

Multidimensional reactive transport modeling of CO₂ mineral sequestration in basalts at the Hellisheidi geothermal field, Iceland

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Two and three-dimensional field scale reservoir models of CO₂ mineral sequestration in basalts were developed and calibrated against a large set of field data. Resulting principal hydrological properties are lateral and vertical intrinsic permeabilities of 300 and 1700×10^{-15} m², respectively, effective matrix porosity of 8.5% and a 25 m/year estimate for regional groundwater flow velocity.

Reactive chemistry was coupled to calibrated models and predictive mass transport and reactive transport simulations carried out for both a 1200-tonnes pilot CO₂ injection and a full-scale 400,000-tonnes CO₂ injection scenario. Reactive transport simulations of the pilot injection predict 100% CO₂ mineral capture within 10 years and cumulative fixation per unit surface area of 5000 tonnes/km. Corresponding values for the full-scale scenario are 80% CO₂ mineral capture after 100 years and cumulative fixation of 35,000 tonnes/km². CO₂ sequestration rate is predicted to range between 1200 and 22,000 tonnes/year in both scenarios.

The predictive value of mass transport simulations was found to be considerably lower than that of reactive transport simulations. Results from three-dimensional simulations were also in significantly better agreement with field observations than equivalent two-dimensional results.

Despite only being indicative, it is concluded from this study that fresh basalts may comprise ideal geological CO₂ storage formations.

1. Introduction

Reducing anthropogenic CO₂ in the atmosphere is among the great challenges of this century (e.g. Broecker, 2007; Hoffert et al., 2002; Lackner, 2003; Pacala and Socolow, 2004; Oelkers and Schott, 2005, and references therein). Large scale capture and sequestration of atmospheric CO₂ has been proposed in an attempt to solve this problem and in situ mineral sequestration is among the suggested methods (Metz et al., 2005; Schrag, 2007). In situ mineral carbonation, injected CO₂ is combined with dissolved metals to form carbonate minerals, such as calcite (CaCO₃), dolomite (CaMg(CO₃)₂), magnesite (MgCO₃) and siderite (FeCO₃), that are stable and environmentally benign over geological time scales. Formation of these minerals can enhance the stability of injected CO₂ substantially (Metz et al., 2005; Oelkers et al., 2008).

Great advances have been made in geochemical computer simulations in recent years and decades. Numerous geochemical

Computer codes have been developed, enabling scientists to peer underground and study the complex interplay of dissolved constituents and fluids by other means than are common in the laboratory. Initially, geochemical simulations involved time-independent equilibrium speciation of multi-component systems (e.g. Wolery, 1979; Reed, 1982; Arnórsson et al., 1982). Reaction path models that capture the sequence of chemical/mineralogical states followed the time-independent equilibrium calculations, providing a way for quantitatively and sequentially interpreting irreversible processes such as chemical weathering and hydrothermal alteration (e.g. Helgeson, 1968; Helgeson et al., 1969). Lichtner (1988) linked mass transport and travel time to the reaction path approach. This approach has now been generalized to heterogeneous multi-dimensional and multi-component flow fields. Resulting reactive transport models can be applied to fundamental Earth Sciences research, as they provide an integrated way to look at the big picture in the complex interplay of material flow, transport and reactions that characterizes subsurface processes, along with assessing the relative importance of individual sub-processes (Steeffel et al., 2005). Such complex interplay is to be expected in association with CO₂ mineral sequestration in basaltic rocks and reactive transport models should therefore be ideal for simulating

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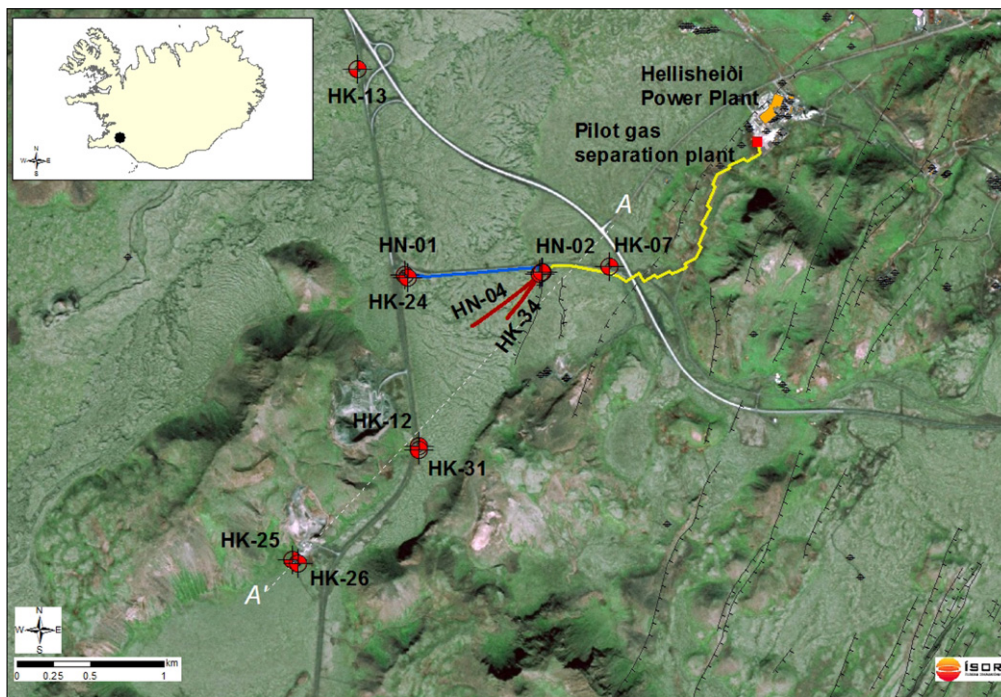


Fig. 1. Aerial map of the CarbFix injection site at Hellisheiði, SW Iceland. Fresh 25 °C groundwater, used for dissolving CO₂, will be pumped from well HN-01 and transported to well HN-02 in a pipeline (blue line). CO₂ captured at the pilot gas separation plant will be transported in a pipeline (yellow line) to well HN-02 where it will be injected along with water from HN-01. Deviation of wells HN-04 and HK-34 is depicted by red lines. Combed lines represent faults. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

its effect in the subsurface. In addition, the models provide tools to predict and optimize long-term management of injection sites, to quantify the amount of CO₂ that can be mineralized, and identify secondary minerals that compete with carbonates for cations leached from the rock matrix.

In this article, we present development of two and three-dimensional field scale reactive transport models that simulate mass and reactive transport, along with the mineral alteration associated with injecting dissolved CO₂ into basalts. The models were built using the TOUGH2 (Pruess, 1991), iTOUGH2 (Finsterle, 1999) and TOUGHREACT (Xu et al., 2006, 2011) simulators. These simulators have been extensively used in simulations of processes associated with geothermal systems and geological storage of CO₂.

2. The CarbFix project

CarbFix (e.g. Aradóttir et al., 2011; Gíslason et al., 2010; Matter et al., 2008; Oelkers et al., 2008) is a combined industrial/academic collaboration project between Reykjavík Energy, the Institute of Earth Science at the University of Iceland, Earth Institute-Lamont-Doherty Earth Observatory at Columbia University in New York and the Centre National de la Recherche Scientifique/Universite Paul Sabatier in Toulouse, that was developed in order to assess the feasibility of in situ CO₂ mineral sequestration in basaltic rocks in Iceland. The project consists of a CO₂ pilot injection, laboratory based experiments, study of natural analogs, predictive model development, numerical modeling and model validation, as well as cost analysis. Rather than injecting CO₂ directly into geological formations, CarbFix is developing a technology to dissolve CO₂ into formation fluids and well water during injection. Once dissolved, CO₂ is no longer buoyant compared to pore fluids, improving considerably security due to decreased leakage risks. This approach of solubility trapping also promotes carbonation of the host rock and thus facilitates the safe long-term sequestration of CO₂ in the subsurface.

Fig. 1 shows an aerial map of the CarbFix injection site. The site is located ca. 3 km SW of the Hellisheiði geothermal power plant, owned and operated by Reykjavík Energy. The power plant is currently generating 303 MWe and 133 MWth by tapping water and steam from a high temperature reservoir at 800–3000 m depth. The power plant annually captures some 40,000 tonnes CO₂ of volcanic origin as a by-product of the geothermal energy production. Reykjavík Energy has already invested substantially in a comprehensive network of deep and shallow wells in the area, in piping and pumping systems and in a pilot plant that separates CO₂ from other volcanic gases.

The wellfield shown in Fig. 1 comprises ideal conditions for studying the long-term effect of mineral CO₂ sequestration in basaltic rocks. Connection to Hellisheiði power plant provides access to a concentrated source of CO₂. Additionally, field data, such as drill cuttings and a calcite rich cap-rock overlying the high-temperature reservoir, suggests that mineral CO₂ sequestration already plays an important role in the evolution of the Hellisheiði geothermal system (see, e.g. Nielsson and Franzson, 2010; Gebrehiwot et al., 2010). Injecting and precipitating CO₂ into nearby formations with the objective of imitating and accelerating this natural CO₂ sequestration process should therefore be considered as an environmentally benign process. Wiese et al. (2008) carried out a study of CO₂ fixation in high-temperature geothermal systems in Iceland and found natural CO₂ sequestration in Hellisheiði to be in the range of 4100–23,500 tonnes/year or about 650,000 tonnes/km². The study showed sequestration to be most intense from 550 to 800 m below surface.

2.1. Process description

A pilot gas separation station has been built next to Hellisheiði geothermal power plant. The pilot station will separate CO₂ from the geothermal gas coming from the condensers of the power plant by sequential extraction (Aradóttir et al., 2011). In the pilot study,

approximately 5% of the total geothermal gas coming from the power station will be separated, resulting in the capture of approximately 2000 tonnes CO₂ for CarbFix on an annual basis. After its capture, CO₂ will be transported in a 3 km pipeline to the pilot injection site as high pressurized gas. There, CO₂ will be injected at depth into well HN-02. Co-injected water will divert injected CO₂ further downward, resulting in a single phase fluid entering the 400–800 m targeted storage formation (Aradóttir et al., 2011).

CO₂ injection was started in Hellisheidi in January 2012. Environmental authorities in Iceland have granted licenses for the injection. It is anticipated that the results of the CarbFix project will be used to optimize the in situ carbon mineralization process, enabling it to also work at other basalt rich sites worldwide.

2.2. Injection site infrastructure

The injection site in Hellisheidi is outfitted with ten wells, whose location is shown in Figs. 1 and 3. Fresh groundwater, used for dissolving CO₂, will be pumped from well HN-01 and transported to well HN-02 in a 1 km pipeline. There it will be injected along with CO₂ from the pilot gas separation plant. The main permeable section in HN-02 is located close to 500 m depth and most of the injected fluid should enter the reservoir there. The static water level in HN-02 is at about 100 m depth and its injectivity by gravity only is about 50 kg/s.

Wells HK-12, HK-25, HK-07 and HK-13 serve as monitoring wells and only penetrate the 5–10 °C, shallow, free surface and fresh groundwater system in the topmost 200 m. Wells HN-04, HK-34, HK-31 and HK-26, however, are drilled down through the storage formation and serve as deep monitoring wells. This reservoir is acting as confined in the short term and extensive well-logging indicates it to follow an approximately linear temperature gradient of 100 °C/km (see, e.g. Björnsson et al., 2006; Gunnarsson et al., 2011; Franzson et al., 2004). Wells HN-04, HK-34, HK-31 and HK-26 are all situated downstream from the injection well. Wells HK-31 and HK-26 are located about 1750 and 3000 m downstream from well HN-02, while the main permeable zones in the directionally drilled HN-04 and HK-34 are at about 70 and 300 m lateral distance from injection well HN-02, respectively. Well HN-04 is most permeable around 400 m depth, while the main permeable section of well HK-34 is located slightly below 500 m depth.

3. Modeling studies within CarbFix

Numerical modeling plays an important role in the CarbFix project as it provides tools to predict and optimize long-term management of the injection site as well as to quantify the amount of CO₂ that can be mineralized. A series of models have been developed within CarbFix, as summarized in Table 1.

TOUGH2 (Pruess, 1991), iTOUGH2 (Finsterle, 1999) and TOUGHREACT (Xu et al., 2006, 2011) have mainly been used for developing the field scale models presented in this article. The workflow of the model development was divided into the following five interrelated stages: (1) development of a conceptual model of regional hydrological flow conditions and reactive transport with respect to ongoing water-rock interaction, (2) compilation and validation of thermodynamic and kinetic databases, (3) parameters in hydrological model constrained by forward and inverse calibration to field data, (4) forward predictive model simulations with a fully coupled reactive mass transport model, and (5) model validation by comparison to laboratory and/or field data. The workflow stages are correlated to the individual model studies in Table 1.

Aradóttir et al. (2011) and Aradóttir et al. (in press) discuss in detail the conceptual model of the pilot CO₂ injection, and the compilation and validation of a thermodynamic database suitable for

this study. This article, therefore, only gives a brief overview of stages (1) and (2) in the model development, while focusing on stages (3)–(5).

4. Overview of conceptual and simple reservoir models

Figs. 2 and 3 show, respectively, in a cross-section, a schematic representation of the CarbFix CO₂ injection site and the conceptual reservoir model of the sequestration site (model II in Table 1). Fresh flood basalts govern the bedrock from surface down to 200 m depth. These basalts host a cold (5–10 °C) groundwater system, with the static water table located about 100 m below surface. Below lies a 200 m thick hyaloclastite layer that separates the cold groundwater system from the storage formation below due to significantly lower vertical permeability. The storage formation extends from about 400 to 800 m depth. It is composed of relatively fresh basaltic lavas interbedded with minor hyaloclastite layers/intrusions. Highest permeability is encountered around 500 m depth.

Two slug type tracer tests have been carried out at the injection site since 2007 with the objective of studying regional hydrology in the area and seeing if tracer dispersivity could identify reservoir flow type (fracture or matrix) within the storage formation. Fig. 4 shows tracer breakthrough curves in well HN-04 from both tracer tests. Rezvani Khalilabad et al. (2008) used a one dimensional and multipath tracer dispersion model to study recovery of the first tracer test (model I in Table 1). The model predicted most of the storage formation to consist of a large volume of relatively homogeneous porous media. A low volume and fast breakthrough path was found to channel only about 3% of the tracer flow between wells HN-02 and HN-04, and thus only plays a minor role in the storage formation.

When the primary rock comes in contact with the injected weakly acidic, carbonated fluid, dissolution reactions occur, leaching cations out of the rock matrix. As dissolution reactions proceed, protons are consumed, the pH of formation fluids increases, and concentration of leached cations builds up. At certain concentrations, the water becomes supersaturated with respect to secondary minerals like carbonates, which begin to precipitate. The efficiency of this reactive chemical transport is highly sensitive to the surface area available for water-rock interaction.

Natural analogs provide important insight into the secondary mineralogy associated with CO₂ rich and CO₂ depleted water-basalt interaction and hence, which minerals are likely to compete with carbonates for dissolved cations in in situ CO₂ mineral sequestration (model III in Table 1). Aradóttir et al. (2011) cover in detail the alteration mineralogy associated with basalt weathering under low and elevated CO₂ conditions. Simple oxides and hydroxides, clay minerals, carbonates and zeolites are among common alteration minerals.

5. Model setup

5.1. Flow conditions

In this study, two and three-dimensional models of the CarbFix injection site were developed (models V, VI, VII, VIII and IX in Table 1). The models are aligned in the direction of regional groundwater flow, which is from north-northeast to south-southwest. Figs. 3 and 5 show layer distribution in the three-dimensional model and the geographical location of the CarbFix numerical models, respectively. The three-dimensional model consists of eight layers, from 100 to 900 m depth, whereas the two-dimensional model is a simplified version of the three-dimensional model, consisting of a single horizontal layer between 425 and 525 m depth. Each layer consists of 6198 elements. Apart from varying

Table 1

Overview of modeling studies within the CarbFix project and correlation to workflow stages.

Model study	Stage	Description	Simulator	D ^a	RT ^b	Temperature	Features	Reference
I	(1)	Tracer dispersion model	TRINV ^c	1D	No	Constant	Multipath dispersion model used for studying reservoir flow paths	Rezvani Khalilabad et al. (2008)
II	(1)	Conceptual, site specific, hydrological model					Extensive research on hydrology of CarbFix site based on continuous well-logging and two tracer tests	Aradóttir et al. (2009, 2011) and references therein
III	(1)	Conceptual, site specific, sequestration model					Extensive literature review on natural analogs of CO ₂ -water-basalt interaction to identify potential alteration minerals	Aradóttir et al. (2011) and references therein
IV	(2)	CO ₂ -water-basalt interaction at Hellisheidi	TOUGHREACT	1D	Yes	Gradient	Batch simulations at T, P corresponding to 100–1000 m depth at Hellisheidi to assess quality of thermodynamic database	Aradóttir et al. (in press)
V	(3)	Hydrological reservoir model	iTOUGH2	3D	No	Gradient	Hydrological properties calibrated using seven datasets on water level fluctuations and downstream tracer recovery	This study, Aradóttir et al. (2009) and Aradóttir (2010)
VI	(4)	Injection/production during pilot CO ₂ injection	TOUGH2	3D	No	Gradient	Simulations of different injections and production scenarios aimed at accelerating groundwater flow during the pilot injection	Aradóttir (2010)
VII	(4)	Pilot CO ₂ injection	TOUGHREACT	2D	Yes ^d	Constant (30 °C)	Fully coupled simulations of CO ₂ -water-basalt interaction associated with pilot injection at Hellisheidi	This study
VIII	(4)	Pilot CO ₂ injection	TOUGHREACT	3D	Yes ^d	Gradient	Fully coupled simulations of CO ₂ -water-basalt interaction associated with pilot injection at Hellisheidi	This study
IX	(4)	Full scale CO ₂ injection	TOUGHREACT	2D	Yes ^d	Constant (30 °C)	Fully coupled simulations of CO ₂ -water-basalt interaction associated with full scale injection at Hellisheidi	This study

^a Dimensionality of model.^b Reactive transport simulations.^c Arason et al. (2004).^d Equivalent mass transport simulations were also carried out.

thickness, all layers have identical element geometry. Generally, layers containing feed zones in the three-dimensional model are set narrow but those that do not contain feed zones are thicker. Elements at the north-northeast and south-southwest boundaries are set at constant temperature, pressure and chemical composition, as well as the top and base layers in the three-dimensional model. The mesh is rectangular, except around injection well HN-02, where it is radial in order to get reasonable distribution of injected fluids and dissolved constituents in the near vicinity of the well. Elements close to the injection well are small (<1 m) but their size gradually increases further away from the well. The horizontal hydraulic gradient that drives natural groundwater flow through the bedrock has been estimated to be 5 m/1000 m, making pressure at the models north-northeast boundary 3.36 bar higher than at the south-southwest boundary. Estimation of the hydraulic gradient is based on water table measurements in wells HN-01, HN-02, HN-04, HK-31 and HK-26 that extend over several years.

Stratigraphy of the CarbFix model is based on the conceptual model shown in Fig. 3. In the topmost 800 m, rock formations mainly consist of relatively fresh basaltic hyaloclastite formations and lava flows. Hydrological parameters of the lavas were

calibrated using field data, but porosity and permeability values for the hyaloclastite formations were taken from Frolova et al. (2005), and set to be 10% and 30 mD, respectively.

Reaction induced matrix permeability changes are calculated from changes in porosity using ratios of permeabilities calculated from the Carman-Kozeny relation (Bear, 1988), ignoring changes in grain size, tortuosity and specific surface area.

5.2. Geochemical system

Aradóttir et al. (2011) carried out an extensive literature review of water-basalt interaction at low and elevated CO₂ conditions with the objective of predicting which minerals are likely to dissolve and precipitate in situ CO₂ mineral sequestration. Subsequently, Aradóttir et al. (in press), developed and validated a thermodynamic database describing mineral reactions of interest for CO₂-water-basalt interaction. All primary and secondary minerals in the database were added to the reactive transport models and their dissolution reactions written in terms of the same basis species as used by Aradóttir et al. (in press) and Gysi and Stefánsson (2011), i.e. H₂O, H⁺, HCO₃⁻, Al(OH)₄⁻, Fe²⁺, Mg²⁺, Na⁺, K⁺, SiO_{2(aq)},

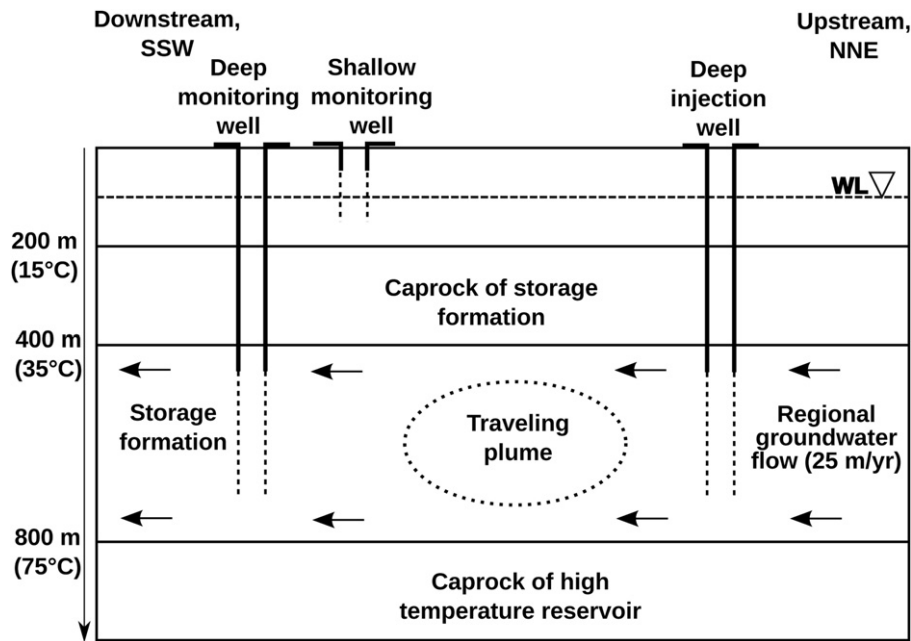


Fig. 2. A simplified schematic representation of the CarbFix pilot CO₂ injection in Hellisheidi, Iceland. Regional groundwater flow will carry injected carbonated water downstream. Cations leached from the rock matrix by the weakly acidic carbonated plume will react with CO_{2(aq)} and form carbonates.

Cl⁻ and O_{2(aq)}. The minerals are listed in Table 2. Selection of basis species is primarily based on the findings of Stefánsson et al. (2001), who studied dominant aqueous species in natural waters in the basaltic terrain of Iceland.

Initial water composition of all model elements at time zero is that of well HN-01 in Hellisheidi, which is given in Table 3. Carbonated injection water has the same composition, except that CO₂ concentration correspond to about Henry's law saturation at 30 bar CO₂ partial pressure, resulting in approximately 1 M total concentration of H₂CO₃, HCO₃⁻ and CO₃²⁻. As a result of acidification, pH of the boundary water drops down to 3.6. Initial rock is composed of plagioclase, pyroxene, olivine and basaltic glass. Basaltic glass composition is that of Oelkers and Gíslason (2001), whereas the specific composition of plagioclase, pyroxene and olivine are those given by Stefánsson (2001). Table 4 shows the initial mineral volume composition used in model simulations.

5.2.1. Kinetics of mineral dissolution and precipitation

Dissolution and precipitation of all minerals except allophane, Al(OH)_{3(am)}, *p*-antigorite, calcite, Fe(II) and Fe(III) hydroxides, imogolite and kaolinite is kinetically controlled. Kinetic rates are a product of the rate constant and reactive surface area, according to the following general rate expression, which is transition state theory based (Lasaga et al., 1994; Steefel and Lasaga, 1994):

$$r = kA \left[1 - \left(\frac{Q}{K} \right)^\theta \right]^\eta \quad (1)$$

where r is rate of dissolution or precipitation, k is the temperature dependent rate constant, A is the specific reactive surface area, K is the equilibrium constant for the dissolution/precipitation reaction taking place and Q is the reaction quotient. θ and η must be determined by experiment but are often set to unity. As

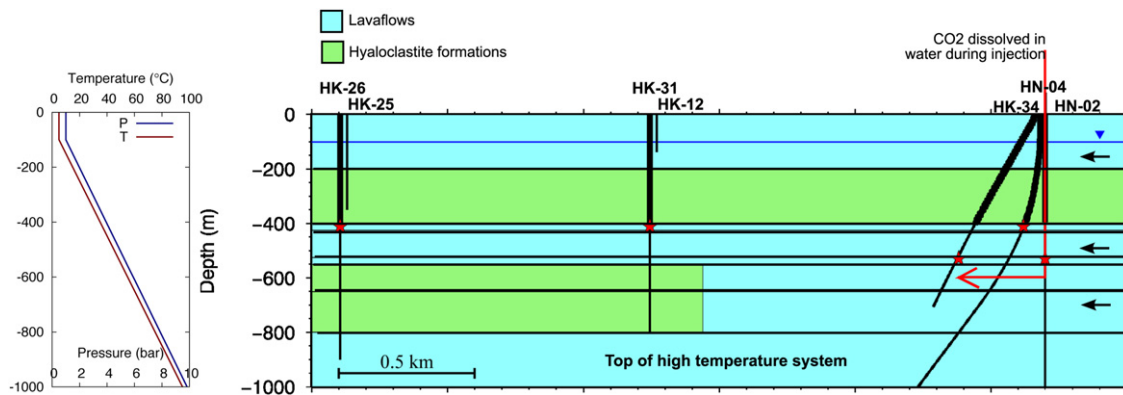


Fig. 3. The CarbFix conceptual model, drawn along A-A' in Fig. 1. CO₂ is dissolved in water during injection into HN-02, resulting in a single phase fluid entering the storage formation (400–800 m depth). Black arrows show direction of regional groundwater flow and red stars main feed-zones in wells. Layers in the three dimensional CarbFix model are designated with horizontal black lines. Layers that contain feed zones are narrower than others. Pressure at the north-northeast boundary is 3.36 bar higher due to the horizontal hydraulic gradient that drives natural groundwater flow through the bedrock. Temperature and pressure gradients are based on extensive well-logging, which show a near-linear increase of temperature and pressure from surface and down to the top (cap-rock) of the high temperature geothermal system.

Table 2

Chemical composition of the minerals and aqueous species considered in this study. See Aradóttir et al. (2011) and Aradóttir et al. (in press) for further information on mineral selection and thermodynamic data.

Group	Mineral	Chemical composition
Primary minerals and glasses	Basaltic glass	SiAl _{0.36} Fe(II) _{0.19} Mg _{0.28} Ca _{0.26} Na _{0.08} O _{3.31}
	Plagioclase	Ca _{0.70} Na _{0.30} Al _{1.70} Si _{2.30} O ₈
	Pyroxene	Ca _{0.35} Mg _{0.42} Fe _{0.23} SiO ₃
	Olivine	(Mg _{0.80} Fe _{0.20}) ₂ SiO ₄
Silicates	Allophane	Al ₂ O ₃ (SiO ₂) _{1.22} (H ₂ O) _{2.5}
	Amorphous silica	SiO ₂
	<i>p</i> -Antigorite ^a	Mg ₃ Si ₂ O ₅ (OH) ₄
	Ca-montmorillonite ^b	Ca _{0.167} Al _{1.67} Mg _{0.33} Si ₄ O ₁₀ (OH) ₂
	Celadonite ^c	KMgAlSi ₄ O ₁₀ (OH) ₂
	Fe-Celadonite ^c	KFeAlSi ₄ O ₁₀ (OH) ₂
	Fe-chlorite ^c	Fe ₅ Al ₂ Si ₃ O ₁₀ (OH) ₈
	Imogolite	Al ₂ SiO ₃ (OH) ₄
	Kaolinite	Al ₂ Si ₂ O ₅ (OH) ₄
	K-montmorillonite ^b	K _{0.33} Al _{1.67} Mg _{0.33} Si ₄ O ₁₀ (OH) ₂
	Mg-chlorite ^c	Mg ₅ Al ₂ Si ₃ O ₁₀ (OH) ₈
	Mg-montmorillonite ^b	Al _{1.67} Mg _{0.5} Si ₄ O ₁₀ (OH) ₂
	Moganite	SiO ₂
Na-montmorillonite ^b	Na _{0.33} Al _{1.67} Mg _{0.33} Si ₄ O ₁₀ (OH) ₂	
Quartz	SiO ₂	
Hydroxides	Amorphous Al hydroxide	Al(OH) ₃
	Fe(II) hydroxide	Fe(OH) ₂
	Fe(III) hydroxide	Fe(OH) ₃
Carbonates	Calcite	CaCO ₃
	Dolomite	CaMg(CO ₃) ₂
	Magnesite ^d	MgCO ₃
	Siderite ^d	FeCO ₃
Zeolites	Analcime	Na _{0.96} Al _{0.96} Si _{2.04} O ₆ ·H ₂ O
	Ca-chabazite ^e	CaAl ₂ Si ₄ O ₁₂ ·6H ₂ O
	Ca-heulandite ^e	CaAl ₂ Si ₇ O ₁₈ ·6H ₂ O
	Laumontite	CaAl ₂ Si ₄ O ₁₂ ·4.5H ₂ O
	Mesolite	Ca _{0.666} Na _{0.666} Al ₂ Si ₃ O ₁₀ ·2.667H ₂ O
	Na-chabazite ^e	Na ₂ Al ₂ Si ₄ O ₁₂ ·6H ₂ O
	Na-heulandite ^e	Na ₂ Al ₂ Si ₇ O ₁₈ ·5H ₂ O
	Natrolite	Na ₂ Al ₂ Si ₃ O ₁₀ ·2H ₂ O
	Stellerite ^f	Ca ₂ Al ₄ Si ₁₄ O ₃₆ ·14H ₂ O
	Stilbite ^f	Ca ₂ NaAl ₅ Si ₁₃ O ₃₆ ·16H ₂ O

^a See Gunnarsson et al. (2005).

^b Forms a solid solution between Ca, K, Mg and Na end-members.

^c Forms a solid solution between Mg and Fe end-members.

^d Magnesite and siderite form a solid solution.

^e Forms a solid solution between Ca and Na end-members.

^f Stellerite and stilbite form a solid solution.

dissolution and precipitation of minerals are often catalyzed by H⁺ (acid mechanism) or OH⁻ (base mechanism), the rate constant in Eq. (1) is often written as the sum of three mechanisms:

$$\begin{aligned}
 r = & k_{25}^{\text{nu}} \exp \left[\frac{-E_A^{\text{nu}}}{R} \left(\frac{1}{T} - \frac{1}{298.15} \right) \right] \\
 & + k_{25}^{\text{H}} \exp \left[\frac{-E_A^{\text{H}}}{R} \left(\frac{1}{T} - \frac{1}{298.15} \right) \right] a_{\text{H}}^{\text{nuH}} \\
 & + k_{25}^{\text{OH}} \exp \left[\frac{-E_A^{\text{OH}}}{R} \left(\frac{1}{T} - \frac{1}{298.15} \right) \right] a_{\text{OH}}^{\text{nuOH}} \quad (2)
 \end{aligned}$$

where nu, H and OH denote neutral, acid and base mechanisms, respectively. E_A is activation energy, k_{25} the rate constant at 25 °C, R is the gas constant, T absolute temperature and a activity of a species. Gíslason and Oelkers (2003) measured and summarized previously measured basaltic glass dissolution kinetics far from equilibrium at temperature ranging from 6 to 300 °C and pH between 1 and 11. The resulting rate law describing basalt glass dissolution is:

$$r_{\text{geo}} = kA \exp \left[\frac{-E_A}{RT} \right] \left(\frac{a_{\text{H}^+}^3}{a_{\text{Al}^{3+}}} \right)^{1/3} \left(1 - \frac{Q}{K} \right) \quad (3)$$

Table 3

Total dissolved component concentration^a (in mmol/L) in initial water.

Component	Concentration (mmol/L)
pH/ ^o C	9.26/21.6
CO ₂	2.122
Si	0.503
Na	1.890
O ₂	0.007
Cl	0.276
Ca	0.134
Mg	0.184
K	0.018
Fe	0.0001
Al	0.0006

^a Well HN-01, Hellisheidi, Reykjavik Energy database.

The rate law was implemented in TOUGHREACT using a general form of a species dependent rate constant that is coded in TOUGHREACT:

$$\begin{aligned}
 r = & k_{25}^{\text{nu}} \exp \left[\frac{-E_a^i}{R} \left(\frac{1}{T} - \frac{1}{298.15} \right) \right] \\
 & + \sum_i k_{25}^i \exp \left[\frac{-E_a^{\text{H}}}{R} \left(\frac{1}{T} - \frac{1}{298.15} \right) \right] a_{\text{H}}^{\text{nuH}} \prod_j a_{ij}^j \quad (4)
 \end{aligned}$$

Table 4
Initial rock mineral composition and potential alteration minerals along with parameters for calculating kinetic rates for mineral dissolution and precipitation.

Mineral ^a	Vol% of solid	A (cm ² /g)	Parameters for kinetic rate law								
			Neutral mechanism		Acid mechanism			Base mechanism			
			k_{25} (mol/m ² /s)	E_a (kJ/mol)	k_{25} (mol/m ² /s)	E_a (kJ/mol)	$n(H^+)$	k_{25} (mol/m ² /s)	E_a (kJ/mol)	$n(H^+)$	
<i>Primary</i>											
Olivine ^b	13	20	2.291×10^{-14}	79.0							
Plagioclase ^b	43	20	1.377×10^{-15}	69.8	3.140×10^{-13}	65.0	0.457	1.256×10^{-18}	71.0	-0.572	
Pyroxene ^b	39	20	7.762×10^{-15}	40.6	4.365×10^{-10}	96.1	0.710				
Basaltic glass ^c	5	20			$k_{25} = 2.730 \times 10^{-11}$						
					$E_a = 25.5$						
					$n(H^+) = 1.00, n(Al^{+3}) = -0.33$						
<i>Secondary</i>											
Celadonite ^d		10,000	1.660×10^{-13}	35.0	1.047×10^{-11}	23.6	0.34	3.020×10^{-17}	58.9	-0.40	
Montmorillonite ^d		10,000	3.890×10^{-15}	48.0	1.950×10^{-12}	48.0	0.220	3.890×10^{-12}	78.0	-0.130	
Moganite ^e		1000	1.653×10^{-13}	70.5							
Quartz ^e		1000	1.223×10^{-12}	80.5							
SiO _{2(am)} ^f		1000	7.32×10^{-13}	60.9							
Dolomite ^d		500	2.951×10^{-8}	52.2	6.457×10^{-4}	14.4	1.0				
Magnesite ^d		500	4.571×10^{-10}	23.5	4.169×10^{-10}	14.4	1.0				
Siderite ^g		500	1.259×10^{-9}	62.76	1.590×10^{-4}	45.0	0.9	2.512×10^{-16}	71.0	-0.572	
Zeolites ^h		1000	1.585×10^{-12}	58.0	1.995×10^{-8}	58.0	0.70	5.495×10^{-15}	58.0	-0.30	

^a Allophane, Al(OH)_{3(am)}, antigorite, calcite, Fe(II) and Fe(III) hydroxides, imogolite and kaolinite precipitate under equilibrium.

^b Forsterite, albite and diopside data from Palandri and Kharaka (2004) used for olivine, plagioclase and pyroxene. Rate constants reduced by 3 orders of magnitude.

^c From Gíslason and Oelkers (2003). Rate constant was renormalized with respect to BET surface area and reduced by 3 orders of magnitude.

^d From Palandri and Kharaka (2004).

^e From Gíslason et al. (1997).

^f From Rimstidt and Barnes (1980). Precipitation occurs under the free energy rate law of Carroll et al. (1998).

^g From Knauss et al. (2005).

^h All zeolites assumed to have the same rate law as heulandite from Palandri and Kharaka (2004).

where i denotes the species dependent mechanism and j specific species to which the rate constant depends on. Parameters used for the kinetic rate expression of different minerals are given in Table 4.

Forsterite, albite and diopside dissolution data from Palandri and Kharaka (2004), based mostly on data from Pokrovsky and Schott (2000), Chou and Wollast (1985) and Knauss et al. (1993), were used for olivine, plagioclase and pyroxene, respectively. Rate constants for albite were multiplied by a factor of five to account for the effect of anorthite in the plagioclase solid solution as anorthite and albite have very different kinetic behaviors. The same does not apply to olivine and pyroxene end-members. The rate law of

the most abundant end-member was therefore chosen to be representative of olivine and pyroxene solid solutions. The rate law of Gíslason and Oelkers (2003) was used for basaltic glass but the geometric rate constant reported by Gíslason and Oelkers (see Eq. (3)) was renormalized with respect to BET surface area.

Rate constants of all primary minerals and glasses were reduced by 3 orders of magnitude, as simulations with the reported rate constants gave way too rapid dissolution during preconfiguration of model VI in Table 1. Simulations of rain water interacting

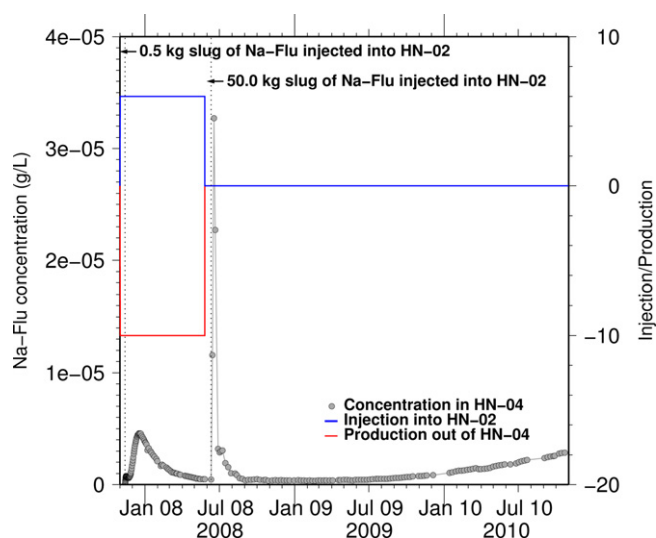


Fig. 4. Recovery of Na-fluorescein injected into HN-02 in observation well HN-04. The preliminary tracer test was launched on November 13, 2007 and ran until May 26, 2008. The natural gradient tracer test was launched on June 10, 2008 and is ongoing. The first hump in each of the tests most likely represents contribution of a small fracture channeling about 3% of the injected tracer while the rest is transported by matrix flow.

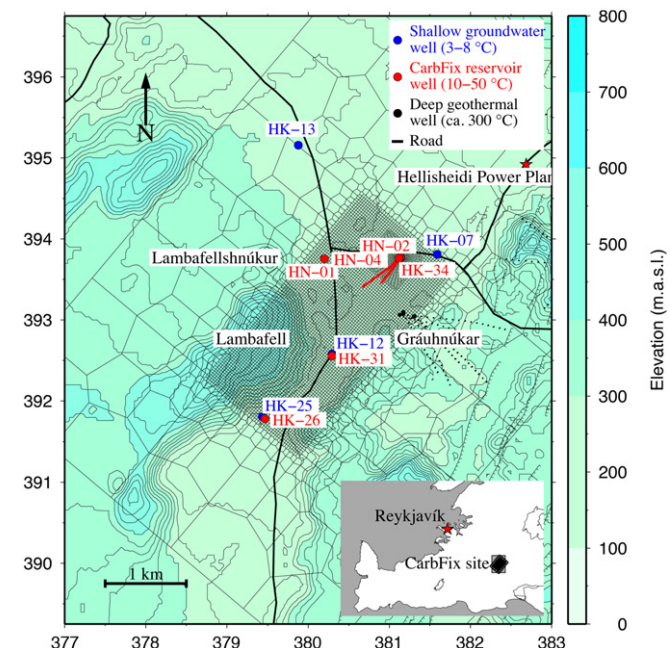


Fig. 5. Geographic location of the CarbFix field model and distribution of elements in the model's mesh. The model is aligned with the regional groundwater flow direction (NNE-SSW) and covers all CarbFix wells.

with the primary rock composition given in Table 4 were carried out with different rate reduction factors and resulting water composition compared to the chemical composition of groundwater of a known age after corresponding simulation times. Simulations using 3 orders of magnitude reduction factor gave excellent match with field observations and the same rate reduction factor was therefore implemented to the rate constants of the primary minerals in Table 4. An equivalent way of reducing simulation rates would have been to reduce mineral reactive surface areas by three orders of magnitude. Use of a rate reduction factor of this sort is justifiable in light of the fact that experimentally determined weathering rates are commonly two to four orders of magnitude faster than field-derived rates as has, e.g. been shown by White and Brantley (2003), White et al. (1996) and Brantley (1992).

Rate law parameters for moganite and quartz were taken from Gíslason et al. (1997) but from Rimstidt and Barnes (1980) for $\text{SiO}_{2(\text{am})}$. Rate law parameters for siderite are those reported by Knauss et al. (2005) but rate law parameters for other minerals are from Palandri and Kharaka (2004) who summarized previous rate measurements from the literature and refitted the data. All zeolites were assumed to have the same rate law as heulandite.

Mineral reactive surface areas in the subsurface are generally unknown. In the current study, reactive surface area of primary minerals and glasses was assumed to be $20 \text{ cm}^2/\text{g}$, which is in agreement with the work of Sonnenthal et al. (2005), who calculated reactive surface areas of several minerals assuming a cubic array of truncated spheres constituting the rock framework. Resulting values were roughly in the range of $10\text{--}100 \text{ cm}^2/\text{g}$. Surface areas of precipitating minerals are generally much higher than those of the primary rock. In the current study, surface areas of secondary clay minerals, zeolites and carbonates were assumed to be 10,000, 1000 and $500 \text{ cm}^2/\text{g}$, respectively. Secondary $\text{SiO}_{2(\text{s})}$ minerals were assumed to have a surface area of $1000 \text{ cm}^2/\text{g}$.

When the aqueous phase supersaturates with respect to a certain secondary mineral, a small volume fraction of 1×10^{-6} is used for calculating a seed surface area for the new phase to grow (Xu et al., 2010). The precipitation of secondary minerals is represented using the same kinetic expression as that for dissolution, except for $\text{SiO}_{2(\text{am})}$ which precipitates under the free energy rate law of Carroll et al. (1998). As precipitation rate data for most minerals are unavailable, parameters for neutral pH rates were employed to describe precipitation. This is a critical but necessary assumption because of lack of data on precipitation kinetics.

6. Simulations

Simulations carried out in this study can roughly be divided into two categories. First, is pure hydrological and tracer recovery modeling aimed at determining primary reservoir properties such as porosity and permeability (model VI in Table 1). Substantial volume of initial state and transient field data was available for constraining the model parameters. Reactive chemistry was then coupled to the calibrated hydrological model and forward predictive simulations carried out on CO_2 -water-basalt interaction associated with pilot and full scale CO_2 injection at Hellisheidi (models VII, VIII and IX).

The numerical models were allowed to reach natural state flow by running no injection/production simulations for 10,000–100,000 years or until pressure and temperature changes between time-steps became negligible. Steady state output was used as initial conditions for other simulations. Batch geochemical simulations of water-rock interaction (model IV) were also carried out before starting fully coupled reactive mass transport simulations in order to pre-react the measured water composition of well HN-01 to a steady state with the primary minerals listed in Table 4.

6.1. Model calibration and well production scheme design

Hydrological properties of the storage formation were calibrated using iTOUGH2 (Finsterle, 1999) to simulate tracer tests that have been ongoing in Hellisheidi since 2007 (model V). The iTOUGH2 simulator is well suited for such simulations as it is capable of solving fluid and heat flow both in forward and inverse mode. The calibration was built on seven datasets on water level fluctuations due to sudden injection and/or production stops as well as downstream tracer breakthrough and recovery curves. The EOS1 equation of state was applied along with the 'two water' option of TOUGH2. This approach was chosen as it involves the simplest way of adding a conservative tracer to a flow problem. The resulting model is based on an integrated form of the mass and energy conservation equations (Lichtner, 1996; Björnsson et al., 2006).

6.2. Predictive mass transport and reactive transport simulations

Reactive chemistry was coupled to the calibrated hydrological model and TOUGHREACT used for predictive reactive transport simulations. The TOUGHREACT simulator solves the same fluid and heat flow equations as iTOUGH2 but also solves equations describing chemical reactions. Simulations were carried out in two (models VII and IX) and three dimensions (model VIII) and involved the pilot injection test as well as a full-scale injection of all CO_2 currently emitted from Hellisheidi geothermal power plant.

In simulations of the pilot CO_2 injection, 3 kg/s of carbonated water ($p_{\text{CO}_2} \sim 30 \text{ bar}$) were modeled to be injected continuously into well HN-02 for 6 months. Well HN-04 was modeled to produce 1 kg/s continuously for the first 12 months of the pilot test, whereas 1 kg/s production out of well HK-34 was turned on 6 months into the pilot test and went continuously on for 18 months (see Fig. 7). In simulations of a full-scale CO_2 sequestration, 40 kg/s of carbonated water ($p_{\text{CO}_2} \sim 30 \text{ bar}$) were injected continuously into HN-02 for 10 years. No downstream wells were produced in this model. Well HN-01 provided water for injection in both modeling scenarios. CO_2 concentration of injected water was set to be at Henry's law saturation at indicated CO_2 pressure as all gas injected into well HN-02 dissolves during injection.

Mass transport simulations, equivalent to the reactive transport simulations described above, were carried out in order to identify the difference between aqueous concentrations in mass transport and reactive transport simulations. The mass transport simulations only involve mass and heat transport but do not take chemical reactions into account. Dissolved CO_2 is expected to be retained by solid phases in CO_2 mineral sequestration, causing downstream concentrations to be lower in reactive transport simulations. Comparing results of equivalent mass transport and reactive transport simulations provides an assessment of how reliable predictive mass transport simulations are for a reactive system. Such information is important because fully coupled reactive transport simulations on large, complex systems can be impractical to run due to heavy computational requirements. In such cases, mass transport simulations have to suffice.

7. Results

7.1. Model calibration and well production scheme design

Calibration of the hydrological field model (V) with field data resulted in horizontal permeability of lava flows to be 300 mD and a vertical permeability of 1700 mD (mD is milli Darcy, $1 \times 10^{-15} \text{ m}^2$). Active matrix porosity was estimated to be 8.5% . The fact that basalt flows in Iceland are commonly composed of vertical basalt columns surrounded by vertical fracture networks could explain why higher

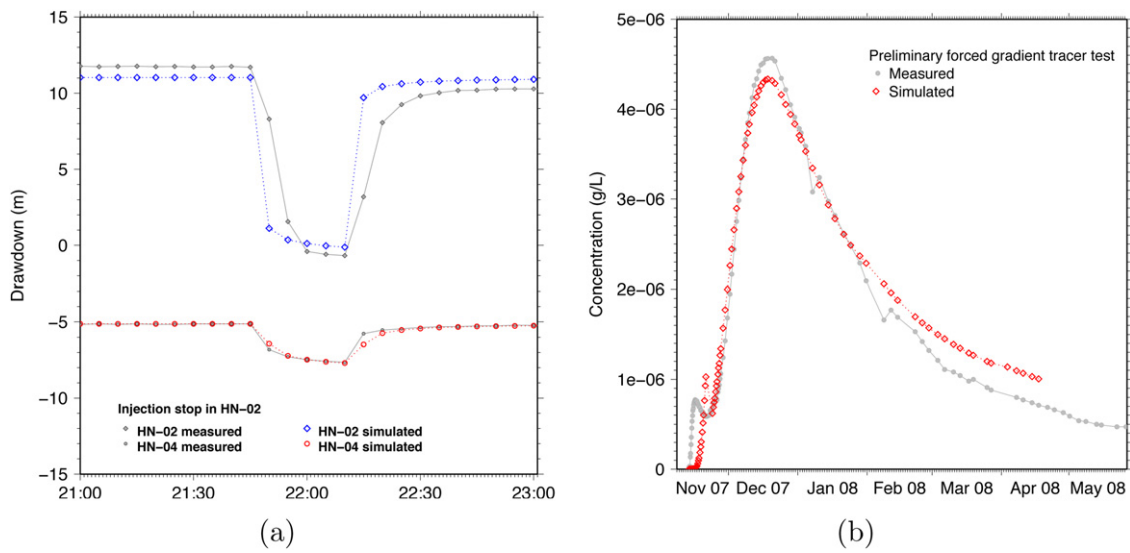


Fig. 6. (a) Comparison between simulated and measured drawdown during a sudden injection stop in HN-02 on January 8, 2008. (b) Comparison of simulated and measured tracer breakthrough in well HN-04 during the first tracer test in Hellisheidi.

vertical permeability was obtained in calibration of hydrological parameters. One way to incorporate this heterogeneity into the numerical model involves using an idealized dual-porosity model that considers fractures in a porous medium (Xu and Pruess, 2001). The method was, however, not used in the current study due to its heavy computational requirements.

Fig. 6 shows comparison between measured and simulated values for two of the calibration datasets used in inverse simulations. Match between measured and simulated values is satisfactory considering that the single porosity model used cannot account for heterogeneities due to, e.g. fractures providing fast flowing but low volume paths between injection and observation wells.

As calibrated hydrological parameters of the storage formation predict groundwater velocity to be low (25 m/year), plans call for accelerating groundwater flow during the pilot CO₂ injection by producing downstream wells. The calibrated numerical model is a valuable tool for designing optimal injection and production schemes (model VI). Simulations showed that production out of HN-04 significantly speeds up downstream recovery of non-reactive tracers, whereas producing HK-34 does not have the same effect. Therefore, additional expenses due to production out of HK-34 are hardly justifiable until injected dissolved constituents have reached well HN-04. Darcy's law is fully applicable due to the low groundwater flow in the storage formation. Its Reynolds number is orders of magnitude lower than 1 (Fetter, 2001).

7.2. Predictive mass transport and reactive transport simulations

The current version of the TOUGHREACT code only allows for single central processing unit (CPU) simulations, which causes difficulties in field scale reactive transport simulations due to the computational requirements of such simulations. Three dimensional mass transport and reactive transport simulations of the 10 year long full-scale CO₂ injection turned out to be too computationally intensive for the single CPU version of TOUGHREACT to handle, as the continuous 40 kg/s injection of carbonated water required small time steps for reasonable distribution of injected species. Three-dimensional reactive transport simulations of the pilot CO₂ injection took several months to run, and were thus only barely practical.

Fig. 7 summarizes predicted HCO₃⁻ concentrations (total dissolved DIC) in wells HN-02, HN-04, HK-34, HK-31, HK-26, and

HN-01 for both the pilot and the full-scale CO₂ injections. Kinks in curves belonging to the pilot injection are due to start of production out of HK-34 after 6 months and stop of production out of HN-04 after 1 year. In the case of the full-scale injection, injected CO₂ reaches water supply well HN-01 after 4 years of continuous production. Fig. 7 will be discussed in detail in the subsequent sections.

7.2.1. Pilot injection – 3D

In the case of the pilot injection, three-dimensional reactive transport simulations predict the CO₂ contaminated plume to reach well HK-34 after 3–4 years. No plume effects are observed in other observation wells.

Fig. 7 shows that only a slight difference exists between HCO₃⁻ concentrations predicted by mass transport and reactive transport simulations in the near vicinity of the injection well (HN-02), but that the difference increases as the CO₂ plume travels downstream. This indicates CO₂ to be retained by solid phases. The predictive value of mass transport simulations of a reactive system therefore decreases as plumes are carried downstream.

Dimensionality effects were studied by comparing HCO₃⁻ concentrations predicted by equivalent two and three-dimensional simulations. Fig. 7 exhibits that a considerable difference exists between concentrations due to dimensionality effects. The three-dimensional field model predicts much earlier plume arrival times in downstream wells than the two-dimensional one. Comparison with measured tracer breakthrough curves shows that good agreement exists between measured tracer arrival times and the plume arrival times predicted by the three-dimensional model, which hence are more realistic than those predicted by the two-dimensional model. Vertical permeability, which was calibrated to be five times higher than the horizontal permeability, is not accounted for in the two-dimensional model, as it consists of a single horizontal slice. This is likely the main reason for slower breakthrough in the two-dimensional model.

Fig. 8 shows cross sectional view of reservoir HCO₃⁻ concentration after 1 year, 5 years, and 10 years in the CO₂ pilot injection, as predicted by three-dimensional reactive transport simulations. The figure shows that the CO₂ plume travels faster deeper in the reservoir (below 500 m depth) than at shallower depths. This is, e.g. due to production out of HN-01 which supplies solvent water for CO₂ injected into HN-02. Increased diffusion due to higher random

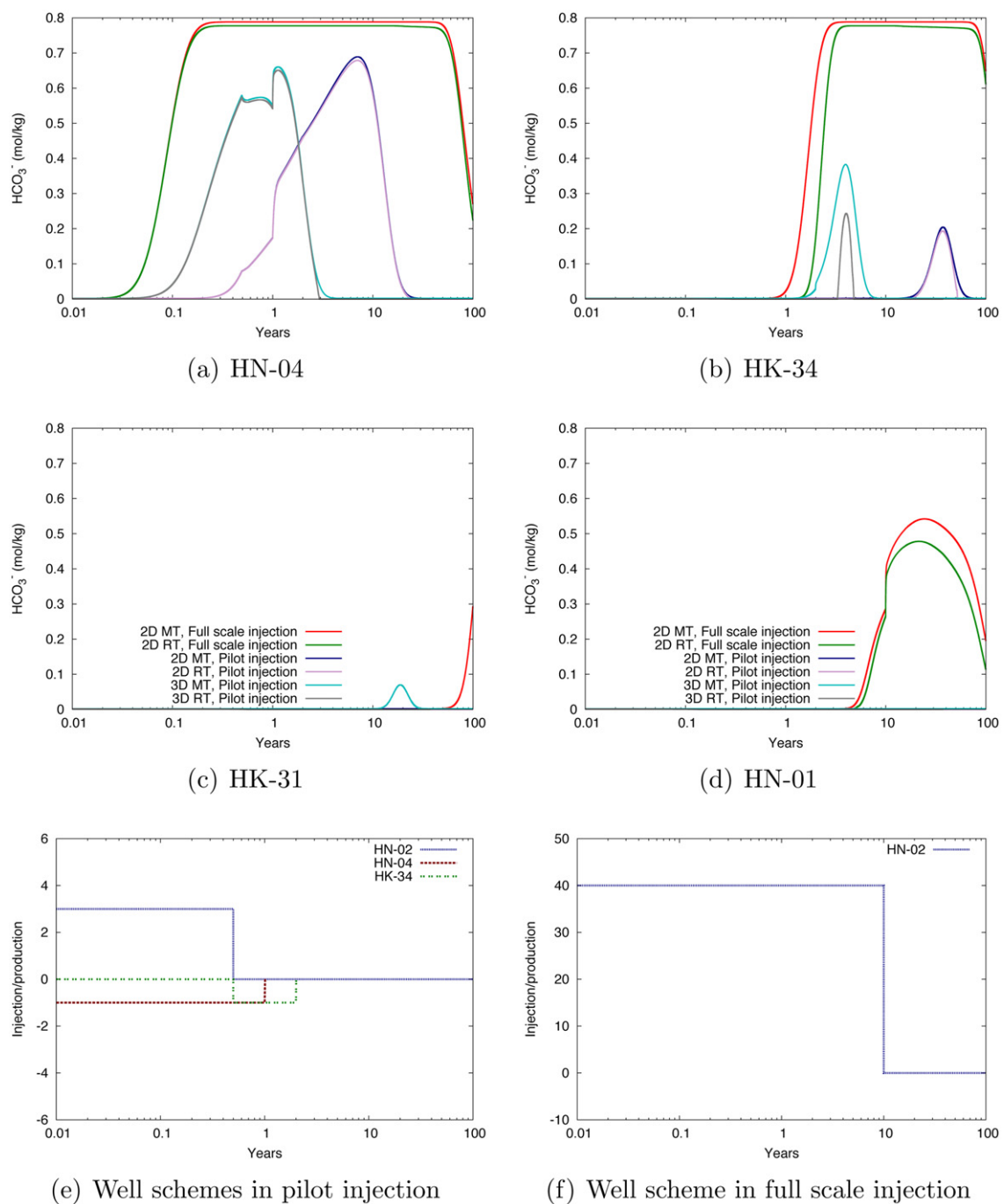


Fig. 7. Predicted HCO_3^- concentrations (total dissolved DIC) in wells in two and three-dimensional mass transport (MT) and reactive transport (RT) simulations of the pilot and full-scale CO_2 injection. Kinks in curves belonging to the pilot injection are due to start of production out of HK-34 after 6 months and stop of production out of HN-04 after 1 year. Subfigures (e) and (f) show well production and injection schemes for the pilot and full scale injection, respectively. (a) HN-04. (b) HK-34. (c) HK-31. (d) HN-01. (e) Well schemes in pilot injection. (f) Well scheme in full scale injection.

kinetic energy at higher reservoir temperature could also play a role. Plume concentration decreases after CO_2 injection is stopped due to dilution, diffusion and chemical reactions, resulting in the plume to be disappearing after 10 years.

Fig. 9 shows primary and secondary mineral abundance predicted by three-dimensional reactive transport simulation of the pilot CO_2 injection along with estimated reservoir sequestering efficiency (model VIII). Initially, the primary rock constituted of 43 vol%, 39 vol%, 13 vol%, and 5 vol% plagioclase, pyroxene, olivine and basaltic glass, respectively, of the reservoir (ca. $6.5 \text{ km} \times 4 \text{ km} \times 0.8 \text{ km}$). The volume percentages correspond to

1.0×10^{13} , 1.5×10^{13} , 7.2×10^{12} , and 2.8×10^{12} moles, respectively. The reservoir bedrock is relatively fresh, but not free of secondary minerals. Clays, simple hydroxides, calcite, quartz, and zeolites are among commonly found alteration minerals in well cuttings collected in the reservoir (e.g. Alfredsson et al., 2007; Helgadóttir et al., 2009). Secondary minerals were not included in the primary rock composition, as quantitative data was not available. Initial interaction between the primary rock minerals and pore water does however, lead to a ‘correction’ of the primary rock composition toward a more inherent state, which includes clay minerals, simple hydroxides and oxyhydroxides, calcite and SiO_2 phases ($\text{SiO}_2(\text{am})$,

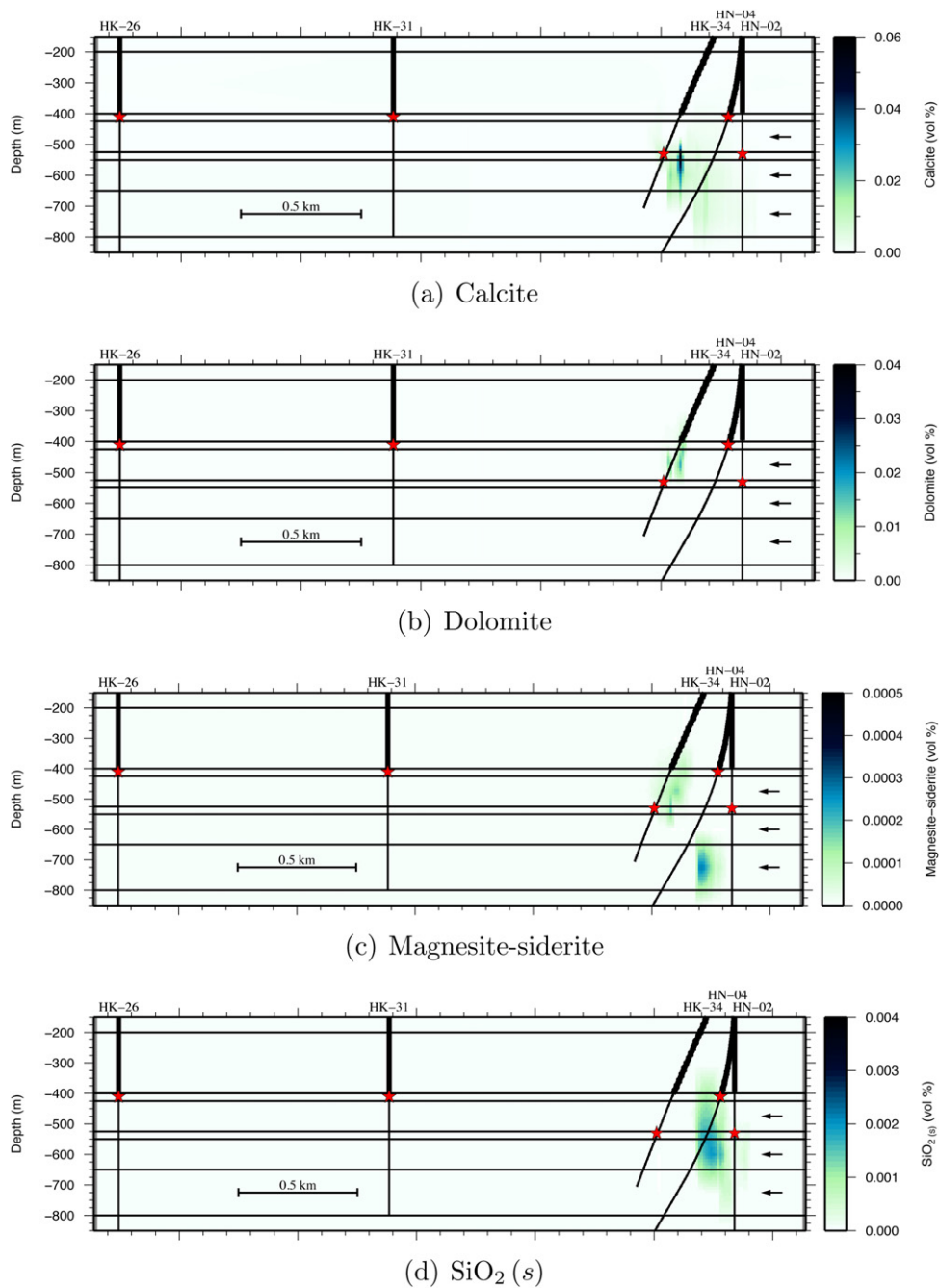


Fig. 8. Cross sectional view of modeled reservoir HCO_3^- concentration in the CO_2 pilot injection after (a) 1 year, (b) 5 years, and (c) 10 years, as predicted by three-dimensional reactive transport simulations. The cross section is drawn along A–A' in Fig. 1. The figure shows how injected carbonated water travels downstream as a plume with regional groundwater flow. Carbonate precipitation and molecular diffusion lead to significant dilution of the plume within 5 years and its disappearance within 10 years. (a) 1 year. (b) 5 years. (c) 10 years.

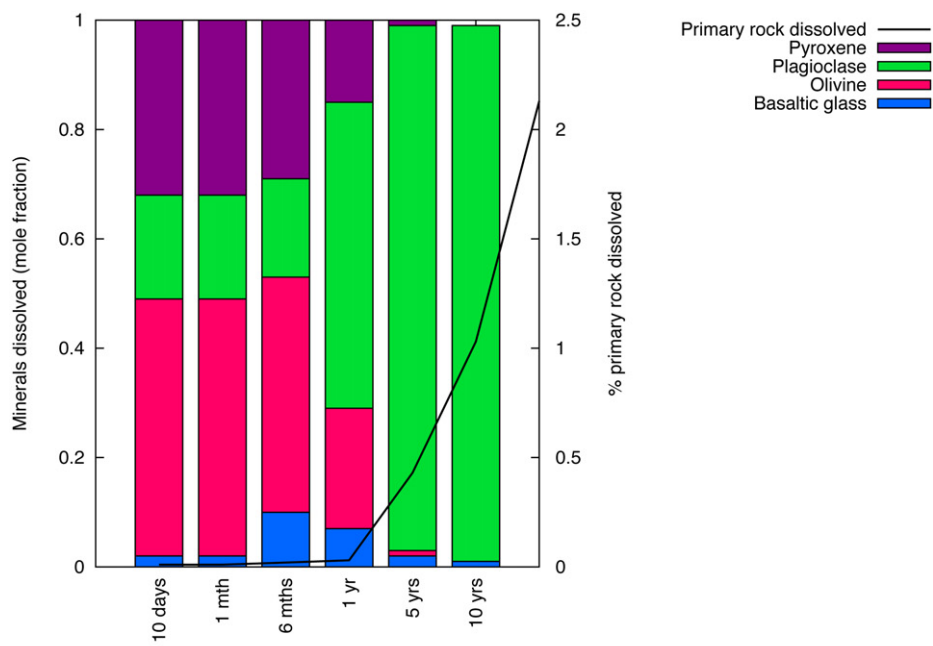
quartz and moganite). The fact that the primary rock evolves to its inherent state, suggests both the model and its thermodynamic database to be of proper quality.

Initially, all primary phases dissolve and hydroxides, carbonates and silicon oxides precipitate as the system evolves to its inherent state. Plagioclase and basaltic glass dissolution gradually increases throughout the simulation time, whereas dissolution of pyroxene and olivine levels out. After 5 years, plagioclase dissolution has become considerably greater than dissolution of other primary phases.

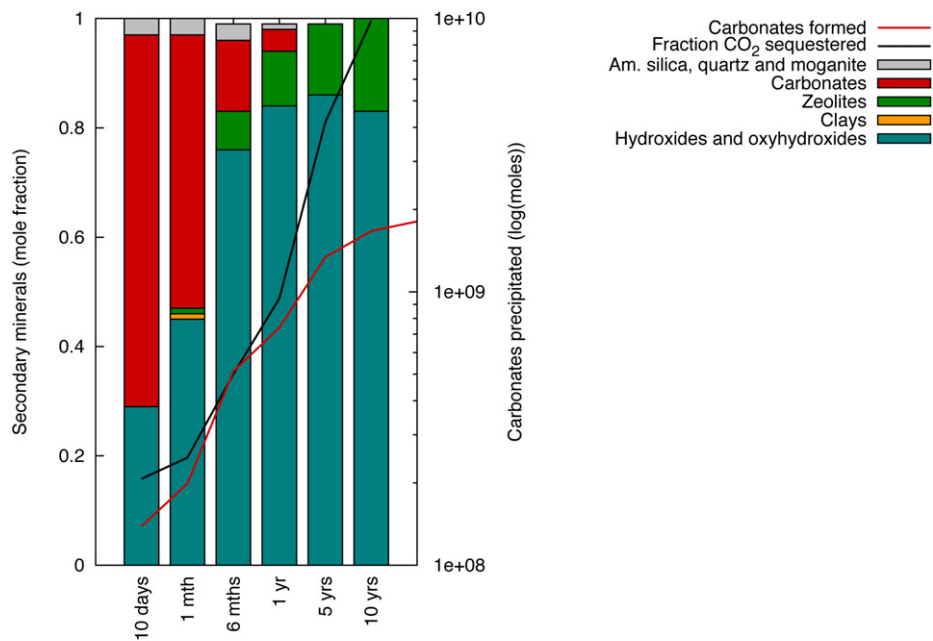
The abundance of simple hydroxides and SiO_2 phases ($\text{SiO}_{2(\text{am})}$, quartz and moganite) increases steadily throughout the whole simulation. Zeolites begin to form after 1 month and rapidly become the

second most abundant alteration mineral group in the reservoir. Carbonate formation increases gradually and all the 1100 tonnes of CO_2 injected are predicted to have been sequestered after 10 years. Calcite is most predominant among carbonates but dolomite and a Fe-rich magnesite–siderite solid solution also form to a lesser degree.

Calcite precipitation is most abundant below 500 m depth for the first 5 years but then it spreads out toward 400 m depth. Highest calcite concentrations are observed between wells HN-04 and HK-34. Dolomite precipitates in large quantities at the front of traveling CO_2 plume but dissolves again as the plume passes by. Magnesite–siderite solid solution precipitates within the CO_2 plume. Calcite and magnesite–siderite phases remain



(a) Primary mineral abundance



(b) Secondary mineral abundance

Fig. 9. Primary and secondary mineral abundance based on a full model mole balance in three-dimensional reactive transport simulation of the pilot CO₂ injection. (a) Primary mineral abundance. (b) Secondary mineral abundance.

stable thereafter. *p*-Antigorite (Gunnarsson et al., 2005) precipitation is generally associated with calcite precipitation in the simulations. Hydroxides form equally through lava flows in the reservoir and aluminum hydroxide is most abundant. Increased silica precipitation is observed behind the moving CO₂ plume. Amorphous silica is most abundant but quartz and moganite also form. The horizontal zeolite zones so commonly found in Iceland and initially described by Walker (1960) are reproduced in the simulations. Mesolite and heulandite are the most abundant zeolites. Fig. 10 shows a cross sectional reservoir view of selected

precipitated minerals in the pilot injection 5 years from injection, as predicted by three-dimensional reactive transport simulations (model VIII).

7.2.2. Full scale injection – 2D

Fig. 11 shows primary and secondary mineral abundance predicted by two-dimensional reactive transport simulation of a 10 year long full-scale CO₂ injection of 40,000 tonnes/year (model IX). Initial interaction between the primary rock and pore water result in the primary rock to evolve to its inherent state, as

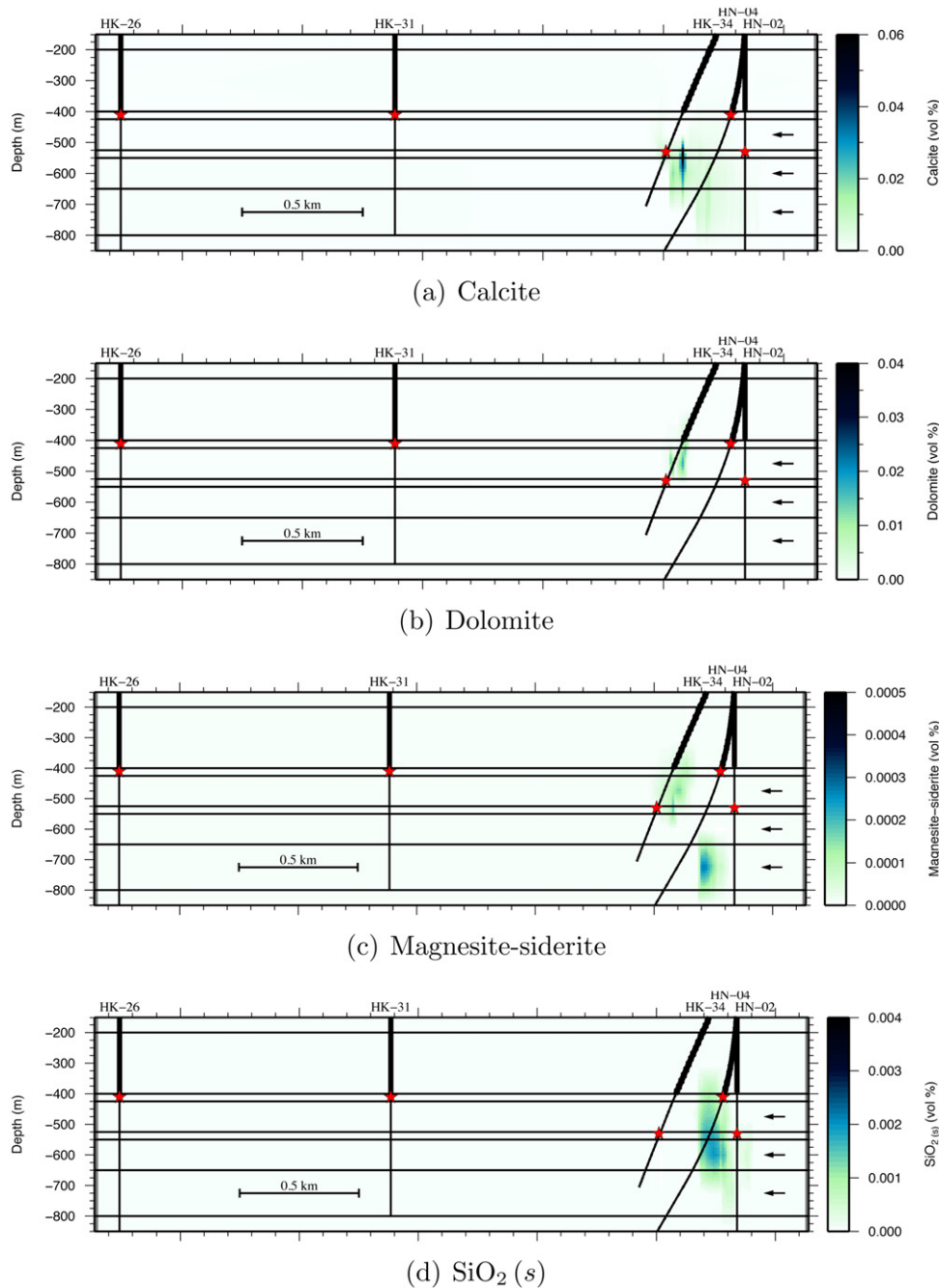
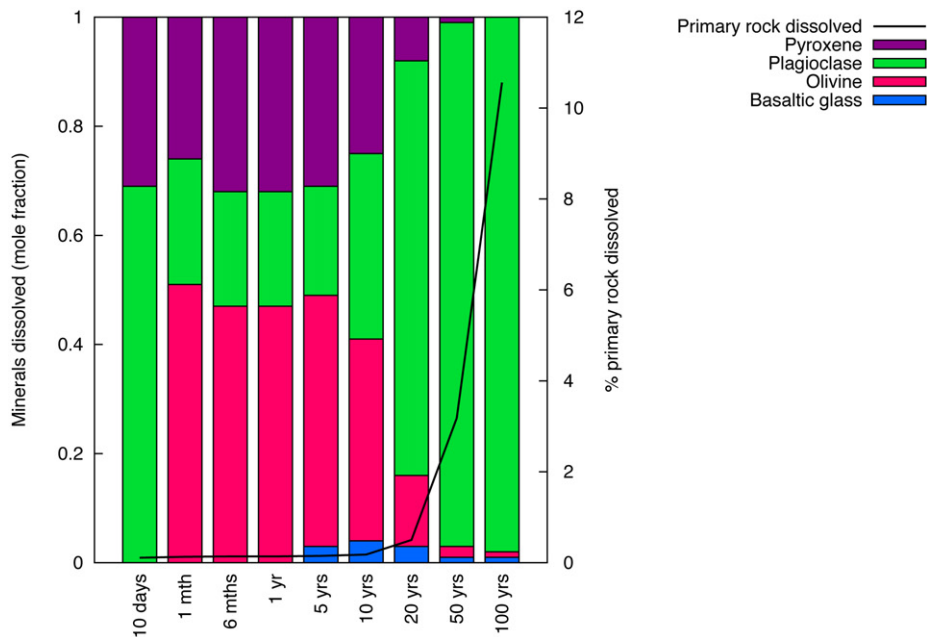


Fig. 10. Cross sectional reservoir view of selected precipitated minerals in the CO₂ pilot injection 5 years from injection, as predicted by three-dimensional reactive transport simulations. The cross section is drawn along A–A' in Fig. 1. The figure shows different minerals forming at different temperature and pressure conditions. Dolomite eventually dissolves again, whereas other carbonates remain stable. (a) Calcite. (b) Dolomite. (c) Magnesite-siderite. (d) SiO_{2(s)}.

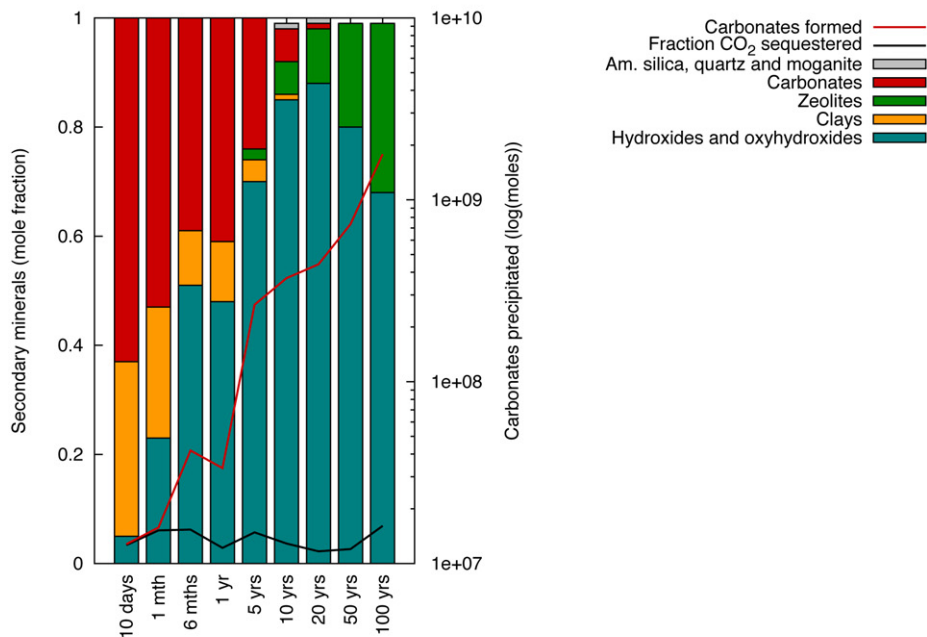
happened in the pilot simulation discussed above. Results from two-dimensional simulations of the full scale injection somewhat differ the three-dimensional ones from the pilot injection, e.g. due to different CO₂ abundance in the reservoir concentrations. Temperature effects most likely also play a role, as the two-dimensional model only consists of a single slice between 400 and 500 m depth, and thus involves lower temperature than are found at greater depth in the three-dimensional model. This could, e.g. explain the higher fractional abundance of clay minerals in two-dimensional simulations than in three-dimensional runs. Celadonite and other clay minerals are typically more common above 500 m depth in the CarbFix reservoir. Another possible explanation for higher fractional clay abundance in two-dimensional simulations

compared to three-dimensional simulations is the initial dominant plagioclase dissolution, which is likely to result in high Al activity and clay formation.

Basaltic glass dissolution is not observed until after 1 year. Delay of glass dissolution compared to three-dimensional simulations are due to lower simulation temperature in the two-dimensional layer. Plagioclase dissolution gradually becomes most extensive, as was also observed in the three-dimensional pilot injection simulations. Clay minerals are abundant and zeolite precipitation becomes extensive after approximately 10 years. Clays are initially Mg-rich (Mg-chlorite and celadonite), but their Fe-content gradually increases until clay solid solutions become equally Fe- and Mg-rich. Carbonates precipitation involves



(a) Primary mineral abundance



(b) Secondary mineral abundance

Fig. 11. Primary and secondary mineral abundance in a two-dimensional reactive transport simulation of a 10 year full-scale CO₂ injection from Hellisheidi power plant. (a) Primary mineral abundance. (b) Secondary mineral abundance.

calcite, dolomite and magnesite–siderite solid solution. Calcite and dolomite are the most and least abundant carbonates, respectively. About 80,000 tonnes CO₂ have been sequestered into carbonates after 100 years. As the two-dimensional model only considers a conservative sequestration volume consisting of a 100 m thick slice between 400 and 500 m depth, it is likely to underestimate the sequestering capacity of the reservoir. The storage formation is four times thicker, so optimally it can be assumed to sequester four times more CO₂ or 320,000 tonnes CO₂ in 100 years.

8. Discussion

The reactive transport simulations carried out in this study indicate in situ CO₂ mineral sequestration in basalt to be among viable solutions to manage anthropogenic CO₂ emissions. The models predict precipitation of alteration minerals to occur at a considerable distance from the injection well, which indicates that clogging in its near vicinity should not be a problem. CO₂ sequestration rate is modeled to range between 1200 and 22,000 tonnes/year, a value in good agreement with the natural sequestration rate

of the Hellisheidi geothermal system, which [Wiese et al. \(2008\)](#) estimated to be 4100–23,500 tonnes/year. Furthermore, models predict cumulative mass of CO₂ fixed in the storage formation per unit surface area in the pilot and full scale injections to be approximately 5000 tonnes/km² and 35,000 tonnes/km², respectively. Natural fixation per unit storage area in Hellisheidi was estimated by [Wiese et al. \(2008\)](#) to have accumulated to 650,000 tonnes/km² over the past 70,000–140,000 years. The sequestering capacity of the storage formation in Hellisheidi is therefore likely to become significantly greater with time than predicted by 100 year long simulations.

Anthropogenic CO₂ emissions in 2010 were projected to be 35 Gtonnes by [Friedlingstein et al. \(2010\)](#). In global context the results of the current study thus indicate that on an annual basis some 50,000–100,000 km² of a few hundred meter thick basaltic formations are needed in order to sequester all anthropogenic CO₂ emissions. When put in perspective to the very large volumes of dry land basalts accessible worldwide, it is not unrealistic to presume that mineral sequestration of CO₂ can develop into being one of the industrially viable solutions available in mitigating the atmospheric CO₂ levels. However, as sequestration was found to be most efficient between approximately 40–80 °C, injection wells should be drilled to a corresponding depth. The storage formation in Hellisheidi also is able to disperse the carbonated water injected, thus maximizing surface area of the reactive transport. Such favorable conditions are harder to find as basalts get buried and older.

Simulated alterations are in agreement with observations from natural analogs. Initial alteration minerals include those commonly found in the CarbFix reservoir at Hellisheidi, which are typical for water–basalt interaction at low CO₂ conditions. Carbonates that form are predominantly calcite – the only carbonate commonly found in Icelandic basalts. In reservoir subdomains containing high CO₂ concentration and low pH, a Fe-rich magnesite–siderite solid solution also forms along with dolomite at the plume front, and these results are in agreement with findings from the basalt hosted petroleum reservoir in Greenland ([Rogers et al., 2006](#)). Clay minerals, simple oxides and oxyhydroxides are predicted to compete with magnesite–siderite solid solution for Mg and Fe leached from primary minerals, whereas zeolites compete with calcite for dissolved Ca.

[Figs. 9 and 11](#) show that plagioclase dissolves in largest quantities of primary minerals, and in fact its dissolution becomes orders of magnitude greater than for other minerals with increasing simulation time. Recent work by [Gudbrandsson et al. \(2011\)](#) shows that at low pH, olivine dissolution dominates plagioclase dissolution, which was not the case in the lower temperature two-dimensional simulations carried out in the current study. Therefore, indications are that modeled plagioclase dissolution is too extensive, and as a result thermodynamic and kinetic relations for plagioclase need to be re-evaluated. The excessive plagioclase dissolution leads to a rise in pH, and a build-up of Al, Si, Ca, and Na ions within the aqueous phase. As a result, Al-hydroxide, zeolites and potentially also clays, become supersaturated and precipitate in what is likely to be too massive amounts.

Extensive monitoring during and after the CO₂ pilot injection will provide the basis for the testing, validating and further calibrating the two- and three-dimensional mass transport and reactive transport models discussed in this contribution. Plans call for drilling an exploratory well into the storage formation after the pilot CO₂ injection. The results of reactive transport simulations suggest that it would be wise to wait for about 3–5 years to drill such a well as then collected core samples will be more likely to contain larger amounts of sequestered carbonates and associated alteration minerals. Furthermore the simulations indicate that the exploratory well should be drilled

in the north-eastern vicinity of the main feed zone of well HK-34.

The current study indicates how reactive transport models can be used for deepening understanding of subsurface processes associated with CO₂ injection and sequestration as well as designing management schemes for injection sites. However, it is clear that if reactive transport simulations of complex subsurface processes are to become more practical, a parallel version of codes such as TOUGHREACT is essential. Such tools are also necessary if complex calibration of, e.g. reactive surface areas, rate-law parameters, and stochastic permeability is to become feasible.

9. Uncertainties associated with reactive transport modeling

Large uncertainties are generally associated with reactive transport modeling, in part due to uncertainties in laboratory measured values, but also due to significant differences between values measured in the field and those measured in the laboratory. Often one must thus accept significant uncertainties in parameters describing permeability, diffusivity, reactive surface area and mineral dissolution/precipitation rates. Hence, it follows that the results of reactive transport calculations can have uncertainties as high as several orders of magnitude. Much of this uncertainty can, however, be overcome by obtaining extensive and specific system and/or site specific physical and chemical parameters, as was done in the current study.

A better understanding of reaction mechanisms, and in particular precipitation mechanisms, is an important factor in decreasing uncertainties associated with reactive transport modeling. Discrepancies between rates measured in the laboratory and those measured in the field also need to be further identified. Information on reactive surface areas in the field is also scarce, and often difficult to measure, as it changes with time due to chemical reactions. [Xu et al. \(2005\)](#) performed sensitivity simulations regarding ankerite and dawsonite precipitation kinetics, which indicated changes in reaction rates to result in small changes in precipitated amounts. The reason for this is that precipitation requires reactants whose availability is controlled by the slow dissolution of aluminosilicate minerals.

10. Conclusions

A series of modeling studies has been carried out with the objective of quantitatively studying CO₂ mineral sequestration in basalts. Two and three-dimensional field scale reservoir models of a proposed CO₂ sequestration formation at Hellisheidi geothermal field were developed through a multistage numerical modeling approach. The models were calibrated against a large set of field and laboratory data. Resulting principal hydrological properties are lateral and vertical intrinsic permeabilities of 300 and 1700 × 10⁻¹⁵ m², respectively, effective matrix porosity of 8.5% and a 25 m/year estimate for the regional groundwater flow velocity.

Predictive numerical modeling results guide in decision making, indicating, e.g. that it is feasible to pump downstream observation wells in order to expedite recovery of the injected carbonated water. Thereby, the planned pilot CO₂ injection study will generate urgently needed field data to validate reactive numerical models. Predictive reactive transport simulations for both a 1200-tonnes pilot CO₂ injection and a full-scale 400,000-tonnes CO₂ injection scenario show that injected carbonated water is capable of leaching cations from the basaltic rock matrix. This eventually results in pore water becoming super-saturated with respect to carbonates, into which CO₂ is sequestered. Reactive transport simulations of the pilot injection predict 100% mineral capture of injected CO₂ within

10 years, whereas two-dimensional simulations of the full-scale injection predict as much as 80% capture within 100 years.

The predictive value of mass transport simulations was found to be considerably lower than that of equivalent reactive transport simulations as injected CO₂ is retained by solid phases. Furthermore, results from three-dimensional simulations were in significantly better agreement with field observations than results from equivalent two-dimensional simulations. Simulated alterations are in agreement with observations from natural analogs, providing confidence in the developed models and some verification of their accuracy. Clay minerals, simple oxides and oxyhydroxides are predicted to compete with magnesite–siderite solid solution for Mg and Fe leached from primary minerals, whereas zeolites compete with calcite for dissolved Ca.

Despite being only indicative it is concluded from this study that relatively fresh and glassy basaltic formation of high porosity may comprise ideal conditions for geological storage of CO₂.

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References

Alfredsson, H.A., Hardarson, B.S., Franzson, H., 2007. Well HK-31 in Threngslí, Helliðshéidi – Strata, Alteration and Permeability. Report ISOR-2007/030.

Aradóttir, E., Sonnenthal, E., Björnsson, G., Gunnlaugsson, E., Jónsson, H., 2009. Development of a coupled reactive fluid flow model for mineral CO₂ capture in Helliðshéidi, Iceland. In: Proceedings, TOUGH Symposium 2009, Lawrence Berkeley National Laboratory, Berkeley, CA, September 14–16.

Aradóttir, E.S.P., 2010. Carbfix Reservoir Model – Hydrological Parameters and Pumping Schemes for Experimental CO₂ Injection. Report 2010-03. Reykjavík. Report available from author.

Aradóttir, E.S.P., Sigurdardóttir, H., Sigfússon, B., Gunnlaugsson, E., 2011. CarbFix – a CCS pilot project imitating and accelerating natural CO₂ sequestration. *Greenhouse Gas.: Sci. Technol.* 1, 105–118.

Aradóttir, E.S.P., Sonnenthal, E.L., Jónsson, H.I., 2012. Development and evaluation of a thermodynamic dataset for phases of interest in CO₂ sequestration in basaltic rocks. *Chem. Geol.* 304–305, 26–38.

Arason, T., Björnsson, G., Axelsson, G., Bjarnason, J.O., Helgason, P., 2004. ICEBOX. Geothermal Reservoir Engineering Software for Windows. A User's Manual. Iceland GeoSurvey Report ISOR-2004/014. Prepared for the United Nations University Geothermal Training Programme, Reykjavík, Iceland.

Arnórsson, S., Sigurdsson, S., Svavarsson, H., 1982. The chemistry of geothermal waters in Iceland. I. Calculation of aqueous speciation from 0 to 370 °C. *Geochim. Cosmochim. Acta* 46, 1513–1532.

Bear, J., 1988. *Dynamics of Fluids in Porous Media*. Dover Publications, New York.

Björnsson, G., Gunnlaugsson, E., Hjartarson, A., 2006. Applying Hengill geothermal model in power plant decision making an environmental impact studies. In: Proceedings, TOUGH Symposium 2006, Berkeley, CA, 15–17 May.

Brantley, S.L., 1992. Kinetics of dissolution and precipitation - experimental and field results. In: Kharaka, Y., Maest, A. (Eds.), *Water-Rock Interaction, Proceedings of the 7th International Symposium*, Park City, Utah, July 13–18, A.A. Balkema, Rotterdam.

Broecker, W., 2007. Climate change: CO₂ arithmetic. *Science* 315, 1371.

Carroll, S., Mroczek, E., Alai, M., Ebert, M., 1998. Amorphous silica precipitation (60 to 120 °C): comparison of laboratory and field rates. *Geochim. Cosmochim. Acta* 62, 1379–1396.

Chou, L., Wollast, R., 1985. Steady-state kinetics and dissolution mechanisms of albite. *Am. J. Sci.* 285, 963–993.

Fetter, C.W., 2001. *Applied Hydrogeology*. Prentice-Hall.

Finsterle, S., 1999. *ITOUGH2 User's Guide*. LBNL-40040. Lawrence Berkeley National Laboratory, Berkeley, CA.

Franzson, H., Björnsson, G., Hafstad, T., Sigurdsson, O., Thordarson, S., Blischke, A., 2004. Helliðshéidi Power Plant: Well HN-1. Internal Report in Icelandic.

Friedlingstein, P., Houghton, R.A., Marland, G., Hackler, J., Boden, T.A., Conway, T.J., Canadell, J.G., Raupach, M.R., Ciais, P., Le Quééré, C., 2010. Update on CO₂ emissions. *Nat. Geosci.* 3, 811–812.

Frolova, J., Ladygin, V., Franzson, H., Sigurdsson, O., Stefánsson, V., Shustrov, V., 2005. Petrophysical properties of fresh to mildly altered hyaloclastite tuffs. In: *World Geothermal Congress 2005*, Anatolia, Turkey.

Gebrehiwot, K., Koestono, H., Franzson, H., Mortensen, A.K., 2010. Borehole geology and hydrothermal alteration of well HE-24, Helliðshéidi geothermal field, SW-Iceland. In: *Proceedings World Geothermal Congress 2010*, Bali, Indonesia, 25–29 April.

Gíslason, S., Heaney, P., Oelkers, E., Schott, J., 1997. Kinetic and thermodynamic properties of moganite, a novel silica polymorph. *Geochim. Cosmochim. Acta* 61, 1193–1204.

Gíslason, S., Oelkers, E., 2003. Mechanisms, rates and consequences of basaltic glass dissolution: II. An experimental study of the dissolution rates of basaltic glass as a function of temperature. *Geochim. Cosmochim. Acta* 67, 3817–3832.

Gíslason, S.R., Wolff-Boenisch, D., Stefánsson, A., Oelkers, E.H., Gunnlaugsson, E., Sigurdardóttir, H., Sigfússon, B., Broecker, W.S., Matter, J.M., Stute, M., Axelsson, G., Fridriksson, T., 2010. Mineral sequestration of carbon dioxide in basalt: a pre-injection overview of the CarbFix project. *Int. J. Greenhouse Gas Control* 4, 537–545.

Gudbrandsson, S., Wolff-Boenisch, D., Gíslason, S., Oelkers, E., 2011. An experimental study of crystalline basalt dissolution from 2 ≤ pH ≤ 11 and temperatures from 5 to 75 °C. *Geochim. Cosmochim. Acta* 75, 5496–5509.

Gunnarsson, G., Arnaldsson, A., Oddsdóttir, A.L., 2011. Model simulations of the Hengill area, Southwestern Iceland. *Transp. Porous Med.* 90, 3–22, doi:10.1007/s11242-010-9629-1.

Gunnarsson, I., Arnórsson, S., Jakobsson, S., 2005. Precipitation of poorly crystalline antigorite under hydrothermal conditions. *Geochim. Cosmochim. Acta* 69, 2813–2828.

Gysi, A.P., Stefánsson, A., 2011. CO₂–water–basalt interaction. Numerical simulation of low temperature CO₂ sequestration into basalts. *Geochim. Cosmochim. Acta* 75, 4728–4751.

Helgadóttir, H.M., Franzson, H., Sigurdsson, O., Ásmundsson, R.K., 2009. Helliðshéidi – Well HN-2. Report ISOR-2009/031. Internal Report in Icelandic.

Helgeson, H.C., 1968. Evaluation of irreversible reactions in geochemical processes involving minerals and aqueous solutions: I. Thermodynamic relations. *Geochim. Cosmochim. Acta* 32, 853–877.

Helgeson, H.C., Garrels, R.M., MacKenzie, F.T., 1969. Evaluation of irreversible reactions in geochemical processes involving minerals and aqueous solutions: II. Applications. *Geochim. Cosmochim. Acta* 33, 455–482.

Hoffert, M., Caldeira, K., Benford, G., Criswell, D., Green, C., Herzog, H., Jain, A., Ksheshig, H., Lackner, K., Lewis, J., Lightfoot, H., Manheimer, W., Mankins, J., Mauel, M., Perkins, L., Schlesinger, M., Volk, T., Wigley, T., 2002. Advanced technology paths to global climate stability: energy for a greenhouse planet. *Science* 298, 981–987.

Knauss, K., Nguyen, S., Wee, 1993. Diopside dissolution kinetics as a function of pH, CO₂, temperature and time. *Geochim. Cosmochim. Acta* 57, 285–294.

Knauss, K.G., Johnson, J.W., Steefel, C.I., 2005. Evaluation of the impact of CO₂, co-contaminant gas, aqueous fluid and reservoir rock interactions on the geologic sequestration of CO₂. *Chem. Geol.* 217, 339–350.

Lackner, K., 2003. A guide to CO₂ sequestration. *Science* 300, 1677–1678.

Lasaga, A., Soler, J., Ganor, J., Burch, T., Nagy, L., 1994. Chemical weathering rate laws and global geochemical cycles. *Geochim. Cosmochim. Acta* 58, 2361–2386.

Lichtner, P.C., 1988. The quasi-stationary state approximation to coupled mass transport and fluid–rock interaction in a porous medium. *Geochim. Cosmochim. Acta* 52, 143–165.

Lichtner, P.C., 1996. Continuum formulation of multicomponent-multiphase reactive transport. *Rev. Mineral. Geochem.* 34, 1–81.

Matter, J., Broecker, W., Stute, M., Gíslason, S., Oelkers, E., Stefánsson, A., Wolff-Boenisch, D., Gunnlaugsson, E., Axelsson, G., Björnsson, G., 2008. Permanent carbon dioxide storage into basalt: the CarbFix pilot project, Iceland. In: *Proceedings of the 9th International Conference on Greenhouse Gas Control Technologies*, Washington, DC, USA, 16–20 November.

Metz, B., Davidson, O., de Coninck, H., Loos, M., Meyer, L. (Eds.), 2005. *IPCC Special Report on Carbon Dioxide Capture and Storage*. Cambridge University Press, New York.

Nielsen, S., Franzson, H., 2010. Geology and hydrothermal alteration of the Hverahlíð HT-system, SW-Iceland. In: *Proceedings World Geothermal Congress 2010*, Bali, Indonesia, 25–29 April 2010.

Oelkers, E., Gíslason, S., 2001. The mechanism, rates and consequences of basaltic glass dissolution: I. An experimental study of the dissolution rates of basaltic glass as a function of aqueous Al, Si and oxalic acid concentrations at 25 °C and pH = 3 and 11. *Geochim. Cosmochim. Acta* 65, 3671–3681.

Oelkers, E., Gíslason, S., Matter, J., 2008. Mineral carbonation of CO₂. *Elements* 4, 331–335.

Oelkers, E., Schott, J., 2005. Geochemical aspects of CO₂ sequestration. *Chem. Geol.* 217, 183–186.

Pacala, S., Socolow, R., 2004. Stabilization wedges: solving the climate problem for the next 50 years with current technologies. *Science* 305, 968–971.

- Palandri, J., Kharaka, Y., 2004. A Compilation of Rate Parameters of Water–Mineral Interaction Kinetics for Application to Geochemical Modeling. Report 2004-1068.
- Pokrovsky, O.S., Schott, J., 2000. Kinetics and mechanisms of forsterite dissolution at 25 °C and pH from 1 to 12. *Geochim. Cosmochim. Acta* 64, 3313–3325.
- Pruess, K., 1991. TOUGH2: A General Numerical Simulator for Multiphase Fluid and Heat Flow. Report LBL-29400.
- Reed, M., 1982. Calculation of multicomponent chemical equilibria and reaction processes in systems involving minerals, gases and an aqueous phase. *Geochim. Cosmochim. Acta* 46 (513–528), 513–528.
- Rezvani Khalilabad, M., Axelsson, G., Gíslason, S., 2008. Aquifer characterization with tracer test technique; permanent CO₂ sequestration into basalt, SW Iceland. *Mineral. Mag.* 72 (1), 121–125.
- Rimstidt, J.D., Barnes, H.L., 1980. The kinetics of silica–water reactions. *Geochim. Cosmochim. Acta* 44, 1683–1699.
- Rogers, K., Neuhoﬀ, P., Pedersen, A., Bird, D., 2006. CO₂ metasomatism in a basalt hosted petroleum reservoir, Nuussuaq, West-Greenland. *Lithos* 92, 55–85.
- Schrag, D., 2007. Preparing to capture carbon. *Science* 315, 812–813.
- Sonnenthal, E., Ito, A., Spycher, N., Yui, M., Apps, J., Sughita, Y., Conrad, M., Kawakami, S., 2005. Approaches to modeling coupled thermal hydrological and chemical processes in the Drift Scale Heater Test at Yucca Mountain. *Int. J. Rock Mech. Min. Sci.* 42, 684–719.
- Steefel, C., Lasaga, A., 1994. A coupled model for transport of multiple chemical species and kinetic precipitation/dissolution reactions with applications of reactive flow in single phase hydrothermal systems. *Am. J. Sci.* 294, 529–592.
- Steefel, C.I., DePaolo, D.J., Lichtner, P.C., 2005. Reactive transport modeling: an essential tool and a new research approach for the Earth sciences. *Earth Planet. Sci. Lett.* 240, 539–558.
- Stefánsson, A., 2001. Dissolution of primary minerals of basalt in natural waters I. Calculation of mineral solubilities from 0 °C to 350 °C. *Chem. Geol.* 172, 225–250.
- Stefánsson, A., Gíslason, S., Arnórsson, S., 2001. Dissolution of primary minerals in natural waters II. Mineral saturation state. *Chem. Geol.* 172, 251–276.
- Walker, G., 1960. Zeolite zones and dike distribution in relation to the structure of the basalts of eastern Iceland. *J. Geol.* 68, 515–528.
- White, A.F., Blum, A.E., Schulz, M.S., Bullen, T.D., Harden, J.W., Peterson, M.L., 1996. Chemical weathering of a soil chronosequence on granitic alluvium: 1. Reaction rates based on changes in soil mineralogy. *Geochim. Cosmochim. Acta* 60, 2533–2550.
- White, A.F., Brantley, S.L., 2003. The effect of time on the weathering of silicate minerals: why do weathering rates differ in the laboratory and field. *Chem. Geol.* 202, 479–506.
- Wiese, F., Fridriksson, T., Armannsson, H., 2008. CO₂ Fixation by Calcite in High-temperature Geothermal Systems in Iceland. Tech. Rep., ISOR-2008/003, Iceland Geosurvey, Report available at: www.os.is/gogn/Skyrslur/ISOR-2008/ISOR-2008-003.pdf.
- Wolery, T.J., 1979. Calculation of Chemical Equilibrium between Aqueous Solution and Minerals: The EQ3/6 Software Package.
- Xu, T., Apps, J.A., Pruess, K., 2005. Mineral sequestration of carbon dioxide in a sandstone–shale system. *Chem. Geol.* 217, 295–318.
- Xu, T., Kharaka, Y.K., Doughty, C., Freifeld, B.M., Daley, T.M., 2010. Reactive transport modeling to study changes in water chemistry induced by CO₂ injection at the Frio-I Brine Pilot. *Chem. Geol.* 271, 153–164.
- Xu, T., Pruess, K., 2001. On fluid flow and mineral alteration in fractured caprock of magmatic hydrothermal systems. *J. Geophys. Res.* 106, 2121–2138.
- Xu, T., Sonnenthal, E., Spycher, N., Pruess, K., 2006. TOUGHREACT – a simulation program for non-isothermal multiphase reactive geochemical transport in variably saturated geologic media: applications to geothermal injectivity and CO₂ geologic sequestration. *Comput. Geosci.* 32, 146–165.
- Xu, T., Spycher, N., Sonnenthal, E., Zhang, G., Zheng, L., Pruess, K., 2011. TOUGHREACT version 2.0: a simulator for subsurface reactive transport under non-isothermal multiphase flow conditions. *Comput. Geosci.* 37, 763–774.

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