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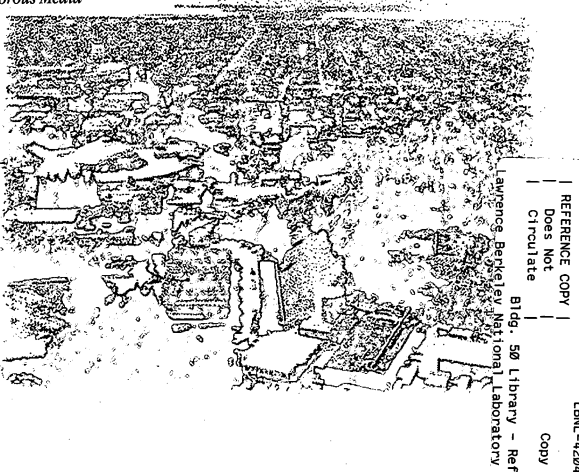
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Submitted to Transport in Porous Media



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¹ Earth Sciences Division, Lawrence Berkeley National Laboratory, University of California, Berkeley, CA 94720.

² Applied Mathematics Group, Industrial Research Ltd., P.O. Box 31310, Lower Hutt, New Zealand

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Tianfu Xu1, Stephen P. White2, and Karsten Pruess1

¹ Earth Sciences Division, Lawrence Berkeley National Laboratory, University of California, Berkeley, CA 94720.

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Abstract. Pyrite (FeS₂) is one of the most common naturally occurring minerals that is present in many subsurface environments. It plays an important role in the genesis of enriched ore deposits through weathering reactions, is the most abundant sulfide mineral in many mine tailings, and is the primary source of acid drainage from mines and waste rock piles. The pyrite oxidation reaction serves as a prototype for oxidative weathering processes with broad significance for geoscientific, geological engineering, and environmental applications. Mathematical modeling of these processes is extremely challenging because aqueous concentrations of key species vary over an enormous range, oxygen inventory and supply are typically small in comparison to pyrite inventory, and chemical reactions are complex, involving kinetic control and microbial catalysis. We present the mathematical formulation of a general multiphase advective-diffusive reactive transport model for redox processes. Two alternative implementations were made in the TOUGHREACT and TOUGH2-CHEM simulation codes which use sequential iteration and simultaneous solution, respectively. The simulators are applied to reactive consumption of pyrite in (1) saturated flow of oxidizing water, and (2) saturated-unsaturated flow in which oxygen

transport occurs in both aqueous and gas phases. Geochemical evolutions predicted from different process models are compared, and issues of numerical efficiency are discussed.

Key words: Reactive chemical transport, Pyrite oxidation, Acid mine drainage, Saturated-unsaturated media, oxygen transport.

1. Introduction

Pyrite oxidation generates sulfuric acid which gives rise to intense alteration of primary rock minerals, the development of secondary mineral assemblages, and the mobilization of metals. The process has been studied under laboratory and field conditions by many investigators. Lowson (1982) has reviewed the literature on abiotic chemical oxidation, while Nordstrom (1982) reviewed the literature on both abiotic and biotic geochemical oxidation. More recent studies include McKibben and Barnes (1986), Ague and Brimhall (1989), Welch *et al.* (1990), Walter *et al.*, (1994), Wunderly *et al.*, (1996), Nordstrom and Alpers, (1997), Lefebvre (1995), and Lefebvre *et al.* (1998).

In the unsaturated zone, pyrite oxidation occurs through a complex interplay of multiphase flow and transport processes. The ultimate oxidant is atmospheric oxygen, although the actual pyrite oxidation step may involve the ferric ion, Fe³⁺ (Lefebvre *et al*, 1998). In natural subsurface environments, bacterial catalysis plays an important role in the kinetics of pyrite oxidation (Kleinmann *et al.*, 1981). The oxygen consumed during pyrite oxidation is supplied from gaseous oxygen dissolved in the aqueous phase. Dissolved oxygen is replenished by recharge of oxygenated rainwater, and by dissolution of oxygen that is present in the gas-filled portion of the

pore space. The dissolution of gaseous oxygen reduces oxygen partial pressure, as well as the total pressure of the gas phase. Gaseous oxygen is replenished by diffusion and advection in the gas phase from the land surface boundary. As will be seen below, under conditions of low liquid saturation in the unsaturated zone, gas diffusion can be a more important mechanism for oxygen supply than gas or liquid advection. Dissolution of gaseous oxygen and subsequent transport to the reactive mineral surfaces may involve diffusive transport processes on a 'local' (pore level) scale. Additional process complexities arise in fractured porous media, where global transport of oxygen occurs primarily through a network of interconnected fractures, while oxygen may penetrate into tight matrix blocks primarily through relatively slow diffusive transport in gas and liquid phases. Depending on the spatial scale of interest, the 'kinetics' of pyrite oxidation may involve not just 'true' chemical kinetics, but may be affected by small-scale transport processes.

Redox reactions provide severe challenges for mathematical modeling because the concentrations of important chemical species such as O₂(aq), SO₄²⁻ and HS⁻ may change by as much as 70 orders of magnitude between oxidizing and reducing conditions (Xu *et al.*, 1998a). The purpose of the present paper is to present and partially evaluate alternative approaches for modeling redox processes in reactive flow systems. Modeling approaches for coupled processes of mass transport and chemical reactions in subsurface systems have been presented by a number of investigators (Yeh and Tripathi, 1991; Simunek and Suares, 1994; Steefel and Lasaga, 1994; Walter *et al.*, 1994; Zysset *et al.*, 1994; White, 1995; and Lichtner, 1996). These differ in their emphasis on accuracy and comprehensiveness of flow and transport modeling on the one hand, and chemical interactions between rocks and fluids on the other. Three major approaches have been used to couple mass transport and chemical reaction: (1) the direct substitution approach (DSA), which substitutes the chemical reaction equations directly into the transport equations; (2)

the sequential iteration approach (SIA), which solves the transport and the reaction equations separately in a sequential manner with an iterative procedure; and (3) the sequential non-iterative approach (SNIA), which is similar to the SIA but without the iterative procedure. Owing to the enormous diversity of transport and chemical reaction phenomena, no universally accepted methodology for modeling such processes has emerged, nor is likely to emerge. Instead, simulation methodology must be carefully chosen and tailored to the specific physical and chemical properties of the system being investigated, and to the objectives of the modeling study.

In this paper, we begin with a description and specification of the chemical system under study, and summarize two alternative modeling approaches as implemented in the TOUGH2-CHEM (White, 1995) and TOUGHREACT (Xu et al., 1997) codes. Then we describe the fully and variably saturated test problems examined in this study. Results from both codes are compared with respect to predicted geochemical evolution, and with respect to numerical accuracy and efficiency. The paper concludes with a discussion of possibilities and limitations for modeling redox-driven reactive chemical transport in multi-dimensional, heterogeneous, and field-scale systems over extended time periods.

2. Reaction mechanisms and kinetic rates

Recent studies have shown reasonable agreement on pyrite oxidation rates. Nevertheless, it is often difficult to make a direct comparison between experiments as initial pH, preparation methods, grain size, temperature, etc. may be different. Also reaction rates measured in the field may differ by orders of magnitude from those measured in the laboratory.

Another problem is determining the dominant reaction responsible for pyrite oxidation. Two candidates (see, for example, McKibben and Barnes, 1986) are

$$FeS_2 + \frac{7}{2}O_2 + H_2O \Leftrightarrow Fe^{2+} + 2SO_4^{2-} + 2H^+$$
 (1)

and the pair

$$FeS_2 + 14Fe^{3+} + 8H_2O \Leftrightarrow 15Fe^{2+} + 2SO_4^{2-} + 16H^+$$
 (2a)

$$Fe^{2+} + \frac{1}{4}O_2 + H^+ \Leftrightarrow Fe^{3+} + \frac{1}{2}H_2O$$
 (2b)

Reaction (2) is known to be much faster than reaction (1). Nordstrom and Alpers (1997) summarized the pyrite oxidation rate constants from several studies including those of McKibben and Barnes (1986), Brown and Jurinak (1989), and Nicholson (1994) (cited for a pH close to 2, a Fe³⁺ concentration of 10⁻³ mol Γ^1 , temperature close to 25 °C, and oxygen in equilibrium with the atmosphere). The rate constants of pyrite oxidation by O₂ range from 1.1×10⁻¹⁰ to 5.3×10⁻¹⁰ mol m⁻²s⁻¹ (moles of pyrite oxidation per square meter reactive surface area per second), while those by Fe³⁺ range from 9.6×10⁻⁹ to 1.9×10⁻⁸ mol m⁻²s⁻¹. Because the pyrite oxidation rate by Fe³⁺ is much faster than that by O₂, it is important to know the oxidation rate for Fe²⁺ to Fe³⁺ according to equation (2b). Under acid conditions, the rate becomes very slow. Singer and Stumm (1968) reported an abiotic rate of 2.7×10⁻¹² mol Γ^1 s⁻¹ (moles per liter solution per second) at pH values below 4. After normalizing this rate in terms of square meter reactive surface area it is equal to

1.53×10⁻¹⁰ mol m²s⁻¹ using a porosity of 0.1, a water saturation of 0.3, and a specific surface area of 586.7 m²m⁻³ (square meter per cubic meter medium). Such rates are considerably slower than the pyrite oxidation rate by Fe³⁺, hence equation (2b) would be the rate-limiting step were it not for the catalytic effect of the bacteria *Thiobacillus ferrooxidans* (Singer and Stumm, 1968; Nordstrom and Alpers, 1997). This effect on the oxidation of Fe²⁺ to Fe³⁺ may increase the rate by a factor of 10⁵ from about 2.7×10⁻¹² mol 1⁻¹s⁻¹ to about 2.7×10⁻⁷ mol 1⁻¹s⁻¹ according to Singer and Stumm (1968, 1970). The catalytic effect on the oxidation rate of pyrite is not clear. Nordstrom and Alpers (1997) note that results of Olson (1991) suggest the oxidation rate of pyrite in the presence of *Thiobacillus ferrooxidans* is little changed from that for the oxidation of pyrite by Fe³⁺. However they argue that the rate determining step is the production of Fe³⁺ by oxidation of Fe²⁺ (2b) rather than the heterogeneous reaction (2a).

Estimates of field pyrite oxidation rate cover a wide range of values. We cannot use field measurements of pyrite oxidation rates in this work, as the rate determining process is often transport of oxygen or other reactants to the reaction site. These are exactly the processes we wish to model in this work and so it is more appropriate to use the laboratory rates. The important parameter is not the pyrite oxidation rate constant but an effective rate given by the product (oxidation rate constant) × (reactive surface area). There is also great uncertainty as to the appropriate reactive surface area to use for field simulations. The effective value may be much smaller than those documented in Nordstrom and Alpers (1997). If flow is predominantly in fractures, then only a small portion of the pyrite will be exposed to oxygenated groundwater. Field pyrite oxidation rates are also complicated by many other factors such as microbial growth kinetics, organic compounds, secondary mineral formation, and climatic patterns, all of which are beyond the scope of this study.

Regardless of whether pyrite oxidation is dominated by reaction (1) or (2), oxygen availability is a very important factor. Reaction (1) directly uses oxygen, while (2) involves oxygen only indirectly. Extensive oxidation is limited to the unsaturated zone, where gas phase O₂ transport may be rapid. As aqueous phase oxygen is depleted through reaction with pyrite, it is replenished by dissolution from the gas phase,

$$O_2(g) \Leftrightarrow O_2(aq)$$
 (3)

and by diffusive transport in the gas phase from the atmospheric boundary at the land surface. As a result, the water can have high concentrations of Fe²⁺, SO₄²⁻, and H⁺. All reactions considered are listed in Table 1. Aqueous dissociation and gas dissolution are assumed to proceed according to the local equilibrium assumption. Pyrite oxidation is subject to kinetic control.

Table 1. List of reactions involved in pyrite oxidation. Thermodynamic equilibrium constants are from the EQ3/6 database (Wolery, 1992).

Reaction	log ₁₀ K at 25
$O_2(g) = O_2(aq)$	- 2.898
$pyrite(FeS_2) + 7/2O_2(aq) + H_2O = 2H^+ + 2SO_4^{2-} + Fe^{2+}$	217.4
$HS^- + 2O_2(aq) = H^+ + SO_4^{2-}$	138.32
$H_2S(aq) + 2O_2(aq) = 2H^+ + SO_4^{2-}$	131.33
$Fe^{2+} + 1/4O_2(aq) + H^+ = Fe^{3+} + 1/2H_2O$	8.475
$OH^- + H^+ = H_2O$	13.995
$HSO_4^- = H^+ + SO_4^{2-}$	-1.9791
$FeSO_4(aq) = SO_4^{2-} + Fe^{2+}$	-2.2

Lasaga *et al.* (1994) give the following general expression to describe the kinetic reaction rate for mineral dissolution or precipitation

$$r = Aka_{H+}^{n} (1 - \Omega^{\theta})^{\eta}$$
(4)

where k is the rate constant (moles per unit mineral surface area and unit time) which is a function of temperature, A is the specific reactive surface area per unit liquid volume, Ω is the mineral saturation ratio (i.e., the ratio between the ion activity product and the equilibrium constant). The parameters θ and η must be determined from experiments; in this work, they are assumed equal to one. The term $a_{H^+}^n$ accounts for the catalytic effect of H^+ where the value of n is determined experimentally; in this simulation this term is not considered (i.e., $a_{H^+}^n = 1$). We use a rate expression based on (4) but include a dependence on liquid saturation in variably saturated media. The rate given in (4) is multiplied by a function of liquid saturation $f(S_1)$ (see Figure 1) derived from laboratory measurements by Welch *et al.* (1990). The fractional factor $f(S_1)$ accounts for the effective sizes of the mineral pyrite surface that are actively available to adsorb and chemically bond with aqueous species.

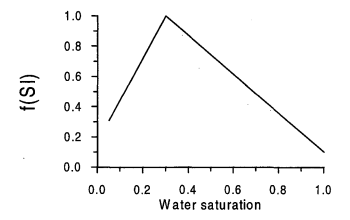


Figure 1. Liquid saturation dependent factor used together with kinetic rate expression (Eq. 4).

3. Mathematical model and numerical approaches

3.1 Governing equations

Two types of primary governing equations must be solved: (1) non-isothermal multiphase flow, and (2) multi-component reactive chemical transport.

Non-isothermal multiphase flow

Non-isothermal multiphase flow consists of fluid flow in both liquid and gas phases, and heat transport. The processes occurring in multiphase flow were discussed in detail by Pruess (1987, 1991). In this work the major assumptions made are: (1) air is treated as an ideal gas; and additivity of partial pressure is assumed for air/vapor mixtures; (2) porosity and permeability changes from mineral dissolution/precipitation are not considered; (3) concentration changes of aqueous and gaseous chemical species do not influence fluid thermophysical properties such as density and viscosity; and (4) heat generation due to chemical reactions such as pyrite oxidation is

neglected. Although in this study we assume isothermal conditions (constant temperature, 25 °C), our model is applicable to non-isothermal conditions as well; an application to sharply non-isothermal flows is presented in Sonnenthal *et al.* (1998). The conservation equations for multiphase flow of water, air and heat can be written as

for water

$$\frac{\partial}{\partial t} \left[\phi(S_1 \rho_1 X_{wl} + S_g \rho_g X_{wg}) \right] = -\nabla \cdot (X_{wl} \rho_1 \mathbf{u}_1 + X_{wg} \rho_g \mathbf{u}_g) + q_w$$
 (5)

air

$$\frac{\partial}{\partial t} \left[\phi(S_1 \rho_1 X_{al} + S_g \rho_g X_{ag}) \right] = -\nabla \cdot (X_{al} \rho_1 \mathbf{u}_1 + X_{ag} \rho_g \mathbf{u}_g) + q_a \tag{6}$$

and heat

$$\frac{\partial}{\partial t} \left[\phi(S_1 \rho_1 U_1 + S_g \rho_g U_g) + (1 - \phi) \rho_s U_s \right] = -\nabla \cdot \left[\sum_{\beta = 1, g} h_{\beta} \rho_{\beta} \mathbf{u}_{\beta} - \lambda \nabla T \right] + q_H$$
 (7)

where

$$\mathbf{u}_{\beta} = -\mathbf{k} \frac{\mathbf{k}_{r\beta}}{\mu_{\beta}} \left(\nabla \mathbf{P}_{\beta} - \rho_{\beta} \mathbf{g} \right) \qquad \beta = 1, \mathbf{g}$$
 (8)

is volumetric flux (Darcy velocity); the subscripts l, g, and s indicate the liquid, gas and solid phases, respectively; the subscripts w, a, and H indicate water, air and heat respectively; S_{β} is the

saturation of phase β ; ϕ the porosity; ρ_{β} the phase density; μ_{β} the phase viscosity; $X_{a\beta}$ and $X_{w\beta}$ are mass fraction of air and water in phase β ; k is the permeability, $k_{r\beta}$ is the relative permeability for phase β ; P_{β} is the pressure in phase β ; g is the gravitational acceleration; u_{β} is Darcy velocity for phase β ; U_{β} is internal energy of phase β ; h_{β} is the specific enthalpy of phase β ; λ is the heat conductivity of the rock-fluid mixture; T is the temperature; and q_{w} , q_{a} , q_{H} are source/sink terms for water, air and heat, respectively. The non-isothermal multiphase flow equations must be complemented with constitutive relationships that express all parameters as functions of the primary thermophysical variables. These expressions are given by Pruess (1987).

Multi-component reactive chemical transport

Aqueous (dissolved) species are subject to transport in the liquid phase, as well as to local chemical interactions with other aqueous species, and solid and gaseous species. Gaseous species are transported in the gas phase and react with aqueous species. Solid species only participate in local chemical reactions. Based on mass conservation, the transport equations for chemical components present in the liquid and gas phases can be expressed as

for the liquid phase

$$\frac{\partial}{\partial t}(\phi S_{l}C_{jl}) = -\nabla \cdot (\mathbf{u}_{l}C_{jl} - D_{l}\nabla C_{jl}) + q_{jl} + r_{js} + r_{jg} \quad j = 1, 2, ..., N_{l}$$
 (9)

and for the gas phase

$$\frac{\partial}{\partial t}(\phi S_g C_{jg}) = -\nabla \cdot (\mathbf{u}_g C_{jg} - D_g \nabla C_{jg}) + q_{jg} - r_{jg} \qquad j = 1, 2, ..., N_g$$
 (10)

where subscript j is the component index; N_1 and N_g are the numbers of chemical components present in the liquid and gas phases; $C_{j\beta}$ is the concentration of component j in phase β ; D_{β} is the molecular diffusion coefficient in phase β and assumed to be not species-dependent; $q_{j\beta}$ is external source/sink rate for component j in phase β (=l or g); r_{js} is the mass transfer rate from the solid to the liquid phase; and r_{jg} is the gas dissolution rate.

Assuming the gas phase acts as an ideal gas, gaseous concentrations (in moles per liter) can be calculated from their partial pressures using

$$C_{jg} = \kappa P_{jg} \tag{11a}$$

where

$$\kappa = \frac{10^2}{RT_s} \tag{11b}$$

 P_{jg} is the partial pressure (in bar) of component j, R is the gas constant (8.314 J mol⁻¹K⁻¹), T_s is the absolute temperature, and C_{jg} is the concentration (in moles per liter) of component j in the gas phase. For use in TOUGHREACT, by substituting (11a) into 10), the transport equations for chemical components present in the gas phase can be written in terms of partial pressures

$$\frac{\partial}{\partial t}(\phi S_g \kappa P_{jg}) = -\nabla \cdot \left[\kappa(\mathbf{u}_g P_{jg} - D_g \nabla P_{jg})\right] + q_{jg} - r_{jg} \qquad j = 1, 2, ..., N_g \quad (12)$$

Terms r_{js} and r_{jg} in (9) and (12) are functions of dissolved concentrations which render the reactive chemical transport equations highly nonlinear. TOUGH2-CHEM incorporates equation (10) directly into the transport equations and need not assume perfect gas behavior or Henry's law (although it does for this work). These primary governing equations must be complemented with constitutive chemical relationships. These relationships, including mass-balance equations for equilibrium reactions and rate expressions for kinetic reactions, are given in Xu *et al.* (1997).

3.2 Numerical approaches

Two alternative modeling approaches are implemented in the TOUGH2-CHEM (White, 1995) and TOUGHREACT (Xu et al., 1997) codes. Both codes are based on the framework of the non-isothermal multiphase flow simulator TOUGH2 (Pruess, 1991). The numerical solution of the multiphase flow and reactive chemical transport equations employs space discretization by means of integral finite differences (Narasimhan and Witherspoon, 1976) and fully implicit first-order finite differences in time. However, the coupled approach between flow and reactive transport is different in the two codes. TOUGH2-CHEM employs a fully coupled approach where all flow and reactive transport equations are solved simultaneously, which is originally designed for reactive flow and transport in geothermal reservoir systems. TOUGHREACT uses a sequential iteration approach similar to Yeh and Tripathi (1991), Simunek and Suares (1994),

Steefel and Lasaga (1994), Walter et al. (1994), and Lichtner (1996). The multiphase flow and reactive chemical transport equations are solved sequentially. After solution of the flow equations, the fluid velocities and phase saturations are used for chemical transport simulation. The chemical transport is solved on a component basis by taking chemical reaction source/sink terms from the previous reaction iteration. The resulting concentrations obtained from the transport are substituted into the chemical reaction model. The chemical reactions are solved on a grid-block basis. The chemical transport and reactions are iteratively solved until convergence.

TOUGHREACT can work with two different fluid flow (equation of state) modules, known as EOS9 and EOS3. Using EOS9, only saturated-unsaturated liquid phase flow is considered (Richards' equation). In this case, the only gaseous species transport process considered is oxygen diffusion. Using EOS3, the full non-isothermal multiphase flow equations (for liquid, gas and heat) are solved. The dissolution of gaseous oxygen induced by pyrite oxidation reduces oxygen partial pressure, as well as the total pressure of the gas phase. As a result, the gas flow is modified, which in turn modifies the liquid phase flow. This reaction effect on the fluid flow is not considered, although advection and diffusion are considered for oxygen transport processes. On the other hand, TOUGH2-CHEM (using the EOS3 flow module) solves the multi-component chemical transport simultaneously with the heat and mass flow problem so that equations of flow and multi-component reactive transport are fully coupled and the reaction effect on the fluid flow is taken into account. This coupling of flow to chemical reaction is not significant in the examples presented in this paper; however, when fluid flow and chemical reactions are strongly coupled, such as when boiling with strong precipitation and dissolution effects takes place in geothermal reservoirs, this may be essential.

Due to the large inventory of the solid mineral (pyrite) and the comparatively small concentrations of aqueous reactants such as dissolved oxygen, the mineral (pyrite) dissolution requires on the order of 10⁴-10⁵ pore volumes of fluid. For these reasons, a number of numerical improvements for convergence and efficiency, such as modification of the Jacobian matrix, use of quasi-stationary approximation, and automatic time stepping were made (see Xu *et al.*, 1998).

4. Flow systems

TOUGHREACT and TOUGH2-CHEM are capable of modeling non-isothermal reactive transport in one, two, or three dimensions. Due to the complexity of reactive chemical transport under multiphase flow conditions, we demonstrate the capabilities discussed here through isothermal modeling in two one-dimensional flow systems. In the first example (Figure 2), a horizontal porous column 2 m long with a cross-sectional area of 1 m² contains the mineral pyrite; at the left boundary water is injected at 0.02 m day⁻¹ and is withdrawn from the right boundary at the same rate. Porosity of the column is 0.4 which, with the prescribed injection rate, leads to a pore velocity of 0.05 m day⁻¹. The column is discretized with a grid spacing of 0.05 m.



Figure 2. Schematic representation of the horizontal saturated flow system used for pyrite oxidation.

The second example models a vertical column extending from the atmosphere through an unsaturated and a saturated zone and extending below the water table (see Figure 3). Oxygen is supplied to the top of the column as a dissolved species in infiltrating rainwaters and may also be transported by gaseous diffusion and advection from the land surface. Because TOUGH2-CHEM includes full coupling between the gas and liquid phase flows, and chemical reactions, it is not possible to solve completely identical problems with the two codes when the gas phase is included. For the unsaturated example TOUGH2-CHEM must include an extra component, nitrogen, to correctly model the gas phase. The initial flow conditions are set by specifying the rate of water infiltration into the top of the model (0.07 m yr⁻¹, variations in infiltration is not considered in this study) and a constant pressure of 3.5 bar at the bottom. The steady-state water saturations obtained by ignoring chemical reactions are used as initial conditions for the calculation of reactive chemical transport. Parameters for this unsaturated-saturated medium are listed in Table 2. The steady-state water saturation profile is given in Figure 4.

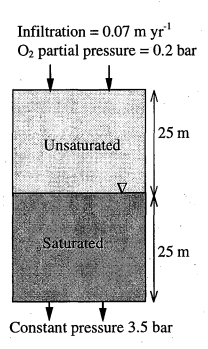


Figure 3. Schematic representation of the unsaturated-saturated flow system used for pyrite oxidation. The system can be considered as part of mine tailings or waste pile, or as a mineralized site that contains the mineral pyrite (FeS₂).

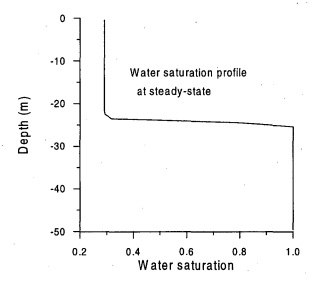


Figure 4. The steady-state water saturation distribution in the unsaturated-saturated medium.

Table 2. Physical and chemical parameters used for the unsaturated-saturated flow system.

Parameter	Value
Infiltration (m yr ⁻¹)	0.07
Depth (m)	50
Grid spacing (m)	1
O ₂ partial pressure at the land surface (bar)	0.2
Permeability (m ²)	7×10^{-12}
gas O ₂ diffusivity (m ² s ⁻¹)	4.4×10^{-5}
Tortuosity	0.1
Porosity	0.1
Relative permeability and capillary pressure	
(van Genuchten, 1980):	
$^{+}$ λ	0.457
$S_{ m lr}$	0.05
${f S}_{ m ls}$	1.0
$P_0(Pa)$	1.96×10^3
Kinetic rate constant, k, (mol m ⁻² s ⁻¹)	2×10^{-13} - 2×10^{-10}
Specific surface area (m ² m ⁻³)	58.67

For both flow systems, the initial water composition corresponds to a dilute reducing water with aqueous oxygen concentration, $c_{O_2(aq)}$, of 1.0×10^{-70} mol I^{-1} . The infiltration water compositions correspond to a dilute oxidizing water with $c_{O_2(aq)} = 2.53\times10^{-4}$ mol I^{-1} , which is at equilibrium with an atmospheric O_2 partial pressure of 0.2 bar. Initial pyrite abundance is 9% by volume.

Five cases were simulated for the two flow systems that involved different combinations of chemical transport by liquid advection, oxygen gas transport by diffusion and advection, and pyrite oxidation controlled by local equilibrium or kinetics. These cases are summarized in Table 3, in order of increasing complexity.

Table 3. List of simulation cases with different combinations of flow system, transport process, and reaction type (equilibrium/kinetics). Symbol k represents the kinetic rate constant (units: mol m⁻²s⁻¹)

Flow	Transport process Pyrite oxidation (equilibrium or kinetics)	Case	Codes used
Fully saturated	Aqueous advection		
	Equilibrium	1	Both
Unsaturated-	Aqueous advection		
saturated	Equilibrium	2	Both
	Kinetics (k = 2×10^{-13} , 2×10^{-10})	3	Both
	Aqueous advection and gas diffusion		
	Kinetics (k = 2×10^{-11} , 2×10^{-10})	4	TOUGHREACT
	Aqueous advection, gas diffusion and		
	advection		
	Kinetics (k = 2×10^{-11} , 2×10^{-10})	5	TOUGH2-CHEM

5. Results

5.1. Horizontal saturated medium

Case 1 (aqueous advection and equilibrium pyrite oxidation)

This is a relatively simple case for which the gross behavior of the system can be determined without detailed calculation. Thus it provides a check on the simulation codes as it allows comparison of calculated dissolution front speeds with theory, and a check on mass conservation. Pyrite oxidative dissolution occurs only in the first grid block, and the $O_2(aq)$ is consumed completely at this location. Dissolution of one mole of pyrite consumes 3.5 moles of aqueous $O_2(aq)$, and creates 2 moles of H^+ , 2 moles of SO_4^{2-} , and 1 mole of H^{2-} . Consequently, total dissolved S and Fe concentrations increase and pH decreases. After one day (note that time in days is equivalent to grid block pore volume, since it takes one day to move one grid block with pore velocity = 0.05m day⁻¹ and the grid spacing = 0.05 m), a quasi-stationary state or QSS

(Lichtner, 1988) is reached. The solution is entirely buffered by pyrite, and dissolved concentrations remain constant along the entire column with the following values: $10^{-64.86}$ mol 1^{-1} for aqueous O_2 , 1.44×10^{-4} for total dissolved S, 7.2×10^{-5} for Fe, and 3.86 for pH. The pH evolution during the early transient period in the first grid block is presented in Figure 5. For this QSS, pyrite abundance in the first block decreases at a constant rate of 2.53×10^{-4} yr⁻¹ by volume, which equals 0.54 moles per year. Pyrite initially has a volume fraction of 9%, corresponding to 188 moles per grid block. It requires 356 yr to dissolve all the pyrite in a grid block. At a pore velocity of 365 grid blocks per year, the retardation factor for the oxidative dissolution front is, and is: $356 \text{ yr} \times 365 \text{ yr}^{-1} = 129,940$. The total amount of aqueous O_2 injected during a one year period is: 2.53×10^{-4} mol $1^{-1} \times 7300$ l yr⁻¹=1.847 mol yr⁻¹. The ratio of total oxygen consumption and pyrite dissolution obtained in the simulation is: 1.847/0.54 = 3.42, which is very close to the stoichiometric coefficient (3.5) in the pyrite oxidation reaction (1). This shows that most of the oxygen is consumed by pyrite oxidation.

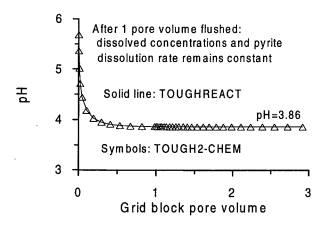


Figure 5. pH evolution during the early transient period (note that grid block pore volume equals time in days) in the first grid block for the horizontal saturated column.

5.2. Unsaturated-saturated medium

Case 2 (aqueous advection and equilibrium pyrite oxidation)

In this scenario, pyrite oxidation and dissolution take place only in the topmost (first) grid block, and aqueous oxygen is consumed completely. The chemical evolution is similar to Case 1. However, the transient period lasts much longer than in Case 1 because of the low infiltration rate of 0.07 m yr⁻¹ (vs. 7.3 m yr⁻¹ in Case 1) and larger grid spacing (1 m vs. 0.05 m). After approximately 150 yr, a quasi-stationary state is reached throughout the entire flow domain. The steady-state pH and dissolved component concentrations are the same as those of Case 1 (see Figure 6).

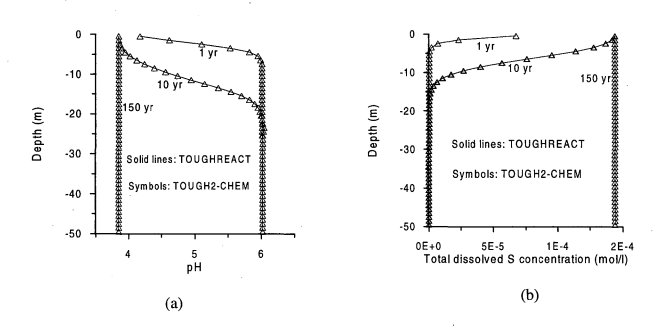


Figure 6. pH (a) and dissolved S concentration (b) at different times for Case 2 (unsaturated-saturated flow, aqueous advection, and equilibrium pyrite oxidation). Total dissolved Fe

concentration equals half of S according to the stoichiometric coefficients of pyrite (FeS₂). Case 3 results (obtained with a rate constant of $k = 2 \times 10^{-10}$ mol m⁻²s⁻¹) are indistinguishable from the equilibrium case.

Case 3 (aqueous advection and kinetic pyrite oxidation)

We first used a rate constant of $k = 2 \times 10^{-10}$ mol m⁻²s⁻¹. The result is the same as for the equilibrium Case 2 (Figure 6), indicating that the reaction rate is limited by oxygen supply, not chemical kinetics. As mentioned above, field kinetic rate constants cover a wide range of values. To illustrate a kinetically-controlled process, we reduced the rate constant by three orders of magnitude to $k = 2 \times 10^{-13}$ mol m⁻²s⁻¹. The result is presented in Figure 7. We can observe a step change in pH from Figure 7b. This may be explained from the reaction processes taking place in the system. Pyrite is first oxidized creating H⁺ and Fe²⁺, and consuming O₂(aq). Fe²⁺ is oxidized to Fe^{3+} by the excess $O_2(\text{aq})$, and the ratio $\text{Fe}^{2+}/\text{Fe}^{3+}$ is controlled by the $O_2(\text{aq})$ concentration. Then aqueous species such as O₂(aq), H⁺, Fe²⁺, and Fe³⁺ are transported downward. At approximately 14 m depth $O_2(aq)$ becomes depleted, and the reaction: $4Fe^{2+} + O_2(aq) + 4H^+ = 4Fe^{3+} + 2H_2O$ is driven to the left-hand side. This reaction provides both H⁺ to lower the pH and O₂(aq) to oxidize some more pyrite and lower the pH further. Because we treat the aqueous reactions at local equilibrium, almost all the Fe³⁺ is reacted in a single grid block and there is a step change in pH. Figure 8 shows the concentration distribution of Fe²⁺ and Fe³⁺ obtained with the code TOUGHREACT. We can see that in the top 13 blocks Fe³⁺ is dominant. At the 14th grid block, Fe³⁺ is dramatically reduced and Fe²⁺ becomes dominant. From Figure 7 we can see that results from both codes agree very well.

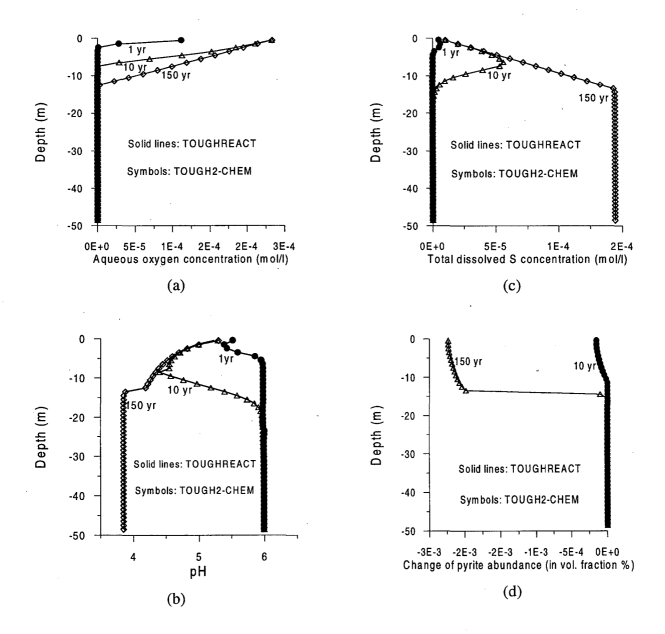


Figure 7. Aqueous oxygen (a), pH (b), total dissolved S (c) and change of pyrite abundance (d) after 150 yr (steady-state) considering only chemical advective transport in the liquid phase (Case 3). Kinetic rate constant is 2×10^{-13} mol m⁻²s⁻¹. Change of pyrite abundance is expressed as change in pyrite volume per unit medium volume (pyrite density = 4.177×10^4 mol m⁻³)

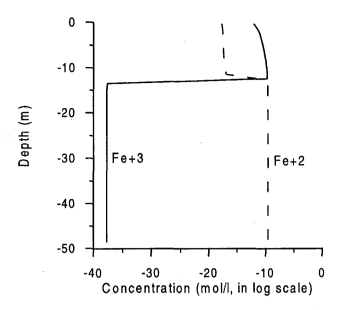


Figure 8. Concentration distribution of species Fe^{2+} and Fe^{3+} after 150 yr (steady-state) for Case 3 with a kinetic rate constant of $k = 2 \times 10^{-13}$ mol m⁻²s⁻¹. The results are obtained with TOUGHREACT.

Cases 4 and 5 (aqueous advection, gaseous transport, and kinetic pyrite oxidation)

The TOUGHREACT calculation includes oxygen diffusion in the gas phase in addition to the processes considered in Case 3. Using an active gas phase in TOUGH2-CHEM not only invokes oxygen diffusion in the gas phase, but also includes an overall advective gas flow, in response to reduction of gas phase pressures from oxygen consumption. Because of these slightly different process descriptions, some differences are expected between Cases 4 and 5.

Results for a rate constant of $k = 2 \times 10^{-10}$ mol m⁻²s⁻¹ are shown in Figure 9. Comparing with results that did not include gaseous transport (Case 3, Figure 6), it is seen that the pH drops much more rapidly to lower values, and total dissolved S concentrations are several orders of magnitude larger. This indicates that gas diffusion provides a much stronger mechanism for oxygen supply than aqueous advection. Results for a tenfold smaller kinetic rate constant ($k = 2 \times 10^{-11}$ mol m⁻²s⁻¹) shown in Figure 10 demonstrate that gaseous oxygen supply maintains aqueous oxygen

concentrations at high levels throughout the unsaturated zone. Quasi-stationary concentrations of the dissolved S at t = 150 yr have similar shapes as for the larger rate constant, but are approximately ten times smaller, again indicating that oxygen consumption is limited entirely by reaction kinetics, not by transport.

The TOUGHREACT and TOUGH2-CHEM simulations agree well for the quasi-stationary states (QSS), but show significant differences during the transient approach to the QSS. This is attributed to the somewhat different process descriptions, with TOUGHREACT neglecting effects of changes in oxygen partial pressure on gas and liquid advection.

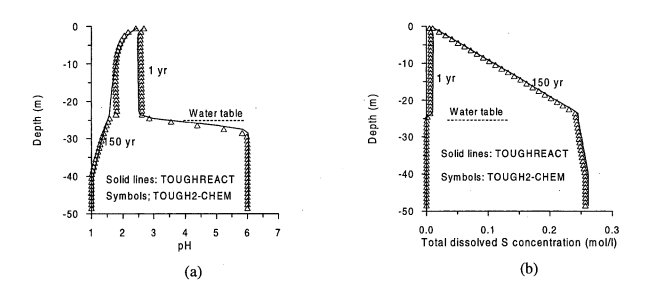


Figure 9. pH (a) and total dissolved S (b) for Case 4 (TOUGHREACT) and Case 5 (TOUGH2-CHEM), using a kinetic rate constant of $k = 2 \times 10^{-10}$ mol m⁻²s⁻¹. Solid lines: TOUGHREACT results considering oxygen gas diffusion in addition to chemical transport in the liquid phase. Symbols: TOUGH2-CHEM results considering reaction effect on the fluid flow, gas advection and diffusion.

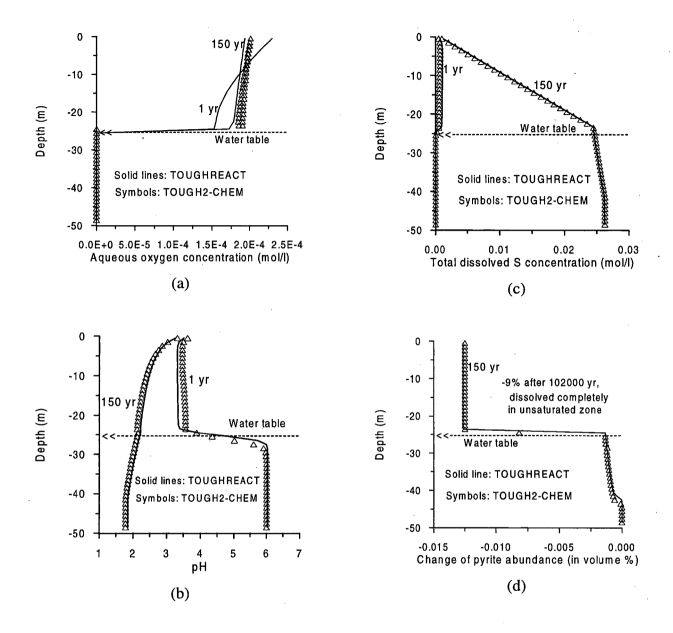


Figure 10. Aqueous oxygen (a), pH (b), total dissolved S (c) and change of pyrite abundance (d) using a kinetic rate constant of $k = 2 \times 10^{-11}$ mol m⁻²s⁻¹.

6. Discussion

6.1 Mass balance

The saturated column (Case 1) was employed to examine global mass conservation. Three runs were performed with TOUGHREACT using three different time steps: 1, 3 and 6 days. Results from the three runs agree well and are in fact indistinguishable on the scale of Figure 5. However, the mass balance errors are slightly different. A chemical component initially present (aqueous and solid phases) in the system plus that entering the system (called input), should be equal to the component currently present in the system plus that leaving the system (output). A relative error (E_r) for the mass balance, is defined as

$$E_{r} = \frac{|(current + output) - (initial + input)|}{initial + input} \times 100\%$$
 (13)

 E_r for S and Fe is presented in Figure 11. It can be seen that the error increases as the time step size decreases. This is because a smaller time step requires more computing operations for the same period of simulation. The error for all three runs is linearly increasing with time. Errors for a short simulation period are very small (on the order of 10^{-8}). However, for simulations over geological time the error can become significant. Extrapolating from the data given in Figure 11, for a time step of 1 day the error can reach 1% after 1.71×10^6 yr.

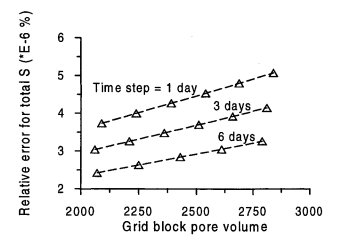


Figure 11. Relative mass balance error (E_r) for total sulfur versus grid block pore volume (equals time in days) with different time steps.

6.2 Advancement of pyrite oxidation front in liquid-advection dominated flow systems

From Figure 7, we can see that in the top blocks, free aqueous oxygen concentration decreases linearly (Figure 7a), and total dissolved S increases linearly (Figure 7c). When aqueous oxygen concentration is high, pyrite oxidation is far from the equilibrium state (Ague and Brimhall, 1989). Under these conditions, Ω <<1 in Eq. 4, so that the dissolution rate takes on an approximate constant or maximum value, $r_m = Ak$ (note that chemical calculation is in terms of kg H_2O or liter solution if density is 1 kg I^{-1}). For case 3 using a rate constant of $k=2\times10^{-13}$ mol m⁻²s⁻¹, the actual rate over the maximum rate r/r_m at different times is presented in Figure 12. The pyrite oxidation front moves as the water is moving. Behind the front, pyrite oxidative dissolution has a constant rate. After 20 years the rate distribution remains unchanged.

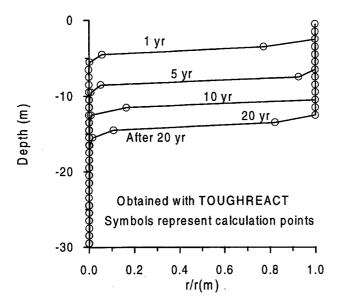


Figure 12 Relative dissolution rate (divided by the maximum rate) distribution at different times for Case 3 using a rate constant of $k = 2 \times 10^{-13}$ mol m⁻²s⁻¹.

The advancement of the front can be estimated from the oxygen mass balance. We first start with

$$\int_{0}^{t} P_{O2} + \int_{0}^{t} E_{O2} = \int_{0}^{t} I_{O2}$$
 (14)

where P_{O2} is the rate of oxygen consumption by pyrite oxidation in the entire system per unit time (in units of mol s⁻¹), E_{O2} is the rate of oxygen consumption (negative for generation) by all other reactions except pyrite oxidation (including free aqueous oxygen) per unit time, and I_{O2} is the oxygen supply rate by infiltration from the land surface. The contribution of the E_{O2} term decreases with time (see Figure 13), and amounts to less than 3% after a quasi-stationary state is reached. Neglecting E_{O2} , we have

$$3.5 \cdot 10^{3} \cdot \phi S_{1} \int_{0}^{z_{b}} r(z) dz = I_{O2}$$
 (15)

where z_b is the depth beyond which r=0 (note that dissolution of 1 mole of pyrite requires 3.5 moles of oxygen, and the unit of r must be in mol $1^{-1}s^{-1}$). For most parts, r takes on a constant value of r_m (see Figure 12), so we have

$$3.5 \cdot 10^{3} \cdot \phi S_{1} \left[r_{m} z_{p} + \int_{z_{p}}^{z_{b}} r(z) dz \right] = I_{O2}$$
 (16)

where z_p is the depth of pyrite oxidation with the maximum rate. The second term of Eq. 16 is small (see Figure 12). For estimation purposes, we neglect this term, and then we have

$$z_{p} = \frac{I_{O_{2}}}{3500\phi S_{1} r_{m}} \tag{17}$$

In the simulations for the unsaturated-saturated flow system, $I_{O2} = 5.6126 \times 10^{-10}$ mol m⁻²s⁻¹, $\phi = 0.1$, $S_1 = 0.3$ (unsaturated zone), so $z_p = 5.345 \times 10^{-12}/r_m$. For different values of the rate constants k, the values of z_p calculated from Eq. 17 are listed in Table 4. This estimation is confirmed by the computer simulation (see Figure 12).

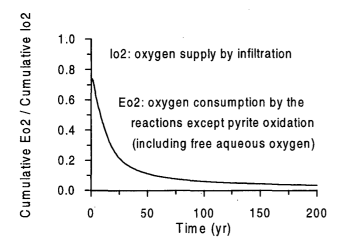


Figure 13. Relative oxygen consumption by reactions except pyrite oxidation (including free aqueous oxygen) versus time for Case 3 using a rate constant of $k = 2 \times 10^{-13}$ mol m⁻²s⁻¹.

Table 4. The estimation of the depth (z_p) of pyrite oxidation front having the maximum rate during a quasi-stationary state using different kinetic rate constant k.

k (in mol m ⁻² s ⁻¹)	r _m (in mol I ⁻¹ s ⁻¹)	z _p (in m)
2×10 ⁻¹⁰	3.91×10 ⁻¹⁰	1.367×10 ⁻²
2×10 ⁻¹¹	3.91×10^{-11}	1.367×10 ⁻¹
2×10 ⁻¹²	3.91×10^{-12}	1.367
2×10 ⁻¹³	3.91×10 ⁻¹³	13.67

6.3. Kinetics versus equilibrium

Whether a particular reaction should be described as governed by the local equilibrium approximation (LEA) or by kinetic rates depends not only on the reaction itself but also on the rates of hydrodynamic processes (Domenico and Schwartz, 1990). A reaction is kinetically controlled when the time required to reach equilibrium is significant in comparison to characteristic times for transport of reactants. In the above simulations for reactive chemical

transport only in the liquid phase (Cases 2 and 3), the choice of equilibrium depends mainly on three factors: (1) kinetic rate, (2) space discretization, and (3) oxygen availability.

At a large rate of $k = 2 \times 10^{-10}$ mol m⁻²s⁻¹ (Case 3), the results become indistinguishable from those for equilibrium (Case 2). The lower limit for applicability of the LEA is obtained by requiring the depth to the reaction front to be less than the grid spacing, $z_p \le \Delta z$. From Eq. 17, we have

$$\frac{I_{O_2}}{3500\phi S_1 r_m} \le \Delta z \tag{18}$$

For the grid spacing size $\Delta z = 1$ m used in our simulations, the LEA is applicable when $r_m \geq 5.345 \times 10^{-12}$ mol $\Gamma^1 s^{-1}$ (rate constant $k \geq 2.734 \times 10^{-12}$ mol $m^{-2} s^{-1}$). Note that the constraint for applicability of the LEA becomes more stringent for finer space discretization. If we use a large rate constant of $k = 2 \times 10^{-10}$ mol $m^{-2} s^{-1}$ ($r_m = 3.91 \times 10^{-10}$ mol $\Gamma^1 s^{-1}$, see Table 4), applicability of the LEA requires $I_{O_2} \leq 4.11 \times 10^{-8}$ mol $m^{-2} s^{-1}$. With the same aqueous oxygen concentration used in the simulations, the corresponding infiltration rate limit is $q \leq 5.126$ m yr⁻¹.

If oxygen transport by gas diffusion processes is considered, the LEA cannot be used. For cases 4 and 5, the liquid advection time for a grid block is 0.43 yr (157 days), while the gas diffusion time is 1.32 days, so the gas diffusion process is several orders of magnitude faster than the liquid advection.

6.4. Importance of gaseous oxygen transport

When oxygen gas transport processes are considered in addition to chemical transport in the liquid phase, a maximum rate of pyrite oxidation is reached very quickly throughout the entire unsaturated zone (see the curve after 0.1 yr in Figure 14). In the top of the saturated zone, pyrite is still oxidized by oxygen supply through liquid advection but the rate is much smaller. With the same $k = 2 \times 10^{-10}$ mol m⁻²s⁻¹ and by neglecting oxygen gas transport processes, pyrite oxidation occurs only at the first grid block and the maximum cannot be reached (see Table 4). Much lower pH and higher dissolved concentrations are obtained by considering gas transport processes. The oxygen gas diffusion plays a dominant role in the chemical evolution for pyrite oxidation in unsaturated-saturated conditions.

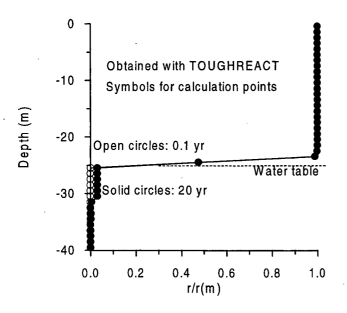


Figure 14. Relative dissolution rate (divided by the maximum rate) distribution at different times for Case 4 using a rate constant of $k = 2 \times 10^{-11}$ mol m⁻²s⁻¹.

6.5 Comparison of the two codes

For the Cases (1, 2 and 3) considering only chemical transport in the liquid phase, the results obtained with both codes match very well. If gaseous species transport processes are considered in addition to transport in the liquid phase, results are slightly different. TOUGH2-CHEM provides a more complete process description, which includes the effects of partial pressure reduction due to reactions on the fluid flow. On the other hand, TOUGHREACT is more efficient than TOUGH2-CHEM (see Table 5). TOUGH2-CHEM solves all flow and multi-component transport and reaction equations simultaneously. TOUGHREACT solves them sequentially, and transport and reaction iteratively. The transport is solved on a component basis, while the reactions are modeled on a grid block basis. The number of equations solved by the two approaches are summarized in Table 6. For pyrite oxidation only by rainwater infiltration, we know the steady-state flow distribution but TOUGH2-CHEM still needs to solve the flow. In this condition, TOUGHREACT solves the flow only once at the first time step.

Table 5. CPU times (minutes on a Pentium 200 PC) required by TOUGHREACT and TOUGH2-CHEM for the unsaturated-saturated cases (a simulation time of 150 yr).

	TOUGHREACT	TOUGH2-CHEM
Considering gas transport (cases 4, 5)	3.0	90
No gas transport (case 3)	1.9	15

Table 6. Number of equations per iteration required to be solved simultaneously by TOUGHREACT and TOUGH2-CHEM for the pyrite oxidation problem in the unsaturated-saturated medium. N_B=number of grid blocks (50), N_C=number of chemical components (5), and N_S=number of chemical species (13).

Approach	Number of equations
TOUGHREACT	Flow (EOS3): $3\times N_B=150$ for one time
	Transport: $N_B=50$ for $N_C=5$ times
	Reaction: N_C+1 (pyrite) for $N_B=50$ times
TOUGH2-CHEM	Flow+transport+reaction
	$2 \times N_B + N_C \times N_B = 350$ for one time

7. Conclusions

Oxidative weathering of pyrite, as well as of other metal sulfides, is an extremely difficult process to model numerically because (1) it is driven by oxygen, whose concentrations in ambient soil gas and in pore water in the unsaturated zone are very small, so that of the order of 10⁵ pore volumes of fluid are required to consume 1 vol.-% of solid mineral; (2) the chemical activities and total elemental solubilities of key species vary over an enormous range of tens of orders of magnitude during redox processes; and (3) the oxidation involves several reaction steps that are partly kinetically controlled, and catalyzed by microbial acitivity.

This paper has presented the formulation of a general multiphase advective-diffusive reactive transport model for redox processes. Two alternative numerical implementations were developed, using either a sequential iteration approach for transport and reactions, or a fully-

coupled simultaneous solution. Simulations of a fully and a variably-saturated pyrite oxidation problem showed reasonable geochemical behavior, and satisfactory agreement between the two approaches. In the unsaturated zone, gas diffusion was found to be the dominant mechanism for oxygen supply. Although simple one-dimensional domains were used in the present study, both TOUGHREACT and TOUGH2-CHEM can deal with two- and three-dimensional problems in hydrologically and geochemically heterogeneous media.

The fully-coupled solution approach allows larger time steps, but the numerical work per time step is also larger. For the problem considered in this study, time steps had to be limited to resolve adequately highly transient changes, and the sequential iteration approach was found to be more efficient numerically. The relative efficiency of different approaches appears to be highly problem dependent.

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Notation

- a air index (subscript)
- A specific reactive surface area, m²m⁻³
- c concentration of chemical species, mol I⁻¹.

 \mathbf{C} concentration of component, mol 1⁻¹. molecular diffusion coefficient in phase β (1 and g). $\mathbf{D}_{\mathbf{B}}$ $\mathbf{E}_{\mathbf{r}}$ relative mass balance error oxygen consumption rate by pyrite, mol s⁻¹ P_{02} oxygen supply rate by infiltration, mol s⁻¹ I_{02} gas phase index (subscript) g gravitational acceleration vector, m s⁻² g H heat index (subscript) specific enthalpy of phase β (l and g), J kg⁻¹. h_{β} chemical component index (subscript or superscript). time step number. k medium permeability, m². k Mineral dissolution rate constant, mol m⁻² s⁻¹ $\mathbf{k}_{\mathbf{m}}$ $K_{r\beta}$ relative permeability of phase β (1 and g). K Thermodynamic equilibrium constant. 1 liquid phase index (subscript). N_l number of aqueous components N_g number of gaseous components. P_{β} pressure of phase β , Pa. P_{ig} partial pressure of gaseous component j, Pa. source/sink terms for water, air and heat, kg m⁻³s⁻¹ for mass or J m⁻³ s⁻¹ for heat q_w , q_a , q_H source/sink terms of dissolved component j, mol m⁻³s⁻¹. q_{jl} source/sink terms of gaseous component j, mol m⁻³s⁻¹. \mathbf{q}_{jg} gas constant, 8.314 kgm²s⁻²K⁻¹mol⁻¹. R dissolution rate of gaseous component i, mol m⁻³ s⁻¹ rjg mass transfer rate of component j from the solid to liquid phase, mol m⁻³ s⁻¹ \mathbf{r}_{js} solid phase index (subscript) S saturation of phase β (1 and g). S_{β} T temperature. time. t

u Darcy velocity, m s⁻¹.

 U_{β} specific internal energy of phase β (1, g and s), J kg⁻¹.

 $X_{j\beta}$ mass fraction of component j in phase β .

w water index (subscript).

Greek Symbols

β phase index, β = g: gas; 1: liquid; s: solid phase.

 Δ increment of a variable

 λ overall medium thermal conductivity, W m⁻¹ K⁻¹.

φ porosity

 ρ density, kg m⁻³.

μ dynamic viscosity, Pa s.

 $\Omega_{\rm m}$ saturation ratio of m-th mineral.

 ∇ .() divergence operator.

 ∇ () gradient operator.

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Ernest Orlando Lawrence Berkeley National Laboratory One Dyglotron Road | Berkeley, Galifornia 94720