure 9 illustrates the isotherms of As(III) and As(V) adsorption onto KIT-Fe 8 % and KIT-Al 8 %. The As(III) is removed more efficiently by KIT-Fe 8 % than by KIT-Al 8 % while the opposite trend is displayed for As(V) removal. As(III) isotherm is closer to a Freundlich type isotherm than a Langmuir type for As(III) removal while it is the opposite case for As(V) isotherms as per the respective $r^2$ values from Table 4. The isotherm shape of As(III) removal by KIT-Al 8 % shows the very unfavourable nature of the adsorption process involved. The values reported in Table 4 for $q_m$ for As(III) removal are very high but the related $b_L$ value is very low. This combination gives a very slow increase of the $q_e$ value at low concentration but implies a continuous increase until reaching a $q_m$ value of nearly 81 mg.g$^{-1}$. Arsenic removal efficiency of coated OMS materials at low concentration is very low, especially for arsenite adsorption as presented by $q_e$ values at 0.1 ppm ($q_e$-0.1). Moreover, only As(V) adsorption onto KIT-Al 8 % is a relatively favourable process as the $R_L$-0.1 value is lower than 0.8. It is worth mentioning that the As(III) adsorption capacity of KIT-Fe 8 % is higher than KIT-Al 8 % especially at high concentration and this difference decreases at low concentration as seen in Table 5. The tests performed in order to modify the coated OMS materials for specific adsorption are then very dependent on the concentration range used in the test. The optimisation study was carried out at a very high initial concentration (~50 ppm).

Table 6 gives a comparison of the observed materials removal capacity with data from the literature for equivalent materials having high surface area and high porosity. The Fe and Al
coated OMS materials developed in this work display high As(III) and As(V) removal capacities which are generally higher than equivalent coated mesoporous silica materials.

3.5 *Kinetic removal of arsenic in 2D, 3D and unorganised sorbents*

Figure 10 presents the kinetic removal of arsenite by unorganised, 2D and 3D organised materials coated by iron (a) or aluminium oxides (b) and the kinetic removal of arsenate by the same materials in Figure 10 (c) and (d) respectively. From Figure 10 (a), the As(III) removal by Al oxides is very limited compared to the removal by Fe oxides (Figure 10 (b)). The As(III) removal by KIT-Al 8 % is higher than that of pure Al oxide at equilibrium which is in opposition to the total weigh of aluminium oxide present in each sample. When Fe oxides are used, the unorganised Fe materials display a higher capacity than the organised 3D material which itself present a higher capacity than the 2D organised material. The As(III) removal capacity at equilibrium of the Fe oxide material can actually be related to the total iron oxide content by weight of the materials: Fe oxide (100 %), KIT-Fe 8 % (20 %), SBA-Fe 8 % (15.4 %).

The As(V) removal presents very similar trends as seen in Figure 10 (c) and (d). The main difference is the much higher removal capacity of the Al oxides materials when compared to the Fe oxides materials. It was also noticed that the KIT-Al 8 % material has a higher removal capacity than the unorganised Al-oxides. The Fe oxide materials show a very high removal capacity for the unorganised materials when compared to the 2D and 3D organised materials, a difference that can be again linked to the total weight of iron oxides in the materials.

In terms of removal kinetic, minor differences can be noticed, but the differences of $q_e$ reached make the comparison difficult. Relative kinetic curves are thus presented in Figure 11. In Figure 11 (a) a small difference in removing As(III) by Al oxides materials being
organised or not was found, while in Figure 11 (b) differences are visible. The kinetic performances of the iron oxides based materials toward As(III) follow the trend: unorganised > 2D > 3D. In the case of As(V) removal, little effect of the organisation of Fe oxides materials is noticed (Figure 11 (c)) while diffusion of As(V) in Al materials is affected by the organisation of the materials (Figure 11 (d)). The materials can be classified from the quickest to the slowest in removing As(III): SBA-Al 8 % > KIT-Al 8 % > Al-oxides. The As(III) diffusion is more affected by the pore size of Fe based materials while As(V) is more affected by the pore size of Al based materials. Generally, the organisation of the materials in term of kinetic performances has little impact on the removal of As(III) and As(V). The experiments were carried out with material of small particle size (Ø < 75 µm); in the case of bigger particle, an organised structure might have bigger impact than unorganised materials, especially because of a better access to the adsorption sites. Table 7 gives the pseudo first and pseudo second order model parameters obtained. In general the $k_1$ and $k_2$ values obtained for As(V) are higher than that obtained for As(III) removal for both Fe and Al based materials. This difference was already noticed in the various previous figures. The data obtained for the pseudo first and pseudo second order models, $k_1$ and $k_2$ values, are relatively higher than equivalent data found in the literature. For example Liu et al. found $k_1$ and $k_2$ values of $5.50 \times 10^{-4}$ min$^{-1}$ and $8.80 \times 10^{-3}$ g.mg$^{-1}$.min$^{-1}$ for As(III) and $5.33 \times 10^{-4}$ min$^{-1}$ and $6.83 \times 10^{-4}$ g.mg$^{-1}$.min$^{-1}$ for As(V) removal by iron coated bamboo charcoal at starting arsenic concentration of 32 ppm [51].

3.6 Diffusion and adsorption investigation

Generally, the intraparticle diffusion rate is constant over a period of time at initial condition only. In fact several regions can usually be noticed in the diffusion of pollutants in porous materials [34]. The presence of different linear regions in the plot of $q_t$ versus $t$ is usually
assumed to show the predominance of the resistance to diffusion of one process over another [34]. The diffusion of As(III) and As(V) in unorganised and organised materials was studied using both the intraparticle diffusion model and the Boyd diffusion model.

### 3.6.1 Intraparticle diffusion model

Figure 12 [(a) and (b)] presents the intraparticle diffusion modelling applied to As(III) and As(V) adsorption onto unorganised Al oxides and Al oxides coated OMS while Figure 12 [(c) and (d)] presents the same data obtained for Fe oxides and Fe oxides coated OMS. The same trends noticed in the kinetic experiments can be noticed for the diffusion modelling. Moreover it can be observed that As(III) and As(V) adsorption in Al oxides is mainly following a three-step diffusion process while the diffusion of these species in Fe oxides based materials seems to follow only a two-step diffusion process.

The intraparticle model does not follow a straight line which implies that intraparticle resistance is not the only one present or is evolving under the experimental conditions. The film diffusion cannot be limiting the arsenic adsorption as the particles are very small and the stirring speed sufficiently high. The presence of surface diffusion resistance is then an explanation for the profiles noticed. The decrease of the percentage of sorption sites available can also reduce significantly the diffusion of pollutant in the materials. In the present case it seems that, when the adsorbent coverage reaches a certain percentage, the diffusion nature changes thus decreasing significantly. As the adsorption is taking place, pore diffusion is initially the main resistance to As adsorption, but it is then replaced by surface resistance.

There is no real correlation between the organisation dimension of the materials and the number of diffusion steps and/or the steepness of these steps. The intraparticle diffusion coefficients obtained are presented in Table 8.

In general, higher values were obtained for As(V) diffusion coefficients in the OMS materials studied. However, it should be noted that As(III) diffuses more quickly in Fe unorganised
Intraparticle diffusion seems to be the limiting step in the diffusion of As(III) and As(V) in the coated OMS materials.

3.6.2 Boyd model

The Boyd model plotted for the different materials studied can be found in the supporting information Figure S2. The Boyd model is not a straight line passing thought the origin, though, in most cases, the Boyd model can be approximated to a straight line. The intraparticle diffusion remains the limiting diffusion step. Intraparticle diffusion can be divided into pore diffusion, surface diffusion and surface adsorption. In the materials studied, the second segment of the Boyd model is characteristic of surface diffusion being the limiting step in the adsorption process. No clear difference can be noticed for the Boyd models applied to unorganised, 2D or 3D organised materials in terms of diffusion profile for the intraparticle model. In the case of As(III) adsorption onto Al based sorbents, the Boyd model is a straight line for all materials and the diffusion pattern is similar for all materials, (Figure S2 (a)). For As(V) adsorption with these same materials, 2D organised materials perform better, followed by 3D organised materials and unorganised materials, (Figure S2 (b)). The Fe based materials behaviour toward As(III) and As(V) adsorption is the opposite of that of Al based materials. For As(III) adsorption a clear difference can be noticed between unorganised and organised materials, while this difference is less clear for As(V) adsorption. The diffusion of As(III) and As(V) in Fe based materials follows 2 steps. This trend underlines the presence of surface diffusion being more of a resistance toward As adsorption in Fe based materials than in Al based materials. In general 2 steps are identified in the case of 2D or 3D organised materials. The presence of different steps in the diffusion of As inside the pores in the case of organised materials could reveal the presence of a resistance to diffusion due to the presence of channels and the effect of the walls onto the diffusion of the
species in these materials. Surface diffusion is decreased, in organised materials due to the close presence of other adsorption sites and the effect of the wall. The resistance toward As diffusion in unorganised materials comes from the low porosity of these materials and a low accessibility of the sorption sites.

The B values and associated $D_{\text{eff}}$ values obtained for the As(III) and As(V) adsorption onto the coated OMS sorbents are presented in Table S2. The difference in particle diameter explains the difference in the diffusion coefficients. The small particle size of the materials used in this study is mainly responsible for the low diffusion coefficient of As(III) and As(V) inside the unorganised and coated OMS material. The lower the $D_{\text{eff}}$ values the better the diffusion. In general the diffusion coefficients obtained are higher for As(V) adsorption than for As(III) adsorption. This is in accordance with the kinetic parameters obtained, As(III) being removed more quickly by coated OMS than As(V).

3.7 Adsorption processes mechanisms

Figure 5 (c) presents the FTIR spectra of KIT-Al 8 % materials before and after adsorption with As(III) and As(V) while Figure 5 (d) focuses onto the iron coated OMS. It can be noticed that the peak at 1390 cm$^{-1}$ corresponding to the $-\text{NO}_3$ bonds disappears after adsorption experiments. Remaining $-\text{NO}_3$ groups dissolve in water during the adsorption experiments and can also react quickly with both As(III) and As(V) providing the possibility of chemisorption and the formation of precipitate (Al-As precipitate) at the surface or inside the pores.

The increase of the 890 cm$^{-1}$ peak for As(V) adsorption shows the formation of Al-As bonds and the presence of chemisorption or ion-exchange in the removal process of As(V) by KIT-Al 8 %. In the case of As(III) adsorption, an increase in the 716 cm$^{-1}$ peak is observed; this increase is also present for the As(V) treated sample. The peaks noticed for As(V) adsorption
are in concordance with the results from Myneni et al. who noticed two peaks for $\text{H}_2\text{AsO}_4^{4-}$ bonding onto Ettringite at 760 and 908 cm$^{-1}$ and 2 peaks for the bonding of $\text{HAsO}_4^{2-}$ at 690 and 865 cm$^{-1}$ approximately [52]. As these two As(V) species are present at pH 7 the peaks are probably an overlapping of the two species respective peaks. In the case of arsenic adsorption onto KIT-Fe 8 % very little difference between the material before and after adsorption of both As(III) and As(V) can be noticed. This shows the predominance of physisorption processes over chemisorption in the removal of As by this material. The lower level of removal capacity of the material can also account for the difficulty in identifying clear peaks. Table 9 shows the DR isotherms parameters obtained for the adsorption of As(III) and As(V) onto KIT-Al 8 % and KIT-Fe 8 %. The energy of adsorption associated with the different processes shows that physisorption is governing all these adsorption reactions ($E < 8 \text{ kJ/mol}$). Only the adsorption of As(V) by KIT-Al 8 % can be associated with ion-exchange or chemisorption to some extent. This is usually observed in the literature, for example the removal of As(V) by Al coated mesoporous silica made from silica rice was governed by chemisorption according to Wantala et al. [53].

4 Conclusion

This work revealed the possibility of using specific iron and aluminium oxides coated 3D Organised Mesoporous Silica materials simultaneously in order to adsorb specific amounts of As(III) and As(V). This procedure enables the newly developed materials to perfectly match specific groundwater characteristics. The materials produced were fully characterised and tested in batch experiments, showing a very high removal capacity, around 7.38 mg.g$^{-1}$ for As(V) and 0.21 mg.g$^{-1}$ for As(III) at 100 ppb. The pH effect showed very similar behaviour of As(V) adsorption by KIT-Al 8 % or KIT-Fe 8 % with a maximum adsorption capacity noticed at pHs closed the respective materials PZC. Very significant increase of As(V)
adsorption capacity was noticed at low pH: for KIT-Al 8 %, a maximum of 55 mg.g⁻¹ at pH 5 and, for KIT-Fe 8 %, a maximum of 35 mg.g⁻¹ at pH 4. The possibility to tailor the coating of the OMS materials toward As(III) or As(V) using both iron and aluminium oxides will then be very dependent on the groundwater pH. Moreover the kinetic and diffusion studies demonstrated an insufficient effect of the 3D or 2D organisation structure compared to unorganised materials. Unorganised materials reached equilibrium more quickly than organised materials. Granulation of materials will be carried out to ascertain the potential higher diffusion properties of organised materials. Finally, the study showed that the main adsorption process was physisorption, with the presence of minor chemisorption mechanisms due to remaining -NO₃⁻ groups from unoxidized nitrate compounds.

**Acknowledgement**

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5 References

6 FIGURES

Figure 1 Pore size distribution of the KIT serie of materials produced at different temperature (a) and N\textsubscript{2} adsorption isotherm of KIT-100 °C and SBA-100 °C (b)

Figure 2 Full range XRD analysis (5 to 75 °θ) (a) with spectra shifted by 500 cps and low angle XRD analysis (0.1 to 5 °θ) (b) with spectra shifted by 5x10\textsuperscript{5} cps of OMS and selected coated OMS materials
Figure 3 XRD analysis of selected coated OMS and raw oxides. Full scan (5 to 75 2θ) with identified Fe and Al oxides phase XRD main peaks
Figure 4 TEM images of selected OMS and coated OMS, KIT-100 x38000 (a) and x43000 (d), SBA-100 x38000 (b) and x71000 (c) and SBA-Fe 8% x71000 (c) and x97000 (f)
Figure 5 FTIR analysis of KIT-100 °C and SBA-100 °C, spectrum un-shifted (a) and comparison of KIT-100 °C and KIT coated with 8 % Al or 8 % Fe, spectrum shifted by 0.2 A.U. FTIR analysis of KIT coated with 8 % Al (c) or Fe (d) oxides before and after adsorption of As(III) and As(V)
Figure 6 Arsenic adsorption onto KIT-Fe (a) and KIT-Al (b) adsorbents loaded with increasing metal oxide content (adsorbent concentration is 1 g.L\(^{-1}\); initial corresponding arsenic concentration is ~50 ppm and equilibrium pH is ~8).

Figure 7 As(III) and As(V) single compound removal study (a) and simultaneous As(III) and As(V) removal study (b) using KIT coated with 8% mixed Al and Fe oxides (adsorbent concentration is 1 g.L\(^{-1}\); initial corresponding arsenic concentration is ~50 ppm and equilibrium pH is ~8).
Figure 8 pH effect onto As(III) and As(V) adsorption onto KIT-Al 8 % (a) and KIT-Fe 8 % (b). Starting corresponding arsenic concentration is 55 ppm and adsorbent dosage is 1 g.L\(^{-1}\).

Figure 9 Adsorption isotherms of As(III) and As(V) onto KIT-Fe 8 % (a) and KIT-Al 8 % (b). Equilibrium pH is 8.
Figure 10 Kinetic experiments using Al and Al coated OMS materials for removing As(III) (a) and As(V) (c) or using Fe and Fe coated OMS materials in removing As(III) (b) and As(V) (d). Starting concentration is ~ 50 ppm.
Figure 11 Relative kinetic removal of As(III) by Al and Al coated OMS (a) or Fe and Fe coated OMS (b) or As(V) by Al and Al coated OMS (c) or Fe and Fe coated OMS (d)

Al-oxides
KIT-Al 8%
SBA-Al 8 %
Pseudo 2nd order models

Fe-oxides
KIT-Fe 8%
SBA-Fe 8 %
Pseudo 2nd order models

As(III)
As(V)
Figure 12 Intraparticle diffusion plot of As(III) (a) and As(V) (b) onto unorganised Al oxides, Al oxides coated 2D and 3D OMS and As(III) (c) and As(V) (d) onto unorganised Fe oxides, Fe oxides coated 2D and 3D OMS
7 TABLES

Table 1 BET analysis results for the Fe oxides coating and Al oxides coating procedures applied to KIT-100 °C materials. These materials were used in the adsorption experiments related to the optimisation of the materials coating.

<table>
<thead>
<tr>
<th>Silica</th>
<th>Fe loading</th>
<th>Surface area</th>
<th>Porosity</th>
<th>Average pore size</th>
<th>Al loading</th>
<th>Surface area</th>
<th>Porosity</th>
<th>Average pore size</th>
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<td>6.36</td>
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<td>1.356</td>
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<td>8</td>
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<td>0.118</td>
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</table>

Table 2 BET analysis summary of materials used in adsorption experiments.

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<tr>
<th>Name</th>
<th>Silica</th>
<th>Surface area</th>
<th>Porosity</th>
<th>Average pore size</th>
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<tr>
<td>KIT-Fe 8 %</td>
<td>KIT</td>
<td>520.9</td>
<td>0.575</td>
<td>4.42</td>
</tr>
<tr>
<td>KIT-Al 8%</td>
<td>KIT</td>
<td>207.8</td>
<td>0.434</td>
<td>8.36</td>
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<td>SBA-Fe 8 %</td>
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<td>643.2</td>
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<td>SBA-Al 8%</td>
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<tr>
<td>Fe oxide</td>
<td>-</td>
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<td>0.231</td>
<td>4.54</td>
</tr>
<tr>
<td>Al oxide</td>
<td>-</td>
<td>11.3</td>
<td>0.001</td>
<td>0.49</td>
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Table 3 Adsorption capacity of tailored coated OMS sorbents toward As(III) and As(V).

<table>
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<tr>
<th>OMS coating content</th>
<th>Relative removal capacity for As(III)</th>
<th>As(V)</th>
<th>As(III) + As(V)</th>
<th>Absolute As removal</th>
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</thead>
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<tr>
<td>Fe oxides</td>
<td>Al oxides</td>
<td>%</td>
<td>%</td>
<td>%</td>
</tr>
<tr>
<td>0</td>
<td>100</td>
<td>16.19</td>
<td>83.81</td>
<td>100</td>
</tr>
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<td>0</td>
<td>86.78</td>
<td>13.22</td>
<td>100</td>
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Table 4 Langmuir and Freundlich models constants for As(III) and As(V) adsorption isotherms onto KIT-Fe 8 % and KIT-Al 8 %.

39
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<tr>
<th>Adsorbent</th>
<th>As</th>
<th>(q_m) mg.g(^{-1})</th>
<th>(b_1\times10)</th>
<th>(r^2)</th>
<th>(K_F) mg.(^1).g(^{-1}).L(^r)</th>
<th>(n)</th>
<th>(r^2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>KIT-Al 8 %</td>
<td>As(III)</td>
<td>80.77</td>
<td>0.06</td>
<td>0.989</td>
<td>0.54</td>
<td>0.91</td>
<td>0.988</td>
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<tr>
<td>KIT-Al 8 %</td>
<td>As(V)</td>
<td>31.49</td>
<td>30.59</td>
<td>0.954</td>
<td>18.48</td>
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<td>0.797</td>
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<tr>
<td>KIT-Fe 8 %</td>
<td>As(III)</td>
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<td>0.962</td>
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<td>0.983</td>
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<td>KIT-Fe 8 %</td>
<td>As(V)</td>
<td>6.52</td>
<td>2.97</td>
<td>0.989</td>
<td>2.30</td>
<td>0.28</td>
<td>0.942</td>
</tr>
</tbody>
</table>

Table 5 Separation factor and adsorption capacity at 100 ppb using Langmuir model for As(III) and As(V) adsorption isotherms onto KIT-Fe 8 % and KIT-Al 8 %

<table>
<thead>
<tr>
<th>Adsorbent</th>
<th>Pollutant</th>
<th>(R_{-0.1})</th>
<th>(q_{-0.1}) mg.g(^{-1})</th>
</tr>
</thead>
<tbody>
<tr>
<td>KIT-Al 8 %</td>
<td>As(III)</td>
<td>0.999</td>
<td>0.045</td>
</tr>
<tr>
<td>KIT-Al 8 %</td>
<td>As(V)</td>
<td>0.766</td>
<td>7.377</td>
</tr>
<tr>
<td>KIT-Fe 8 %</td>
<td>As(III)</td>
<td>0.984</td>
<td>0.206</td>
</tr>
<tr>
<td>KIT-Fe 8 %</td>
<td>As(V)</td>
<td>0.971</td>
<td>0.188</td>
</tr>
</tbody>
</table>

Table 6 A comparison of the produced materials adsorption capacity with data from the literature. MAA (Mesoporous Activated Alumina), MS (Mesoporous Silica), OMA (Organised Mesoporous Alumina), ZVI (Zero Valent Iron), AC (Activated Carbon)

<table>
<thead>
<tr>
<th>Adsorbents</th>
<th>As ppm</th>
<th>qm in mg.g(^{-1})</th>
<th>qe-0.1 in mg.g(^{-1})</th>
<th>Best fit isotherm</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>CuO - MAA</td>
<td>0 - 1</td>
<td>2.16</td>
<td>2.02</td>
<td>0.11</td>
<td>0.14</td>
</tr>
<tr>
<td>Fe(^{3+}) and amino functioned MS</td>
<td>0 - 200</td>
<td>142.60</td>
<td>101.74</td>
<td>0.16</td>
<td>0.94</td>
</tr>
<tr>
<td>Iron oxide coated sponge</td>
<td>0 - 5</td>
<td>3.85</td>
<td>4.50</td>
<td>0.33</td>
<td>0.29</td>
</tr>
<tr>
<td>Fe 10 % - MCM-41</td>
<td>0 - 10</td>
<td>-</td>
<td>26.25</td>
<td>-</td>
<td>0.72</td>
</tr>
<tr>
<td>Highly OMA</td>
<td>0 - 0.4</td>
<td>-</td>
<td>-</td>
<td>5.00</td>
<td>19.80</td>
</tr>
<tr>
<td>ZVI on AC</td>
<td>0 - 2</td>
<td>18.19</td>
<td>12.02</td>
<td>8.59</td>
<td>10.54</td>
</tr>
<tr>
<td>CeO(_2)-ZrO(_2) nanospheres</td>
<td>0.5 - 60</td>
<td>74.13</td>
<td>133.50</td>
<td>16.49</td>
<td>57.70</td>
</tr>
<tr>
<td>KIT-Al 8 %</td>
<td>0 - 50</td>
<td>80.80</td>
<td>31.50</td>
<td>0.05</td>
<td>7.38</td>
</tr>
<tr>
<td>KIT-Fe 8 %</td>
<td>0 - 50</td>
<td>13.30</td>
<td>30.60</td>
<td>0.21</td>
<td>0.19</td>
</tr>
</tbody>
</table>
Table 7 Kinetic models parameters for As(III) and As(V) removal onto Al/Fe oxides and Al/Fe coated OMS materials obtained at C₀ ~ 50 ppm

<table>
<thead>
<tr>
<th>Arsenic</th>
<th>Adsorbents</th>
<th>Pseudo-1ˢᵗ order model</th>
<th>Pseudo-2ⁿᵈ order model</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>qₑ</td>
<td>kₑ×10⁷</td>
</tr>
<tr>
<td>As(III)</td>
<td>Al</td>
<td>6.75</td>
<td>1.17</td>
</tr>
<tr>
<td></td>
<td>KIT-Al 8 %</td>
<td>9.65</td>
<td>1.12</td>
</tr>
<tr>
<td></td>
<td>SBA-Al 8 %</td>
<td>1.82</td>
<td>1.48</td>
</tr>
<tr>
<td></td>
<td>Fe</td>
<td>31.07</td>
<td><strong>4.03</strong></td>
</tr>
<tr>
<td></td>
<td>KIT-Fe 8 %</td>
<td>9.33</td>
<td>1.97</td>
</tr>
<tr>
<td></td>
<td>SBA-Fe 8 %</td>
<td>4.72</td>
<td>3.87</td>
</tr>
<tr>
<td>As(V)</td>
<td>Al</td>
<td>14.70</td>
<td>1.96</td>
</tr>
<tr>
<td></td>
<td>KIT-Al 8 %</td>
<td>28.12</td>
<td>7.67</td>
</tr>
<tr>
<td></td>
<td>SBA-Al 8 %</td>
<td>12.68</td>
<td>23.50</td>
</tr>
<tr>
<td></td>
<td>Fe</td>
<td>28.70</td>
<td>2.28</td>
</tr>
<tr>
<td></td>
<td>KIT-Fe 8 %</td>
<td>7.26</td>
<td>13.30</td>
</tr>
<tr>
<td></td>
<td>SBA-Fe 8 %</td>
<td>8.53</td>
<td>34.40</td>
</tr>
</tbody>
</table>

Table 8 Intraparticle diffusion coefficients for As(III) and As(V) adsorption onto Al and Fe unorganised oxides and Al and Fe coated OMS materials

<table>
<thead>
<tr>
<th>Materials</th>
<th>Pollutant</th>
<th>Intraparticle diffusion coefficients in mg·g⁻¹·min⁻⁰·⁵</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>As(III)</td>
<td>kₐ₁×10⁻⁰·⁵</td>
</tr>
<tr>
<td>Al</td>
<td>4.45</td>
<td>2.65</td>
</tr>
<tr>
<td>KIT-Al 8 %</td>
<td>6.98</td>
<td>3.01</td>
</tr>
<tr>
<td>SBA-Al 8 %</td>
<td>1.74</td>
<td>0.56</td>
</tr>
<tr>
<td>Fe</td>
<td>41.61</td>
<td>6.38</td>
</tr>
<tr>
<td>KIT-Fe 8 %</td>
<td>9.41</td>
<td>2.67</td>
</tr>
<tr>
<td>SBA-Fe 8 %</td>
<td>4.94</td>
<td>0.79</td>
</tr>
<tr>
<td>Al</td>
<td>16.64</td>
<td>5.65</td>
</tr>
<tr>
<td>KIT-Al 8 %</td>
<td>70.44</td>
<td>12.23</td>
</tr>
<tr>
<td>SBA-Al 8 %</td>
<td>41.63</td>
<td>2.37</td>
</tr>
<tr>
<td>Fe</td>
<td>26.94</td>
<td>0.94</td>
</tr>
<tr>
<td>KIT-Fe 8 %</td>
<td>20.57</td>
<td>2.77</td>
</tr>
<tr>
<td>SBA-Fe 8 %</td>
<td>32.52</td>
<td>1.54</td>
</tr>
</tbody>
</table>

Table 9 Dubinin–Radushkevich isotherm parameters and adsorption energy associated to As removal by adsorption onto KIT-Fe 8 % and KIT-Al 8 %

<table>
<thead>
<tr>
<th>Adsorbent</th>
<th>Pollutant</th>
<th>pHₑ</th>
<th>qₑ</th>
<th>kₐₒ×10⁸</th>
<th>r²</th>
<th>E</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>mol⁻²·J⁻¹</td>
<td></td>
<td>J·mol⁻¹</td>
</tr>
<tr>
<td>KIT-Fe 8 %</td>
<td>As(III)</td>
<td>8</td>
<td>10.33</td>
<td>263.00</td>
<td>0.817</td>
<td>435.81</td>
</tr>
<tr>
<td>KIT-Fe 8 %</td>
<td>As(V)</td>
<td>8</td>
<td>5.60</td>
<td>104.00</td>
<td>0.903</td>
<td>692.72</td>
</tr>
<tr>
<td>KIT-Al 8 %</td>
<td>As(III)</td>
<td>8</td>
<td>12.73</td>
<td>2380.00</td>
<td>0.920</td>
<td>144.95</td>
</tr>
<tr>
<td>KIT-Al 8 %</td>
<td>As(V)</td>
<td>8</td>
<td>29.52</td>
<td>4.75</td>
<td>0.969</td>
<td>3243.64</td>
</tr>
</tbody>
</table>
Highlights:

- 3D and 2D Organised Mesoporous Silica were efficiently coated with Al to Fe oxides
- 8 % metal oxide loading was the most appropriate content for efficient As removal
- Tailoring of adsorbent for As(III) or As(V) was achieved by varying Al to Fe ratio
- Efficiency of tailored absorbent is highly pH and concentration dependent
- No superior diffusion into organised materials over unorganised materials was shown