Interaction of gypsum with As(V)-bearing aqueous solutions: Surface precipitation of guerinite, sainfeldite, and Ca$_2$NaH(AsO$_4$)$_2$·6H$_2$O, a synthetic arsenate

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ABSTRACT

The interaction of arsenate-bearing aqueous solutions with gypsum at a starting pH of 9 and 25 °C results in surface precipitation of guerinite, Ca$_2$(HAsO$_4$)$_2$(AsO$_4$)$_2$·9H$_2$O, sainfeldite, Ca$_2$(HAsO$_4$)$_2$(AsO$_4$)$_2$·4H$_2$O, and occasionally Ca$_2$NaH(AsO$_4$)$_2$(AsO$_4$)$_2$·6H$_2$O, a new arsenate. These three solid phases are characterized by the simultaneous presence of HAsO$_4^{2–}$ and AsO$_4^{3–}$ groups in their structure, which is explainable since crystallization occurs within a pH range in which both HAsO$_4^{2–}$ and AsO$_4^{3–}$ are available in the aqueous solution. The interaction leads to a decrease in the As(V) concentration in the aqueous phase to reach values controlled by the solubility of these solid phases. The study combines several macroscopic experiments, in which changes in the solution chemistry are monitored as a function of time, with the characterization of solid phases by SEM-EDS and XRD. The crystal morphologies of the precipitating phases are interpreted on the basis of their respective structures. The thermodynamic solubility products of both guerinite and the new arsenate have been determined, being 10$^{-31.17}$±0.03 and 10$^{-13.83}$±0.03, respectively. The reaction paths followed by the system and the equilibrium endpoints have been modeled using the geochemical code PHREEQC.

Keywords: Crystal growth, calcium arsenate, analysis chemical, sainfeldite, phase equilibrium, guerinite, gypsum

INTRODUCTION

Although the mobility and toxicity of As in the environment have been studied extensively (Smedley and Kinniburgh 2002; Vaughan 2006), numerous issues remain unresolved. On the one hand, it is well established that under oxidizing conditions, As(V) is the most common form of As in natural waters, with AsO$_4^{3–}$, HAsO$_4^{2–}$, H$_2$AsO$_4^{–}$, and H$_3$AsO$_4$ being the prevailing aqueous species in different pH ranges. On the other hand, the crystal chemistry and thermodynamic properties of arsenates remain largely unknown. Moreover, with some significant exceptions (Bothe and Brown 1999a; Lee and Nriagu 2007), the determination of thermodynamic solubility products of arsenates has received little attention, perhaps as a consequence of the complicated crystallization behavior of these compounds in aqueous environments, which involves formation of an enormous diversity of hydrates, double salts, the presence of arsenate groups with different protonation degrees, etc. This lack of data is an important handicap because an in-depth study of arsenate mobility in natural systems requires a precise knowledge of the solid phases that may or may not precipitate their crystal chemistry and their solubility.

Immobilization of arsenate in the environment can occur by precipitation of low-solubility salts or by sorption on mineral surfaces in soils, sediments, and aquifers. Artificial remedia-
for 11 of these compounds. The most complete research on the crystal chemistry of these arsenates was carried out during the seventies by Ferraris and co-workers, who studied the structures of CaHAsO₄ (weilite), CaHAsO₄·2H₂O (pharmacolite), CaHAsO₄·H₂O (haideringite), Ca₃(AsO₄)₂(SO₄)·9H₂O (guerinite and ferrarisite), Ca₃(HAsO₄)₂(AsO₄)·4H₂O (sainfeldite), and other arsenates and H arsenates of Ca (Ferraris et al. 1971, 1972; Ferraris and Chiari 1970; Catti and Ferraris 1974; Catti et al. 1980; Ferraris and Abbona 1972). Bothe and Brown (1999a) identified and determined the solubility products of several arsenates of low solubility, including Ca₃(AsO₄)₂OH, 4H₂O, Ca₃(AsO₄)₂·2H₂O, and Ca₃(AsO₄)₆·2H₂O. Unfortunately, these insoluble Ca arsenates crystallize at high pH values. At a pH < 8, there is a tendency to form more-soluble, hydrated acidic arsenates, which are unlikely to produce aqueous solutions with As concentrations below the guideline values proposed for As dissolved in potable water (Magalhães 2002). Moreover, there is also strong evidence that Ca-arsenate compounds of low solubility are not stable in the presence of air at a pH > 8 (Robins 1981; Nishimura et al. 1983). Carbon dioxide in air will convert the Ca arsenate to Ca carbonate with the subsequent release of As back into the aqueous phase.

In addition to Ca arsenates, several metal-arsenates and mineral-like arsenates (e.g., Jiménez et al. 2004; Twidwell et al. 1999 and references therein) have been proposed as possible immobilizing phases, but the solubility and the precise identity and structure of these phases is in many cases not known. The obvious conclusion is that research on the structure, solubility, and crystallization behavior of new arsenate-containing materials has to be conducted. Roman-Ross and co-workers (Roman-Ross et al. 2003; Fernández-Martinez et al. 2006) have recently studied the precipitation of gypsum by mixing CaCl₂ and Na₂SO₄ aqueous solutions in the presence of dissolved Na₃HAsO₄, and have detected the incorporation of some arsenate (substituting for sulfate) into the gypsum structure. This incorporation supposes an increase of the unit-cell parameters, as AsO₄³⁻ (or HAsO₄⁻) is greater than SO₄²⁻. In conclusion, these authors suggested that As(V) could be sorbed onto gypsum by forming a limited Ca-arsenate-sulfate solid solution. In fact, surface precipitation of pollutant-bearing solid solutions has been found to be a significant, long-term mechanism of sorption not only of cations but also of oxyanions (Prieto et al. 2002; Andara et al. 2005; Fernández-González et al. 2006) on carbonates and sulfates. The existence of a certain connection between arsenate and gypsum has also been documented by Juillot et al. (1999), who, in studying the mobilization of As from buried wastes at an industrial site, observed an intimate association of gypsum and Ca arsenates, detecting that gypsum may contain some As and that Ca arsenates may contain some S. However, to our knowledge there are no studies in the literature quantifying the uptake behavior of arsenate on gypsum.

In this work, we have studied the interaction of Na₃HAsO₄ aqueous solutions with gypsum, to check the effectiveness of this mineral in removing As from water within a moderately basic pH range (< 9). To this aim, we have combined several macroscopic techniques, in which changes in the solution chemistry have been monitored as a function of time, with microscopic observations and X-ray diffraction (XRD). We find that the formation of arsenate-bearing gypsum (with arsenate substituting for sulfate in the structure) has a negligible effect on the solution composition, with the removal of As mostly occurring by surface precipitation of guerinite, sainfeldite, and a new arsenate, Ca₃Na₃(HAsO₄)(AsO₄)·6H₂O. Although gypsum does not seem to be a suitable remediation tool at pH < 9, the nature, solubility, and crystallization behavior of the solid phases that precipitate in this pH range need to be investigated. In this framework, our study proposes an estimation of the solubility product at 25 °C and 1 atm for both guerinite and the new arsenate, as well as a model for the reaction paths followed by the system.

**EXPERIMENTAL METHODS**

**Interaction experiments**

Interaction experiments were carried out at 25 ± 0.1 °C and at atmospheric pressure by reacting gypsum grains with Na₃HAsO₄ aqueous solutions in thermostated and continuously stirred (100 rpm) polypropylene vessels. In most of the experiments, 2 g of mineral grains with diameters ranging from 1.0 to 1.5 mm were added to 100 cm³ of reacting solution. Moreover, to check the grain size effect, one set of experiments was performed with solids ranging from 0.5 to 1.0 mm in diameter. Gypsum grains were obtained by crushing natural single crystals. The resultant fragments were sieved to the selected size range and ultrasonically cleaned in an ethanol bath. This starting material was confirmed to be gypsum by X-ray diffraction (XRD, Philips-X’Per-PRO) and analyzed by X-ray fluorescence spectrometry (XRF, Philips-PW2404). XRF analyses yielded less than 0.2 wt% foreign elements, with Na, Sr, and P as major impurities. The grains are essentially cleavage fragments dominated by (010) crystal faces, with a specific N₂-BET (Micrometrics-ASAP-2010) surface area of 0.16 m²/g (0.22 m²/g for the grains with diameters in the range 0.5–1.0 mm). The use of cleavage fragments with smooth surfaces within this size range minimized the significance of adsorption in comparison with surface precipitation.

For each starting solution (100, 50, 25, 20, 10, 5, and 2 mM of arsenate), a set of experiments with increasing reaction times was carried out. The experiments were performed by placing 100 cm³ (weighed to 0.001 g) of solution and 2.000 ± 0.005 g of gypsum grains in the reacting vessel. The starting solutions were prepared with reagent grade Na₃HAsO₄·7H₂O (Merek) and deionized water (MilliQ System) at ambient CO₂ partial pressure. The vessels were then closed with polypropylene caps to avoid contact with the atmosphere and minimize evaporation during the experiments. After the corresponding reaction period, the experiment was stopped and the solution was assayed for dissolved Ca, sodium, sulfate, As, and pH. Concentrations and pH evolution were monitored in this way for 50 days. Each experiment was repeated three times. Thus, the reported results are the average value of three analogous experiments. The solution pH was measured with a combination electrode (Ross-Thermo-Orion-810200) and a digital pH-meter (CyberScan-pH-2100). Multipoint calibration of the electrode was performed with 4.01, 7.00, and 10.01 NIST-traceable buffers (Thermo-Orion). Dissolved As was determined by Inductively Coupled Plasma Optical Emission Spectroscopy (ICP-OES, Perkin-Elmer-Optima-3300-DV). Calcium and Na were measured by Atomic Absorption Spectrometry (AAS, Pye-Unicam-SP-90); dissolved sulfate was analyzed by ion chromatography (IC, Metrohm Compact IC 761). The results are reproducible, particularly in the case of As. The variation in As concentration from replicate experiments at the same reaction time was always less than ±4% (RSD). For Ca, Na, and sulfate this variation was found to be within ±8% (RSD), ±6%, and ±5%, respectively. Speciation and aqueous solution modeling was carried out using the geochemical code PHREEQC (Parkhurst and Appelio 1999), a geochemical modeling program based on the ion-association aqueous model with a diversity of capabilities.

**Characterization of the solid phases**

After the experimental runs, representative individuals of the gypsum crystals were selected from the aqueous solution to verify the incorporation of As-bearing phases on their surfaces. For this purpose, the samples (the gypsum grains with the corresponding overgrowth) were carbon-coated and then examined with a scanning electron microscope (SEM, JEOL-JSM-6100). This SEM was also used to estimate the composition of the precipitates using an INCA Energy 200 microanalysis system (EDS) with a silicon detector (PentaFET, Oxford Instruments) fitted
with an ultra-thin window that allows the detection of O. SEM-observations were combined with an XRD study (Philips-X-Pert-PRO) to complete the characterization of the precipitate layer. To avoid the inclusion of peaks from the substrate, the diffraction patterns were obtained by carrying out 20 scans (from 3 to 45°) at a fixed small angle of incidence (3 or 4° depending on the sample) on the substrate surface. The diffractograms obtained in this way involve essentially the precipitate overgrowth and only in some cases include the most intense peaks corresponding to the underlying gypsum.

Starting from a 100 mM As(V) parent solution, some tiny crystals of an unknown phase appeared in the early stages of the experiment (see the section of results). To obtain samples of this phase suitable for characterization by single-crystal XRD, specific crystallization experiments were carried out by mixing CaCl₂ (0.01 mM) and Na₂HAsO₄ (1 M) aqueous solutions in a thermostated (25 ± 0.1 °C) polypropylene vessel. 50 cm² of each parent solution were placed in the reacting vessel, which was then sealed with a polypropylene cap to avoid evaporation. Following mixing, the solution pH reached a value of ~8.90. After several days of growth, the resultant crystals (~100 µm) were extracted and studied by single-crystal XRD. The structure was solved by Patterson methods, using the program DIRDIF-99 (Beurskens et al. 1992). Experimental details on the crystal-structure determination of this new phase can be seen in Jiménez et al. (2006). Moreover, to determine the solubility of this unknown phase, the evolution of both solution pH and composition was monitored over 1800 h.

**RESULTS**

**Uptake of As, release of Ca, and pH evolution**

Figure 1 shows the evolution of As and Ca concentrations as a function of time for the experimental series carried out with a 25 mM parent solution and grain sizes in the range 1–1.5 mm. As can be observed, the As concentration decreases quickly during the first 24 h. This decrease goes with an increase in the Ca and sulfate concentration and a reduction of pH (see inset in Fig. 1). The result is similar in the experiments carried out starting from 100, 50, 20, and 10 mM Na₂HAsO₄ parent solutions, in which two consecutive stages were observed: the As concentration falls rapidly in the early hours of the experiment and then evolves slowly toward an asymptotic value (As∞) mea-

**TABLE 1. Uptake of As (asymptotic values)**

<table>
<thead>
<tr>
<th>Asᵢ</th>
<th>As∞</th>
<th>U∞</th>
<th>U/Sᵦₜₑᵗ</th>
<th>U/Sᵦₚ₆</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>100</td>
<td>62.5</td>
<td>1.88</td>
<td>11.8</td>
<td>89.5</td>
<td></td>
</tr>
<tr>
<td>50</td>
<td>31.2</td>
<td>0.94</td>
<td>5.88</td>
<td>44.5</td>
<td></td>
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<tr>
<td>25</td>
<td>16.2</td>
<td>0.44</td>
<td>2.75</td>
<td>21.0</td>
<td></td>
</tr>
<tr>
<td>25(small size)</td>
<td>16.1</td>
<td>0.45</td>
<td>2.05</td>
<td>12.9</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>13.9</td>
<td>0.31</td>
<td>1.94</td>
<td>14.8</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>9.4</td>
<td>0.03</td>
<td>0.19</td>
<td>1.43</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>5</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
course of the interaction with the gypsum crystals, the solution pH slowly decreases toward an asymptotic value, following a trend parallel to the decrease of the As concentration (see Fig. 1). Table 2 shows the initial solution pH (pH0) and the value measured at the end of the experiments (pH∞). For the experiments carried out with 100, 50, 25, 20, and 10 mM solutions, the pH decreases during the first 24 h from an initial value of ~9.0 to a value <7.5, approaching a lower asymptotic value for prolonged reaction times. Starting from less-concentrated solutions (5 and 2 mM), the pH decreases a few tenths in the beginning of the experiment, then remains virtually constant.

The increase in Ca concentration is obviously a consequence of the gypsum dissolution, which is common in all the experiments. At the beginning of every experiment, the aqueous solution is free of Ca, but the concentration of this element increases rapidly during the first ~24 h and rise slowly afterward, approaching an asymptotic value for prolonged reaction times (see Ca∞ in Table 2). The concentration of sulfate (S∞ in Table 2) increases in a similar way, but it is always higher than that of Ca. As noted below, this feature is due to the fact that a part of the released Ca is consumed during the process by precipitation of Ca-bearing solid phases. In contrast, the concentration of Na remains virtually constant (deviations are in the rank of the experimental uncertainties) during the experiments. Only in the case of the experiments carried out in 100 mM parent solution, the isolated aggregates grow more slowly, and cover less than 20% of the surface after prolonged reaction times. In this case, a detailed observation of the precipitate already shows the existence of a transitory but very reproducible decrease in As concentration during the first 6–8 h of reaction. The data-points represent the mean of three replicate experiments and clearly confirm the existence of a transitory but very reproducible decrease in As concentrations.

**Microscopic characterization of the solids**

Figure 4a shows the surface of a typical gypsum grain after 18 h of reaction with a 50 mM parent solution. The surface shows dissolution signs and is covered by a precipitate consisting of aggregates of small (~30 µm) crystals with laminar morphologies (Fig. 4b). The precipitate begins to grow at the cleavage steps of the grain perimeters and extends to quickly cover the (010) gypsum surface, which becomes completely covered by a crust of precipitate after 24 h of reaction. The precipitate features are similar in the experimental series carried out with a 25 mM parent solution, but in this case the nucleation density is lesser and the crystallography of the highest order is observed. This observation is in good agreement with the presence of the main reflections of guerinite (at d ~14.1, 3.90, 2.91 Å, etc.) in the diffractograms.

Guerinite is a Ca arsenate in which unprotonated AsO4 3− and monoprotonated arsenate HAsO4 2− groups coexist to form a nonahydrate, Ca4(HAsO4)4(AsO4)2 ·9H2O, with monoclinic, P21/n, structure. The crystal structure of guerinite is characterized by (T01) layers built up by [AsO4] tetrahedra linked to Ca polyhedra (N.C. 7 or 8) by sharing vertices and edges (Catti and Ferraris 1974). An H-bonding system connects these (T01) layers between them, both the water H atoms and the anionic H atoms being involved in the H bridges. Two of the five independent Ca polyhedra share faces and form chains along the [010] direction. These structural features are consistent with the morphological characteristics observed in the precipitated guerinite crystals. Figure 5 shows the typical morphology, which consists of thin lamellar sheets elongated on the a axis. These sheets correspond to {T01} pinacoids, defined by the crystallographic planes with the highest d-spacing (14.1 Å).

Using a 20 mM parent solution, the isolated aggregates grow and cover less than 20% of the surface after prolonged reaction times. In this case, a detailed observation of the precipitate al-

### Table 2. Concentration of Ca, sulfate and pH level after prolonged reaction times

<table>
<thead>
<tr>
<th>Parent solution</th>
<th>Ca∞ (mM)</th>
<th>S∞ (mM)</th>
<th>pH0</th>
<th>pH∞</th>
</tr>
</thead>
<tbody>
<tr>
<td>100</td>
<td>15.5</td>
<td>62.3</td>
<td>8.9</td>
<td>7.0</td>
</tr>
<tr>
<td>50</td>
<td>16.4</td>
<td>40.0</td>
<td>8.9</td>
<td>7.1</td>
</tr>
<tr>
<td>25</td>
<td>17.9</td>
<td>29.1</td>
<td>9.0</td>
<td>7.2</td>
</tr>
<tr>
<td>25(small size)</td>
<td>17.8</td>
<td>28.9</td>
<td>9.0</td>
<td>7.2</td>
</tr>
<tr>
<td>20</td>
<td>18.5</td>
<td>25.7</td>
<td>9.0</td>
<td>7.2</td>
</tr>
<tr>
<td>10</td>
<td>18.6</td>
<td>19.3</td>
<td>9.0</td>
<td>7.5</td>
</tr>
<tr>
<td>5</td>
<td>16.1</td>
<td>16.2</td>
<td>9.0</td>
<td>8.7</td>
</tr>
<tr>
<td>2</td>
<td>15.3</td>
<td>15.2</td>
<td>8.9</td>
<td>8.8</td>
</tr>
</tbody>
</table>
discerning between two different kinds of crystals, G and S (Fig. 6), with laminar and prismatic shapes, respectively. The laminar crystals are analogous (morphology and EDS analyses) to those shown in Figures 4b and 5, which were identified as guerinite. This analogy is in agreement with the presence of the main reflections of guerinite in the diffractograms, which firmly supports the identification of the crystals labeled G as this mineral. The crystals labeled S show an elongated shape with well-defined crystal faces. EDS analyses yield Ca/As atomic ratios similar to that observed in the laminar crystals identified as guerinite. However, both O/Ca and O/As atomic ratios were systematically determined to be significantly lower than that of guerinite, within the unavoidable experimental error. Both ratios point to the sainfeldite stoichiometry, \( \text{Ca}_5(\text{HAsO}_4)_2(\text{AsO}_4)_2\cdot4\text{H}_2\text{O} \), which is in agreement with the presence of the main reflections of this mineral (at \( d \AA \approx 8.7, 4.67, 3.88, \) and \( 3.18 \)) in the diffractograms.

In a similar way to guerinite, AsO\(_4\)\(^3–\) and HAsO\(_4\)\(^2–\) groups coexist in sainfeldite to form a Ca-arsenate hydrate with monoclinic (\( C2/c \)) structure (Ferraris and Abbona 1972). The arsenate groups are linked to irregular [CaO\(_6\)] octahedra to form a three-dimensional framework with channels along [001], where the water molecules, which are Ca-coordinated, are located. These molecules, as well as the acidic H atoms of the HAsO\(_4\)\(^2–\) groups, are involved in H bonds that reinforce the structure, especially along [001]. Moreover, different periodic bond chains (PBCs) involving As-O-Ca links can be discerned within the polyhedral framework, of which those along [001] seem to be particularly strong. An in-depth study of PBCs is, however, beyond the scope of the present work.

Figure 6 illustrates the typical morphology of these crystals, which consists of two four-faced prisms, \{11\(\bar{1}\)\} and \{110\}, and the front \{100\} pinacoid. These three forms correspond to the crystallographic planes with the highest \( d \)-spacings (6.78, 8.69, and 9.32 \( \AA \), respectively) and were expected to be among the most important growth forms of sainfeldite. In fact, the crystal habit in Figure 6c has been simulated using the Donnay-Harker (D-H) tool of the computer code SHAPE (V7.1.2, Shape Software 2004), according to the following equation (Dowty 1980):

\[
RG_{hkl} = d_{hkl} \exp(-kd_{hkl})
\]

where \( k \) is a semiempirical constant that depends on a set of different crystal-growth parameters, including surface energy and supersaturation. \( RG_{hkl} \) is the linear growth rate of the \((hkl)\) crystal.
several trial-and-error runs using different initial concentrations. The crystallization conditions were optimized after gypsum grains simply by mixing CaCl$_2$ solutions. The crystallization conditions were optimized after gypsum grains simply by mixing CaCl$_2$ solutions. The crystallization experiment has been performed (see experimental section). The idea was to reproduce the conditions at the early stages of the 100 mM interaction experiment, i.e., a basic pH (around 9), a very high concentration of Na$_2$HAsO$_4$, and a low concentration of Ca. To avoid the subsequent evolution of the system, the experiment was performed without the presence of gypsum grains simply by mixing CaCl$_2$ and Na$_2$HAsO$_4$ aqueous solutions. The crystallization conditions were optimized after several trial-and-error runs using different initial concentrations.

After a while, a precipitate of microcrystals began to develop and was left to grow over a period of 10 days. The resultant crystals exhibit tabular shapes (~150 µm) with well-defined polygonal contours and unknown powder diffraction patterns. Single-crystal XRD indicates that this phase is Ca$_2$Na(HAsO$_4$)(AsO$_4$)$_2$·6H$_2$O (Jiménez et al. 2006). Moreover, EDS-analyses of several suitable crystals (over 10 µm thick) indicate the presence of Na, Ca, As, and O. These crystals exhibit the same habit and essentially the same composition as those observed in the interaction experiments. Therefore, it is reasonable to assume that both crystal families correspond to the same phase. The mean Ca/As and Na/As atomic ratios are, respectively, close to 1 and 0.5. Finally, within the unavoidable experimental error, the O/As ratio has a mean value around 7. These mean ratios point to a hydrated double-arsenate of Ca and Na, Ca$_2$Na(HAsO$_4$)(AsO$_4$)$_2$·6H$_2$O, which is in agreement with the data obtained from the structural determination (Jiménez et al. 2006). The coexistence of acidic HAsO$_4^-$ and plain arsenate AsO$_4^{3-}$ groups is analogous to that observed in guerinite and sainfeldite, and is explainable if one considers that crystallization occurs within a pH range in which both HAsO$_4^-$ and AsO$_4^{3-}$ ions are available in the aqueous solution. As the crystal structure of this new arsenate has not yet been described in a detailed way, we include here a description of the different types of polyhedra, their links to neighboring polyhedra, and other structural peculiarities. The crystal morphology is interpreted on this basis.

Structure and morphology of Ca$_2$Na(HAsO$_4$)(AsO$_4$)$_2$·6H$_2$O, a new arsenate

This new compound crystallizes in the triclinic space group $PT$, with $a = 6.680$ Å, $b = 8.223$ Å, $c = 12.537$ Å, $\alpha = 73.46^\circ$, $\beta = 78.89^\circ$, and $\gamma = 87.47^\circ$, and two formula units per unit cell (Jiménez et al. 2006). Selected interatomic distances and bond angles derived for this phase are shown in Tables 3 and 4. Figure 7 shows a projection of the structure onto (010). The structure is formed by layers parallel to (001) consisting of [AsO$_4$] tetrahedra and [CaO$_6$] polyhedra. These layers are linked to each other by

![Figure 6](image-url) Surface of a typical gypsum grain after several days of reaction with a 20 mM Na$_2$HAsO$_4$ parent solution. (a) Guerinite (G) and sainfeldite (S) crystals on the gypsum surface. (b) Detail at a higher magnification. (c) Habit of a sainfeldite crystal simulated according to Equation 3.
Ca-O distances within these octahedra show significant differences among them (Table 3) and the O-Ca-O angles (Table 4) differ significantly from 90° and 180°. These features confirm that both [CaO₆] octahedra are quite irregular, [CaO₆] being more distorted. Figure 8 shows the connection of these [CaO₆] octahedra with adjacent polyhedra in the structure. As can be observed, two [CaO₆] octahedra are edge-sharing through the O atoms O¹⁰ and O¹¹ no. 4. In the same way, two [CaO₆] octahedra are edge-sharing via O¹⁰ no. 3 and O¹¹ no. 5. The O⁷ atom is simultaneously shared by [AsO₄]³⁻, [CaO₆] and [CaO₆]. Similarly, O⁷ is shared by two octahedra (centered in Ca³⁺ and Ca⁶⁺) and one neighboring As²⁺ tetrahedron. Moreover, O¹² links [CaO₆] with an adjacent [AsO₄]³⁻ tetrahedron and O¹³ links [CaO₆] with an adjacent [AsO₄]³⁻ tetrahedron. Finally, the O atom furthest from Ca in each octahedron, O¹⁷ in [CaO₆] and O¹⁷ in [CaO₆], belongs to a water molecule.

Sodium atoms are surrounded by six O atoms to form [NaO₆] octahedra, which connect the layers formed by [AsO₄]³⁻ and [CaO₆] polyhedra. A scrutiny of Na-O distances and O-Na-O angles (Tables 3 and 4) reveals that these octahedra are closer to a regular octahedron than those coordinating the Ca atoms. The O atoms farthest from the central Na (O¹⁶ and O¹⁷) connect two [NaO₆] octahedra, which thus form edge-shared chains along [100]. Five of the six O atoms surrounding each Na belong to H₂O molecules (O¹⁵, O¹⁶, O¹⁷, O¹⁸ no. 6, and O¹⁹ no. 7 in Table 3). The remaining O, O⁰, is shared with one [AsO₄]³⁻ tetrahedron, thereby connecting the [NaO₆] chains with the (001) layers formed by [CaO₆] and [AsO₄] polyhedra.

Figure 9 shows a representation of some typical morphologies exhibited by the Ca₂NaH(AsO₄)₂·6H₂O crystals. The crystal faces have been indexed according to the unit-cell parameters.

### Table 3. Atomic distances (Å) for Ca₃NaH(AsO₄)₂·6H₂O

<table>
<thead>
<tr>
<th>AsO₄ tetrahedra</th>
<th>Calcium octahedra</th>
<th>Sodium octahedra</th>
<th>Water molecules</th>
</tr>
</thead>
<tbody>
<tr>
<td>As₁-O₁ = 1.715</td>
<td>Ca₁-O₁ = 2.298</td>
<td>Na₁-O₁ = 2.395</td>
<td>O₁-H₆ = 1.000</td>
</tr>
<tr>
<td>As₁-O₂ = 1.652</td>
<td>Ca₁-O₂ = 2.352</td>
<td>Na₁-O₂ = 2.425</td>
<td>O₁-H₆ = 0.998</td>
</tr>
<tr>
<td>As₁-O₃ = 1.656</td>
<td>Ca₁-O₃ = 2.367</td>
<td>Na₁-O₃ no. 7 = 2.478</td>
<td>O₁-H₆ = 1.002</td>
</tr>
<tr>
<td>As₁-O₄ = 1.664</td>
<td>Ca₁-O₄ no. 3 = 2.385</td>
<td>Na₁-O₄ = 2.478</td>
<td>O₁-H₆ = 1.002</td>
</tr>
<tr>
<td>As₁-O₅ = 1.684</td>
<td>Ca₁-O₅ no. 4 = 2.412</td>
<td>Na₁-O₅ no. 6 = 2.517</td>
<td>O₁-H₆ = 1.002</td>
</tr>
<tr>
<td>As₁-O₆ = 1.689</td>
<td>Ca₁-O₆ no. 1 = 2.435</td>
<td>O₁-O₁ no. 1 = 2.481</td>
<td>O₁-H₆ = 0.999</td>
</tr>
<tr>
<td>As₁-O₇ = 1.665</td>
<td>Ca₁-O₇ = 2.275</td>
<td>O₁-O₁ no. 2 = 2.417</td>
<td>O₁-H₆ = 1.002</td>
</tr>
<tr>
<td>As₁-O₈ = 1.672</td>
<td>Ca₁-O₈ = 2.331</td>
<td>O₁-O₁ no. 3 = 2.360</td>
<td>O₁-H₆ = 1.002</td>
</tr>
<tr>
<td>As₁-O₉ = 1.672</td>
<td>Ca₁-O₉ no. 5 = 2.417</td>
<td>O₁-O₁ no. 5 = 2.447</td>
<td>O₁-H₆ = 1.002</td>
</tr>
<tr>
<td>O₁-H₆ = 0.62</td>
<td>Ca₁-O₁ no. 6 = 2.517</td>
<td>O₁-O₁ no. 7 = 2.478</td>
<td>O₁-H₆ = 1.002</td>
</tr>
</tbody>
</table>

Notes: no. 1 = x + 1, y + z; no. 2 = x + y - 1, + z; no. 3 = - x + 1, y - z + 1; no. 4 = - x + 1, - y + 1, - z + 1; no. 5 = - x, - y, - z + 1; no. 6 = - x + 1, - y, - z + 2; no. 7 = - x, + y, - z + 2.

### Table 4. Selected bond angles (°) for Ca₃NaH(AsO₄)₂·6H₂O

<table>
<thead>
<tr>
<th>AsO₄ tetrahedra</th>
<th>Calcium octahedra</th>
<th>Sodium octahedra</th>
</tr>
</thead>
<tbody>
<tr>
<td>O₁-Ca₁-O₆ no. 2 = 108.2</td>
<td>O₁-Ca₁-O₆ no. 3 = 170.3</td>
<td>O₁-Na₁-O₆ = 96.4</td>
</tr>
<tr>
<td>O₁-Ca₁-O₆ = 112.8</td>
<td>O₁-Ca₁-O₆ = 152.7</td>
<td>O₁-Na₁-O₆ = 96.25</td>
</tr>
<tr>
<td>O₁-Ca₁-O₆ = 110.1</td>
<td>O₁-Ca₁-O₆ = 95.5</td>
<td>O₁-Na₁-O₆ = 166.3</td>
</tr>
<tr>
<td>O₁-Ca₁-O₆ = 114.3</td>
<td>O₁-Ca₁-O₆ = 82.2</td>
<td>O₁-Na₁-O₆ = 169.6</td>
</tr>
<tr>
<td>O₁-Ca₁-O₆ = 107.3</td>
<td>O₁-Ca₁-O₆ no. 3 = 85.9</td>
<td>O₁-Na₁-O₆ = 84.4</td>
</tr>
<tr>
<td>O₁-Ca₁-O₆ = 105.1</td>
<td>O₁-Ca₁-O₆ = 105.4</td>
<td>O₁-Na₁-O₆ = 84.2</td>
</tr>
</tbody>
</table>

Notes: no. 1 = x + 1, y + z; no. 2 = x, + y - 1, + z; no. 3 = - x + 1, - y, - z + 1; no. 4 = - x + 1, - y + 1, - z + 1; no. 5 = - x, - y, - z + 1;
previously determined. In all cases, the crystals show a tabular habit parallel to (001) with a polygonal contour delimited by [100], [010], [110], and [110] crystal edges that correspond to the crystallographic directions with the smallest translations (6.68, 8.22, 10.36, and 10.62 Å, respectively). The fact that the crystal habit is dominated by (001) is easy to understand if one considers the layered structure of this compound, which also explains the perfect cleavage and the frequent twinning on (001). Due to the small crystal thickness, the side forms are difficult to index but the described polygonal contour could correspond to the intersection of (001) with minor (010) or (011), (100) or (101), (111), and (110) faces. All these faces represent the most important structural planes according to the D-H rules. The D-H tool of the computer code SHAPE ranks [001] first, with a reticular area ($V/d_{\text{cell}}$) of 54.88 Å². The forms [010], [101], [010], [110], [011], [012], [111], and [110] follow, but the reticular areas of the corresponding structural planes are significantly larger (82.18, 85.00, 98.83, 103.63, 110.93, 117.13, 123.04, and 127.82 Å², respectively).

**FIGURE 8.** Connection between two [CaO₄] octahedra with adjacent polyhedra in the Ca₃Na(AsO₄)(AsO₄)·6H₂O structure. As can be observed, two [CaO₄] polyhedra are edge-sharing through two O⁰⁺ O atoms. In the same way, other two [CaO₄] octahedral are edge-sharing via two O⁰⁻ O atoms.

**FIGURE 7.** Projection of the Ca₃Na(AsO₄)(AsO₄)·6H₂O structure onto (010). The structure is formed by layers parallel to (001) consisting of [AsO₄] tetrahedra and [CaO₆] polyhedra. These layers are linked to each other by [NaO₆] octahedra that form chains along [100].

**FIGURE 9.** (a) SEM image of an aggregate of Ca₃Na(AsO₄)(AsO₄)·6H₂O. (b) Habit of a Ca₃Na(AsO₄)(AsO₄)·6H₂O crystal simulated using SHAPE.

**DISCUSSION**

This work shows that the interaction of arsenate-bearing aqueous solutions with gypsum at a starting pH of ~9 leads to surface precipitation of guerinite, sainfeldite, and occasionally Ca₃Na(AsO₄)(AsO₄)·6H₂O. The process involves the release of Ca²⁺ and SO₄²⁻ ions from the gypsum surface to the aqueous solution and the reaction of these ions with As(V) aqueous species to produce the nucleation and growth of Ca arsenates. In the case of guerinite, the process could be described according to the reaction:

$$2\text{AsO}_4^{2-}(\text{aq}) + 2\text{AsO}_4^{2-}(\text{aq}) + 5\text{CaSO}_4 \rightarrow 2\text{HAsO}_4^{2-}(\text{aq}) + 7\text{H}_2\text{O}$$

where the subscripts aq, l, Gy, and G stand for aqueous species, liquid, gypsum, and guerinite, respectively. An analogous reaction would describe the formation of the tetrahydrate (sainfeldite). The three phases—guerinite, sainfeldite, and Ca₃Na(AsO₄)(AsO₄)·6H₂O—are characterized by the simultaneous presence of HAsO₄₂⁻ and AsO₄³⁻ groups in the structure. Obviously, an explanation of the precipitation behavior of these solid phases requires consideration of the relevant aqueous species of As(V) in the studied pH range. Here, to determine the driving forces operating during the experiments, the activities ($a_i$) of the aqueous species (free ions and complexes) have been calculated by applying PHREEQC to the experimental data (solution pH and analytical concentrations of As, Ca, Na, and sulfate). To this aim, the PHREEQC database was completed for the aqueous species AsO₄³⁻, HAsO₄₂⁻, H₂AsO₄⁻, H₃AsO₄³⁻, CaAsO₄, CaHAsO₄₂⁻, and CaH₂AsO₄ using the stability constants compiled by Bothe and Brown (1999a). In the output, PHREEQC reports concentrations and activities of all the aqueous species and the saturation index with respect to all the relevant solid phases. Moreover, for each experimental data set, PHREEQC calculates a charge balance from which the data quality can be assessed. Here, the charge-balance errors were always less than 3% and predominantly less than 2%, which supports the quality of both the analytical data and the model. For instance, after 50 days of reaction with a 20 mM parent solution, the solution pH was 7.2 and the concentrations of As, Ca, Na, and sulfate were 139, 185, 394, and 257 mM, respectively (see Tables 1 and 2). Under these
conditions, the electrical balance was –1.362 meq/kg, which corresponds to an error of –1.21%. Tables 1 and 2 only display the analytical data corresponding to the end of the experiments (50 days). The complete data set, including all the concentrations and pHs measured at intermediate reaction times, was reported in Rodriguez-Blanco (2006).

**Determination of solubility products and saturation indices**

The saturation index (SI) is a measure of the supersaturation and is given by

\[ SI = \log \frac{\Pi a_i}{K_{sp}} \]  

(5)

where \( \Pi a_i \) stands for the product of ion activities in the actual aqueous solution, \( v_i \) is the stoichiometric number of the ion \( i \) in the solid formula, and \( K_{sp} \) stands for the thermodynamic solubility product of the solid phase, i.e., the equilibrium value of \( \Pi a_i \). Obviously, \( SI = 0 \) reflects equilibrium, \( SI < 0 \) subsaturation, and \( SI > 0 \) supersaturation. Therefore, to assess the saturation index with respect to a specific solid phase, one needs to know not only the activities of the participating aqueous ions, but also the solubility product of that solid phase. Unfortunately, the determination of thermodynamic solubility products of Ca arsenates has received little attention and many published data are still open to debate. This occurs because many Ca arsenates are difficult to produce as pure phases. Instead, they are most readily formed in conjunction with other phases such as a mixture of different hydrates, different protonation degrees, etc. Bothe and Brown (1999a) have determined the solubilities of several Ca arsenates including haidingerite (CaHAsO$_4$·H$_2$O) and the two polymorphs, guerinite and ferrarisite, of the nonahydrate Ca$_3$((HAsO$_4$)$_2$(AsO$_4$)$_2$·9H$_2$O. Nevertheless, to our knowledge, no data about the solubility products of sainfeldite and, obviously, about Ca$_3$Na$_4$(HAsO$_4$)(AsO$_4$)$_2$·6H$_2$O have been reported. Moreover, the solubility product of the two polymorphs, guerinite and ferrarisite, were determined by far-from-equilibrium precipitation experiments in which an assemblage of two solid phases formed. In such experiments, equilibrium was assumed to occur when, after a while, no change in the aqueous solution composition was observed and the solid phase assemblage remains unaltered. Nevertheless, reaching a true equilibrium situation (at an invariant point) requires a dissolution-recrystallization process involving extremely slow changes, which are difficult to detect at least for short reaction times. In the case of ferrarisite, Bothe and Brown also reported problems of reproducibility in the XRD patterns of the solid phase assemblage. Finally, using the solubility product as determined by Bothe and Brown, the aqueous solution was shown to be undersaturated with respect to guerinite during most of the experiments, which is incompatible with our experimental observations.

Here, the solubility product of guerinite has been estimated assuming that the aqueous solution is at equilibrium with guerinite at the end of the experiments (50 days), which seems to be reasonable if one considers the asymptotic shape of the curve of the analytical data (see Fig. 1). Both the solid phase assemblage and the composition of the aqueous solution remains virtually unchanged during the last 30 days of the experiment. Moreover, the final concentration is virtually the same in the three replicate experiments, the relative standard deviation being less than 8%. We have interpreted this to mean that the saturation index with respect to guerinite has to approach zero at the end of the experiments, and that the solubility product can be determined by calculating the asymptotic value of \( \Pi a_i \) for prolonged reaction times. For guerinite, the ionic activity product (IAP) is given by the expression:

\[ IAP = \sum \Pi a_i = a_{Ca^{2+}} \cdot a_{HAsO_4^{-}} \cdot a_{AsO_4^{3-}} \]  

(6)

Figure 10 shows the evolution of the logarithm of IAP$_G$ during the experiment performed with 1.0–1.5 mm gypsum grains and a 25 mM Na$_2$HAsO$_4$ parent solution. The data points have been fitted to an exponential decay function. The asymptotic limit is \(-31.17 ± 0.05\). The result corresponds to a solubility product \( K_{sp} \) (guerinite) = \( 10^{-31.17} \), which is slightly lower than that estimated by Bothe and Brown (10$^{-30.69}$). The fact that analogous values have been obtained in the case of the experiments performed with 100, 50, and 20 mM Na$_2$HAsO$_4$ parent solutions strongly supports the present estimate. Moreover, this method has proved to be a reliable tool to determine the solubility products of different Ca arsenates at specific pHs. Such is the case of pharmacolite, CaHAsO$_4$·2H$_2$O, whose solubility product was determined at pH 7 (Rodríguez-Blanco et al. 2007) in similar interaction experiments with gypsum.

In the case of sainfeldite, Ca$_3$(HAsO$_4$)$_3$(AsO$_4$)$_2$·4H$_2$O, the determination of the solubility product is more problematic. This mineral has only been observed to form in the experiments carried out with 20 mM Na$_2$HAsO$_4$ parent solutions, always in coexistence with the nonahydrate guerinite. In the three replicate experiments, sainfeldite was the last phase that precipitated, i.e., sainfeldite crystallized when the aqueous solution approached saturation with respect to guerinite. Both minerals are different hydrates of the same compound, so that the expression for the two ionic activity products is the same, that is, Equation 6. Obviously, the tetrahydrate can be expected to be stable at a higher temperature than the nonahydrate. Therefore, the fact that both hydrates form together in this experiment indicates that (1) one of the two hydrates is crystallizing in a metastable way, and/or
(2) the transition temperature is very close to 25 °C. On the other hand, the fact that guerinite is the only phase formed in most of the experiments seems to indicate that guerinite is the stable hydrate at 25 °C. Another possibility is that a comparatively lower Ca/As(V) proportion in the aqueous solution favors (from a kinetic point of view) the formation of guerinite or vice versa. In any case, it is reasonable to think that the sainfeldite solubility should be quite similar to the solubility of guerinite and thus the evolution of the saturation index for sainfeldite has to be parallel and probably very close to the evolution of the saturation index for guerinite.

A similar method has been applied to determine the solubility product of Ca₂Na(H₃AsO₄)(AsO₄)·6H₂O, but in this case, crystallization was carried out by mixing CaCl₂ and Na₂H₂AsO₄ parent solutions (see experimental section). Equilibrium was assumed to occur after 1500 h of reaction, when the composition of the aqueous solution showed no change with time. The ionic activity product for this new phase is given by the expression:

\[
IAP = a_{Ca}^{2} \cdot a_{Na} \cdot a_{HAsO_{4}^{2-}} \cdot a_{AsO_{4}^{3-}}.
\]

Again, the activities have been calculated by applying PHREEQC to the experimental data (solution pH and analytical concentrations of As, Ca, Na, and Cl). In this case, the equilibrium \(IAP\) was \(K_{sp} = 10^{3.8 \pm 0.3}\). The newly obtained solubility products for guerinite and Ca₂Na(H₃AsO₄)(AsO₄)·6H₂O can now be incorporated into PHREEQC to calculate the evolution of the saturation indices with respect to these phases during the experiments.

**Reaction paths and equilibrium endpoints**

Figure 11 shows the evolution of the saturation indices of gypsum and guerinite for the experiments carried out using a 25 mM parent solution and a gypsum grain size in the range 1.0 < \(\varnothing\) < 1.5 mm. These indices have been obtained using a \(pK_{sp}\) of 4.58 for gypsum (PHREEQC database) and of 3.17 (this work) and 30.69 (Bothe and Brown 1999a) for guerinite. Moreover, taking into account that in the present experiments the final pH values were around 7.2, the saturation indices of pharmacolite and haidingerite, which usually form below a pH of 7, have also been considered. At the very beginning of the experiment, the aqueous solution is undersaturated with respect to any solid phase, but the fast dissolution of the gypsum grains leads to a dramatic increase of all these saturation indices. As can be observed, the fluid becomes quickly supersaturated with respect to guerinite (note that the first values on the left of the graph do not correspond to 0 h but to 0.5 h of interaction), reaching a maximum (\(SI_{g} = 2\)) during the first 12 h of the experiment. Afterward, the uninterrupted growth of guerinite leads to a decrease of As(V) concentration in the fluid, and \(SI_{g}\) gradually approaches zero. The saturation index of gypsum evolves differently, approaching equilibrium from undersaturation. As the gypsum grains dissolve, the concentration of Ca and sulfate in the aqueous solution increases, and the gypsum saturation index tends to reach a value close to zero after about 300 h of reaction.

Figure 11 also shows that the fluid is undersaturated with respect to both haidingerite and pharmacolite during the whole reaction time, so that the nucleation of these phases is not possible. The only mineral that precipitates is guerinite because the aqueous solution becomes highly supersaturated with respect to this phase in the early hours of reaction. The subsequent nucleation and growth of this mineral leads to the removal of Ca and As(V) from the aqueous solution in an uneven proportion, five Ca²⁺ ions for each four arsenate ions, in agreement with solid stoichiometry. As a consequence, the Ca/As(V) ratio in the aqueous solution decreases, which is consistent with a reduction of pH from 9 to ~7.2.

The evolution of the guerinite saturation index observed with a 25 mM solution is quite similar in the experiments performed with 10, 20, 50, and 100 mM Na₂H₂AsO₄ parent solutions. In all these cases, the saturation index for guerinite is close to zero for prolonged reaction times, the deviations being within the experimental error. The existence of this common endpoint supports the suitability of the solubility product estimated in this paper and confirms that the aqueous solution reaches equilibrium with guerinite at the end of the experiments. In contrast, the end point reached by the gypsum saturation index depends on the parent solution concentration. In the experiments carried out with 2, 5, 10, 20, and 25 mM As(V) parent solutions, equilibrium with gypsum is reached after a more-or-less prolonged reaction time. In contrast, when the initial concentration is higher, the aqueous solution remains undersaturated with respect to gypsum during the whole reaction time (Table 5). This scenario could be explained in view of the SEM observations, which show that, using 50 or 100 mM parent solutions, the gypsum grains become completely covered by guerinite in less than 48 h of reaction. As a consequence, the guerinite precipitate protects the substrate from further dissolution, so that the process stops, at least on the experimental timescale. Obviously, at this stage the system is not at equilibrium, but the term “partial equilibrium” (Helgeson 1968; Prieto et al. 2003) could be used to describe a situation in which the reactive solid (gypsum) becomes isolated from the aqueous solution by a coating of secondary solids that maintain equilibrium with the aqueous phase. In contrast, using 25, 20, or 10 mM parent solutions, the gypsum grains do not become completely covered by the overgrowing crystals and continue to dissolve until equilibrium with respect to both gypsum and guerinite is attained.
Relationship between the evolution of the Na concentration and the saturation index of Ca$_2$Na(HAsO$_4$)$_2$(AsO$_4$)$_2$·6H$_2$O. Parent solution: 100 mM Na$_2$HAsO$_4$. Grain size: 1.0 < O < 1.5 mm.

In the case of the 100 mM parent solution, the evolution of the saturation index of Ca$_2$Na(HAsO$_4$)$_2$(AsO$_4$)$_2$·6H$_2$O requires a separate consideration. Figure 12 shows the relationship between the evolution of the Na concentration and the saturation index of this new phase. There is a quick rise of the saturation index during the first 2–8 h of the experiment, which is followed by a simultaneous decrease in Na concentration. The aqueous solution only becomes slightly supersaturated in Ca$_2$Na(HAsO$_4$)$_2$(AsO$_4$)$_2$·6H$_2$O during a very short period of time (~6 h), and afterward becomes undersaturated in this phase. This evolution could be explained by the precipitation and subsequent dissolution of Ca$_2$Na(HAsO$_4$)$_2$(AsO$_4$)$_2$·6H$_2$O crystals and the corresponding removal-release of Na during the first hours of reaction. Moreover, this result supports the assumption that the solids obtained by mixing of CaCl$_2$ and Na$_2$HAsO$_4$ aqueous solutions correspond to the transitory phase formed using a 100 mM parent solution.

As a final point, it is interesting to discuss the maximum aqueous As(V) concentration that can exist in the presence of gypsum within this pH range. The equilibrium end point would imply simultaneous equilibrium of both guerinite (G) and gypsum (Gy) with the aqueous solution, according to Equation 4. The equilibrium constant for this reaction can be easily derived from the solubility products of both minerals:

$$K_{eq}(G/G) = \frac{a_{SO_4}^2}{a_{HAsO_4}^4 \cdot a_{AsO_4}^2} = \frac{a_{Ca}^2 \cdot a_{SO_4}^2}{a_{Ca}^2 \cdot a_{HAsO_4}^4 \cdot a_{AsO_4}^2} = \frac{K_{sp}(Gy)}{K_{sp}(G)}$$  

(8)

This result means that, at equilibrium, the aqueous activities of SO$_4^{2-}$, HAsO$_4^{2-}$, and AsO$_4^{3-}$ are related in such a way that the ratio $K_{sp}(Gy/G)$ has to be $\approx 1.9 \times 10^4$. This condition is reasonably satisfied (within the experimental error) at the end of the experiments carried out with 25, 20, and 10 mM parent solutions (see Table 5). However, in the case of experiments carried out with 100 and 50 mM parent solutions, the actual ratio differs by one order of magnitude with respect to the equilibrium value, which confirms the existence of a partial equilibrium situation. Equation 8 also shows that if by some other process, the SO$_4^{2-}$ concentration decreases while equilibrium is maintained for both guerinite and gypsum, then the total concentration of arsenate also decreases due to the solvent-mediated conversion of gypsum to guerinite. For instance, it is easy to calculate using PHREEQC that, if at pH = 7.2 the concentration of sulfate decreases from 30 to 15 mM, then the total concentration of arsenate has to decrease from ~14.6 to ~7.8 mM to maintain equilibrium for both guerinite and gypsum.

Another key factor in determining the removal capacity of the reaction in Equation 4 is the solution pH. At the pH = 7 attained at the end of the experiments, the prevailing arsenate species in the aqueous solution is HAsO$_4^{3-}$, followed by the doubly protonated arsenate ion, H$_2$AsO$_4$. The unprotonated arsenate ion, AsO$_4^{3-}$, follows but its concentration is several orders of magnitude lower than that of HAsO$_4^{3-}$. In these conditions, a rise of pH would have the effect of increasing the relative concentration of AsO$_4^{3-}$ at the expense of the concentration of H$_2$AsO$_4$. This redistribution, in turn, would lead to an increase in the saturation index of guerinite, so that this phase should precipitate to restore equilibrium. For instance, if the pH increases from 7 to 9, the total concentration of arsenate has to decrease from ~10 to ~2 mM to maintain equilibrium for both guerinite and gypsum. The removal of arsenate is considerably more effective at pH >9. However, under such conditions, which are beyond the scope of the present work, other Ca arsenates that are less soluble than guerinite would precipitate. Future papers will deal with this matter.

The interaction of As(V)-bearing aqueous solutions with gypsum at a starting pH of ~9 results in surface precipitation of guerinite, sainfeldite, and in some cases Ca$_2$Na(HAsO$_4$)$_2$(AsO$_4$)$_2$·6H$_2$O, a new arsenate. These three solid phases are characterized by the simultaneous presence of HAsO$_4^{3-}$ and AsO$_4^{3-}$ groups in their structures, which is explainable as crystallization occurs within a pH range in which both HAsO$_4^{3-}$ and AsO$_4^{3-}$ ions are available in the aqueous solution. This phenomenon leads to a decrease in the As(V) concentration in the aqueous phase to reach values (~15 mM) controlled by the solubility of these solid phases at the final pH (~7.2). This concentration could be further reduced by buffering the pH to 8–9 and/or in the presence of a sink for sulfate, but a value of ~1 mM seems to be the minimum value attainable by precipitation of guerinite. As expected, this concentration is extremely high from the perspective of water quality and thus gypsum does not seem to be a suitable remediation tool in this pH range. However, the nature, solubility, and crystallization behavior of the solid phases that precipitate in this pH range needed to be investigated to get a better understanding of the As(V) behavior in aqueous environments.

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