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Use of Synthetic and Natural Zeolites Tailored for As(V) Sorption

Andrey E. Krauklis

Abstract
Arsenic in drinking water poses serious potential health risks in more than 30 countries with total affected population of around 100 million people. Natural and synthetic zeolites can be tailored in order to obtain improved sorption of As(V) making them a relatively cheap and efficient material for water remediation. The chapter is concentrated on the zeolitic materials for water remediation, and reports new findings regarding modification methods and comparison of such materials for the use in As(V) sorption applications. Methods of modification of zeolites are developed and explained. On the experimental and novel scale, using developed methods, 11 novel materials are synthesized and studied. Initial and modified materials are characterized by optical microscopy, SEM and EDX, as well as by metal content in those which are determined using dissolution in acids and FAAS.

Keywords: zeolites, water remediation, As(V), adsorption, sorbents, arsenic, environmental remediation, equilibrium, kinetics, heavy metals, metalloids

1. Introduction
Groundwater contamination with arsenic compounds is an everyday problem for millions of people as water resources are crucial for use as drinking water, as well as in food production and agriculture [1–3]. Accumulation of arsenic in the body poses significant health risks and may lead to arsenicosis [3]. USA Environmental Protection Agency (EPA) in 2006 has diminished acceptable maximum contaminant level (MCL) for arsenic in drinking water from 50 down to 10 μg/L, and stricter norms have been set in headquarters of Peoples Republic of China, United Nations Health Organization, and European Union [4–6]. Arsenic contamination is both natural and due to anthropogenic sources. Weathering of rocks and minerals is a typical example of a natural process, while pesticides, metal waste, fertilizers and fossil fuel combustion are due to industrial and agricultural activities [7–9].
Sorption is considered as one of the most reasonable water remediation techniques because of its lower costs [10–13]. Zeolites are crystalline, microporous, hydrated aluminosilicate minerals containing alkali or alkaline cations. These materials are often praised for their efficient sorptive properties. Zeolite surface modification is a possible route for substantially improving oxyanion sorption [12]. Zeolites generally possess high values of specific surface area, which can be larger than 700 m$^2$/g [14]. This parameter is an extremely important one for adsorptive properties. Most of zeolitic water remediation technologies are based on cation exchange principles; however, there are cases when zeolites are used and useful for anion removal, such as cases of arsenate and arsenite sorption [15, 16]. Advantages of zeolitic sorbents are as follows: Generally, they are environmentally friendly, relatively cheap and can be treated for secondary use. Considering affinity of metalloids to interact with Fe and Mn-containing compounds [17], studies of As(V) sorption can be promoted in direction of using iron- and manganese-modified zeolites.

1.1. Clinoptilolite
Clinoptilolite is a high silicate content heulandite group mineral (Na,K,Ca$_{0.5}$Sr$_{0.5}$Ba$_{0.5}$Mg$_{0.5}$)$_6$[Al$_6$Si$_{30}$O$_{72}$]$_2$O$_2$ – HEU, often main component of natural zeolites. Its effective pore diameter is in range from about 4.5 up to 6.0 Å. Such clinoptilolite’s properties as chemical stability in basic environments, thermostability and high sorption rate make it a useful material in chemical research and industry [15].

1.2. Zeolite A
Zeolite A is one of the most widely used synthetic zeolites. It has a Linde Type A (LTA) structural type as defined by International Zeolite Association (IZA). Most commonly used in its sodium form (4A), its chemical formula in dehydrated form is [Na$_{12}$Al$_{12}$Si$_{12}$O$_{48}$]$_8$ [18]. Potassium (3A) and calcium (5A) forms are also widely used. These zeolite names roughly represent pore opening diameters in ångströms (Å). It is possible to convert one form into another one via ion exchange [18].

1.3. Zeolite X
Zeolites X contain 12 rings or pore openings with a diameter of 7.4 Å (8.1 Å when completely empty). Diameter of the central cavity is 13.7 Å [19].

1.4. Iron oxides
Iron oxides and hydroxides are known for their use in wastewater treatment [20]. Natural iron oxides (without using zeolites as the host material) have affinity for arsenic compounds but show low sorption efficiency of treating aqueous solutions contaminated with As compounds (0.02–0.4 mg/g) due to low specific surface area [5, 21–23]. Thus it is important to introduce a host material with high specific surface area, i.e. zeolite, which can be modified with iron compounds, such as FeOOH. Fe(III) ionic radius is 0.67 Å [24], small enough in order to use it for modification of zeolites.
1.5. Manganese oxides

As(V) is sorbed more easily onto solid surfaces than As(III), and thus oxidation of As(III) followed by adsorption is a potentially effective route for the removal of arsenic compounds [25, 26]. It has been reported that clinoptilolite modification with MnO₂ significantly enhances sorption capacity of As(V) and is believed to be independent of pH [27]. MnO₂ can oxidize As(III) to As(V) based on sources [24, 28], which is backed up with the following red-ox reactions, shown in Eqs. 1 and 2:

\[
\begin{align*}
\text{HAsO}_2^2+ 2\text{H}_2\text{O} & \rightleftharpoons \text{H}_3\text{AsO}_4^+ + 2\text{H}^+ + 2e^- \quad (\phi = -0.56 \text{ e}) \\
\text{MnO}_2 + 4\text{H}^+ + 2e^- & \rightleftharpoons \text{Mn}^{2+} + 2\text{H}_2\text{O} \quad (\phi = +1.23 \text{ e})
\end{align*}
\]  

The summary red-ox reaction is represented by Eq. 3:

\[
\text{HAsO}_2 + \text{MnO}_2 + 2\text{H}^+ \rightleftharpoons \text{H}_3\text{AsO}_4 + \text{Mn}^{2+} \quad (\phi = +0.67 \text{ e})
\]

The value of standard potential (\(\phi = +0.67 \text{ e}\)) is positive, which indicates that equilibrium is shifted towards right of Eq. 3, in direction of As(V) as a product [24].

Mn(II), Mn(IV) and Mn(VII) ionic radii are 0.80–0.91 Å, 0.50–0.52 Å and 0.46 Å, respectively [24]. Mn-O bond in a permanganate ion is 1.629 Å, and Mn is in the tetrahedron’s centre [29]. This is small enough in order to use it for modification of zeolites.

1.6. Sorption models

Langmuir and Freundlich isotherms are the most popular models in describing sorption in water environments; however, other models are also known and widely used such as Dubinin-Radushkevich, Temkin and Redlich-Peterson models [18, 30].

1.7. Langmuir model

The model is based on assumption that each sorption active centre is equivalent, and it is energetically irrelevant whether adjacent sorption centres are empty or already occupied [20].

Langmuir sorption model [31]:

\[
q_e = \frac{q_mK_L C_e}{1 + K_L C_e},
\]

where \(q_e\) [mg/g] is equilibrium concentration in adsorbent which corresponds to initial concentration in solution \(C_0\) [mg/L]; \(q_m\) [mg/g] is maximum monolayer coverage capacity; \(C_e\) [mg/L] is equilibrium concentration in solution; \(K_L\) [L/mg] is Langmuir constant.

Linearized form of Eq. 4 is represented with Eq. 5:

\[
\frac{C_e}{q_e} = \frac{1}{q_m} C_e + \frac{1}{q_mK_L}
\]
In order to compare experimental data’s fit with the model, linearized form is used, plotting data in coordinates $C_e^q - C_e$ and obtaining determination coefficient $R^2$ [30].

### 1.8. Freundlich model

Freundlich model is based on assumption that sorption occurs on nonequivalent sorption centres, which is due to repulsion between the sorbed particles. It is assumed there is an infinite number of sorption centres [20]. Freundlich sorption model [32]:

$$q_e = K_F C_e^\frac{1}{n},$$  \hspace{1cm} (6)

where $K_F$ [mg$^{1-n}$L$/g]$ is Freundlich coefficient; $n$ [-] is Freundlich constant.

Linearized form of Eq. 6 is represented with Eq. 7 [30]:

$$\ln q_e = \ln K_F + \frac{1}{n} \ln C_e$$  \hspace{1cm} (7)

In order to compare experimental data’s fit with the model, linearized form is used, plotting data in coordinates $\ln q_e - \ln C_e$ and obtaining determination coefficient $R^2$ [30].

### 1.9. Dubinin-Radushkevich model

Usually the model is used to differentiate between physisorption and chemisorption. Initially it was used to describe physisorption. Dubinin-Radushkevich sorption model [33]:

$$q_e = q_s \frac{e^{-\varepsilon_{DR}}} {C_e}$$  \hspace{1cm} (8)

$$\varepsilon_{DR} = RT \ln \left(1 + \frac{1}{C_e} \right)$$  \hspace{1cm} (9)

$$E = \frac{1}{\sqrt{2K_{DR}}},$$  \hspace{1cm} (10)

where $\varepsilon_{DR}$ [kJ/mol] is Dubinin–Radushkevich isotherm variable; $E$ [kJ/mol] is mean free energy of adsorption; $q_s$ [mg/g] is theoretical Dubinin–Radushkevich saturation sorption capacity; $K_{DR}$ [mol$^2$/kJ] is Dubinin–Radushkevich constant.

Linearized form of Eq. 8 is represented with Eq. 11 [30]:

$$\ln q_e = \ln q_s - K_{DR} \varepsilon_{DR}^2$$  \hspace{1cm} (11)

In order to compare experimental data’s fit with the model, linearized form is used, plotting data in coordinates $\ln q_e - \varepsilon_{DR}^2$ and obtaining determination coefficient $R^2$ [30].

### 1.10. Temkin model

The model is based on assumption that heat of adsorption (function of temperature) decreases linearly with increasing amount of sorbed particles. Temkin sorption model [33]:

$$q_e = q_s \frac{1}{1 + K_{DR} C_e}$$
\[ q_e = \frac{RT}{b_T} \ln K_T C_e, \quad (12) \]

where \( R \) [8.314 J/(mol·K)] is universal gas constant; \( T \) [K] is temperature; \( K_T \) [mg/L] and \( b_T \) [L/mg] are parameters describing adsorbate-adsorbent interactions.

Linearized form of Eq. 12 is represented with Eq. 13 [30]:

\[ q_e = \frac{RT}{b_T} \ln K_T + \frac{RT}{b_T} \ln C_e \quad (13) \]

In order to compare experimental data’s fit with the model, linearized form is used, plotting data in coordinates \( q_e – \ln C_e \) and obtaining determination coefficient \( R^2 \) [30].

1.11. Redlich-Peterson model

The model is a hybrid between Langmuir and Freundlich models. When value of coefficient \( \beta_{RP} \) is equal to 1, the model becomes equivalent with Langmuir model. Often this three-parameter model is able to explain experimental data more precisely. Redlich–Peterson sorption model [34]:

\[ q_e = \frac{K_{RP} C_e}{1 + \alpha_{RP} C_e^{\beta_{RP}}}, \quad (14) \]

where \( \alpha_{RP} \) [L/mg], \( \beta_{RP} [-] \) and \( K_{RP} \) [L/g] are Redlich–Peterson isotherm parameters.

Linearized form of Eq. 14 is represented with Eq. 15 [30, 34]:

\[ \ln \left( \frac{K_{RP} C_e}{q_e} - 1 \right) = \beta_{RP} \ln C_e + \ln \alpha_{RP} \quad (15) \]

\( K_{RP} \) is optimized by finding the closest fit to the model via the highest determination coefficient. In order to compare experimental data’s fit with the model, linearized form is used, plotting data in coordinates \( \ln \left( \frac{K_{RP} C_e}{q_e} - 1 \right) – \ln C_e \) and obtaining determination coefficient \( R^2 \) [30, 34].

1.12. Purpose and aim of this work

The purpose of this work is to present zeolites as potential sorbents for As(V) sorption for water remediation as well as to present novel modification methods and materials. The aim of the study is to provide and interpret results of As(V) sorption onto raw and modified zeolites.

2. Materials

2.1. Slovakian natural clinoptilolite (Slov)

Clinoptilolite natural zeolite from Slovakian deposit Nižný Hrabovec was used. Its specific weight is 2200–2440 kg/m³; porosity 24–32%; clinoptilolitic content fluctuates in range from 86
to 94%; according to manufacturer, it also contains cristobalite, clay mica, plagioclase, rutile, quartz; Si:Al ratio is in range 4.80–5.40; fraction of 1–2.5 mm was used [35].

2.2. Ukrainian natural clinoptilolite (Ukr)
Clinoptilolite natural zeolite from Ukrainian Zakarpattian deposit Sokimicke (ukr. Сокирницьке родовище). Specific weight 2370 kg/m³; porosity 44%; clinoptilolitic content 77%; fraction 1–3 mm [36].

2.3. Russian natural clinoptilolite (Khol)
Clinoptilolite natural zeolite from Russian Zabaykalsky Krai deposit Kholinskoе (rus. Холинское месторождение). Specific weight 1900–2800 kg/m³; bulk density 1.02–1.20 g/cm³. Porosity is in range of 20–23% [37]. Zeolite was crushed and sieved using vibro sieve FRITSCH analysette 3 SPARTAN (Germany; International), collecting fraction from 0.8 up to 1.4 mm.

2.4. Synthetic zeolite A (4A)
SYLOSIV A 4 was used (purchased from Grace Davison, USA), zeolite 4A in fine powder form (particle size 6–9 μm). According to manufacturer, this material is chemically stable in basic, neutral and weak acidic environments; specific surface area 800 m²/g; effective pore volume 0.25–0.30 cm³/g; specific weight 1900–2300 kg/m³ [38].

2.5. Synthetic zeolite X (13X)
Zeolite 13X was used (purchased from Hong Kong Chemical Corp.). Bulk density 0.601 g/cm³; porosity 0.55%; fraction 4–5 mm.

2.6. Reagents
All compounds used were of analytical grade (≥ 98%) and were used without further purification. Sodium hydroxide, potassium chloride and 65% nitric acid were obtained from Sigma-Aldrich (Riedel-de Haën, Germany). Iron(III) chloride hexahydrate, calcium chloride dihydrate and 30% hydrogen peroxide were obtained from Enola (Riga, Latvia). Manganese(II) chloride tetrahydrate was obtained from Firma Chempur (Piekary Śląskie, Poland). All aqueous solutions were prepared using high purity deionized water (10–15 MΩ·cm), produced via water purification system Millipore Elix 3 (Billerica, USA). Arsenate stock solution was prepared using disodium hydrogenarsenate heptahydrate Na₂HAsO₄·7H₂O obtained from Alfa Aesar (Haverhill, USA).

3. Zeolite modification methods
Zeolites Ukr, Slov, Khol, 4A and 13X were modified using 6 different methods. Altogether, using these methods, 11 novel materials were synthesized and are described in this chapter. Additionally, FeOOH-modified zeolites A were obtained and described in the following source [17]. It should be also noted that another aluminosilicate, clay montmorillonite was modified using a similar approach and described in another work [39].
3.1. First method

First method is based on the method described in source [40]. The basis for FeOOH-modified sorbent synthesis is the iron oxohydroxide precipitation on the raw material. Description of modification is the following: 0.25 mol FeCl$_3$$\cdot$6H$_2$O is dissolved in 250 mL DI water, adding 250 mL 3 M NaOH solution and aged for 3 h. Obtained Fe(OH)$_3$ precipitates are decanted. 100 g of raw material are mixed in the Fe(OH)$_3$ dispersion. The mixture is then carefully mixed and filtered under vacuum and washed with 250 mL DI water. The filtered and washed material is then dried in air atmosphere for 1 h at room temperature and then dried in the oven Gallenkamp Plus II (London, UK) for 4 h at 60°C.

Materials obtained using this method: Fe-Ukr(1); Fe-Slov.

3.2. Second method

Second method is based on the first method, modified based on the idea that the material is first soaked with a respective metal salt, followed by the reaction (FeOOH synthesis) inside the zeolite structure. The developed method is the following: 1 M FeCl$_3$$\cdot$6H$_2$O solution is prepared and 100 g of zeolite are placed into it. The mixture is aged for 24 h. The mixture is then filtered, and without washing, 250 mL of 3 M NaOH solution are added to the soaked material and aged for 24 h. The mixture is filtered under vacuum and washed with 250 mL DI water. The filtered and washed material is dried in air atmosphere for 1 h at room temperature and subsequently dried in the oven Gallenkamp Plus II (London, UK) for 4 h at 60°C.

Materials obtained using this method: Fe-Ukr(2); Fe-13X; Fe-KHol.

3.3. Third method

Third method is based on the first method, adapted to be applicable for efficient modification of powders (as in case of Fe-4A and Fe-5A described in another work [17]). The idea is to conduct a reaction in the wet FeOOH mass, simplifying and accelerating modification process. The developed method is the following: 0.25 mol FeCl$_3$$\cdot$6H$_2$O is dissolved in 250 mL DI water, 250 mL of 3 M NaOH is added and the mixture is aged for at least 3 h. Synthesized Fe(OH)$_3$ precipitates are decanted. The mass is filtered under vacuum, mixing in 100 g of raw material into the Fe(OH)$_3$ precipitates, while carefully mixing. Porridge-consistency mixture is washed with 250 mL DI water. The filtered and washed material is then dried in air atmosphere for 2 h at room temperature, and further dried in the oven Gallenkamp Plus II (London, UK) for 4 h at 110°C.

Materials obtained using this method: Fe-Ukr(3); in combination with other methods, FeMn-Slov with fourth method.

3.4. Fourth method

Fourth method is based on methods described in sources [27, 41] which are natural zeolite modification methods with manganese oxides. Concentrations used are the same as described in source [27]. 100 g of zeolite is weighted in the beaker and dried in air atmosphere in the oven for 1 h at 70°C. 2.5 M MnCl$_2$ solution and 10 M NaOH solution are prepared. 100 mL of prepared 2.5 M MnCl$_2$ solution is added, while
mixing, to the zeolite in the beaker. 1 mL of prepared 10 M NaOH solution is added and mixed. Solution is aged for 24 h. Without filtering, the mixture is placed in the oven for 3 h at 150°C. The result is densified zeolite granule/pellet mass, covered with precipitates. This mass is then placed into the crucibles. Crucibles with the obtained mass are placed in the muffle furnace and are held there for 5 h in air atmosphere at 550°C. Crucibles are then taken out and cooled down in air at room temperature. After cooling, modified zeolite is washed with 300 mL of DI water. Material is dried in air for 1 h at room temperature, and further dried in the oven Gallenkamp Plus II (London, UK) for 4 h at 60°C.

Materials obtained using this method: Mn-Slov(1); in combination with other methods, FeMn-Slov with third method, CaMn-Slov with sixth method.

3.5. Fifth method

Fifth method is based on fourth method, but, while fourth method is rooted in Mn(II) oxidation at elevated temperature, this method is based on Mn(VII) reduction reaction using ethanol. The reaction as represented by Eq. 16 is conducted at room temperature [42]:

\[ 2\text{KMnO}_4 + 3\text{C}_2\text{H}_5\text{OH} \rightarrow 2\text{MnO}_2 + 3\text{CH}_3\text{CHO} + 2\text{KOH} + 2\text{H}_2\text{O} \]  (16)

This modification method is developed and chosen in order to perform modification of zeolites with MnO₂ in softer conditions at room temperature. Concentration and volume of potassium permanganate solution is chosen based on KMnO₄ solubility in water at room temperature (if limited by) and in order to introduce the same amount of Mn atoms as in fourth method (0.25 mol Mn). The amount of ethanol is chosen such so that C₂H₅OH and KMnO₄ are in molar ratio 1:1 (stoichiometrically). 100 g of zeolite is weighted in the beaker. Then, 1.5 L 0.17 M KMnO₄ solution is prepared and added into the beaker with the zeolite. The aging proceeds for 96 h in the solution. Then, 24.3 mL 60% C₂H₅OH is added to the mixture in order to reduce KMnO₄. Acetaldehyde is obtained and characteristic smell can be felt. Modified zeolite is filtered under vacuum and washed with 300 mL DI water. Material is then dried in air atmosphere at room temperature for 1 h, and further dried in the oven Gallenkamp Plus II (London, UK) for 4 h at 60°C.

Materials obtained using this method: Mn-Slov(2), Mn-4A.

3.6. Sixth method

The sixth method is a Ca²⁺ ion-exchange reaction used in combination with previously described methods. It is based on a principle that one can tailor the effective micropore diameter in zeolites by exchanging the cations. Divalent Ca²⁺ cations are used to increase the effective pore opening [18]. 20 g of zeolite are placed in 100 mL glass vessels. 4.5 M CaCl₂ solution is prepared and 50 mL is added to the material. The vessels are then placed on the orbital shaker Biosan Multifunctional Orbital Shaker PSU-20i (Riga Latvia), setting frequency to 150 rpm for 96 h. The mass is decanted and then washed with 100 mL DI water. Afterward, the material is dried for 2 h at room temperature in air atmosphere, and then subsequently dried at 110°C in the oven for 4 h.

Materials obtained using this method: in combination with other methods, CaMn-Slov with fourth method.
4. Material characterization

Modified and raw zeolites were characterized using optical microscopy, scanning electron microscopy (SEM), energy-dispersive X-ray spectroscopy (EDX, EDS), by analysis of Fe and Mn content, which was performed by dissolution in acids followed by flame atomic absorption spectrometry (FAAS). Bulk density of all sorbents was determined.

4.1. Optical microscopy

Optical microscopy allows to study the surface of the material and to evaluate homogeneity. Optical microscope Leica was used with a digital camera Leica DFC480, ocular Leica 10X/21B, which enables magnification from x7.1 up to x115. Additional lighting was used: Leica Fluorescent Ringlight and Leica CLS150X (Germany; International) [43]. Software used for obtaining images was Leica Application Suite v4.1.0. A square of 1 cm was cut out of millimetre paper, which was placed adjacent to the sample as a scale. In order to obtain qualitative images, 11 micrographs were used on average using focus stacking (pyramid) approach. Software used for stacking was Helicon Focus (Kharkiv, Ukraine).

4.2. SEM and EDX

Materials were covered with a thin layer of gold in order to prevent charging due to electron beam. Gold sputtering was performed using Quorum Technologies Emitech K550X (Laughton, UK). Methods were performed using Tescan Mira/LMU (Brno, Czech Republic) in backscattered electron regime with working voltage of 15 kV.

4.3. FAAS

Fe and Mn content was obtained using dissolution in acids followed by flame atomic absorption spectrometry (FAAS). 1 g of each material was weighed in a beaker, using analytical scales (± 0.1 mg). 25 mL 65% HNO₃ and 5 mL 35% H₂O₂ were then added in each beaker. Beakers were placed in a thermostat Biosan MyLab Thermo-Block TDB 400 (Riga, Latvia) and heated up to 160°C. When half of the solution had evaporated, 25 ml 65% HNO₃ was added while heating the sample. The solutions were allowed to cool down in air at room temperature, filtered into a graduated vessel and diluted with DI water to a total volume of 60 mL each. Furthermore, a blank sample was prepared for background correction. 10 mL of each solution were moved in test tubes, using pipettes. Prepared samples were analysed using PerkinElmer AAnalyst 200 (Waltham, USA) with flame atomization. Mn and Fe content measurements were performed using background correction in air-acetylene flame. Fe and Mn content in samples was calculated using Eqs. 17 and 18:

\[
w_{Fe} = \frac{C_{Fe} \cdot V}{m} \tag{17}
\]

\[
w_{Mn} = \frac{C_{Mn} \cdot V}{m} , \tag{18}
\]
where $w_{Fe}$ and $w_{Mn}$ are Fe and Mn mass fractions in materials, respectively [mas%]; $C_{Fe}$ and $C_{Mn}$ are Fe and Mn ion concentrations in solution, respectively [mg/L]; $V$ is volume of the solution (0.060 L) [L]; $m$ is mass of sorbent (1000 mg) [mg].

4.4. Determination of bulk density

The method of bulk density determination is based on a standard method, which is described in literature [44–46]. An empty 250 mL graduated cylinder was weighted, using analytical scales Kern ALJ 220–4 (± 0.1 mg). Approximately, 100 g of powder or granules/pellets were placed in the graduated cylinder, while determining the total mass of material and the cylinder. Deducing cylinder’s mass from the total, the mass of the material was obtained. The cylinder was gently hit on the flat surface until the material became compacted and the change in volume was not observable. The volume of the material was determined using the closest mark on the graduated cylinder. The bulk density was determined as the ratio of obtained mass and volume.

5. Sorption experiments

Sorption experiments were conducted using batch system. $\text{Na}_2\text{HAsO}_4\cdot7\text{H}_2\text{O}$ was used for preparing arsenic stock solutions at various concentrations (300, 200, 100, 50, 25, 10 and 5 mg/L). 0.5000 ± 0.0001 g of each sorbent was weighed in every 100 mL glass vessel using analytical scales Kern ALJ 220–4 (Balingen, Germany). 30.00 ± 0.05 mL of an As(V) solution was then added to every vessel with the adsorbent. Vessels were then shaken for 24 h at room temperature ($23\pm1^\circ\text{C}$) at 150 rpm using orbital shaker Biosan Multi-functional Orbital Shaker PSU-20i (Riga, Latvia) to ensure sorption equilibrium was achieved. Suspensions were filtered into 50 mL test tubes, and concentration of As(V) in the filtrate was then analysed using PerkinElmer AAnalyst 200 with flame atomization. Absorption was measured using background correction in $\text{N}_2\text{O}-\text{C}_2\text{H}_2$ flame. A spectral line of 193.7 nm was used. FAAS spectrometer was calibrated with 1000 mg As/L standard solutions obtained from Scharlau (Barcelona, Spain) (As$_2$O$_3$ in 0.5 M HNO$_3$). Each measurement was performed 3 times. In order to ensure arsenic analysis quality control, experiments were performed systematically; accurate As(V) concentration of respective stock solutions was measured 3 times and standard deviation was determined, which then was taken into account when describing sorption capacity of the materials.

6. Experimental results

6.1. Homogeneity and bulk density

Homogeneity of materials was evaluated using optical microscope. All studied materials were homogeneous except Fe-Ukr(1), Fe-Ukr(2), Fe-Ukr(3) and Fe-Slov. Due to inhomogeneities, these materials were not further analysed with SEM and EDX. Obtained information about bulk density and homogeneity is summarized in Table 1.

6.2. Metal ion content

Fe and Mn content by weight in all materials is summarized in Table 2 as an average of three repeated measurements with standard deviation. Material with highest Fe content is FeMn-Slov.
### Table 1. Material bulk density, form and homogeneity.

<table>
<thead>
<tr>
<th>Material</th>
<th>Bulk density ($g/mL$)</th>
<th>Form</th>
<th>Homogeneity</th>
</tr>
</thead>
<tbody>
<tr>
<td>4A</td>
<td>0.46 ± 0.02</td>
<td>Powder</td>
<td>Homogeneous</td>
</tr>
<tr>
<td>Mn-4A</td>
<td>0.77 ± 0.03</td>
<td>Powder</td>
<td>Homogeneous</td>
</tr>
<tr>
<td>13X</td>
<td>0.75 ± 0.03</td>
<td>Granules/Pellets</td>
<td>Homogeneous</td>
</tr>
<tr>
<td>Fe-13X</td>
<td>0.74 ± 0.03</td>
<td>Granules/Pellets</td>
<td>Homogeneous</td>
</tr>
<tr>
<td>Ukr</td>
<td>1.04 ± 0.04</td>
<td>Granules/Pellets</td>
<td>Homogeneous</td>
</tr>
<tr>
<td>Fe-Ukr(1)</td>
<td>1.05 ± 0.04</td>
<td>Granules/Pellets</td>
<td>Non-homogeneous</td>
</tr>
<tr>
<td>Fe-Ukr(2)</td>
<td>1.01 ± 0.04</td>
<td>Granules/Pellets</td>
<td>Non-homogeneous</td>
</tr>
<tr>
<td>Fe-Ukr(3)</td>
<td>1.00 ± 0.04</td>
<td>Granules/Pellets</td>
<td>Non-homogeneous</td>
</tr>
<tr>
<td>Slov</td>
<td>0.91 ± 0.04</td>
<td>Granules/Pellets</td>
<td>Homogeneous</td>
</tr>
<tr>
<td>Fe-Slov</td>
<td>0.92 ± 0.04</td>
<td>Granules/Pellets</td>
<td>Non-homogeneous</td>
</tr>
<tr>
<td>Mn-Slov(1)</td>
<td>1.12 ± 0.04</td>
<td>Granules/Pellets</td>
<td>Homogeneous</td>
</tr>
<tr>
<td>Mn-Slov(2)</td>
<td>0.97 ± 0.04</td>
<td>Granules/Pellets</td>
<td>Homogeneous</td>
</tr>
<tr>
<td>FeMn-Slov</td>
<td>0.91 ± 0.04</td>
<td>Granules/Pellets</td>
<td>Homogeneous</td>
</tr>
<tr>
<td>CaMn-Slov</td>
<td>1.11 ± 0.04</td>
<td>Granules/Pellets</td>
<td>Homogeneous</td>
</tr>
<tr>
<td>Khol</td>
<td>0.90 ± 0.04</td>
<td>Granules/Pellets</td>
<td>Homogeneous</td>
</tr>
<tr>
<td>Fe-Khol</td>
<td>0.96 ± 0.04</td>
<td>Granules/Pellets</td>
<td>Homogeneous</td>
</tr>
</tbody>
</table>

### Table 2. Fe and Mn content of materials obtained using FAAS.

<table>
<thead>
<tr>
<th>Material</th>
<th>Fe content (mas%)</th>
<th>Mn content (mas%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>4A</td>
<td>0.19 ± 0.01</td>
<td>0.00 ± 0.01</td>
</tr>
<tr>
<td>Mn-4A</td>
<td>0.03 ± 0.00</td>
<td>0.57 ± 0.02</td>
</tr>
<tr>
<td>13X</td>
<td>0.34 ± 0.01</td>
<td>0.01 ± 0.01</td>
</tr>
<tr>
<td>Fe-13X</td>
<td>2.92 ± 0.11</td>
<td>0.02 ± 0.00</td>
</tr>
<tr>
<td>Ukr</td>
<td>0.53 ± 0.02</td>
<td>0.01 ± 0.01</td>
</tr>
<tr>
<td>Fe-Ukr(1)</td>
<td>1.37 ± 0.05</td>
<td>0.02 ± 0.01</td>
</tr>
<tr>
<td>Fe-Ukr(2)</td>
<td>1.67 ± 0.06</td>
<td>0.01 ± 0.01</td>
</tr>
<tr>
<td>Fe-Ukr(3)</td>
<td>1.70 ± 0.06</td>
<td>0.05 ± 0.01</td>
</tr>
<tr>
<td>Slov</td>
<td>0.45 ± 0.02</td>
<td>0.01 ± 0.01</td>
</tr>
<tr>
<td>Fe-Slov</td>
<td>8.75 ± 0.33</td>
<td>0.05 ± 0.01</td>
</tr>
<tr>
<td>Mn-Slov(1)</td>
<td>0.47 ± 0.02</td>
<td>4.58 ± 0.17</td>
</tr>
<tr>
<td>Mn-Slov(2)</td>
<td>0.43 ± 0.02</td>
<td>0.33 ± 0.01</td>
</tr>
<tr>
<td>FeMn-Slov</td>
<td>19.22 ± 0.73</td>
<td>2.80 ± 0.10</td>
</tr>
<tr>
<td>CaMn-Slov</td>
<td>0.35 ± 0.01</td>
<td>3.68 ± 0.13</td>
</tr>
<tr>
<td>Khol</td>
<td>0.33 ± 0.01</td>
<td>0.03 ± 0.01</td>
</tr>
<tr>
<td>Fe-Khol</td>
<td>1.43 ± 0.05</td>
<td>0.06 ± 0.01</td>
</tr>
</tbody>
</table>
(19.22 ± 0.73 mas%). Materials with highest Mn content are Mn-Slov(1) (4.58 ± 0.17 mas%), CaMn-Slov (3.68 ± 0.13 mas%) and FeMn-Slov (2.80 ± 0.10 mas%).

6.3. Optical microscopy

All raw and modified materials were studied using optical microscopy. Optical micrographs were obtained at different magnifications: x7.1, x20 and x80; and x115 for some materials. Optical micrographs with magnification of x80 for all studied materials are summarized in Figure 1.

Optical micrographs (Figure 1) indicate that obtained materials FeMn-Slov, Fe-Khol, Fe-13X, Fe-Ukr(1), Fe-Ukr(2), Fe-Ukr(3) and Fe-Slov are modified with an iron compound. Furthermore, also in x80 magnification materials look homogeneous, except Fe-Ukr(1), Fe-Ukr(2), Fe-Ukr(3) and Fe-Slov, where iron compound covers only parts of the materials’ surface. Optical micrographs (Figure 1) also indicate that obtained materials Mn-4A, Mn-Slov(1), Mn-Slov(2), FeMn-Slov and CaMn-Slov are modified with manganese compounds.

6.4. Scanning electron microscopy

All raw and modified materials, except for nonhomogeneous Fe-Ukr(1), Fe-Ukr(2), Fe-Ukr(3) and Fe-Slov are studied with scanning electron microscopy (SEM). SEM micrographs are summarized in Figure 2.

SEM micrographs (Figure 2) indicate that obtained materials FeMn-Slov, Fe-Khol and Fe-13X are modified with an amorphous compound, which is also further proved using EDX (for more details, also see source [17]). SEM micrographs (Figure 2) indicate that obtained materials Mn-4A, Mn-Slov(1), Mn-Slov(2), FeMn-Slov and CaMn-Slov are modified with manganese compounds. In case of Mn-4A and Mn-Slov(2), manganese compound is amorphous, while for Mn-Slov(1), FeMn-Slov and CaMn-Slov a new crystalline phase (Mn₈O₁₀Cl₃) is obtained (more details can be found in source [17]).

6.5. Energy-dispersive X-ray spectroscopy

All raw and modified materials, except nonhomogeneous Fe-Ukr(1), Fe-Ukr(2), Fe-Ukr(3) and Fe-Slov were studied with energy-dispersive X-ray spectroscopy (EDX). EDX results are summarized in Table 3. Elements with probability ≥95% are shown as an average of 6 repeated measurements at different locations.

Analysing amorphous iron compounds with EDX, it was deduced that it consists of 65.28 ± 3.91 mas% Fe and 34.72 ± 2.08 mas% O. This result is in agreement with elemental content of FeOOH (62.85 mas% Fe, 36.01 mas% O and 1.14 mas% H).

Analysing manganese crystals with EDX, it was deduced that it consists of 54.58 ± 3.27 mas % Mn, 30.27 ± 1.82 mas% O and 15.15 ± 0.91 mas% Cl. This result is in agreement with elemental content of Mn₈O₁₀Cl₃ (62.26 mas% Mn, 22.67 mas% O, 15.07 mas% Cl). Elevated oxygen content can be explained with signal from zeolite oxygen and/or with other manganese oxide presence.
Figure 1. Optical micrographs with x80 magnification of all studied materials: (A) 4A; (B) Mn-4A; (C) Ukr; (D) Slov; (E) Mn-Slov(1); (F) Mn-Slov(2); (G) FeMn-Slov; (H) CaMn-Slov; (I) Khol; (J) Fe-Khol; (K) 13X; (L) Fe-13X; (M) Fe-Ukr(1); (N) Fe-Ukr(2); (O) Fe-Ukr(3); (P) Fe-Slov.
Figure 2. SEM micrographs of studied materials: (A) 4A; (B) Mn-4A; (C) Ukr; (D) Slov; (E) Mn-Slov(1); (F) Mn-Slov(2); (G) FeMn-Slov; (H) CaMn-Slov; (I) Khol; (J) Fe-Khol; (K) 13X; (L) Fe-13X.
6.6. As(V) sorption experiments

Experimental data is compared with sorption models: Langmuir, Freundlich, Dubinin–Radushkevich, Temkin and Redlich–Peterson isotherm linearized forms comparing obtained determination coefficients ($R^2$). As(V) sorption experimental results and determination coefficients of fit with the sorption models are reported in Table 4.

As(V) sorption on raw zeolites is most precisely characterized by Freundlich’s sorption model in all cases. As(V) sorption on iron oxo hydroxides indicates that for these materials physisorption is dominant [3]. As(V) on Fe(III)-modified clinoptilolite and synthetic zeolites is most precisely characterized by Langmuir model [17, 20], which is also consistent with experimental data in this work, except for Fe-Ukr(2). Inconsistency in case of Fe-Ukr(2) can be explained by material’s inhomogeneity, leaving large areas of zeolite unmodified.

As(V) sorption on Mn-modified zeolites is most precisely characterized with Freundlich model in cases of Mn-Slov(1), FeMn-Slov and CaMn-Slov and with Redlich–Peterson model in cases of Mn-4A and Mn-Slov(2).

Obtained As(V) equilibrium sorption capacities and Langmuir monolayer coverage (where applicable) are systematized in Table 5. Improvement shows how much material’s equilibrium sorption capacity (at the highest studied As(V) solution concentration of 300 mg As(V)/L) increased after modification. Each sorption experiment was performed 3 times and obtained results represent average with standard deviation.

As(V) sorption isotherms for each respective material group (raw and modified, from top to bottom: Slov, Ukr, Khol, A and X) are shown in Figure 3.
6.7. As(V) equilibrium sorption capacity correlation with Fe and Mn content

A correlation analysis was performed using obtained sorption data and Fe and Mn content of materials. The analysis was performed using Microsoft Excel Data Analysis toolpack (USA; International). Three different As(V) equilibrium concentrations were chosen – at initial As(V) solution concentration of 300, 100 and 5 mg/L. Correlation coefficients are summarized in Table 6. Correlation analysis was performed using all studied materials, as well as separate material groups (where applicable).

Correlation between material’s As(V) equilibrium sorption capacity $q_e$ and Fe content is strong (correlation strength ≥0.67 by absolute value) for all groups and for all materials altogether at all studied concentrations.

Correlation between material’s As(V) equilibrium sorption capacity $q_e$ and Mn content is strong (correlation strength ≥0.67 by absolute value) only for Ukrainian natural zeolites at initial As(V) solution’s concentration of 5 and 100 mg/L. Medium correlation (correlation strength between 0.33 and 0.67 by absolute value) is observed for Ukrainian and Slovakian natural zeolites at initial As(V) solution’s concentrations of 300 mg/L. For studied materials altogether at all concentrations and for Slovakian zeolite group at initial As(V) solution’s concentration of 5 and 100 mg/L correlation is weak (correlation strength <0.33 by absolute value).

Table 4. As(V) sorption data and comparison with sorption models. Values marked in bold signify a respective model with the highest fit.

<table>
<thead>
<tr>
<th>Material</th>
<th>Determination coefficient $R^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Langmuir</td>
</tr>
<tr>
<td>4A</td>
<td>0.0586</td>
</tr>
<tr>
<td>Mn-4A</td>
<td>0.9911</td>
</tr>
<tr>
<td>Ukr</td>
<td>0.3874</td>
</tr>
<tr>
<td>Fe-Ukr(1)</td>
<td>0.9853</td>
</tr>
<tr>
<td>Fe-Ukr(2)</td>
<td>0.9062</td>
</tr>
<tr>
<td>Fe-Ukr(3)</td>
<td>0.9990</td>
</tr>
<tr>
<td>Slov</td>
<td>0.7607</td>
</tr>
<tr>
<td>Fe-Slov</td>
<td>0.9936</td>
</tr>
<tr>
<td>Mn-Slov(1)</td>
<td>0.9896</td>
</tr>
<tr>
<td>Mn-Slov(2)</td>
<td>0.9625</td>
</tr>
<tr>
<td>FeMn-Slov</td>
<td>0.9704</td>
</tr>
<tr>
<td>CaMn-Slov</td>
<td>0.5771</td>
</tr>
<tr>
<td>KHol</td>
<td>0.6213</td>
</tr>
<tr>
<td>Fe-KHol</td>
<td>0.9968</td>
</tr>
<tr>
<td>13X</td>
<td>0.5236</td>
</tr>
<tr>
<td>Fe-13X</td>
<td>0.7705</td>
</tr>
</tbody>
</table>
7. Conclusion

Natural and synthetic zeolites were modified and tailored for As(V) sorption using novel methods. Based on optical microscopy, SEM, EDX and FAAS results, it was proved that zeolites were modified with amorphous Fe(III) oxohydroxide, amorphous Mn(IV) oxide and crystalline mixed oxidation state manganese oxide-chloride Mn$_8$O$_{10}$Cl$_3$. FeOOH, and Mn$_8$O$_{10}$Cl$_3$-modification improves As(V) sorption capacities of these aluminosilicates. Granulated material with the highest As(V) sorption capacity is Mn$_8$O$_{10}$Cl$_3$-FeOOH-modified Slovakian clinoptilolite natural zeolite (6.96 ± 0.49 mg/g). These materials are effective and potential As(V) sorbents, able to remove >98% As(V) in water environments. Sorption capacity of granulated materials (per unit volume) was improved up to 26.5 times. As(V) sorption on FeOOH-modified zeolites follows Langmuir model most precisely, while sorption on unmodified zeolites is described by Freundlich isotherm. In case of Mn$_8$O$_{10}$Cl$_3$-modified and MnO$_2$-modified zeolites, Freundlich and Redlich–Peterson models show the most precise fit, respectively. There is a strong and positive correlation between As(V) sorption capacity and Fe content among all zeolites. Among studied materials, Mn content showed a strong positive correlation only among Ukrainian clinoptilolite natural zeolites.

<table>
<thead>
<tr>
<th>Material</th>
<th>Equilibrium sorption capacity $q_{e,300}$ [mg/g]</th>
<th>Equilibrium sorption capacity $q_{e,100}$ [mg/g]</th>
<th>Equilibrium sorption capacity $q_{e,5}$ [mg/g]</th>
<th>Langmuir monolayer coverage $q_m$ [mg/g]</th>
<th>Improvement [times]</th>
<th>Water remediation [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>4A</td>
<td>0.15 ± 0.01</td>
<td>0.06 ± 0.00</td>
<td>0.01 ± 0.00</td>
<td>—</td>
<td>4.2</td>
<td>—</td>
</tr>
<tr>
<td>Mn-4A</td>
<td>0.39 ± 0.02</td>
<td>0.30 ± 0.02</td>
<td>0.13 ± 0.01</td>
<td>0.40</td>
<td>2.6</td>
<td>43.0</td>
</tr>
<tr>
<td>Ukr</td>
<td>0.43 ± 0.02</td>
<td>0.14 ± 0.01</td>
<td>0.02 ± 0.00</td>
<td>—</td>
<td>—</td>
<td>6.6</td>
</tr>
<tr>
<td>Fe-Ukr(1)</td>
<td>1.08 ± 0.06</td>
<td>0.96 ± 0.07</td>
<td>0.17 ± 0.02</td>
<td>1.19</td>
<td>2.5</td>
<td>57.2</td>
</tr>
<tr>
<td>Fe-Ukr(2)</td>
<td>0.87 ± 0.05</td>
<td>0.40 ± 0.03</td>
<td>0.05 ± 0.01</td>
<td>—</td>
<td>2.0</td>
<td>17.4</td>
</tr>
<tr>
<td>Fe-Ukr(3)</td>
<td>0.92 ± 0.05</td>
<td>0.83 ± 0.04</td>
<td>0.24 ± 0.01</td>
<td>0.93</td>
<td>2.1</td>
<td>80.2</td>
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<tr>
<td>Slov</td>
<td>0.36 ± 0.02</td>
<td>0.14 ± 0.01</td>
<td>0.02 ± 0.00</td>
<td>—</td>
<td>—</td>
<td>8.2</td>
</tr>
<tr>
<td>Fe-Slov</td>
<td>4.74 ± 0.29</td>
<td>4.08 ± 0.29</td>
<td>0.29 ± 0.03</td>
<td>4.92</td>
<td>13.2</td>
<td>&gt;98.0</td>
</tr>
<tr>
<td>Mn-Slov(1)</td>
<td>4.92 ± 0.30</td>
<td>2.21 ± 0.13</td>
<td>0.12 ± 0.01</td>
<td>10.50</td>
<td>13.7</td>
<td>41.2</td>
</tr>
<tr>
<td>Mn-Slov(2)</td>
<td>0.66 ± 0.04</td>
<td>0.41 ± 0.02</td>
<td>0.13 ± 0.01</td>
<td>0.71</td>
<td>1.8</td>
<td>44.3</td>
</tr>
<tr>
<td>FeMn-Slov</td>
<td>6.96 ± 0.49</td>
<td>3.79 ± 0.27</td>
<td>0.30 ± 0.02</td>
<td>6.82</td>
<td>19.3</td>
<td>&gt;98.0</td>
</tr>
<tr>
<td>CaMn-Slov</td>
<td>1.32 ± 0.10</td>
<td>0.51 ± 0.04</td>
<td>0.06 ± 0.00</td>
<td>—</td>
<td>3.7</td>
<td>21.3</td>
</tr>
<tr>
<td>KHol</td>
<td>0.24 ± 0.02</td>
<td>0.10 ± 0.01</td>
<td>0.02 ± 0.00</td>
<td>—</td>
<td>—</td>
<td>7.9</td>
</tr>
<tr>
<td>Fe-KHol</td>
<td>1.25 ± 0.06</td>
<td>1.18 ± 0.06</td>
<td>0.30 ± 0.02</td>
<td>1.28</td>
<td>5.2</td>
<td>&gt;98.0</td>
</tr>
<tr>
<td>13X</td>
<td>0.30 ± 0.02</td>
<td>0.12 ± 0.01</td>
<td>0.01 ± 0.00</td>
<td>—</td>
<td>—</td>
<td>3.2</td>
</tr>
<tr>
<td>Fe-13X</td>
<td>1.05 ± 0.05</td>
<td>0.42 ± 0.02</td>
<td>0.08 ± 0.00</td>
<td>—</td>
<td>3.5</td>
<td>26.0</td>
</tr>
</tbody>
</table>

Table 5. As(V) equilibrium sorption capacities.

Use of Synthetic and Natural Zeolites Tailored for As(V) Sorption
http://dx.doi.org/10.5772/intechopen.72614
Figure 3. As(V) sorption experimental data and constructed isotherms.
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