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Experimental assessment of CO$_2$-mineral-toxic ion interactions in a simplified freshwater aquifer: Implications for CO$_2$ leakage from deep geological storage

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Abstract

The possible intrusion of CO$_2$ into a given freshwater aquifer due to leakage from deep geological storage involves a decrease in pH, which has been directly associated with the remobilization of hazardous trace elements via mineral dissolution and/or via desorption processes. In an effort to evaluate the potential risks to potable water quality, the present study is devoted to experimental investigation of the effects of CO$_2$ intrusion on the mobility of toxic ions in simplified equilibrated aquifers. We demonstrate that remobilization of trace elements by CO$_2$ intrusion is not a universal physicochemical effect. In fact goethite and calcite, two minerals frequently found in aquifers, could successfully prevent the remobilization of adsorbed Cu(II), Cd(II), Se(IV) and As(V) if CO$_2$ is intruded into a drinking water aquifer. Furthermore, a decrease in pH resulting from CO$_2$ intrusion could reactivate the adsorption of Se(IV) and As(V) if goethite and calcite are sufficiently available in underground layers. Our results also suggest that adsorption of cadmium and copper could be promoted by calcite dissolution. These adsorbed ions on calcite are not remobilized when CO$_2$ is intruded into the system, but it intensifies calcite dissolution. On the other hand, arsenite As(III) is significantly adsorbed on goethite, but is partially remobilized by CO$_2$ intrusion.
Keywords: CO₂ leakage, Freshwater aquifer, Goethite; Calcite; Remobilization, Metalloids; Heavy metals
Introduction

Unregulated CO₂ emissions into the Earth’s atmosphere (about 22x10⁹ ton CO₂/year), caused mainly by fossil fuel combustion, have led to concerns about global warming. To maintain the atmospheric CO₂ level below 500 ppm, CO₂ emissions will have to be stabilized at current levels, although they are forecast to double over the next 50 years (1). Capture from individual industrial sources and long-term geological storage are realistic and available ways of reducing CO₂ emissions because large volumes of this gas can be stored in various deep geological formations (2-5). However, CO₂ injection into porous media (deep geological formations) can significantly modify their storage properties. For example in brine aquifers, nucleation-growth and/or dissolution of minerals are envisaged and these reactions can be enhanced by high in-situ temperatures (80-110°C) and pressures (100-400 bar). Toxic elements (e.g. metalloids or heavy metals) could be liberated or released by native mineral dissolution reactions as measured and/or simulated in some studies (6-8); subsequently, dissolved or released ions could be sequestered by nucleation-growth of secondary mineral phases (9-10). In summary, several physicochemical reactions at fluid-rock interfaces have been envisaged and they need to be studied in detail in the coming years as claimed by various recent studies (5, 8-13).

The deep geological formations considered (e.g. brine aquifer) are always isolated by impermeable layers, so-called caprock formations. Unfortunately, CO₂ leakage from storage sites to higher formations (e.g. drinking water sources) by diffusion through less impermeable zones or transport along (micro-) fractures could be a potential risk in the long term, possibly aggravated by seismic activity and micro-seismicity (14). In this hypothetical case, CO₂ dissolution could influence the mobility and/or retention of hazardous trace elements (e.g. heavy metals and/or metalloids) contained in native aquifer minerals, as recently investigated by reactive transport simulations (7, 15-18), laboratory experiments (19-22) and field tests.
(e.g. MSU-ZERT field site) (23-24). In this way, various trace elements such as Cd, Pb, Zn, U and As, which can be toxic to humans or wildlife at µg/L-mg/L concentrations, have been predicted to be released by dissolution reactions of native aquifer minerals (15-18). All simulations in these studies have assumed that the intrusion of CO₂ into drinking water aquifers decreases their pH, enhancing the dissolution reactions of native aquifer minerals and consequently remobilizing trace metal species if such metals are contained in the minerals. Unfortunately, few experimental data are available in the literature in order to challenge or support such predictions. Moreover, the sorption/desorption processes of trace elements (fast or instantaneous processes at the mineral-fluid interfaces) are also rarely simulated because it is difficult to combine them with dissolution-precipitation reactions in reactive transport simulations (16). It should be noted that desorption/adsorption via surface complexation is possibly the most important process controlling the fate of toxic trace elements mobilized by CO₂ leakage (7, 15).

For the above reasons, the present study is devoted to experimental investigation of the effects of CO₂ intrusion on the mobility/retention of toxic ions in simplified equilibrated aquifers (drinking water sources). In our study, “simplified equilibrated aquifer” means a synthetic solution enriched with one or two metalloid ions equilibrated for 24h with a single or binary mineral system at 20°C and 10 bar of argon prior to CO₂ injection (8 bar). In this context, six metalloid ions ((Cd(II), Cu(II), Se(IV), Se(VI), As(III) and As(V)) and two widespread minerals (goethite, calcite and goethite-calcite composite) were investigated. A schematic illustration of this experimental study with emphasis on CO₂ leakage from deep geological storage is provided in Figure 1. Goethite and calcite were chosen because of their abundance in nature as minerals (abiotic origin) and biominerals (biotic origin). Both minerals can co-exist in several terrestrial environments such as deep geological formations, aquifers (drinking water sources), soils and aerosols, playing an important role in the fate and transport
of several metalloid and heavy metal trace elements and organic molecules at the mineral-
fluid interfaces (25-29). Finally, it should be noted that various suspensions were withdrawn
from the reactor before and after injection of CO_{2} in each experiment or run, in order to
monitor the pH and ion concentration ([Ca], [Fe], [Na], [As], [Se], [Cd] and [Cu]) ex-situ as a
function of time (up to 48 hours). This allows a simple macroscopic evaluation of element
transfer at the mineral-solution interfaces, involving mineral dissolution and/or
retention/mobility of adsorbed toxic ions related to the intrusion of CO_{2} in the system.

Materials and methods

Minerals

Pure goethite, pure calcite and a goethite-calcite composite were synthesized as mineral
analogues for a given simplified potable aquifer. In this experimental study powdered and
high-purity minerals were used in order to increase their surface reactivity and also to
identify/quantify clearly their physicochemical affinity with investigated toxic ions interacting
in CO_{2}-poor and CO_{2}-rich solutions as schematically illustrated in Fig. 1. A brief description
of these mineral syntheses is provided in the following paragraphs.

Goethite: Low acicular goethite with a high specific surface area (130m^{2}/g) was synthesized
by placing 1mol of NaOH and 0.2mol of FeCl_{3}.6H_{2}O in a 2L reaction cell and adding 1L of
high-purity water. Constant agitation (400rpm) of the solution and a moderate temperature
(30°C) for 24h of reaction were required. For more specific details on the synthesis procedure
and goethite characterization refer to Montes-Hernandez et al. (30).

Calcite: Nanosized calcite with a moderate specific surface area (7m^{2}/g) was synthesized via
gas-solid carbonation of Ca(OH)_{2} nanoparticles in a static bed reactor. The specific procedure
and fine calcite characterization have already been reported by Montes-Hernandez et al. (31).
In short, pure calcite was synthesized by placing 74g of Ca(OH)_{2} in a 2L reaction cell slightly
heated to 30°C and by injecting high-purity CO\textsubscript{2} at 20 bar. Complete Ca(OH)\textsubscript{2}-to-CaCO\textsubscript{3} (calcite) transformation was reached after about 24h of reaction.

Goethite-Calcite composite: Low acicular goethite intimately adhering to sub-micrometric calcite - so-called goethite-calcite composite - with a high specific surface area (90m\textsuperscript{2}/g) was synthesized via three sequential precipitation reactions: (1) Fast precipitation of nanosized acicular goethite (α-FeOOH) using a high OH/Fe molar ratio (=5); (2) Instantaneous precipitation of portlandite (Ca(OH)\textsubscript{2}) by adding CaCl\textsubscript{2} salt into a goethite alkaline suspension, and; (3) Sub-micrometric calcite precipitation by injecting CO\textsubscript{2} into a goethite-portlandite alkaline suspension. These sequential precipitation reactions and textural characterization of goethite-calcite composite have been specifically described by Montes-Hernandez et al. (32).

\textbf{Aqueous solutions}

One liter of high-purity water was used to prepare synthetic solutions enriched with single or double metal(loid) ions by using commercial salts of As(III), As(V), Se(IV), Se(VI), Cd(II) and Cu(II). The chemical compositions of all the salts used are summarized in Table 1. For the double metal(loid) ion solutions, two ions were arbitrarily combined as follows: As(III)-Se(IV), As(V)-Se(VI) or Cd(II)-Cu(II). In all cases, a high initial element concentration of about 50mg/L was also arbitrarily considered. This high concentration is not realistic in a given potable water aquifer, but it leads to a high saturation on the surface of the minerals being investigated through a sorption process prior to CO\textsubscript{2} injection. Moreover, it is easy to quantify the release and/or mobilization of adsorbed ion(s) due to CO\textsubscript{2} intrusion into the system.
Experiments: CO$_2$-mineral-toxic ion interactions

One liter of single or double metal(loid) ion solution and 3g of calcite, goethite or goethite-calcite composite were placed in a 2L reaction cell. This suspension was immediately stirred using constant mechanical agitation (400 rpm) and argon was also immediately injected at 10 bar in order to simulate pressures of the kind relating to potable water aquifers. After 24h of mineral-ion solution equilibration at room temperature ($\sim$20°C), i.e. assuming that adsorption of a given ion takes place during the first 24h, high-purity CO$_2$ was injected at 8 bar into the dispersion in order to assess the release of adsorbed toxic ions and also to quantify the mineral dissolution, both processes being possibly intensified by intrusion of CO$_2$ into the system. In all experiments, about 15ml of dispersion were withdrawn from the reactor as a function of time before (from 0 to 24h) and after (from 24 to 48h) injection of CO$_2$. The pH of each sample of suspension was measured at room temperature. The suspensions were then filtered through a 0.2 µm Teflon filter. 10 ml of the resulting filtered solutions were acidified for measurement of [Ca], [Fe], [Na], [As], [Se], [Cd] and [Cu] by inductively coupled plasma optical emission spectrometry (ICP-OES).

Results and discussion

As predicted and/or measured, the intrusion of CO$_2$ within a freshwater aquifer decreases the pH, enhancing dissolution reactions of native aquifer minerals and consequently remobilizing trace metal species if such metals are contained in them. Here, various trace elements such as Cd, Pb, Zn, U and As were predicted to be released by dissolution reactions of native aquifer minerals and/or by fast desorption (7, 15-18). In this experimental study, we provide more specific insights into the remobilization of Se(IV), Se(VI), As(III), As(V), Cd(II) and Cu(II) related to CO$_2$ intrusion into a pre-equilibrated mineral-metal(loid) ion solution system. The chemical speciation of these six ions in solution before and after injection of CO$_2$ was
determined by using the Phreeqc numerical code (33) in order to understand certain observed behaviors more easily (see below). pH data (measured and calculated) and the dominant species in solution are summarized in Table 1.

Goethite as an aquifer mineral analogue

Figure 2 displays a compilation of five different adsorption experiments using pure goethite as adsorbent. Here, the adsorbed/remobilized amount of a given metal(loid) ion at the mineral-solution interfaces was normalized with respect to the available surface area in the system (3g*130m^2/g). In short, Cd(II), Cu(II) and As(III) are significantly adsorbed on the goethite prior to CO_2 intrusion into the system. A lower adsorption capacity was measured for Se(IV) and Se(VI) in the conditions investigated. After CO_2 intrusion, residual dissolved Se(IV) is largely removed from the solution onto the goethite, i.e. the adsorption process is reactivated by CO_2 intrusion. Conversely, adsorbed Cd(II) and Cu(II) are largely removed from the goethite by CO_2 intrusion into the system. Adsorbed arsenite As(III), which is reputed to be a very highly mobile element, was slightly affected by CO_2 intrusion. In general, the so-called oxyanions were re-adsorbed or slightly remobilized (case: As(III)). However, the heavy metals investigated (Cd(II) and Cu(II)) (single cations) were preferentially remobilized from the goethite. This can be explained by a significant change in the surface charge of the goethite after CO_2 intrusion, assuming that surface charge is strongly dependent on pH, as is widely described in the literature (e.g. 27, 34 and references therein) and on the understanding that the Cu and Cd speciation in solution remains unchanged in the presence of goethite when CO_2 is intruded into the system (see Table 1). In our study, the pH (measured ex-situ) changes drastically when CO_2 is intruded into the system (Fig. 2b). In this context, the intrusion of CO_2 could create a positive charge on the goethite, enabling the re-sorption of available dissolved anions in the interacting solution. Finally, a slight release of Fe (<2mg/L) was measured prior to CO_2 intrusion, indicating slight dissolution of the goethite, and Fe was no
longer detected in the interacting solution (<0.1 mg/L, “minimum value in our calibrating curve”) when CO$_2$ was intruded into the system (Fig. 2b). This is in agreement with the literature, in the fact that the goethite is stable over a broad range of pH values (from 3 to 13.5) at ambient temperature (35, 36) as also verified by equilibrium Phreeqc calculations.

Calcite as an aquifer mineral analogue

In a similar way to the goethite scenario, Figure 3 summarizes five different adsorption experiments now using pure calcite as adsorbent. The respective pH profiles and an average Ca-concentration profile can be also consulted in Figure 3b. The adsorbed/remobilized amounts for all the ions investigated were also normalized with respect to the available surface area in the calcite system (3g*7m$^2$/g). Here, the major result is the fast, strong sorption of Cd(II) and Cu(II) onto/into the calcite. The ex-situ monitoring of pH and Ca concentration in the interacting solution (Fig. 3b) suggests that slight dissolution of calcite enhances the sorption of cadmium and copper (heavy metals) as claimed earlier by Limousin (37) for cadmium in particular. We also note that CO$_2$ intrusion into the calcite system has no significant effect on the Cd(II) and Cu(II) already sorbed. This means that dissolution of the calcite, i.e. the release of Ca from the calcite into the interacting solution as a result of CO$_2$ intrusion, was dissociated from potential remobilization of sorbed Cd and Cu as might be speculated at first sight. On the other hand, the oxyanions investigated (Se(IV), Se(VI) and As(III)) were only slightly adsorbed on the calcite, if the values are compared with those for the sorbed amounts of Cd(II) and/or Cu(II) (Fig. 3a). We also note that arsenite As(III) and selenate Se(VI) were completely remobilized from the calcite when CO$_2$ was intruded into the system. Conversely, a more complex behavior was observed in the case of selenite Se(IV); in fact, a given amount of dissolved Se(IV) was re-adsorbed on the calcite by CO$_2$ intrusion. In this way, the decrease in pH resulting from CO$_2$ intrusion plays an important role with regard
to the surface charge of a given mineral, the chemical speciation of dissolved elements (partial
protonation of selenite from $\text{SeO}_3^{2-}$ to $\text{HSeO}_3^-$: see Table 1) and possibly the surface
complexation process (25). It should be noted that the decrease in pH can be significantly
moderated/buffered by the dissolution of carbonate minerals (e.g. calcite) as illustrated in
Figures 2a (goethite system) and 3a (calcite system). This pH buffering capacity can be
envisaged in freshwater aquifers because these underground layers generally contain
carbonate minerals. Numerical simulations have been successfully used to quantify this pH
buffering capacity in groundwater in response to CO$_2$ leakage from deep geological storage
(7).

Goethite-calcite composite as an aquifer mineral analogue

Five supplementary adsorption/release experiments are reported in Figure 4, but, in this case,
a goethite-calcite composite was used as adsorbent. The pH profile for each adsorption/release
experiment and release of Fe and Ca by dissolution of goethite and calcite, respectively, are
also reported in Fig. 4b. As indicated in the previous cases, the adsorbed/remobilized amounts
of the ions investigated were normalized with respect to the available surface area in the
goethite-calcite system ($3\text{g}^\ast 90\text{m}^2/\text{g}$). In a similar way to the calcite system, fast, strong
adsorption of Cd(II) and Cu(II) onto the goethite-calcite was measured. However, the kinetic
behavior was reversed. For example, Cd was adsorbed in the first 10 minutes when pure
calcite was used as adsorbent and its adsorption was considerably retarded in the case of the
goethite-calcite system (see Fig. 3a and 4a). This was probably due to different textural
properties between pure calcite (nanosized) and calcite contained in the composite (sub-
micrometric) (31-32). On the basis of the results obtained with the calcite system, we assume
that such high adsorption is controlled preferentially by the calcite contained in the composite.
Additionally, these two adsorbed heavy metals were scarcely perturbed by CO$_2$ intrusion into
the system; this is in complete agreement with the calcite system (Fig. 3a and 4a). Concerning
the so-called oxyanions, a given proportion of residual dissolved selenite Se(IV) was reabsorbed onto the goethite-calcite composite by CO₂ intrusion. Conversely, adsorbed arsenite As(III) was partially remobilized from the goethite-calcite composite. This is in agreement with the calcite and goethite systems. The selenate Se(VI) ion was slightly adsorbed on the calcite and/or goethite before and/or after CO₂ intrusion into the system (Fig. 2a and 3a). This highly mobile ion had not presented any adsorption affinity with the goethite-calcite composite (Fig. 4a). Finally, mineral dissolution could be intensified by CO₂ intrusion in this way; we assumed that the goethite contained in the composite was extremely stable in the conditions investigated because the Fe concentration measured in the solution was less than 0.1 mg/L, the “minimum value in the calibrating curve”. Conversely, significant Ca concentration was measured in the interacting solution and this was intensified by CO₂ intrusion, reaching about 340 mg/L at the end of the experiment (48h) (Fig. 4b). As mentioned above, calcite dissolution may moderate the decrease in pH and such dissolution is not related directly to the remobilization of adsorbed heavy metals (e.g. Cd and Cu).

Competitive adsorption/remobilization of binary metal(loid) ions

For competitive adsorption/remobilization of binary metal(loid) ions using goethite-calcite composite as adsorbent, two ions were arbitrarily combined as follows: As(III)-Se(IV), As(V)-Se(VI) or Cd(II)-Cu(II). These three independent experiments are reported in Figure 5. 

Cd(II)-Cu(II) competition: All available copper in the solution (about 50 mg/L) was adsorbed more rapidly than the available cadmium (unfilled circle profiles in Figure 5). The same kinetic behavior was observed in the case of single metal(loid) ion solutions (Fig. 4a). This suggests insignificant competitive adsorption between Cu and Cd using goethite-calcite composite. However, CO₂ intrusion into the system created slight remobilization of adsorbed Cu. This is in disagreement with a single metal(loid) ion solution in the case of the goethite-
calcite system. Such disagreement could be explained if all the adsorption sites on the available calcite were saturated with Cd and Cu, and the liberated Cu therefore comes from the goethite, as observed in the pure goethite system when CO$_2$ is intruded (Fig. 2a).

$\textit{Se(IV)-As(III) competition:}$ Clearly, arsenite As(III) is adsorbed on the goethite-calcite composite in preference to the selenite Se(IV) ion (unfilled star profiles in Figure 5).

However, CO$_2$ intrusion into the system reactivates the adsorption of residual dissolved Se(IV). Conversely, adsorbed As(III) is partially remobilized from the goethite-calcite composite. CO$_2$ intrusion was observed to have a similar effect in the case of single Se(IV) and As(III) ion solutions (Figure 4a).

$\textit{Se(VI)-As(V) competition:}$ Arsenate As(V) was strongly adsorbed on the goethite-calcite composite. Conversely, insignificant adsorption of selenate Se(VI) was measured in the conditions investigated (unfilled triangle-down profiles in Figure 5). Surprisingly, CO$_2$ intrusion into the system led to re-adsorption of residual dissolved As(V) on the goethite-calcite composite while As(III) was remobilized in all the other cases investigated. This can be explained in terms of chemical speciation in the solution because arsenate remains partially protonated before and after injection of CO$_2$ (dominant species H$_2$AsO$_4^-$; see also Table 1) while arsenite is completely protonated from H$_2$AsO$_3^{1-}$ to H$_3$AsO$_3^{0}$ when CO$_2$ is intruded into the system.

In conclusion, this simple experimental study provides further specific insights into the remobilization/re-adsorption of Se(IV), Se(VI), Cd(II), Cu(II), As(III) and As(V) caused by CO$_2$ intrusion into an equilibrated mineral-ion solution system. This has direct implications for CO$_2$ leakage from deep geological formations and its possible intrusion into freshwater aquifers. CO$_2$ intrusion into a given aquifer has been directly associated with the release and/or remobilization of trace elements via mineral dissolution and/or desorption. In the present study, we demonstrated that the remobilization of trace elements as a result of CO$_2$
intrusion is not a universal physicochemical effect. In fact, goethite and calcite, two minerals frequently found in aquifers, can successfully prevent the remobilization of adsorbed Cu(II), Cd(II), Se(IV) and As(V) if CO₂ is intruded into a drinking water aquifer. Moreover, a decrease in pH resulting from CO₂ intrusion can reactivate the adsorption of Se(IV) and As(V) if goethite and calcite are sufficiently available in underground layers. Our results also suggest that the adsorption of cadmium and copper could be promoted by calcite dissolution.

In this way, adsorbed Cd(II) and Cu(II) on calcite are not remobilized when CO₂ is intruded into the system, but it intensifies calcite dissolution, which plays a buffering role. On the other hand, arsenite As(III) is significantly adsorbed on goethite, but is partially remobilized by CO₂ intrusion. Finally, our results confirm the low or even insignificant adsorption affinity of selenate Se(VI) with calcite and/or goethite. The latter oxyanion is reputed to be a very highly mobile element in natural oxic environments.
Acknowledgements

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Table 1. Dominant species and pH in suspensions prior to and after CO$_2$ injection into the system. These values were determined by using Phreeqc numerical simulations with an element initial concentration of 50 mg/L and 8 bar of CO$_2$ in the system.

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$^1$ Equilibration between ion solution and mineral prior to CO$_2$ injection; $^a$ pH measured ex-situ at 20°C in the mother solution; $^2$ Equilibration between ion solution and mineral after CO$_2$ injection; $^b$ pH measured ex-situ at 20°C at the end of the experiment (after 48h).
Figure 1. Schematic representation of CO₂ leakage from deep geological storage, focusing in red on the present study: “experimental assessment of CO₂-mineral-toxic ions interactions in a simplified potable aquifer”.
Figure 2. (a) Kinetic adsorption behavior of a single metal(loid) ion on goethite and effect of CO₂ intrusion into the system. (b) pH profiles before and after CO₂ intrusion. In all experiments, a similar initial ion concentration (about 50mg/L) was used and the adsorbed/remobilized amount of each element was normalized with respect to the initial available surface area (3g*130m²/g).
Figure 3. (a) Kinetic adsorption behavior of a single metal(loid) ion on calcite and effect of CO$_2$ intrusion into the system. (b) pH profiles and average [Ca] profile before and after CO$_2$ intrusion. In all experiments, a similar initial ion concentration (about 50mg/L) was used and the adsorbed/remobilized amount of each element was normalized with respect to the initial available surface area (3g*7m$^2$/g).
Figure 4. (a) Kinetic adsorption behavior of a single metalloid ion on goethite-calcite composite and effect of CO$_2$ intrusion into the system. (b) pH profiles and average [Fe] and [Ca] profiles before and after CO$_2$ intrusion. In all experiments, a similar initial ion concentration (about 50mg/L) was used and the adsorbed/remobilized amount of each element was normalized with respect to the initial available surface area (3g*90m$^2$/g).
Figure 5. Competitive adsorption/remobilization of binary metal(loid) ions by using goethite-calcite composite as adsorbent. Two ions were arbitrarily combined as follows: As(III)-Se(IV), As(V)-Se(VI) or Cd(II)-Cu(II). A similar initial ion concentration (about 50mg/L for each element) was used and the adsorbed/remobilized amount was normalized with respect to the initial available surface area (3g*90m^2/g).