Review

Tailored zeolites for the removal of metal oxyanions: Overcoming intrinsic limitations of zeolites

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HIGHLIGHTS

• First review manuscript singly focusing on the interaction of zeolites with metal oxyanions.
• The focus on chromium and arsenic oxyanions offers the covering of a wide array of zeolite modification technologies.
• Detailed description of the zeolite modification technologies that have been successfully employed to date.
• Global overview of the experimental results obtained with the modified supports, with given considerations on the diverse aspects of heavy metal treatment.

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ABSTRACT

This review aims to present a global view of the efforts conducted to convert zeolites into efficient supports for the removal of heavy metal oxyanions. Despite lacking affinity for these species, due to inherent charge repulsion between zeolite framework and anionic species, zeolites have still received considerable attention from the scientific community, since their versatility allowed tailoring them to answer specific requirements. Different processes for the removal and recovery of toxic metals based on zeolites have been presented. These processes resort to modification of the zeolite surface to allow direct adsorption of oxyanions, or by combination with reducing agents for oxyanions that allow ion-exchange with the converted species by the zeolite itself. In order to testify zeolite versatility, as well as covering the wide array of physicochemical constraints that oxyanions offer, chromium and arsenic oxyanions were selected as model compounds for a review of treatment/remediation strategies, based on zeolite modification.

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1. Introduction

Heavy metal contamination of the environment is a well-known concern. The discharge of these pollutants into aqueous streams is a direct result of human activity, mainly industrial [1]. Although some metals are essential to the enzymatic processes of aerobic or anaerobic organisms [2], high concentrations of the same can be toxic to lethal. Also, as heavy metals accumulate in the organisms and persist through their food chain, they ultimately present a risk for human health.

Numerous technologies have been developed for the treatment of heavy metal contamination, in aqueous medium or in soils. Ion-exchange and adsorption are amongst the most referred processes for this treatment [3]. Given the increasing environmental and economical awareness, development efforts for new technologies must bear in mind the potential for recovery-reuse capability, in a bid to decrease demand on processing of mineral ores and ultimately reducing manufacturing costs by allowing reutilization of otherwise discarded spent materials [4].

Zeolites have been proposed by some authors to be an efficient and economical resource for the treatment of general heavy metal contamination of effluents [3–6]. Their ion-exchange and adsorption capacities endure these supports with the ability to recover most cationic species from solution. Conversely, zeolites present very low to negligible affinity for the exchange of anionic compounds. Nevertheless, their physicochemical properties also render zeolites quite versatile materials, and many researchers found interest in these supports and conducted efforts to tailor zeolites in order to allow the removal of anionic compounds. These tailored materials have found application on the removal of toxic metal oxyanions, such as chromate or arsenate [7], radioactive uranyl ions [8], as well as other non-metallic anionic species such as iodides [9], phosphates [10] or nitrates [11].

This paper presents details of the strategies that have been developed to overcome the natural limitation of zeolites to the removal of oxyanions. The focus is given on the treatment of As(III), As(V) and Cr(VI) compounds, some of the most commonly found oxyanions in metal-contaminated sites. The choice on these compounds is justified as they cover some interesting differences between them, which are also transported to some extent to the application and performance of tailored zeolites.

A brief introduction to zeolites and the mentioned oxyanions will be followed by a detailed description of the functionalization procedures, ending on a comparison of achievements of the different tailoring processes on the removal/treatment of chromium and arsenic oxyanions.

1.1. Zeolites

Zeolites are highly crystalline aluminosilicate materials, consisting of either Si or Al atoms interconnected via oxygen bridges. Si and Al are also referred as “T atoms”, as they are the center of tetrahedral SiO4 or AlO4 structures. These tetrahedra are the building blocks forming the several crystalline lattice structures known so far, from either naturally occurring or specifically designed synthetic zeolites. Their intricate structural arrangement is also responsible for the high specific surface area of these materials, which present figures exceeding 700 m²/g [12]. This is a key factor for the excellent adsorption and catalytic properties of these materials.

Since aluminum has a valence of +3 and silicon +4, isomorphous framework substitution of Si by Al originates a charge deficit that is responsible for the need of charge-balancing cations, in order to maintain the neutrality of the zeolitic network [13]. Common charge-balancing cations are alkaline or rare-earth metals like Na+, K+, Mg2+ or Ca2+, which are electrostatically bound to the Al atoms [14]. The mobility of these species promotes the ion-exchange capability of zeolites for cationic species. The ion-exchange capacity of a zeolite is dependent on the Si/Al ratio of the framework, where a higher Al content will correspond to a higher content of charge-balancing cations, which in turn will increase the overall ion-exchange capacity of the zeolite [15]. Naturally occurring zeolites present high Al content, with Si/Al ratios typically ranging from 1 to 6 [16].

1.2. Chromium oxyanions

Chromium is an important industrial compound, mainly used in electric plating, refining, printing and dying industries [17,18]. It is one of the most common inorganic contaminants present in groundwater of hazardous waste sites, in the form of hexavalent chromium, one of the most toxic priority pollutants [18–20].

In aqueous medium, this transition metal can be found in different oxidation states, the most common are the trivalent and hexavalent states [21], being Cr(III) the most stable. Divalent Cr(II) is also known to be stable in oxygen-free conditions, as it is strongly reducing [22]. Trivalent chromium is an essential trace element for the metabolism of glucose and lipids [20]. Most Cr(III) compounds
present very low solubility at neutral or physiological pH, therefore considered to bear low toxicity to cells [23]. Cr(VI) forms much more soluble species, being extremely mobile in soils or groundwater [24,25]. Also, the external cell structure is more permeable to Cr(VI) than for the case of Cr(III) [26]. The carcinogenic and/or mutagenic effects of Cr(VI) arise from the combination of its mobility in cellular structures and high redox potential. Cr(VI) permeates cellular walls and, once in the cellular cytoplasm, is converted into Cr(III), which can bind covalently to biomolecules such as DNA [21].

Cr(VI) is commonly found as chromate or dichromate anions [27]. This speciation is related to pH and concentration [28], where chromate is predominant at very low pH. The negative charge of chromate and dichromate is responsible for their poor adsorption in soils or most mineral structures, which explains the great mobility of Cr(VI) in these media.

Since most contaminant metals form cationic species in aqueous media, technologies designed for the treatment of common metal contamination may suit Cr(III) but lack the flexibility for the anionic Cr(VI) species. The most common treatment procedures for Cr(VI) include reduction–precipitation, liquid–solid transfer, precipitation–floculation or concentration [17]. Most of these technologies are disposal-type, so efforts have been conducted on the research for recovery-reuse procedures, in order to address cost-effectiveness and decrease demand on chromium ore resources.

1.3. Arsenic oxyanions

Arsenic is the 20th most abundant element of the Earth’s crust and contamination of aqueous sources is known to occur naturally, although human activity also contributes to the release of this metal into the environment [29,30]. It has found application in several industries, namely mining, glassmaking, electronics, wood preservation and pharmaceutics [31]. Apart from the natural contamination sources, use of arsenic in agricultural products such as fungicides, herbicides and insecticides has added to the contamination of soils and underground water sources. Arsenic is known to be carcinogenic and its contamination, both natural or anthropogenic, is a known cause for concern, especially as there are widespread reports of populations depending on As-contaminated water sources [29,32].

Arsenic is a metalloid with similar chemistry to phosphor and antimony [33]. In environmental conditions, it is found in the trivalent and pentavalent oxidation states, both forming oxyanions. Oxidized environments favor the As(V) inorganic acid species, ranging from H₃AsO₄ at pH below 2 to the arsenates H₄AsO₄n⁺³ (n ≤ 2), where AsO₄³⁻ is found at pH higher than 10. As(III) compounds are normally found in anoxic and/or reducing environment such as underground waters, typically as H₃AsO₃ at pH below 9, or the arsenite forms H₄AsO₄n⁻³ (n ≤ 2) for higher pH [32,33]. Some bioprocesses also contribute to the speciation of As(V) to As(III), as bacteria can promote reduction and methylation of arsenic as a defense to its toxicity. Also, intra-cellular As(V) reduction into As(III) may give rise to free radicals which promote damage to cellular structures, and arsenite in known to inhibit several enzymes [31]. However, some bacteria are known to use arsenic in their energy metabolism, since arsenate is similar to phosphate and may replace the latest in biomolecules.

Comparatively to Cr(VI), the negative charge of the different arsenite or arsenenate species requires specific treatment processes. As(V) is considered less toxic as it absorbs more strongly to solid surfaces than As(III), so oxidation–precipitation or oxidation–coagulation/flocculation processes are typically employed for the treatment of As-containing wastewaters or for in situ remediation of soil contamination [32,34,35]. Other processes include ion-exchange in specific resins, membrane processes and adsorption on iron-based materials.

2. Experimental aspects of zeolite tailoring

Due to their ion-exchange capacity, zeolites have found widespread use in research studies related to the removal of metal contaminants in general [4,6]. Studies on the removal of Cr and As have also been conducted.

For the removal of Cr(III), zeolites have been widely considered to be good supports for the direct removal of these species, in either natural or synthetic form, and in the presence of other competing metal species [36–39]. Studies on the removal of Cr(VI) present a different nature from other cationic metal ions removal processes. Zeolites are unable to perform direct ion exchange of Cr(VI) compounds, as charge repulsions from the negatively charged framework limits its permeability to anions [40]. This limitation was witnessed by several authors, in both natural and synthetic zeolites [24,41]. Arsenic ions present some differences toward Cr(VI), as unmodified zeolites have some affinity for arsenic oxyanions (detailed in Section 3). Nevertheless, studies on the usage of zeolites for the removal of Cr and As oxyanions generally present indirect methodologies, where zeolites are previously modified in order to overcome their limitation for negatively charged ions.

From the different methodologies proposed, a separation into two distinct processes is possible: processes that involve zeolite surface functionalization for direct oxianion adsorption (discussed in Section 2.1), and processes involving redox reactions of Cr or As and subsequent retention by the zeolite-based system (discussed in Section 2.2).

2.1. Zeolite tailoring for the direct adsorption of oxyanions

The adsorption of various anions on tailored zeolites has been achieved. In order to allow adsorption to occur, tailoring of the zeolite surface is required as to modify the negative zeta potential that repels these compounds. Two main processes have been successfully employed: deposition of surfactants on the zeolite surface (Section 2.1.1), and grafting molecules with affinity for negative ions onto the zeolite (Section 2.1.2). Additionally, the modification of zeolites with metal ions that form insoluble chromates or arsenates has been proposed for the removal of Cr and As (Section 2.1.3).

2.1.1. Surfactant-modified zeolites (SMZ)

One of the strategies for tailoring zeolites in order to allow adsorption of negatively charged species is the modification of the polarity of the surface using surfactant compounds. The surfactancy ability of zeolites is known from the 1980s, when zeolites replaced phosphates as water softeners in detergent formulations. The earliest reports of deliberate modification of the surface polarity of inorganic materials date back to 1988: Boyd et al. presented the modification of soil samples using the surfactant hexadecyltrimethylammonium (HDTMA), to allow the retention of several organic compounds [42].

Surfactant-modified zeolites (SMZ), also named organo-zeolites, are prepared using certain surfactant molecules, mostly quaternary amine salts, which can be ion-exchanged on the surface of the zeolite. If these amines are provided with one long alkyl chain, the hydrophilic section or “head” (where the positively charged nitrogen atom is located) will be attracted by the zeolite, whereas its hydrophobic counterpart (the alkyl “tail”) will oppose the zeolite (Fig. 1). If the surfactant concentration in solution is kept below the respective critical micelle concentration (cmc), a monolayer (hemimicelle) will form on the zeolite surface. If the
concentration of surfactant is further increased beyond cmc, the hydrophobic tails will attract each other and form a double layer or admicelle [43] over the zeolite surface. This double layer will cover the zeolite with one of the N’R surfaces deposited on the zeolite and the other N’R surface facing the solution, therefore endowing the outer surface with positive charges. These can be used to adsorb metal oxanions or other anions, as depicted in Fig. 2.

The Cr(VI) adsorption capacity of most surfactants used in the preparation of SMZs has been demonstrated even on the unsupported form (macelle form) [44]. While this fact may lead to consider this procedure as a more economical process when compared to the process costs of tailoring zeolites, Bowman described SMZs prepared from natural zeolites to be a cost-effective support [45], since natural zeolites are considerably less expensive than the cost of the surfactants used. Moreover, the same author added that SMZs have the potential of being alternative supports to activated carbon or ion-exchange resins. The advantages of being a heterogeneous support (granting ease in recovery) and still having the intrinsic properties of zeolites render SMZs attractive sorbents, especially for multi-component effluents. Zeolites present internal and external cation exchange sites; surfactant molecules exchange mainly in the outer layer sites, as their dimensions limit accessibility to the inner zeolite framework [46]. This allows zeolite to maintain their characteristic cation exchange ability after modification with surfactants, potentially allowing simultaneous retention of anionic species on the modified surface and cationic metal species on internal exchange sites. SMZs have also been found to be resistant to organic solvents, extreme pH conditions or high ionic strength [47], so their application range in liquid phase conditions is not restricted when compared to unmodified zeolites. Several surfactants have been successfully used to date, for the modification of either natural or synthetic zeolites, as well as other mineral compounds such as clays. This process enables not only the adsorption of metal oxanions; sulfate, nitrate and phosphate [48,49], organic and aromatic compounds [45,50], pharmaceutical drugs [51], bacteria [52] or even viruses [53] have been reported to be adsorbed on SMZs.

### 2.1.2. Surface functionalization of zeolites

An alternative route to modify zeolites in order to induce affinity for anions is surface functionalization. This can be achieved by grafting species with adequate anion-affinity on the surface of the zeolite, in a process that resembles to SMZ, as the objective is the inversion of the zeta potential of the surface of zeolites. However, some differences exist between surface functionalization and SMZ. While SMZs are prepared by adsorption of surfactants, grafting involves generation of covalent bonding between the grafting agent and the zeolite.

The most common grafting agents used are hybrid organic–inorganic compounds, such as alkyl silanes. These will bond to the zeolite surface silanol groups (SiOH), which are relatively reactive [54]. A review from Moller and Bain offered considerable detail on the silane-based functionalization of several MCM materials, as well as the respective reactive conditions for their preparation [55]. Fig. 3 illustrates an example of grafting aminopropyl-triethoxy-silane (APTES) into a silica-like surface. The amino group in APTES is useful for the adsorption of anions, as it is easily protonated through pH control [56,57], and therefore becomes able to retain negatively charged ions (Fig. 4).

Since SiOH groups are widely available in ordered silica-like structures, these are the predominant targets for surface functionalization. These groups are also available in Si-rich zeolites or Al-free zeolites (such as the silicalite MFI zeolite [58]), so surface functionalization of high Si/Al ratio zeolites can result in high loading of organic matter on the support. Conversely, Al-rich zeolites will be less prone to grafting silanes, since with the increase in Al content of a zeolite, the lesser available silanol groups will be present at the surface. Nevertheless, their higher capacity for cation exchange renders Al-rich more adequate supports for SMZ preparation. In this regard, surface functionalization and

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**Fig. 1.** Formation of a monolayer or double-layer on the zeolite surface by an alkyl-quatery ammonium surfactant.

**Fig. 2.** Adsorption of a metal oxanion on a SMZ.

**Fig. 3.** Grafting of APTES into a silica-like surface.
surfactant modification of zeolites can be considered two complementary techniques to induce changes on the zeta potential of zeolites according to the Si/Al ratio.

Unlike most large pore mesoporous silica materials, zeolites present two types of available surface area: the internal surface area, only accessible to molecules that are able to diffuse through zeolite pores; and the external surface area [59]. The internal surface area is less prone to functionalization, as it presents less silanol groups as the external surface, being the hydroxyl groups available as the Si-(OH)-Al Brønsted sites [60]. Therefore, zeolites are mainly functionalized on the outer surface area. Nano-crystalline zeolites have been considered more adequate supports for functionalization, as the ratio between external/internal surface area is higher [61].

As it is the case of SMZs, zeolites are able to maintain their intrinsic properties such as ion-exchange and shape selectivity after functionalization, which in turn allows for bi- or multi-functionality of these materials. This property is commonly explored in catalysis [59]; however, a different kind of bifunctional zeolites has been proposed by Barquist and Larsen for the specific removal of hexavalent chromium [62]. The authors selected magnetic Y zeolites as supports for the functionalization with APTES. The magnetization of Y zeolite can be achieved by forming iron oxides in the zeolite, through ion-exchange of Fe(III) and Fe(II) ions and then subjecting to alkaline treatment [63]. This property was chosen to overcome the hydrophobicity of APTES-functionalized zeolites, which tend to form colloidal suspensions in water media, therefore losing easiness of recovery.

2.1.3. Metal modification of zeolites (MMZ)

Anion adsorption on metal-modified zeolites has been studied for some time and the adsorption of Cr or As oxyanions has been successfully achieved.

Metal modification of zeolites can be performed by two processes; simple ion-exchange, using cationic metal ions that present great affinity for the target oxyanions; or through promoting the formation of metal oxide species on the zeolite. While the later allows direct adsorption and further retrieval of sorbates, the first method is usually employed to introduce oxyanion-precipitating agents or oxidant/reductant species in the zeolite that will perform the treatment of As or Cr oxyanions. In this section, only the processes for oxyanion removal without redox reactions will be detailed, while MMZs with reducing/oxidizing-species will be discussed later.

Zeolites most commonly present counter-ions from alkali or alkaline earth metal elements. The replacement of these for metals which form insoluble Cr and/or As salts has been proposed as a process for the removal of Cr(VI) and As(V) from solution. Chromate/dichromate and arsenate salts of metal ions such as Ag(I), Al(III), Ba(II), Fe(II) or Fe(III), Hg(II), Pb(II) and Zn(II) are highly insoluble, so provided zeolites are exchanged with these cations previously, it is possible to immobilize the corresponding chromates/arsenates on the zeolite. The preparation of this kind of MMZs is straightforward. Normally, the zeolite is impregnated with a solution of a salt of the selected metal ion, usually a nitrate or chloride. The solution is stirred at room temperature and the solid is recovered, washed and dried prior to application.

Ion-exchanged MMZs have demonstrated good performance, despite some detrimental re-solubilization of chromates and arsenates taking place. In order to overcome this limitation, covalently bonded metal oxide MMZs were successfully used for metal oxyanion removal. Iron oxides are commonplace for the preparation of metal oxide MMZs for arsenic removal. This is mainly due to the high success of granular ferric hydroxide (GFH) on the treatment of arsenic contamination [64], where the coupling of Fe oxides to zeolites attempts to mimic the specificity of GFH for arsenate-like ions. The preparation of Fe-MMZs normally adds a calcination step following the ion-exchange procedure, where the sample is calcined in alternating oxidizing/reducing atmospheres (ex: N2 and H2) [65]. This class of materials is often termed iron-coated zeolites (ICZ).

The formation of other metal oxides on zeolites bears a similar principle to the previously described, where the zeolite is impregnated with the chosen metal, followed by slow addition of a base, normally NaOH, to promote the formation of metal oxides on the zeolite upon drying or calcination [66]. This process was also successfully used for the preparation of magnetic zeolites, using a mixture of Fe(II)/Fe(III) ions [63,67]. The adsorption of Cr and As in metal-oxide MMZs is reversible, offering an easier recovery of the removed compounds, when compared the other MMZs.

2.2. Zeolite conjugation with reducing/oxidizing species

A different perspective for the removal of metal oxyanions with zeolites is to convert them through redox reactions into other ions which are possible to be exchanged by the zeolite.

This is mostly evident for Cr(VI). Both natural and synthetic zeolites possess limited affinity for hexavalent chromium, but have demonstrated good ion-exchange properties for the trivalent form. This occurs since Cr(III) forms cationic species in aqueous solution. Therefore, provided that Cr(VI) is previously reduced to Cr(III), zeolites would then be able to recover Cr from solution in an indirect process. Overall, the reduction of hexavalent chromium occurs according to Eq. (1) (for acidic pH) or Eq. (2) (for alkaline pH) [22]:

\[
\text{Cr}_2\text{O}_7^{2-} + 14\text{H}^+ + 6\text{e}^- \rightarrow 2\text{Cr}^{3+} + 7\text{H}_2\text{O} \quad E^\circ = +1.33 \text{ eV} \quad (1)
\]

\[
\text{CrO}_4^{2-} + 4\text{H}_2\text{O} + 3\text{e}^- \rightarrow \text{Cr(OH)}_{3+} + 5\text{OH}^- \quad E^\circ = -0.12 \text{ eV} \quad (2)
\]
The inversion of charge in arsenic ions is not so straightforward as between Cr(VI)/Cr(III). Both As(V) and As(III) exist in nature as oxyanions, being the redox properties of the As(V)/As(III) pair described in Eq. (3) [68]. Despite that the redox pair As(III)/As(V) does not yield any cationic species, the oxidation of arsenite to arsenate is still of great importance since the trivalent form is more toxic and more difficult to remove.

$$\text{H}_2\text{AsO}_4^- + 2\text{H}^+ + 2\text{e}^- \rightarrow \text{H}_3\text{AsO}_3^- + \text{H}_2\text{O} \quad E^\circ = +0.56 \text{eV} \quad (3)$$

Different redox-assisted processes for the treatment of Cr and As on modified zeolites have been mentioned in the literature. All consist in endowing the zeolite with redox agents that assist the retention by the system. MMZs have also been proposed for the redox-assisted removal of oxyanions, or as photocatalysts for the treatment of the same. These will be detailed in Sections 2.2.2.1 and 2.2.2 respectively. Other option is the combination of biomass with zeolites (Section 2.2.2.3).

### 2.2.1. Metal-modified zeolites for redox processes

MMZs have also been successfully used in redox processes for the treatment of Cr and As. The previously described methods for the preparation or MMZs are also valid for the incorporation of reducing/oxygenizing particles. Iron is commonly found coupled to zeolites for redox processes of Cr(VI) or As (III), with iron species such as Fe(III), Fe(II) and zerovalent iron. For the particular case of the conversion of arsenite to arsenate, Fe(III) is commonly employed to assist the oxidation, since the formal electrode potential of the Fe(III)/Fe(II) pair is higher than for As(V)/As(III) [68]. For the Cr(VI)/Cr(III) system, Fe(II) salts are used as the reductant in the reduction–precipitation process, which is the common industrial treatment procedure for hexavalent Cr [69]. Apart from iron, manganese species were also successfully combined with zeolites for the oxidation and removal of As(III) [70].

### 2.2.2. Photocatalytic processes

Photocatalysis arises from the generation of electron–hole pairs on a semiconductor material through photon absorption, leading to chemical reactions between the semiconductor surface and molecules in the vicinity [69]. Photocatalysis has been successfully applied to the treatment of Cr and As solutions, by promoting either reduction of Cr(VI) to Cr(III), or oxidation of As(III) to As(V). To this effect, prominent heterogeneous photocatalysts such as TiO$_2$ and ZnO have been successfully used [71].

Zeolite-based photocatalysts have found application in advanced oxidation processes. Despite these advances, only a limited number of papers detail the use of zeolite photocatalysts for the purpose of Cr(VI) treatment, albeit some interest in mesoporous silica is reported.

To the best of our knowledge, no photocatalytic process has been reported for As(III) oxidation and removal using zeolite-supported photocatalysts. This particular reaction has long been reported using suspended TiO$_2$ photocatalyst, and reports of successful application of TiO$_2$ supported in activated carbon or activated alumina have also been presented [71]. Hence, the perspective of zeolites as useful supports could be also considered for the case of arsenic.

### 2.2.3. Zeolites as supports for bioremediation of Cr and As oxyanions

Biosorption of metal contaminants has been thoroughly investigated and is a cost-effective and environmental alternative to less efficient metal treatment processes [2,72,73]. Different kinds of biomass, either live or dead, are able to retain metal ions on the high multiplicity of chemical groups present on cellular structures. Bacteria are well known for their use on the treatment of metal contaminated solutions [2]. The structural polymers on bacterial cell walls present several functional groups such as carboxyl, phosphoryl and amino groups, which are directly responsible for the reactivity of bacterial cells to aggression from metal contaminants [74]. The external surface of bacteria is intrinsically reactive toward dissolved metals, despite the different surface formats between the different types of bacteria.

While the removal and recovery of most metal cations through biosorption has been demonstrated, mechanisms regarding the biosorption of Cr(VI) and As(III)/(V) by live bacteria present interesting differences. Bacteria are able to reduce Cr(VI), owing to the high electronic density on the cell surface. Several microorganisms have been used for the removal of chromium from effluents with very promising results [75–79]. The mechanism of Cr(VI) bioreduction was intensively studied by Park et al., who provided evidence that the process is an “adsorption-coupled-reduction” [80,81]. Many functional groups that are present on cell walls can act as electron donors [82] and the diol-rich polysaccharide content of cell walls was found to coordinate with an intermediary oxidation state, Cr(V), hence assisting the overall reduction process. This unstable intermediary was found to be present on Cr-reducing cells [26]. In terms of bacterial oxidation of As(III), despite being inhibitory to cellular growth, some bacteria are able to use the redox pair As(III)/As(V) in the electron transport during the respiratory cycle. Specific arsenic-reducing or –oxygenizing enzymes have been extracted from bacteria and the As(V) and As(III) uptake and cellular pathways have been reported [31,83,84]. Under the scope of As contamination treatment, As(III)–oxygenizing bacteria can play a helpful role on the preconcentration of the easier to recover As(V) ions.

The combination of biosorbers with a zeolitic support offers some advantages. Solely suspended microorganisms present the disadvantages of being too sensitive to the xenobiotic effects caused by high levels of pollutant concentration. Also, mechanical strength and low density of microorganisms in suspension can hinder the performance of treatment systems [85]. Immobilization of biomass on a support offers the advantages of mechanical resistance and higher packing capacity. To this respect, zeolites and minerals provide a resistant support that can actively take part on the removal of the metal that underwent the redox process, promoted by the supported biomass.

### 3. Results and achievements on oxyanion removal with tailored zeolites

This section presents an overview on how the previously described zeolite tailoring strategies perform on the removal of arsenic and chromium oxyanions. While the treatment of Cr(VI) has been the driving force for the development of most processes, given the lesser natural affinity of zeolites for this case, some were also successfully applied to arsenic remediation as well. It is common to see studies including other oxyanions as well, and in fact, some works present the simultaneous treatment of As and Cr oxyanions with tailored zeolites [7,86,87]. Additionally, some of the tailoring strategies can be combined, as it is the case of MMZs which are also modified with surfactants [88,89]. The efficiency of the different approaches will be presented according to the respective processes and technologies detailed in Section 2. Cr and As uptake values are merely presented for illustration purpose, and should not be considered for a performance comparison between tailoring strategies, or even between results obtained for the same zeolite modification. Operational conditions between cited works are not identical and key parameters, such as initial oxyanion concentration, pH and contact time, are only mentioned in the text where appropriate. Additionally, the presented maximum uptake values
refer to either experimentally observed, or as predicted from the Langmuir adsorption model, at equilibrium state.

Despite having covered the different zeolite tailoring strategies for oxyanions in the previous section, it is worth to note that the removal of oxyanions using unmodified natural or synthetic zeolites has also been reported. This is commonly reported for the removal of arsenate ions, and the highly chemically related phosphate and antimonate oxyanions have also been found to adsorb on natural zeolites [33,90,91]. The general principle ruling zeolite affinity for this compound family is thought to involve the properties of the surface aluminol groups which, at specific pH values, promote the formation of surface inside–sphere complexes with these compounds [91–93]. At very low pH values, below the point of zero charge (pzc) of most zeolites, aluminol groups are protonated (Eq. (4)), while above pzc, they can be deprotonated by hydroxyl ions. Ions such as arsenate and phosphate react with either protonated or neutral aluminol groups, depending on solution pH being below or at near pzc (Eqs. (5) and (6)) [91,94].

**Zeolite-Al(OH) + H2O → Zeolite-Al(OH)2 + H2O**

**Zeolite-Al(OH)2 + H2O4 − → Zeolite-AlO-X(OH)2 + H2O**

X = AsorP

**Zeolite-Al(OH) + H2O4 − → Zeolite-AlO-X(OH)2 + OH−**

X = AsorP

Some authors also defend that the retention of As(V) by natural zeolites is assisted by their composition presenting a diversity of charge-balancing cations, some of which residual amounts of transition metal cations presenting affinity for As(V) in the same perspective as a metal-modified zeolite would perform. Residual iron and manganese ions in the composition of natural zeolites are regarded as useful on both retention of As(V) or oxidation of As(III) [95].

Despite the affinity between arsenic compounds and zeolites, some authors still conduct modification of zeolites for the improvement of specific properties, such as hydrophobicity and mechanical resistance (in SMZs), for promoting oxidation of As(III) to As(V) or to induce multifunctionality. As an example, SMZs offer the possibility of retaining both arsenic and other oxyanions such as chromate [86], while not compromising the zeolite ability to retain As(V). This was demonstrated by Chutia et al., whom published two simultaneous reports on the removal of arsenic, one describing the use of unmodified synthetic H-MFI zeolites and the other, the use of surfactant-modified natural zeolites. The authors observed that both systems were able to remove As(V) and regeneration of the supports was possible, through washing with NaOH or HCl [94,96].

### 3.1. Direct adsorption of Cr(VI) and As(V)/As(III)

#### 3.1.1. Use of SMZs for the removal of Cr(VI) and As oxyanions

The earliest application of SMZs for the removal of Cr(VI) can be traced back to a report by Santiago et al. in 1992 [4], with the use of ethylenediacetyldimethylammonium (EHDDMA) and cetylpyridinium (CPD) to modify the zeolite surface. The authors found the CPD-SMZ support more efficient for chromate removal, with an uptake of 0.65 mg Cr/g zeolite. Arsenic removal using both natural and surfactant-modified zeolites was reported later, in 1997 for the removal with unmodified zeolites, while a year later, Misaelides et al. described the earliest application of SMZs from natural zeolites for the removal of As(V) [95,97].

Soon after the initial studies of Cr(VI) removal by SMZs, Haggerty and Bowman demonstrated that an adsorption process, where reduction of Cr(VI) does not occur, was the global mechanism for the removal of this oxyanion [43]. Also for Cr(VI), Covarrubias et al. further commented that for monolayer SMZs, the global process that takes place is a surface precipitation, while for surfactant bilayer SMZs, ion–exchange with the surfactant counter–ions is the main process involved [98]. Li et al. demonstrated that the release of counter–ion(s) to the solution is directly related to the amount of sorbed Cr(VI) on SMZ packed columns [99]. As further evidence of the adsorption mechanism, the removal of chromate from solution by SMZs is commonly reported to be well fitted by the Langmuir adsorption model. As an example, Ghiaci et al. performed a study on Cr(VI) removal by natural clinoptilolite, ZSM-5 zeolite and MCM-41 molecular sieve, all tailored with HDTMA and CPD surfactants. Both the adsorption of the surfactants by the zeolites and the following adsorption of chromate by the SMZs were well fitted by the Langmuir adsorption model [100].

In terms of recovery-reuse procedures for SMZs, several authors reported the possibility for regeneration of the prepared supports. A recent report by Zeng et al. demonstrated the reusability of their HDTMA-surfactated clinoptilolite and chabazite supports [101]. The regeneration of the SMZs was conducted by rinsing the support with 0.1 M HCl, and the regenerated support was able to achieve 90.6% of the Cr(VI) removal of as–prepared equivalent SMZ. The same principle is valid for As(V) sorption and desorption with natural zeolite SMZs, as previously referred from the works of Chutia et al. [96]. In the latest, the authors employed NaOH or HCl at a concentration of 0.1 M, with the alkaline washing achieving greater recovery of the sorbed As(V) ions, at 70% desorption efficiency vs. less than 50% for HCl.

**Table 1** presents an overview on the achievements of different SMZs in the adsorption of Cr(VI) or As(V). The predominance of works using natural zeolites is noteworthy, which reflects authors’ efforts to address both cost–effective and environmental concerns. Nevertheless, SMZs prepared from synthetic zeolites have also been successfully employed on the removal of Cr(VI) and As(V).

A variation from the previous SMZs was recently reported by Shyaa, Hasan and Abbad. The authors proposed a polyaniline-loaded zeolite nanocomposite for the removal of Cr(VI). The modification of the zeolite was conducted though adsorption of anilinium ions on the zeolite inner and outer surface, being these ions latter polymerized with ammonium persulphate [111]. The system was able to achieve a maximum uptake of 25 mgCr/gzeolite, from an initial solution of 100 mg Cr/L.

#### 3.1.2. Functionalized zeolites

To date, the most studied organic-functionalized materials have been mesoporous, high-Si content supports [112]. The application of functionalized silica to the selective removal of oxyanions has been proposed since the late 1990s, with the work of Fryxell et al. on the usage of mesoporous silica functionalized with ethylenediamine silane, that was later reacted with Cu(II) ions, resulting in a selective adsorbent for As(V) and Cr(VI) in presence of other competing anions [113]. The basic principle is to immobilize coordinated Cu(II) species which are positively charged and possess electrostatic coordination affinity for tetrahedral anions, such as chromate. The authors reported a maximum uptake of 54 mgCr/g support from initial concentrations ranging to 1000 mg Cr/L. Also, chromium sorption was not significantly disturbed by competing sulfate anions present at 150 mg/L. Later, Yukoi et al. studied the functionalization of MCM-41 with coordinated Fe(III) as a selective adsorbent for Cr(VI), As(V), Mo(VI) and Se(VI) [114]. The authors reported a maximum uptake of 95 mgCr/g support for an initial concentration of 230 mg Cr/L. Yoshitake et al. used mono-, di- and tri-amino substituted MCM-41 and SBA-1 mesoporous silica to
remove chromate and arsenate [86]. The trimino-SBA-1 support achieved the highest uptake of chromate (28 mgCr/g support from a 317 mg Cr/L solution). Lam et al. obtained −NH2 and −COO−Na+ functionalized MCM-41 supports for the selective separation of dichromate and Cu(II) ions [57]. Once protonated at pH 2, NH2-MCM-41 was able to completely separate dichromate ions from Cu(II), achieving an uptake of 23 mgCr/g support (initial Cr(VI) concentration was 7 mg/L).

As previously mentioned APTES-functionalized zeolites present hydrophobicity and tend to form colloidal suspensions in aqueous media, hence easing coalescence of recovery. To overcome this limitation, Barquist and Larsen proposed the added magnetization of a APTES-modified Y zeolite for the specific removal of Cr(VI) [62]. The authors proposed the ion-exchange of Fe(III) and Fe(II) ions followed by alkali treatment in order to promote the formation of iron oxides. The functionalized supports were able to remove Cr(VI) up to 44 mgCr/g support (being the maximum initial Cr concentration of 2 mM) and the magnetization of the zeolite was attained. A different approach was presented by Chen et al. [115], with the preparation of magnetic siliceous MCM-41 for the selective removal of arsenic and chromium from solution. For the magnetization of the support, magnetic iron oxide nanoparticles (Fe3O4) were added to the reaction mixture for the synthesis of MCM-41, and the support presented magnetic properties straight after the synthesis. Subsequently, the support was functionalized with aminopropyltrimethoxysilane (APTMS) and exchanged with Fe3+ in order to acquire selectivity for oxyanions. The authors reported a maximum uptake of 91 mgCr/g support and chromate was selectively removed in the presence of Cu2+ ions. The comparison between the APTMS-magnetic MCM-41 and the APTMS-non-magnetic counterpart proved that the presence of the iron oxide nanoparticles did not interfere with the removal of chromate, as the uptakes were similar between the two supports. Moreover, recovery of the support with a magnet proved to be a much easier process than gravity sedimentation. While this support was able to achieve a higher uptake compared to the magnetic NaY support of Barquist and Larsen, it should be pointed out that Y zeolite has a higher amount of Al in its composition, which decreases the capacity for functionalization (authors reported a Si/Al ratio of 2); also, the magnetization of the support was conducted post-synthesis and the magnetic iron oxides were formed in situ.

### 3.1.3. Metal modified zeolites

The application of MMZs in arsenic and chromium oxyanion removal is a rather recent effort. While SMZs gained notoriety in the 1980s, MMZs were only applied in this field at the start of the XXIst century. The main obstacle for direct adsorption with MMZs is the efficiency of the retention of arsenate or chromate on the zeolite. This was very well illustrated in one of the early publications in the field, from Faghrihan and Bowman. It was interesting as these authors have a strong background in chromate removal with SMZ and investigated the potential of MMZs from natural clinoptilolite, exchanged with several mono-, di- and trivalent cations as inducers of surface precipitation of the corresponding insoluble chromates [116]. A correlation was found between the solubility product of the different chromates and the corresponding chromate uptake by the different MMZs. The maximum uptake was achieved with the Pb-loaded sample, which forms the least soluble chromate from the ions in study, at 29 mgCr/gzeolite. This was sequentially followed by the samples loaded with Ag+ (21 mgCr/gzeolite), Hg2+ (19 mgCr/gzeolite) and Ba2+ (18 mgCr/gzeolite). The authors concluded that the prepared MMZs allowed a much higher uptake of chromate than the comparatively obtained using SMZs. However, the authors also observed that, while very insoluble, minimal loss of the adsorbed lead or mercury chromates did occur, and the toxicity associated to these compounds render this approach undesirable for environmental remediation. A compromise between performance and environmental safety can be achieved when the less toxic barium ions are used in MMZs, and this was the choice of Bolotamir and Egashira in their study of Ba-exchanged natural zeolite, bearing mainly clinoptilolite and some mordenite phases [117]. The authors tested several zeolitic fractions with differing Ba2+ loading and achieved uptake figures ranging from 5 to 30 mgCr/gzeolite.

For the removal of arsenic oxyanions, metal ions such as aluminum, iron and manganese found particular success in the formulation of MMZs. Xu, Okhi and Maeda proposed Al-exchanged natural zeolite (Shirazu zeolite) for the removal of arsenate [118,119]. Depending on pH and the arsenate/sorbent ratio, uptakes values of 6 mgAs/gzeolite were achieved. Desorption of arsenate against NaOH was reported and the supports were re-used, showing some loss of arsenate removal performance (up to 30% loss), which was found to be easily compensated by re-generating

### Table 1

Overview of different SMZs and uptake capacities for Cr(VI) and As(III)/As(V).

<table>
<thead>
<tr>
<th>Selected zeolite</th>
<th>Selected surfactant</th>
<th>Maximum uptake observed (mg/g support)</th>
<th>Reference(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Natural clinoptilolite</td>
<td>HDTMA</td>
<td>Cr(VI): 13.22 As(III)/As(V): 3.7</td>
<td>[43,49,96,99–105]</td>
</tr>
<tr>
<td></td>
<td>CPB</td>
<td>Cr(VI): 3.5</td>
<td>[100,106]</td>
</tr>
<tr>
<td></td>
<td>PHMG</td>
<td>Cr(VI): 3.0</td>
<td>[97,107]</td>
</tr>
<tr>
<td></td>
<td>ODYMA</td>
<td>Cr(VI): 2.3</td>
<td>[9]</td>
</tr>
<tr>
<td>Natural mordenite</td>
<td>HDTMA</td>
<td>Cr(VI): 0.023</td>
<td>[96,108]</td>
</tr>
<tr>
<td></td>
<td>EHDDMA</td>
<td>Cr(VI): 0.017</td>
<td>[108]</td>
</tr>
<tr>
<td></td>
<td>CPB</td>
<td>Cr(VI): 6.9</td>
<td>[98]</td>
</tr>
<tr>
<td>Natural chabazite</td>
<td>CB</td>
<td>Cr(VI): 14.0</td>
<td>[106]</td>
</tr>
<tr>
<td>Natural laumontite-stilbite conjugate</td>
<td>HDTMA</td>
<td>Cr(VI): 8.83</td>
<td>[101]</td>
</tr>
<tr>
<td></td>
<td>EHDDMA</td>
<td>Cr(VI): 0.71</td>
<td>[109]</td>
</tr>
<tr>
<td>Natural natrolite</td>
<td>HMDA</td>
<td>Cr(VI): 2.3</td>
<td>[110]</td>
</tr>
<tr>
<td>Synthetic LTA</td>
<td>HDTMA</td>
<td>Cr(VI): 1.2</td>
<td>[87]</td>
</tr>
<tr>
<td>Synthetic FAU type</td>
<td>HDTMA</td>
<td>Cr(VI): 2.0</td>
<td>[7]</td>
</tr>
<tr>
<td></td>
<td>CPB</td>
<td>Cr(VI): 8.0</td>
<td>[98]</td>
</tr>
<tr>
<td>Synthetic LTA type</td>
<td>CB</td>
<td>Cr(VI): 6.0</td>
<td>[98]</td>
</tr>
<tr>
<td>ZSM-5 (MFI)</td>
<td>HDTMA</td>
<td>Cr(VI): 0.66</td>
<td>[100]</td>
</tr>
<tr>
<td></td>
<td>CPB</td>
<td>Cr(VI): 0.53</td>
<td></td>
</tr>
</tbody>
</table>

List of abbreviations: HDTMA, hexadecyltrimethylammonium; EHDDMA, ethylenehexadecyltrimethylammonium; ODYMA, octadecyltrimethylammonium; CPB, cetylpyridinium bromide; PHMG, polyhexamethylene guanidine; HMDA, hexamethyl-1,9-nonanediammonium.
the MMZ with fresh aluminum solution prior to a new arsenic removal cycle. The authors also tested the influence of competing anions such as phosphate, chromate, nitrate and others, and reported that only the chemically related phosphate ion had a significant interference in arsenate removal [120]. A recent publication by Šiljeg et al. reported an iron-modified natural clinoptilolite for the removal of As(III) and As(V) ions. The authors reported a detrimental influence of sulfate and phosphate ions on the overall efficiency, while desorption studies with HCl revealed that the retrieval of As(III) from the support was possible, but As(V) desorption was residual [121]. Previously, Payne and Abdel-Fattah reported that Fe(II)-modified natural chabazite, clinoptilolite and synthetic faujasite and LTA zeolite were able to remove As(III) and As(V) [122]. The individual and combined performance of iron and/or manganese MMZs for arsenate removal was well illustrated in a publication from Jiménez-Cedillo et al. By exchanging FeCl3 and MnCl2 in natural Mexican clinoptilolite, the authors prepared Fe-, Mn- and Fe–Mn-MMZs which were used in the removal of arsenate in concentrations up to 0.1 ppm [123]. Sorption equilibrium was quickly attained and the maximum uptake of 8 μgAs(III)/gzeolite was achieved with the Fe-MMZ sample with higher Fe(II) loading. For comparative metal loadings, it was found that the arsenate removal performance increased in the following sequence: Mn-MMZ, Mn–Fe-MMZ and Fe-MMZ.

In terms of application of metal oxide MMZs for oxyanion adsorption, the main focus between arsenic or chromium compounds appears to lie on the first. For the removal of chromate, the few publications available report the use of iron-coated zeolites (ICZ). Very recently, Du et al. reported the preparation of a ICZ from St. Cloud Mountain (USA) natural zeolite, which achieved a maximum uptake of 82 μgCr(III)/gzeolite from batch studies (initial Cr concentration of 50 mg/L maximum) [124]. Arsenic has received a slightly wider array of MMZs. An example of As(V) removal with natural clinoptilolite ICZ was reported by Jeon et al., whom also studied the effect of competing sulfate ions on the arsenic removal efficiency [125]. Starting from diluted As(V) solutions (2.0 mg/L), the predicted maximum adsorption capacity of the ICZ, through the Langmuir model, was 0.68 mgAs(V)/gzeolite, which decreased to 0.53 in the presence of sulfate (at 400 mg/L) in the same experimental conditions. This support was chosen for field application, on the decontamination of mine drainage containing arsenic, cadmium and lead [126].

Beside iron oxide, MMZs with alumina and MnO2 have also been reported on the removal of arsenate or arsenite, and in some cases bimetallic oxide MMZs have been employed. Fe and Al natural clinoptilolite MMZ was reported to adsorb As(V) by Simsek et al. [66]. The authors claim that the maximum uptake of 5.86 mgAs/gzeolite was achieved with the bimetallic Fe/Al oxide MMZ, from a solution of 9.5 mgAs/L. A MnO2–natural clinoptilolite MMZ was recently evaluated for arsenate removal from dilute solutions by Camacho, Parra and Deng. The authors evaluated pH and As(V) feed concentrations and concluded that the inclusion of MnO2 on the zeolite increased its sorption capacity and heterogeneity, as contrasted by the poor fitting obtained by the Langmuir model on the MnO2–MMZ, compared to parent zeolite [127]. Maximum uptake was observed for a 50 μgAs/L solution, at 2.5 μgAs/gzeolite. The growing interest in zeolite synthesis from fly ash has also resulted in the application of an alumina-loaded cancrinite-type zeolite on the treatment of As(V) by Qiu and Zheng [128]. The authors reported a maximum uptake of 34.5 mgAs/V/L with this support. Table 2 presents an overview on the achievements of different SMZs in the adsorption of Cr(VI) or As(V).

<table>
<thead>
<tr>
<th>Modification</th>
<th>Selected zeolite</th>
<th>Metal modification compounds</th>
<th>Maximum uptake observed (mg/gzeolite)</th>
<th>Reference(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ion exchange</td>
<td>Fe(III)</td>
<td>0.082</td>
<td></td>
<td>[124]</td>
</tr>
<tr>
<td>Atomic layer</td>
<td>Fe oxide</td>
<td>0.69</td>
<td></td>
<td>[125]</td>
</tr>
<tr>
<td>Metal oxide formation</td>
<td>Fe oxide</td>
<td>3.87</td>
<td></td>
<td>[66]</td>
</tr>
<tr>
<td></td>
<td>Al oxide</td>
<td>2.14</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Fe and Al oxide</td>
<td>5.11</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Mn oxide</td>
<td>0.034</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Fly ash cancrinite</td>
<td>Al oxide</td>
<td>34.5</td>
<td></td>
</tr>
</tbody>
</table>

### 3.2. Redox processes

#### 3.2.1. Metal modified zeolites

The modification of zeolites with metal ions that are redox agents for Cr(VI) was reported. Some known reducing agents for Cr(VI) also offer intrinsic capacity for being ion-exchanged in a zeolitic structure. That is the case of Fe(II) ions, which have been used on the modification of natural and synthetic zeolites to promote chromate reduction and removal. Kiser and Manning resorted to a Fe(II)-modified synthetic NaY on the treatment of Cr(VI) solutions and observed that the reduced Cr(III) species, being susceptible for ion-exchange, readily displace additional Fe(II) from the zeolite exchange sites. This cooperative mechanism between Cr(III)/Fe(II) results in cation exchange with Fe(II), hence assisting additional Cr(VI) reduction. Furthermore, the authors commented that the presence of the Al–Si–O and SiO2 surfaces in the zeolite structure may assist the reaction of Cr(VI) with Fe(II) by providing...
a catalytic effect [130]. The reported maximum chromium uptake was 16 mg Cr/g for zeolite. Other applications of Fe(II) MMZs for the treatment of chromate were reported by Huang and Zhang [131] and very recently by Ly et al. [132].

Some interesting variations to the former examples are also reported. The most directly related was presented by Lee et al. The authors started from a similarly prepared Fe(II) natural clinoptilolite (Pohang, Korea), but promoted reduction of the iron ions to the zerovalent state with sodium borohydride [133]. The authors used this support for the simultaneous treatment of diluted Cd(II) and Cr(VI) solutions. This support demonstrated reduction capacity for Cr(VI) while retaining the ion-exchange capacity for cadmium, although sorption equilibrium times differed significantly between the two metals. A more elaborated alternative has been proposed by Li et al., whom combined zerovalent iron particles with natural clinoptilolite via an hydrogel prepared with sodium silicate and sodium aluminate, which is cured, homogenized and later treated with HDTMA to form a combined iron MMZ-SMZ for the reduction and sorption of chromate [89]. The combined reduction/sorption of Cr(VI) resulted in an uptake of 780 μg Cr/g for the iron MMZ-SMZ, compared to 520 μg Cr/g for the equivalent SMZ obtained in the absence of zerovalent iron. The same system was successfully employed to the packing of columns for open-flow treatment of Cr(VI) solutions [88].

In terms of As(III) oxidation, few works report the use of zeolitic supports. A work from Anielak et al. did state that Mn-modified clinoptilolite was able to oxidize As(III) to As(V) [70]. Some other works report that MnO2 is reduced to Mn(II) by As(III), and this oxide has been successfully used to coat minerals such as sand or quartz [68]. Therefore, it is expectable that MnO2 MMZs could also be of use on As(III) oxidation and treatment.

3.2.2. Photocatalytic redox processes

A recent publication by Shao et al. reported the photocatalytic reduction of Cr(VI) to Cr(III) by ZSM-5 zeolite, compared to ZnO and in the presence or absence of oxalate [134]. The authors chose oxalate as a model for organic compounds in the environment. ZSM-5 zeolite showed pH-dependent photocatalytic activity in the reduction of Cr(VI), albeit being outperformed by ZnO in the same experimental conditions.

An interesting approach has been previously described by Kanthasamy and Larsen, whom prepared ZSM-5 zeolite hollow tubular structures doped with Fe(III) nitrate for the reduction of Cr(VI) [135]. The zeolitic tubes were synthesized from mesoporous silica, which acted both as a structural template and reagent, since it is consumed during the synthesis. Doping with Fe(III) was conducted in two distinct procedures: during synthesis, which leads to the encapsulation of Fe(III) ions within the hollow structure; or post-synthetically by ion-exchange. The dispersion of iron achieved with the first process was found to be greater, and ion-exchange promoted aggregation of iron particles on the surface, giving rise to subsequent leaching, which led to a decrease in activity of the ion-exchanged catalyst.

3.2.3. Zeolites as supports for bioremediation of Cr and As oxyanions

Hexavalent chromium reduction by biosorbents with subsequent removal of the formed Cr(III) ions by zeolites has been reported. The concept of combining bacterial capacity for biosorption of heavy metals with a zeolitic support was originally proposed by Neves and Tavares. The Arthrobacter viscous bacterium supported on zeolites was extensively used for the treatment of Cr(VI) solutions, and several zeolites were successfully employed as support (natural clinoptilolite or synthetic FAU, MFI, MOR and BEA zeolites) [24,40,136–143]. This bacterium is able to form a biofilm on the zeolitic surface, as reported by Lameiras et al. The authors found that the removal percentages obtained for the biofilm-laden zeolite sorbent are similar than those previously obtained using instead granular activated carbon (GAC) as support [144]. Typical uptake values for initial Cr(VI) concentration up to 100 mg/L range from 5 to 10 mg Cr/g zeolite to 17.0 mg Cr/g zeolite in later reports, usually achieved with synthetic FAU zeolite and with higher biomass concentration [143]. This approach has already reached the stage of pilot-scale testing [145]. An Escherichia coli biofilm supported on NaY zeolite was also evaluated. The results obtained using this bacterium are similar to the obtained using Arthrobacter viscosus on identical conditions [24,146]. For an initial chromium concentration of 100 mg/L, the uptake values found were around 4 mg/g zeolite for both bacteria. E. coli was also evaluated on natural clinoptilolite by Erdoğan et al., as well as other bacterial strains (B. subtilis, S. aureus, S. epidermidis and P. aeruginosa). Based on the fitting of the Langmuir adsorption model, the highest maximum uptake was achieved by the B. subtilis–clinoptilolite system, with 2.44 mg Cr/g required [147].

Arsenate oxidation with bacteria supported on chabazite was investigated by Liévermont et al. [83]. The authors tested a bacterial strain isolated from an arsenic-contaminated site, later identified as a β-Proteobacteria, in combination with some minerals (the referred chabazite zeolite, quartz sand and kutchahorite mineral) and concluded that the zeolite or quartz sand could be used in a first instance to support the biooxidation of As(III) on large-scale remediation of arsenic contamination. An indirect study reporting bio-oxidation of organic As(III) compounds on zeolitic supports was presented by Matusch et al. The authors evaluated the decomposition of arsenobetaine, which is formed mainly by metabolism of arsenate in marine organisms such as fish, mollusks and crustaceans. While studying the decomposition of arsenobetaine with natural and synthetic clinoptilolite, the authors noticed the formation of degradation products from carbon-arsenic bond cleavage pathways using natural clinoptilolite, while the synthetic counterpart only revealed sorption properties. The authors traced back the decomposition to bacterial contamination of the natural clinoptilolite samples and referred possible presence of Leptospirillum ferrooxidans and Thiothrix thiooxidans, while other species would be likely present. Although the purpose of this work was different, it showed the possibility of zeolite-bacteria systems to be also exploited on arsenic removal from organoarsenic compounds [148].

4. Conclusion

As it was presented in this paper, the versatility of zeolites is so ample that it offers interesting options to overcome their own natural limitation for anions, by conducting chemical modification or by tailoring of the matrix. The comparison between chromium and arsenic oxyanions also brings to light some interesting subtle physicochemical differences concerning their interaction with the environment and for the development of treatment/remediation procedures. Their differences are also highlighted through the interaction of Cr(VI), As(III) and As(V) with zeolites. To this respect, designing zeolite-based technologies for the treatment of each of these compounds further stresses the versatility of these materials.

As adsorption processes are concerned, surfactant modification of the zeolite surface proved to be a straightforward process to obtain efficient supports, whereas surface functionalization, despite being a slightly more elaborated methodology, allows for considerable higher uptake values (notably for Cr). These techniques are somewhat complementary in terms of achieving the same purpose from different zeolites, as SMZ require Al-rich zeolites while Al-poor zeolites are preferred for surface functionalization. Despite the special requirements of surface
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