In situ redeposition of trace metals mobilized by CO$_2$-charged brines

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[1] Mobilization of contaminants by CO$_2$-charged brines is one concern relating to injection of CO$_2$ as part of carbon capture and storage projects. This study monitors the mobility of trace metals in an exhumed CO$_2$-charged aquifer near the town of Green River, Utah (USA), where CO$_2$-charged brines have bleached red sandstones, and concentrated trace metals at the bleaching reaction front. Mass balance calculations on the trace metal enrichments are used to calculate time-integrated fluid fluxes and show that a significant fraction of the metals mobilized by the CO$_2$-rich brines are redeposited locally. A sequential extraction procedure on metal-enriched samples shows that these metals are incorporated into secondary carbonate and oxide phases which have been shown to grow at the CO$_2$-promoted bleaching reaction front. We argue that while CO$_2$-charged brines are capable of mobilizing trace metals, local metal redeposition implies that the potential for contamination of overlying freshwater aquifers is low.

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1. Introduction

[2] Geological CO2 sequestration is proposed as one method for mitigating the effects of fossil fuel burning on global climate [e.g., IPCC, 2007]. One of the outstanding scientific questions relates to the potential for injected CO2 to leak and contaminate overlying freshwater aquifers [Gale, 2004; Smyth et al., 2009]. CO2, injected into geological formations in a gaseous or supercritical state, is likely to dissolve into formation fluids at the gas-water interface, forming acid brines [Bickle, 2009; Dubacq et al., 2012]. These brines can mobilize trace metals via desorption of metals bound to mineral surfaces, or by dissolution of trace metal-bearing phases [Bradl, 2004; Yin et al., 1996]. If these brines were to migrate into overlying freshwater aquifers, then there is concern that drinking water supplies could be affected.

Models [Wilkin and DiGulio, 2010; Zheng et al., 2009] and experiments [Little and Jackson, 2010; Lu et al., 2010] have shown that CO2 can mobilize trace metals; however, field experiments have not shown hazardous levels of contamination [Keating et al., 2010; Kharaka et al., 2010]. Of critical importance is whether the mobilized metals remain in solution or are redeposited locally.

[3] CO2 has been escaping along the Little Grand Wash and Salt Wash grabens for at least 400,000 years, based on U-Th dating of surface carbonate deposits. [Kampman et al., 2012; Dockrill and Shipton, 2010]. Near the town of Green River, Utah, the red-bed Entrada sandstone, exposed by the Salt Wash graben, exhibits spectacular bleaching in which hematite grain coatings are removed and trace metals released from the grain coatings are concentrated at the bleaching reaction front (Figure 1). Wigley et al. [2012] argue that the bleaching results from inputs of CO2-charged brines along the Salt Wash fault, distinguishing the site from hydrocarbon-related bleached sandstones elsewhere on the Colorado Plateau [Wigley et al., 2012; Loope et al., 2010; Beittler et al., 2005, 2003; Garden et al., 2001; Chan et al., 2000]. The CO2-promoted reaction fronts are used as an analogue for processes occurring where CO2 leaks from injection sites into overlying aquifers. Here we quantify trace metal enrichments and corresponding time-integrated fluid fluxes [Bickle and Baker, 1990] at the bleaching reaction fronts, and use these to calculate the fraction of mobilized metals which have been redeposited. This allows an estimate to be made of the concentration of metals in the solution leaving the CO2-promoted reaction front.

1.1. Green River Natural Analogue Site

[4] Within the field area, the Jurassic Entrada sandstone is a well-sorted sediment of uniform grain size and mineralogy. Regional burial diagenesis includes the development of Fe-oxide grain coatings [Cullers, 1995; Trimble, 1978], giving the rock its red color. Along the axis of the Green River anticline, the Entrada sandstone has been bleached by diagenetic fluids which have dissolved the Fe-oxide grain coatings [Wigley et al., 2012]. Bleaching occurs at the base of the formation in a broad domal structure and exposed over a 1–2 km east-west section along the Salt Wash graben (Figure 2a).

[5] Bleaching of red bed sandstones is a common worldwide occurrence, with the bleaching being a result of acid-reductive dissolution of hematite grain coatings, and mobilization of otherwise insoluble Fe3+ to soluble Fe2+. Such fluid-rock reaction may be promoted by a wide range of common volatile species (CH4, CO2, H2S, hydrocarbons, and organic acids) present in geological fluids that modify fluid Eh-pH. Several lines of evidence suggest that the fluid causing the bleaching of the Entrada sandstone in the Salt Wash graben was a CO2-charged brine with minor methane.

[6] Calcite cements and veins associated with the Green River bleaching have a heavier δ13C than hydrocarbon-related bleaching, consistent with deposition from isotopically heavy C-rich fluids, similar to those in the modern, actively leaking CO2 system at Green River that underlies the study site [Assayag et al., 2009; Kampman et al., 2009], but inconsistent with isotopically light methane-saturated fluids. Mobilization of low pH-soluble trace metals is inconsistent with methane-rich fluids which have pH ≈7.6 when saturated with Fe-oxides (Figure S1 in the supporting information).1 Regionally, such bleaching is typically localized to structural highs, implicating a low density methane-saturated bleaching fluid. At Green River, bleaching is restricted to the base of the formation, and the morphology of bleached zone is consistent with the flow of a dense fluid as a gravity current, discrediting the role of buoyant methane-saturated bleaching fluids. If the bleaching fluids contained significant H2S, then hematite would be immediately reprecipitated in situ as pyrite and would not

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1 All supporting information may be found in the online version of this article.
Figure 1. (A) Google earth image of the area of bleached sandstones near the town of Green River, Utah. Dotted gray line shows the extent of the bleaching. Black dashed lines show faults, and solid black arrows mark the Green River anticline. (B) Map of Utah describing the location of the field area. Grey rectangle shows the area in Figure 1A.

Figure 2. Illustration and outcrop photos of the exhumed CO₂ reservoir. (a) Bleaching at the crest of the anticline, facing north. (b) Schematic development of bleached sandstone (yellow) in original sandstones (red). CO₂ (or CO₂-rich brine) migrated into the formation near the crest of the Green River anticline along faults and flowed laterally as a gravity current. A CO₂-promoted bleaching front (dotted lines) propagated laterally by advection and vertically by diffusion. (d) The preserved reaction front is subhorizontal and exhibits cusps regions associated with open fractures and granulation seams. (c) Bleached halos develop around vertical fractures. (e) Sampling across subhorizontal reaction fronts.
result in the observed Fe depletion. While some pyrite is observed in fractures, the lack of pyrite (or its Fe-bearing weathering products) throughout the bleached sandstones precludes involvement of significant H2S.

[7] Microthermometry of single-phase aqueous and two-phase liquid-vapor fluid inclusions hosted in quartz overgrowths and gypsum, petrographically linked to the bleaching, suggests that the bleaching fluid was a low temperature (≈ 27°C) brine (salinity 2.5–7 wt%) [Wigley et al., 2012]. Raman spectroscopic analysis of successfully targeted vapor bubbles reveals gas-phase compositions dominated by CO2, with minor methane (11–27 vol%) present in three inclusions (Figure 3) [Wigley et al., 2012]. The presence of a free-phase CO2 bubble suggests the aqueous phase is saturated with CO2 at the inclusion pressure. The magnitude of the Fermi-diad separation in the Raman spectra of CO2 is sensitive to the CO2 density, and this separation has been experimentally calibrated for CO2 density in the vapor phase, for gaseous CO2 to liquid CO2, in the density range of 0.01–1.20 g/cm3 [Rosso and Bodnar, 1995]. Similarly, the Raman peak position of the symmetric C–H stretching mode in CH4 is dependent on pressure, in CH4 bearing inclusions [Lu et al., 2007]. Measurements of the Fermi-diad separation in CO2 spectra reveals CO2 densities of 0.04–0.09 ±0.02 g/cm3 which equates to CO2 pressures of 2.28–4.56 ±1.03 MPa and formation depths of 202–413 ±97 m, assuming hydrostatic pressures. Inclusion pressures based on Raman peak position in CH4 bearing inclusions [Lu et al., 2007] reveal inclusion pressures of 2.59–3.98 ±1.34 MPa. At these inclusion, pressures dissolved CO2 concentrations in equilibrium with a CO2 vapor phase are 0.65–1.30 mol/L, based on the solubility models of Duan et al. [2006]. Based on burial curves for the Entrada Sandstone in the area surrounding Green River [Nuccio, 2000], modified for local variations in burial depth due to the structural high formed by the Green River anticline, the interval from which these samples were taken passed through the burial depth range 200–400 ±97 m between 2.1 and 5.5 ±1.5 Ma, and this gives an approximate age of the bleaching event.

[8] The upper bleached-red contact is sharp and subhorizontal except in regions of fractures and granulation seams, where differences in porosity and tortuosity enhance or retard vertical transport of the reaction front, respectively (Figures 2b–2c). To the east, the bleached zone terminates as a series of meter scale fingers into red sandstones, while the western extent is not exposed. It is thought that the CO2-rich brine migrated as a gravity current, with the main components of flow to the east, down dip and in the direction of regional groundwater flow [Kampman et al., 2009], with vertical transport dominated by diffusion except in areas containing faults and open fractures (Figure 2b) [Wigley et al. 2012].

[9] The contact between bleached and unbleached sandstone is characterized by stepped profiles in bulk rock Fe-concentration, hematite and K-feldspar abundances, which occur over 10–15 cm, and demarc the final position of the acid-reductive
Figure 4. (A–I) Concentration profiles for selected metals across horizontal reaction fronts (distances from the red-bleached transition) and (J–L) fracture-associated reaction fronts (distances from the core of the fracture). Red color denotes unaltered rock, yellow is bleached sandstone. (A–C) Cobalt from transects H1, H4, and H2 showing systematic enrichment within 10 cm of the bleaching reaction front. (C–H) Enrichments in metals along profile H2; (D) Cu, (E) Zn, (F) Ni, (G) Pb, (H) Sn. (I–L) Enrichments across fracture-associated bleaching reaction fronts. (I) Cu, F1; (J) Co, F1; (K) Sn, F1; (L) Cu, F2 and F3. Background concentrations in red and bleached rock are indicated. Error bars show 2σ propagated uncertainties including recovery relative to U.S. Geological Survey (USGS) standard Sco-1, repeat averages, and instrument uncertainties. Raw data and sampling localities are given in Tables S3 and S4.

A fine intergrowth of oxide and carbonate is deposited in a 2–3 cm region spanning the reaction front [Wigley et al., 2012]. Bulk rock trace metal enrichments occur systematically in a narrow zone within 5–10 cm of the upper subhorizontal contact (Figure 4a–4h), due to release of trace metals as grain coating Fe-oxides were dissolved, with subsequent redeposition at the reaction front [Wigley et al., 2012]. The enriched metals are characteristic in having high solubilities at low pH [e.g., Benjamin and Leckie, 1981], and this mobilization-redeposition is thought to be fundamentally driven...
by acid-reductive dissolution of the Fe-oxide grain coatings at low pH, with subsequent transport of the trace metals in solution and redeposition at the reaction front as pH rises due to a combination of pH buffering silicate dissolution reactions and secondary mineralization. The scale of enrichments is determined by the volume of fluid that interacted with the reaction front, which had previously reacted with unaltered rock.

[10] Bleaching also occurs around open fractures (Figure 2c), which are mineralized with carbonate and Fe-oxides, and exhibits the same geochemical and petrological signatures as the subhorizontal bleaching contacts, indicating a common parent fluid and continuity in reactive-transport processes. The fracture-associated bleaching reaction front initiated at the plane of the fracture and migrated laterally by diffusion and advection of the fluid. The asymmetry of the bleached halos is due to the transport away from the fractures being imposed on unidirectional horizontal regional flow in the aquifer [Wigley et al., 2013]. Spikes in trace-metal concentration are observed within 5–10 cm of the red-bleached transition adjacent to the fractures (Figures 4i–4l), similar to those across subhorizontal contacts. These are assumed to be related to deposition of metals released by the bleaching reaction front as it propagates from the fracture, in addition to any metals present in the bleaching fluid entering the fracture, due to the increase in pH across the reaction front.

[11] Key questions include the following: (1) What was the original distribution of the metals? (2) How, and in which phases, are the metals redeposited? (3) What fraction of the mobilized metals is redeposited? (4) Does the concentration of metals in the fluid exceed drinking water guidelines?

2. Methods

2.1. Sampling and Analytical Methods

[12] Profiles from bleached to unbleached sandstones were cored using an electric diamond-edged 40 mm core plug drill, with samples spaced at 2–15 cm intervals. Samples near to the bleached-red contact were subsampled every 4 mm using a 2 mm tungsten-carbide drill bit. Between drilling, the bit and core were cleaned with compressed air. Samples were dissolved in HF and HNO₃ using an open beaker method, and analyzed by inductively coupled plasma (ICP)-atomic emission spectroscopy (major elements) and ICP-mass spectrometry (trace elements) in Cambridge, UK.

2.2. Mass of Metals in Enriched Zones

[13] The mass of a metal in the enriched region is determined by integrating across the enriched zone, assuming a stepped concentration profile. Uncertainties are estimated by propagating the uncertainties on the background and enriched compositions. The background compositions are taken as the average within the bleached zone (bleached sandstone), and the average composition of red sandstone near to the bleached fracture (unaltered sandstone; 1σ uncertainty; Table S1). The uncertainty in peak height is propagated 1σ instrument uncertainty, 1σ of repeat analyses, and recovery relative to USGS standard Sco-1 (Table S2). The range in the fraction of each metal redeposited is calculated from a Monte-Carlo simulation (n =1000) where each parameter listed above is allowed to vary within its range in uncertainty, and the range is given by 1σ about the mean of the resulting distribution.

2.3. Sequential Extraction

[14] The nature of the minerals hosting the redeposited metals was determined by a sequential extraction procedure (modified from Tessier et al., 1979 and Förstner, 1982) applied to a selection of unaltered and bleached samples and to metal-enriched samples from subhorizontal reaction fronts. Leaching was performed at room temperature in five steps: (1) Water rinse: 1.5 mL deionized 18.2 MΩ water with continuous agitation for 2 h. (2) Exchangeable fraction: 1.5 mL 1 M sodium acetate solution (NaOAc) at pH 8.2, with continuous agitation for 3 h. (3) Carbonates: 1.5 mL 1 M NaOAc adjusted to pH 5 with acetic acid (HOAc), with continuous agitation for 7 h, repeated three times. (4) Oxides: 1.5 mL 0.1 M oxalate buffer adjusted to pH 3 with oxalic acid. Occasional agitation for 54 h, repeated three times. (5) HCl rinse (remaining carbonate/oxides): 1.5 mL 16 M HCl with continuous agitation for 2.5 h.

[15] After each leach, the sample was centrifuged for 30 min at 3000 rpm, and the liquid pipetted carefully into a cleaned centrifuge tube. The sample was then rinsed with 18.2 MΩ water, recentrifuged, and the rinse added to the leachate solution. After the final leaching step, the residue was dried down and dissolved in a combination of HF and HNO₃ in an open beaker method. The leachate solutions were evaporated to dryness and redissolved in a dilute nitric acid solution.

[16] The number of moles of a mineral phase dissolved in each sequential leaching step was
Table 1. Fraction of Each Mineral Released in Sequential Leaching Steps\(^{a}\)

<table>
<thead>
<tr>
<th></th>
<th>Average Fraction Released</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>18.2 MΩ Water Rinse</td>
</tr>
<tr>
<td>Quartz</td>
<td>0.00 (0)</td>
</tr>
<tr>
<td>K-feldspar</td>
<td>0.10 (2)</td>
</tr>
<tr>
<td>Plagioclase</td>
<td>0.00 (0)</td>
</tr>
<tr>
<td>Clay</td>
<td>0.07 (1)</td>
</tr>
<tr>
<td>Calcite</td>
<td>0.04 (1)</td>
</tr>
<tr>
<td>Dolomite</td>
<td>0.05 (1)</td>
</tr>
<tr>
<td>Fe-dolomite</td>
<td>0.00 (0)</td>
</tr>
<tr>
<td>Hematite</td>
<td>0.00 (0)</td>
</tr>
<tr>
<td>Pyrite</td>
<td>0.00 (0)</td>
</tr>
<tr>
<td>Gypsum</td>
<td>0.73 (12)</td>
</tr>
<tr>
<td>TiO₂</td>
<td>0.00 (0)</td>
</tr>
</tbody>
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\(^{a}\)Numbers in parentheses are 1σ uncertainties for the last digit, calculated from the magnitude of the misfit to bulk rock values.

determined by a modal decomposition of the compositions of the leachate solutions. This was done by minimizing the difference between theoretical and observed compositions by varying the moles of each mineral dissolved. Mineral compositions of plagioclase, K-feldspar, and carbonates are taken from electron microprobe analyses (Table S5). An average clay composition is used based on the major element chemistry of the < 2µm fraction (\(\text{Al}_1.4\text{Ca}_0.2\text{Fe}_0.5\text{Mg}_0.1\text{Na}_0.2\text{Si}_3.8\text{O}_{10}(\text{OH})_2\); recalculated to a 12 oxygen formula). Fe and Ti-oxides, pyrite and quartz are assumed to exist as pure phases.

3. Results and Discussion

3.1. Phases Dissolved During Sequential Extraction

The results of the modal decomposition of sequential leaching solutions are shown in Table 1. The water rinse predominantly dissolves soluble salts, releasing up to 86% of available S with a Ca/S ratio of ≈1. Modal analysis on water rinse solutions shows that 75% of all minerals dissolved in this step are gypsum (Table 1). The exchangeable leach removes residual soluble salts and minor amounts of calcite and silicates. It also removes loosely adsorbed cations: in some samples, up to 60% of total Sr is removed in this step. Up to 80% total Ca and 60% total Mg are removed in the carbonate leach indicating dissolution of calcite and dolomite. Approximately 95% of minerals dissolved are carbonates, along with some clays and residual dolomite. A moderate correlation (\(R^2 = 0.34\)) between Fe and S suggests dissolution of minor pyrite; however, an Fe/S ratio of ≈100 precludes pyrite dissolution as the dominant control on the Fe content of the leachate solution. The hydrochloric acid leach dissolves residual carbonate and oxide phases, and minor silicates. A larger fraction of total Fe is released in unaltered samples in this step, reflecting partial dissolution of the grain coating oxy-hydroxide phases.

3.1.1. Location of Redeposited Metals

The results of the sequential extraction experiments are typified by cobalt (Co; Figure 5). In both bleached and unbleached samples where no enrichment is observed, Co is released from carbonate, oxides, and by dissolution of the residue (Figure 5a), indicating an equal distribution between carbonate, coarse oxides, and grain-coating phases. In samples with bulk rock Co enrichment, the Co released in the steps dissolving carbonate and oxide phases is two orders of magnitude greater than background values (Figure 5b). Redeposited trace metals are therefore incorporated into the secondary oxides and carbonates growing at the bleaching reaction front [Wigley et al., 2012].

3.2. Fraction of Mobilized Metals Redeposited

The minimum fraction of each trace metal redeposited can be calculated from bleached halos surrounding fractures where the displacement distance of the reaction front is given by the perpendicular distance of the front from the fracture.
The theoretical mass of a metal mobilized along a one-dimensional path is given by the difference between the background concentrations in the red and bleached sandstones, and the displacement distance. The ratio of the predicted to observed mass of metal in the enriched zone gives a measure of the fraction of that metal redeposited at the reaction front.

Some metals analyzed (Cu, V1; Zn, V2 and V3) exhibit redeposited fractions close to unity, with the adsorbed mass being within error of the total mass dissolved (Table 2). Others (Cu, V2 and V3; Co, V1) show enrichment relative to the mass dissolved by propagation of the reaction front from the fracture. The true value lies in the upper range of calculated results as the sampling may have missed peaks in concentration, thus underestimating the mass of metal that has been redeposited. Fluids entering the fractures have previously reacted with unaltered rock and may therefore be enriched in metals and these, along with metals mobilized by the reaction front moving away from the fracture, are all redeposited.

Metals in solution exist either as free-metal cations or metal-anion complexes (e.g., with OH⁻, HCO₃⁻, or Cl⁻). Speciation modeling using the geochemical speciation program PHREEQC [Parkhurst and Appelo, 1999] with the Lawrence Livermore National Laboratory database [Johnson et al., 2000], using the calculated range in P_{CO₂} and salinity and assuming equilibrium with calcite, suggests that Sn, Zn, and Co exist as uncomplexed species, with a minor fraction of Co and Zn complexing with chloride and Zn also forming carbonate complexes (Figure S2); Cu speciation is dominated by chloride species. Realistic variations in methane concentrations do not significantly change metal speciation profiles. This shows that metal complexation does not play a major role in the redeposition process, as metals complexing with hydroxyl, carbonate, and chloride anions as well as free metal cations are all redeposited at the CO₂-promoted reaction fronts.
Transport of mobilized metals is thus controlled by the distance moved by CO2-promoted reaction fronts, rather than by the distance moved by the fluid. The difference between these length scales is controlled by the stoichiometry and kinetics of the fluid-rock reactions, and therefore the mineralogy of the host formation. In nonreactive reservoirs, mobilized metals will be transported further; however, such reservoirs are less likely to contain significant quantities of contaminants.

3.3. Vertical Transport Distances Across Subhorizontal Reaction Fronts

The observation that the metals removed from the bleached sandstone are quantitatively redeposited can be used to calculate transport distances perpendicular to the subhorizontal reaction fronts. The length of rock bleached by the fluid interacting with horizontal reaction fronts is calculated using the difference in background concentration in bleached and unbleached samples and the mass of each metal deposited at the reaction front, assuming 100% redeposition ($l = M/\Delta C$; where $l$ is the length of rock bleached (m), $M$ is the mass of the metal in the enrichment (mol m$^{-2}$), and $\Delta C$ (mol m$^{-3}$) is the difference in background concentrations between red and bleached sandstone). Some metals may have been removed by the subhorizontal advection of the gravity current although this flux may decrease near the margin of a gravity current. If so the calculation underestimates the transport distance. Four transects across horizontal reaction fronts were analyzed (H1–4), and the length of rock bleached for each unit area of reaction front is low (<0.35 m; Figure 6), consistent with the CO2-rich brine migrating as a gravity current with vertical motion dominated by diffusive processes except near open fractures.

3.4. Potential Impact on Water Quality

The enrichment of metals in the fluid is calculated from the mass of metal deposited at the reaction front and the volume of fluid required to transport this mass. The time integrated fluid flux (TIFF, the volume of fluid that has interacted with the reaction front) is calculated from the retardation factor ($R$) and the distance the front is displaced (Table 2). $R$ is defined as the volume of fluid required to displace the reaction front through a unit volume of rock, and is estimated from the reaction stoichiometry and change in brine-CO2 concentration across the reaction front. TIFF is then given by $R \times$ distance, i.e., the volume of fluid required to displace the reaction front through the observed volume of rock.

Metal concentrations in the fluid ($C_m$; mol m$^{-3}$) as the reaction front propagates are given by the following:

$$C_m = \frac{F_m}{F_f}$$

where $F_m$ is the metal flux (mol m$^{-2}$ s$^{-1}$), and $F_f$ is the fluid flux (m s$^{-1}$). The time-averaged metal flux is given by the mass of metal in the enrichment (mol/m$^2$ reaction front); and the TIFF is the corresponding $F_f$ (m$^3$/m$^2$; Table 2).

Calculated concentrations of Cu and Zn are an order of magnitude below the World Health Organization (WHO) drinking water limit (Cu: 2 mg/L, Zn: 5 mg/L [WHO, 2011], and Sn concentrations are well below acceptable limits (150 mg/L [WHO, 2003]). No guidelines exist for Co; however, calculated Co concentrations are 10 times the average composition of U.S. freshwater [ATSDR, 2004] (a worked example is presented in Appendix A).
4. Concluding Remarks

[27] The reaction fronts studied here, where fluid velocities are low and diffusive processes are important, have a high Damköhler number ($N_D$; the ratio of reaction rate to rate of advection) and a low Peclet number ($Pe$; the ratio of advective to diffusive transport). The high injection pressures in CO$_2$-storage reservoirs will result in high rates of advection, and consequently CO$_2$-promoted reaction fronts will have a lower $N_D$ and a higher $Pe$ than those documented here. However, much of the concern over contaminant mobilization relates to migration of CO$_2$ into overlying aquifers where the groundwater flow rates are likely to be considerably lower (Figure 7), being dominated by regional gradients in hydraulic head, such that reaction fronts have a transport characteristic comparable to those studied here. In this context, our results show that mobilized metals are redeposited over small length scales in secondary phases precipitating at the CO$_2$-promoted reaction fronts. Fundamentally, this redeposition process is thought to be driven by buffering of pH by fluid-rock reactions at the reaction front, although a full reactive-transport description of the system is required to fully evaluate the redeposition process. Fluxes of CO$_2$-charged waters of approximately 25 times the volume of an aquifer of the composition of the Entrada sandstone would be required for reaction fronts to pass through the aquifer and for mobile trace metals to be extracted.

Acknowledgments

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Appendix A: Calculating Metal Concentrations in the Fluid: A Worked Example

[29] Calculation of the concentration of a metal in the fluid leaving the reaction front requires an estimate of the retardation factor of the formation and knowledge of the distance moved by the reaction front and the mass of the metal under consideration deposited at the reaction front.

[30] The retardation factor, $R$, is given by the following:

$$ R = \frac{M_{\text{hem}} \cdot S}{\Delta CO_2} \quad (A1) $$

where $M_{\text{hem}}$ is the number of moles of hematite per m$^3$ formation ($\approx$40 moles/m$^3$), $S$ is the number of moles of CO$_2$ needed to dissolve 1 mole of hematite ($\approx$3.2) [Wigley et al., 2012], and $\Delta CO_2$ is the change in CO$_2$ concentration in the fluid across the reaction front ($\approx$5 mol/m$^3$) [Wigley et al., 2012]. This gives a retardation factor of approximately 25.6.

[31] The metal concentration is given by the metal flux divided by the fluid flux (equation (1)). By integrating over the duration of the bleaching event, this reduces to the following:

$$ \text{Concentration} = \frac{M}{\text{TIFT}} \quad (A2) $$

where $M$ is the mass of metal at the reaction front (kg/m$^2$) and TIFT is the time integrated fluid flux (m$^3$/m$^2$), which is given by: $R \times X$, where $X$ is the distance moved by the reaction front (m).
The mass of copper deposited at reaction front V1 is 0.00152 kg/m², and the reaction front has moved 0.506 m. The corresponding TIFF is therefore 13 m³/m², and the concentration of copper in the fluid as the reaction front propagates is 0.000117 kg/m³ or 0.117 mg/L.

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