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Kinetics of low-temperature H₂ production in ultramafic rocks by ferroan brucite oxidation

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T = 378

20 40 60 Duration (days)

Abstract

(Inmol)

Ferroan brucite, (Mg,Fe)(OH)₂, is among the potential mineral candidates for low temperature (<423 K) abiotic H₂ production in ultramafic rocks. To verify this assumption, synthetic ferroan brucite with grain size similar to that observed in natural samples (40–100 nm) was reacted with pure water at temperatures ranging from 348 to 573 K. Experimental products are consistent with the reaction 3 Fe(OH)₂^{brucite} = Fe₃O₄ + H₂ + 2 H₂O. This reaction reached completion in ~2 months at 378 K and is thermally activated with an activation energy of 145 ± 1 kJ/mol. The standard state formation enthalpy and the third law entropy of amakinite, Fe(OH)₂, were refined from the experimental dataset. The new thermodynamic parameters imply that ferroan brucite is stable at significantly lower hydro-

gen activity than previously calculated. The alteration of Fe-brucite produces H_2 at rates compatible with present day observations of H_2 emissions in natural settings (ophiolite and mid-oceanic ridges). However, efficient fluid renewal is required, as opposed to H_2 production through olivine serpentinisation, which can proceed in static hydraulic conditions.

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Introduction

Hydrothermal circulation across upper mantle rocks at midocean ridges promotes serpentinisation reactions. In the course of these reactions, olivine reacts with water to form serpentine, magnetite and ferroan brucite, (Mg,Fe)(OH)₂, along with abiotic hydrogen (Moody, 1976). As observed in ophiolites (Neal and Stanger, 1983; Abrajano *et al.*, 1990; Leong *et al.*, 2023), ultramafic rocks can still produce H₂ at low temperature (*i.e.* at T < 423 K), even if they are extensively serpentinised. The extrapolation of experimental kinetic data collected in the 473–623 K range (*e.g.*, McCollom *et al.*, 2016) indicates that serpentinisation of olivine with a grain size of 500 µm should reach a reaction progress above 90 % in at least 10,000 yr at temperatures below 423 K.

The serpentinisation of olivine produces secondary minerals, including ferroan brucite, which are Fe^{2+} -rich and which can further react to produce H_2 + magnetite at low temperature:

$$3[Fe(OH)_2]^{brucite} = Fe_3O_4 + H_2 + 2H_2O$$
 Eq. 1

where [Fe(OH)₂]^{brucite} represents the Fe component of ferroan brucite.

Petrographic data on ophiolite and dredge seafloor samples (Jöns *et al.*, 2017; Klein *et al.*, 2020; Ellison *et al.*, 2021) seem to indicate that Reaction 1 could proceed at sub-surface conditions in partly serpentinised ultramafic rocks. H₂ production was achieved in hydrothermal experiments carried out

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on serpentinised peridotite at 373 K and was attributed to magnetite formation at the expense of ferroan brucite (Miller *et al.*, 2017).

In order to test the potential of Reaction 1 to produce H₂ at temperatures below 423 K in ultramafic rocks, the kinetics and thermodynamics of Reaction 1 were investigated experimentally here using synthetic (Mg_{1-x}Fe_x)(OH)₂ of grain size (40–100 nm) and composition (*x* from 0.156 to 0.205) relevant to natural ferroan brucite (Malvoisin *et al.*, 2020).

Materials and Methods

Ferroan brucite $(Mg_{1-x}, Fe_x)(OH)_2$, with *x* ranging from 0.156 to 0.205, was synthesised under ambient conditions from a stoichiometric solution of dissolved Fe(II) and Mg chlorides, as described in Carlin *et al.* (2023). The ferroan brucite obtained by this method formed platelets 40 to 100 nm across (Fig. 1a). It was loaded under an Ar atmosphere with degassed ultrapure water either in welded shut gold capsules ('caps' experiments) or in 50 mL Parr 5500 series titanium reactors ('SP' experiments; see details in Tables S-2 and S-3). The capsules were run either in horizontal cold seal pressure vessels at temperatures from 348 to 573 K at 20 MPa (caps#1 to #15) or in an oven at temperatures of 378 and 423 K at the liquid-vapour equilibrium pressure (P_{satr} caps#1 to #12). Titanium reactor experiments, SP#3 to SP#5, were conducted in the 423 to 473 K range, also at P_{sat} . SP#6 was run at 378 K with an initial Ar pressure of ~5 MPa.

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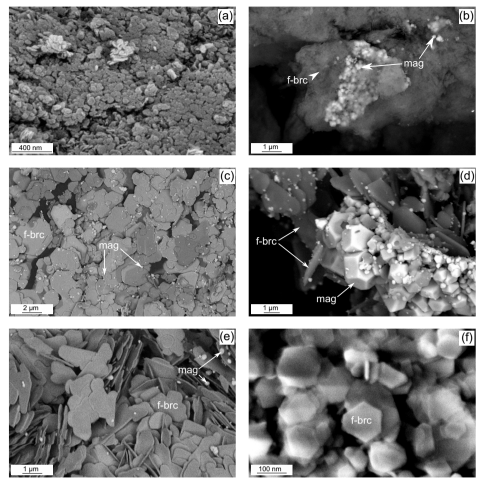


Figure 1 Back-scattered images of (a) the starting ferroan brucite and (b–f) its reaction products. (b) Caps#9; 378 K, 2.5 days. (c, d) SP#3; 473 K, 8 days. Euhedral micrometre-sized magnetite is visible. (e) Caps#15; 573 K, 2 days. Note that ferroan brucite platelets have recrystallised. (f) SP#6; 378 K, 69 days. ~100 nm wide euhedral ferroan brucite platelets are observed. f-brc, ferroan brucite; mag, magnetite.

Pressure evolution in SP experiments was monitored at ±2 kPa with a Keller pressure sensor PA-33X. After the experiments, the H₂ produced and trapped in the gold capsule was sampled using the protocol described in Malvoisin et al. (2013) and analysed by gas chromatography. H₂ leakage through the gold capsule walls is negligible at the temperatures investigated here (Malvoisin et al., 2013). The amount of H₂ produced in titanium reactors was quantified either on the gas sampled at the run conditions, and/or at the end of the experiment. No H₂ was detected in two blank experiments performed at 378 and 473 K over more than 40 days with capsules containing only degassed ultrapure water (100 µL). Before experiments, the titanium reactors were heated to 523 K during one day in air to extract any H₂ potentially solubilised in the reactor wall (Louthan and Derrick, 1975), and to ensure Ti surface oxidation prior to reaction. No hydrogen production was detected in these blank experiments. Details of the sample characterisation techniques are provided in the Supplementary Information.

Results and Discussion

Ferroan brucite oxidation reaction. In all experiments, ferroan brucite partly decomposed into magnetite + H₂ (Table S-2). The H₂ yield increased from 4×10^{-3} to 220 µmol of H₂ *per* gram of starting material when the temperature increased from 348 to 573 K. In eight experiments (Table S-2), pyroaurite was detected (<20 wt. %), indicating the presence of minor CO₂ in the reacting

medium. Pyroaurite does not involve significant H₂ production (Carlin et al., 2023). Its impact on H₂ production rate is, thus, mainly associated with the lowering of the amount of ferroan brucite available for H₂ production. However, for experiments used for determining kinetic and thermodynamic parameters, we estimated that a maximum of 3.4 wt. % of the initial ferroan brucite was consumed to form pyroaurite. This leads to an error on the H₂ production rate that is small compared to the error associated with H₂ measurement of ~11 %. The Fe content $(X_{\rm Fe(OH)_2})$ of ferroan brucite was determined from the refined unit cell parameters (Table S-2) based on Vegard's law (Carlin et al., 2023). It was found to vary from 0.1 to 0.2, i.e. equal or slightly below $X_{Fe(OH)_2}$ in the ferroan brucite starting material. Recrystallisation of ferroan brucite as platelets was observed in the highest temperature experiments (Fig. 1). Unless metastable growth occurred, this observation suggests that ferroan brucite with $X_{Fe(OH)_2} < 0.2$ is a stable reaction product in these experiments.

Altogether, the experimental results revealed that brucite partly reacted during the experiments according to Reaction 1, leading to the overall reaction:

$$(1 - x + y)Mg_{(1-x)}Fe_{x}(OH)_{2} = (1 - x)Mg_{(1-x+y)}Fe_{(x-y)}(OH)_{2}$$
$$+ \frac{y}{3}Fe_{3}O_{4} + \frac{2y}{3}H_{2}O + \frac{y}{3}H_{2}$$
Eq. 2

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where *x* is the initial $X_{\text{Fe}(\text{OH})_2}$ and *y* is a parameter related to reaction progress such as $0 \le y \le x$, which reflects that only a fraction, *y*/*x*, of the Fe(OH)₂ component in ferroan brucite has reacted. Indeed, thermodynamic equilibrium may well be achieved for *y* < *x*.

*H*₂ *production rate and brucite oxidation rate.* The amount of produced H₂ was used to retrieve the progress of Reaction 2. Magnetite was not used to infer reaction progress because, due to the presence of minor Fe³⁺ in the starting material, part of the magnetite product may form independently of Reaction 2, i.e. without H₂ production (Carlin et al., 2023). The amount of H₂ produced in the SP experiments did not induce pressure changes significant enough to retrieve isothermal H₂-production rate laws directly from pressure monitoring. However, the qualitative pressure evolution was used to determine the overall duration of the H₂ production stage, which, in turn, was used to model the reaction kinetics (see Supplementary Information for details). H₂ production kinetics at 378 K could be accurately retrieved from Run SP#6, where gas was regularly sampled (Fig. 2a). In addition, a series of experiments in gold capsules (caps#t5 to #t12) were stopped after different durations and H₂ was analysed in order to further constrain the kinetics of Reaction 2 (Fig. 2b). These experimental data were fitted to a kinetic law (Lasaga, 1998; see Supplementary Information for details) with the reaction rate (r) as:

$$r = r_0 \times \left(1 - \frac{Q}{K}\right)$$
 Eq. 3

where $r_0 = k_0 \times A \times \exp(\frac{-E_a}{RT})$ with k_0 a kinetic constant, A the $Fe(OH)_2$ specific surface area, E_a the activation energy, R the gas constant and T the temperature. Q and K are the reaction quotient and the equilibrium constant of Reaction 1, respectively. The standard state is defined here with unit activity for pure minerals and water at any temperature and pressure, as well as unit fugacity for ideal gas at 1 bar of pressure and any temperature. Q was approximated to $\frac{P_{\rm H_2}}{X_{\rm Fe(OH)_2}^3}$ with $P_{\rm H_2}$ the H₂ partial pressure at the conditions of the experiment and $X_{Fe(OH)_2}$ the molar fraction of Fe(OH)₂ in ferroan brucite, by assuming ideal behaviour for H2 and Fe(OH)2 in the gas phase and in the brucite solid-solution, respectively. X_{Fe(OH)2} was calculated from Equation 2, based on the number of moles of produced H₂ ($n_{\rm H_2}$). $P_{\rm H_2}$ was derived from $n_{\rm H_2}$ considering the amount of H₂ dissolved in the solution as calculated using PHREEQC (Parkhurst and Appelo, 2013). K was estimated with the same procedure as Q_{r} by considering that equilibrium H₂ pressure equals the H₂ partial pressure in the gas at the last measurement multiplied by a factor, λ , slightly above 1 (see Supplementary Information for details on this factor). Fitted r_0 values are displayed in Figure 2c as a function of reciprocal temperature. The slope in the linear fit corresponds to an activation energy of 145 ± 1 kJ/mol. The intercept of the fit provides a $k_0 \times A$ quotient of 8.97×10^7 mol s⁻¹ g⁻¹.

Experimental constraints on thermodynamic properties of ferroan brucite. Based on the kinetic law derived in the previous section, 17 experiments reached equilibrium (Table S-2). These experiments were, thus, used to constrain the equilibrium constant (*K*) of Reaction 1.

As discussed above, a set of *K* values can be calculated based on n_{H_2} , measured at the end of each experiment. A pair of $\Delta_{\text{f}}H^\circ$ and *S*° values for the Fe(OH)₂ end member was retrieved by least square regression through this set of *K* values (see Supplementary Information for thermodynamic calculation details). $\Delta_{\text{f}}H^\circ_{\text{Fe}(\text{OH})_2}$ and $S^\circ_{\text{Fe}(\text{OH})_2}$ of -581.3 ± 2.9 kJ/mol and

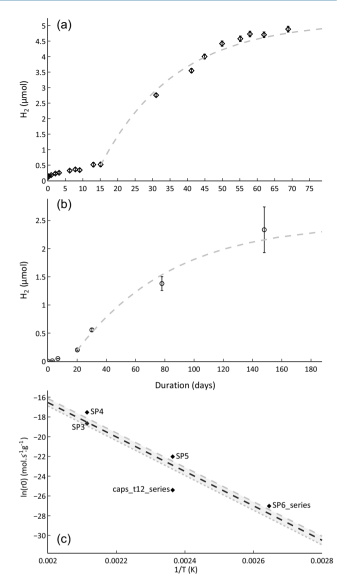


Figure 2 Kinetics of ferroan brucite alteration. **(a)** Number of moles of produced H₂ as a function of time at 378 K (Run SP#6; black diamonds). **(b)** Number of moles of produced H₂ as a function of time at 423 K (Runs caps#t5 to caps#t12; black circles). The dashed lines in **(a)** and **(b)** correspond to a fit of the data with Equation 3. **(c)** $\ln(r_0)$ vs. 1/*T*. The linear regression of r_0 is displayed with a black dashed line ($R^2 = 0.87$) and the grey dashed lines are linear regressions for r_0 with λ values of 1.01 and 1.20.

86.4 ± 6.3 J/mol/K were obtained, respectively. These thermodynamic values are only relevant for calculations using the same standard states as those used here, as well as the same assumption of unit activity and fugacity coefficients for ferroan brucite solid solution and H₂ in the gas phase, respectively. They fall in the range of published values for Fe(OH)₂ (Table S-4, Fig. S-3). The $\Delta_f H^\circ_{Fe(OH)_2}$ value is consistent with the value by Ziemniak *et al.* (1995) and departs by 1.2 % from $\Delta_f H^\circ_{Fe(OH)_2'}$ tabulated in the NIST-JANAF database (Chase, 1998). The values of the NIST-JANAF database only differ by 0.1 % from those commonly used for thermodynamic modelling of fluid–rock interactions in ultramafic rocks (McCollom and Bach, 2009). H₂ production prediction for our experiments is overestimated by more than one order of magnitude with the McCollom and Bach's (2009) database (Fig. S-4).

Implications for low-T H₂ *production in ultramafic systems.* Olivine serpentinisation (*e.g.*, McCollom *et al.*, 2016)



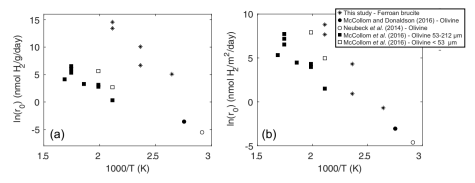


Figure 3 Comparison of experimentally determined rate of H_2 production for ferroan brucite (r_0 , this study) and olivine serpentinisation (Neubeck *et al.*, 2014; McCollom and Donaldson, 2016; McCollom *et al.*, 2016). (a) Reaction rate *per* mass of starting material (r_0). (b) Reaction rate *per* reactive surface area (r_0/A) with A the specific surface area either measured with the BET method or calculated with the relationship provided in Brantley and Mellott (2000).

and ferroan brucite alteration (Miller et al., 2017; Ellison et al., 2021) are the two main processes that have been proposed to account for H₂ production at low temperature (T < 423 K) in ultramafic rocks. Excluding kinetic experiments interpreted as being distorted by artefacts (McCollom and Donaldson, 2016), a maximum of 0.028 nmol H₂/g olivine/day was proposed for H₂ production by olivine serpentinisation at 363 K (Fig. 3a). In comparison, the rate of H₂ production measured here during ferroan brucite reaction (r_0) is approximately three orders of magnitude higher (Fig. 3a). When weighted by the specific surface area of the powders used in the various experiments, however, the reaction rates are rather well aligned in an Arrhenius plot with an activation energy of ~81 kJ/mol (Fig. 3b). The difference in grain sizes between olivine used in the experiments depicted in Figure 3 (38-212 µm) and the synthetic ferroan brucite used here (~50 nm) probably plays a key role in their difference of reactivity.

The respective contribution of olivine and Fe-brucite alteration to the H₂ production rate in natural systems was evaluated using a numerical approach. A fluid–serpentinised dunite system composed of olivine (grain size of 500 μ m; Malvoisin *et al.*, 2017), ferroan brucite (grain size of 50 nm; Malvoisin *et al.*, 2021) and water was modelled with H₂ being only produced by alteration of the latter minerals. A range of escape rates of H₂ (advection and/or diffusion) was defined in order to simulate hydrothermal activity at mid-ocean ridges, water infiltration in an ophiolitic unit or sub-stagnant hydraulic conditions in a deep aquifer. The temperature was set to 363 K, relevant to low-*T* serpentinisation (Fig. 4; see model details in Supplementary Information).

The model shows that ferroan brucite is the first mineral to react with a rate that is three orders of magnitude faster than that of olivine, leading to a rapid H₂ production in the first year of the reaction (Fig. 4b). In a closed system, ferroan brucite reaction rapidly stops due to the attainment of thermodynamic equilibrium (Q/K = 1). The adjustment of the thermodynamic parameters of Fe(OH)₂ proposed here (1.8 % and 1.2 % for $S^{\circ}_{Fe(OH)_2}$ and $\Delta_{\rm f} H^{\circ}_{\rm Fe(OH)_2}$, respectively, compared to the database of McCollom and Bach, 2009) has a strong impact on the predicted equilibrium H₂ partial pressure and, thus, on the amount of H₂ that is produced. At 363 K, equilibrium is achieved for a H₂ molality of 2.2×10^{-5} mol/kg, while it is two orders of magnitude higher $(5.0 \times 10^{-3} \text{ mol/kg})$ with the database of McCollom and Bach (2009). At 313 K, a difference of three orders of magnitude for H₂ molality between the two database is predicted. The oxygen fugacity (fO_2) lies on the H₂O/H₂(g) equilibrium with the database of McCollom and Bach (2009) and five orders of magnitude above with the thermodynamic data derived here. Interestingly, this latter fO_2 is consistent with the fO_2 measured at the bottom of Holes BA1A, BA1D and BA4A during the Oman Drilling Project (Kelemen et al., 2021). After ferroan brucite reaction, olivine completely reacts in the model in approximately 3 Myr and ultimately produces 3000 times more H₂ than ferroan brucite (Fig. 4a).

Ferroan brucite can further react if, at the same time, H_2 escapes from the system at a rate exceeding that of H_2 production associated with olivine serpentinisation (Fig. 4c). This corresponds to minimum H_2 escape rates of 10^{-5} and 7×10^{-4} mol $H_2/day/g$ rock at 313 and 363 K, respectively. The estimated

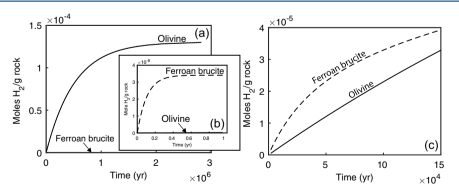


Figure 4 Numerical modelling of the contribution of ferroan brucite alteration and olivine serpentinisation to H_2 production as a function of time in a partly altered peridotite (see Supplementary Information for details). (a) General view and (b) incipient stage simulation in a closed system (no H_2 leak associated with diffusion or fluid advection). (c) Simulation by considering advection of a fluid at a rate of 1.4×10^{-4} kg water/day/kg rock. Dashed line: H_2 production associated with ferroan brucite alteration. Solid line: H_2 production associated with olivine serpentinisation.

maximum escape rate by vertical diffusion is three orders of magnitude lower than this threshold value (see Supplementary Information for details), suggesting that vertical diffusion is not sufficient to drive ferroan brucite reaction. The threshold value is achieved by water renewal at a minimum rate of 7×10^{-6} and 3×10^{-5} kg water/day/kg rock at 313 and 363 K, respectively. The average water-to-rock ratio at mid-ocean ridges is ~1 (Coogan et al., 2019). Considering hydrothermal activity and fluid flow during a minimum of 30,000 yr (Früh-Green et al., 2003), it can be converted into a mean water flux of 10^{-7} kg water/day/kg rock. In the Oman ophiolite, ferroan brucite with x = 0.28 can represent ~50 mol % of the serpentinisation reaction products (Malvoisin et al., 2020). Present day alteration of such ferroan brucite may occur during interaction with rainwater. H2-rich hyperalkaline fluids are found at depths >50 m (Leong et al., 2023), and the recharge rate of the aquifer in Oman is 18 mm/year (Dewandel et al., 2005). Combining these values leads to a mean meteoritic water flux of 3×10^{-7} kg water/day/kg rock. Both in ophiolites and on the seafloor, mean water flux estimates are, thus, approximately one order of magnitude lower than the minimum flux necessary to drive ferroan brucite reaction. However, fluid flow in ultramafic rocks is concentrated in cracks and microcracks (Dewandel et al., 2005; Corre et al., 2023) and is, thus, expected to be, locally, several orders of magnitude higher than the mean water flux. For example, the highest water-to-rock ratio values reported in abyssal peridotites are above 10⁵ (Snow and Reisberg, 1995; Delacour et al., 2008), corresponding to water fluxes $>10^{-2}$ kg water/day/kg rock compatible with H₂ production associated with ferroan brucite oxidation. The measured maximum H₂ production rate in the Oman ophiolite is of 71,000 mol H₂/yr for a minimal volume of altered rock of 0.05 km³ (Leong et al., 2023). This corresponds to a specific flux of 10^{-3} nmol H₂/g rock/day which is consistent with the specific H₂ production rate of 5×10^{-3} nmol H₂/g rock/ day, estimated at 313 K based on the extrapolation of the data acquired here for ferroan brucite alteration (Fig. 3). Actually, the same extrapolation for serpentinisation of olivine having a grain size of 500 µm (Fig. 3b) yields a much lower production rate of 10^{-5} nmol H₂/g rock/day at 313 K. Ferroan brucite oxidation could thus be a main contributor to H₂ production at temperatures below 423 K in ophiolites and on the seafloor. This is consistent with petrographic observations in natural samples showing ferroan brucite oxidation to form magnetite in open systems conditions (Bach et al., 2006; Jöns et al., 2017).

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Additional Information

Supplementary Information accompanies this letter at https://www.geochemicalperspectivesletters.org/article2408.



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Kinetics of low-temperature H₂ production in ultramafic rocks by ferroan brucite oxidation

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Supplementary Information

The Supplementary Information includes:

- Characterisation Techniques
- > Time Needed to Reach Thermodynamic Equilibrium in Titanium Reactors
- ➢ H₂ Production Rate-law
- > Retrieval of $\Delta_f H^\circ$ and S° of the Fe(OH)₂ End Member: Calculation Method
- ▶ Numerical Modelling of H₂ Production During Serpentinised Dunite Alteration
- ➤ Tables S-1 to S-4
- ➢ Figures S-1 and S-4
- Supplementary Information References

An overview of the notation used throughout is given in Table S-1.

Characterisation Techniques

Gas phase contained in the sealed gold capsules was recovered after the experiments using the method described in Malvoisin *et al.* (2013). For titanium reactors, the gas phase was sampled in a syringe connected to the reactor headspace through a valve. For the gas analysis, 250 μ L of gas was injected with gas-tight syringe in a Clarus 500 gas chromatograph (Perkin ElmerTM) equipped with a polymer filled column (Restek ShinCarbonTM) and a thermal conductivity detector (TCD). Argon was used as carrier gas. Each gas sample was analysed at least three times to check reproducibility. Based on these repeated measurements, the uncertainty on H₂ measurement is estimated to be ~11 % of the measured values (one standard deviation).

The recovered sample powder was characterised by X-ray diffraction (XRD) with a Bruker D8 diffractometer. XRD patterns were collected from 10 to 80° (2 θ) using CuK α or CoK α radiation and counting times of 3 s per 0.04° step (Fig. S-1). XRD patterns were analysed using the Rietveld technique with the BGMN software (Doebelin and Kleeberg, 2015). Details of both the refinement strategy and refinement constraints are given in Carlin *et al.* (2023). The results of gas analyses and Rietveld refinement are provided in Tables S-2 and S-3.



Sample powders were mounted on double-sided carbon tape in the glove box for scanning electron microscopy (SEM) imaging. They were then coated under vacuum with a 1 nm thick gold layer and characterised with a field emission gun scanning electron microscope (FEG-SEM; Zeiss Ultra 55) operated at 5 to 10 kV.

Time Needed to Reach Thermodynamic Equilibrium in Titanium Reactors

Pressure evolution in SP experiments was monitored *in situ* (Fig. S-2). The hydrogen partial pressure, P_{H_2} , and thus the amount of produced H_2 , n_{H_2} , were not retrieved from the *in situ* pressure monitoring. Indeed, as illustrated in Figure S-2, a steady state regime is achieved after an initial step of pressure increase but the steady-state pressure cannot be precisely retrieved due to a noise amplitude of ~30 kPa in the measured pressure. These variations have actually the same amplitude as the pressure increase generated by the reaction itself. For example, in experiment SP#5, the pressure oscillation (0.41–0.37 MPa) is equal to 40 kPa, *i.e.* close to the pressure increase generated by the complete reaction (Fig. S-2).

The *in situ* pressure monitoring could however be used to constrain the kinetics of ferroan brucite oxidation by retrieving the approximate reaction time needed to reach steady-state pressure conditions, even though the exact steady-state pressure could not be accurately determined. Steady-state pressure was assumed to represent near equilibrium conditions, and the corresponding reaction time was determined graphically for each SP experiments (Fig. S-2).

H₂ Production Rate-law

In the experiments conducted in gold capsules at \sim 20 MPa, no gas phase is present and H₂ occurs as an aqueous phase only:

$$3[Fe(OH)_2]^{\text{brucite}} = Fe_3O_4 + H_{2,aq} + 2H_2O.$$
 (S-1)

In the other experiments conducted on the liquid/vapour equilibrium, H₂ can be present as a gas phase:

$$3[Fe(OH)_2]^{brucite} = Fe_3O_4 + H_{2,g} + 2H_2O.$$
 (S-2)

We assumed in the following that the kinetics of Reactions S-1 and S-2 are the same. This implies that H_2 transfer between the aqueous and the gaseous phases is not a rate-limiting step (*i.e.* rapid compared to the other processes involved in the reaction). Under this assumption, hydrogen production rate (*r*) according to Reactions S-1 and S-2 can be written, following Lasaga (1998), as:

$$r(t) = \frac{d\bar{n}_{H_2}}{dt} = r_0 \times (1 - \frac{Q_i}{K_i}),$$
(S-3)

with $r_0 = k_0 \times A \times \exp\left(\frac{-E_a}{RT}\right)$. Q_i is computed by assuming that H₂ is an ideal gas and that the activity of Fe(OH)₂ in ferroan brucite is equal to its molar fraction $(X_{Fe(OH)_2})$. This leads for Reaction S-1 to $Q_1 = \frac{[H_{2,aq}]}{X_{Fe(OH)_2}^3}$ with $[H_{2,aq}] = \frac{n_{H_2}}{V_{aq}}$, and for Reaction S-2 to $Q_2 = \frac{P_{H_2}}{X_{Fe(OH)_2}^3}$ with $P_{H_2} = \frac{n_{H_2,g} \times R \times T}{V_g}$. Introducing



Henry's coefficient $(K_{\rm H} = \frac{P_{\rm H_2}}{[{\rm H_2}]})$ allows to relate $n_{{\rm H_2},{\rm g}}$ to $n_{{\rm H_2}}$ as $n_{{\rm H_2},{\rm g}} = \frac{K_{\rm H}V_{\rm g}}{K_{\rm H}V_{\rm g}+RTV_{\rm aq}}n_{{\rm H_2}}$. Assuming that $V_{\rm aq}$ and $V_{\rm g}$ remain constant during the experiment provides the same expression for $\frac{Q_r}{K_r}$ in Reactions S-1 and S-2:

$$\frac{Q_1}{K_1} = \frac{Q_2}{K_2} = \frac{n_{\rm H_2}}{n_{\rm H_2,eq}} \left(\frac{X_{\rm Fe(OH)_2,eq}}{X_{\rm Fe(OH)_2}} \right)^3.$$
 (S-4)

Combining Equation S-4 with an expression of $X_{Fe(OH)_2}$ derived from mass conservation in Reactions S-1 and S-2 $(X_{Fe(OH)_2} = \frac{n^{\circ}_{Fe} - 3n_{H_2}}{n^{\circ}_{Fe} - 3n_{H_2} + n^{\circ}_{Mg}})$ allows to express r(t) as a function of n_{H_2} only as:

$$r(t) = \frac{d\bar{n}_{\rm H_2}}{dt} = \frac{r_0}{f(n_{\rm H_2}(t))},$$
(S-5)

where $f(n_{\text{H}_2}(t)) = \frac{1}{1 - \frac{n_{\text{H}_2}(t)}{n_{\text{H}_2,\text{eq}}} \times (\frac{(n^\circ_{\text{Fe}} - 3n_{\text{H}_2,\text{eq}})(n^\circ_{\text{Fe}} - 3n_{\text{H}_2}(t) + n^\circ_{\text{Mg}})}{(n^\circ_{\text{Fe}} - 3n_{\text{H}_2,\text{eq}} + n^\circ_{\text{Mg}})(n^\circ_{\text{Fe}} - 3n_{\text{H}_2}(t))})^3}.$

We measured n_{H_2} by sampling the gas phase either at the end of the experiment (298 K) or *in situ* (experiment SP#6). During all these measurements, the calculated fraction of H₂ in the aqueous phase is below 3 mol % and was thus neglected. The value of n_{H_2} at equilibrium ($n_{H_2,eq}$) was set to the last measurement of H₂ times a factor (λ) slightly above 1 to account for the fact that n_{H_2} asymptotically tends towards the value at the equilibrium according to Reaction S-3. λ was varied between 1.01 and 1.2 to estimate its impact on the derived kinetic constants (Fig. 2). The parameter r_0 was determined by least-square regression of Equation S-5 through the experimental n_{H_2} dataset. The activation energy and the $k_0 \times A$ product were retrieved with a linear fit in a ln(r_0) vs. 1/T plot.

Retrieval of $\Delta_f H^\circ$ and S° of the Fe(OH)₂ End Member: Calculation Method

For Reactions S-1 and S-2 with H₂ in either the aqueous or gas phase, by definition:

$$\Delta_r G^{P,T} = \left(\sum_i^{\text{phases}} \nu_i \,\Delta_a H_i^{P,T}\right) - T \times \left(\sum_i^{\text{phases}} \nu_i \,S_i^{P,T}\right) + RT \ln \prod_i a_i^{\nu_i}, \quad (S-6)$$

with $\Delta_a H_i^{P,T} = \Delta_f H_i^\circ + \int_{T_r}^T Cp_i dT + V_i^\circ (P - P_r)$ and $S_i^{P,T} = S_i^\circ + \int_{T_r}^T \frac{Cp_i}{T} dT$. Equation S-6 can be simplified assuming equilibrium conditions, *i.e.* $\Delta_r G^{P,T} = 0$ and $\prod_i a_i^{\nu_i} = K$ and

Equation S-6 can be simplified assuming equilibrium conditions, *i.e.* $\Delta_r G^{P,T} = 0$ and $\prod_i a_i^{\nu_i} = K$ and expressed as a function of two unknowns, $\Delta_f H^{\circ}_{Fe(OH)_2}$ and $S^{\circ}_{Fe(OH)_2}$. All the other parameters can be calculated from tabulated data (slop98.dat database and Klein *et al.*, 2009 for other amakinite parameters).

Considering the experimental dataset with all the experiments (*j*) having reached equilibrium at T_j and P_j , a system of equations is obtained:

$$\Delta_{\mathbf{f}} H^{\circ}_{\mathbf{Fe}(\mathrm{OH})_2} - T_j \times S^{\circ}_{\mathbf{Fe}(\mathrm{OH})_2} = f_j(T_j, P_j).$$
(S-7)

 $\Delta_{f}H^{\circ}_{Fe(OH)_{2}}$ and $S^{\circ}_{Fe(OH)_{2}}$ values are then determined by least-squares regression (Fig. S-3).



Numerical Modelling of H₂ Production During Serpentinised Dunite Alteration

A numerical model was developed to determine the contribution of both olivine and ferroan brucite to H_2 production during the reaction of a partly serpentinised dunite at 363 K. The initial composition of the serpentinised dunite was determined by assuming that half of the olivine was already serpentinised according to the following reaction (initial serpentinisation degree of 50 %):

 $24 (Mg_{0.9}Fe_{0.1})_2SiO_4 + 34 H_2O = 12 Mg_3Si_2O_5(OH)_4 + 9 (Mg_{0.8}Fe_{0.2})(OH)_2 + Fe_3O_4 + H_2$ (S-8) Olivine Water Serpentine Ferroan brucite Magnetite

This leads to an initial mineral assemblage with molar fractions in ferroan brucite, olivine, serpentine and magnetite of 20 mol %, 52 mol %, 26 mol % and 2 mol %, respectively.

The production of H_2 during serpentinised dunite alteration at 363 K was considered to depend on three processes: (i) ferroan brucite alteration, (ii) olivine serpentinisation, and (iii) H_2 leakage due to fluid flow or diffusion (also referred to as ' H_2 escape rate' in the main text).

Regarding (i), ferroan brucite was considered to form magnetite according to Reaction 1 at the rate provided in Equation S-3. The value of $n_{H_2,eq}$ required in this latter equation was computed with PHREEQC considering H₂ in equilibrium with an assemblage composed of ferroan brucite + magnetite + chrysotile. The llnl.dat database was used with the thermodynamic data for ferroan brucite derived in the previous section.

Olivine serpentinisation (ii) was modelled according to Reaction S-8. The rate of H₂ production was modelled with the following surface-dependent zero-order rate law:

$$\frac{dn_{\rm H_2,ol}}{dt} = -\left(\frac{1}{24}\right)\frac{dn_{\rm ol}}{\partial t} = k_{\rm ol}SAM_{\rm ol}n_{\rm ol}(t). \tag{S-9}$$

 $M_{\rm ol}$ is the molecular weight of olivine. $k_{\rm ol}$ is calculated as $\frac{k_{\rm MD}}{s_{A_{\rm MD}}}$. One should note that, on the contrary to ferroan brucite alteration rate (Eq. S-3), $n_{\rm H_2,ol}$ does not depend on the total amount of H₂, $n_{\rm H_2}$. This is consistent with thermodynamic predictions of complete olivine reaction even at high activity in H₂ (*e.g.*, McCollom and Bach, 2009). Ferroan brucite formed during olivine serpentinisation (Reaction S-8) is also added to the amount of ferroan brucite available for H₂ production (ferroan brucite alteration (i)).

 H_2 "leakage" (iii) was assumed to be proportional to the H_2 concentration in the fluid expressed as the number of moles of H_2 in the fluid over the mass of water in contact with the rock at each time step:

$$\frac{dn_{\rm H_2,leak}}{dt} = \frac{-n_{\rm H_2}}{m_{\rm water}}J.$$
(S-10)

J is assumed here to be constant. For the fluid flow case, *J* corresponds to the rate of water renewal. In this latter case, a cumulative water to rock ratio can be calculated as the integral of *J* over time. Moreover, fluid flow also induces Mg, Si and Fe transport as aqueous species. The solubility of Mg, Fe and Si in the fluid at the equilibrium with serpentinised dunite were retrieved from the thermodynamic calculation performed with PHREEQC in (i). The predicted pH is 9.7. The solubility in Mg of 6.2×10^{-5} mol/kg at 363 K is approximately 4 and 5 orders of magnitude higher than for Si and Fe, respectively. Ferroan brucite is thus predicted to be the main phase to dissolve during the simulation. The product of Mg solubility by the cumulative water to rock ratio was used to estimate the extent of ferroan brucite dissolution associated with fluid flow.



For the diffusion case, *J* can be expressed as:

$$J = \frac{D\rho_{\text{water}A_{\text{h}}}}{z}.$$
(S-11)

Combining Equations S-10 and S-11 and assuming $n_{\text{H}_2} = 0$ at the surface (z = 0) allow to retrieve Fick's first law of diffusion. A maximum of $\frac{dn_{\text{H}_2,\text{leak}}}{dt}$ for diffusion can be estimated by using $n_{\text{H}_2} = n_{\text{H}_2,\text{eq}}$, *D* corresponding to H₂ diffusion in bulk water (no tortuosity; $5 \times 10^{-9} \text{ m}^2/\text{s}$ at 298 K), z = 2500 m by using a geothermal gradient of 30 K/km, $\rho_{\text{water}} = 1000 \text{ kg/m}^3$ and $A_{\text{h}} = \frac{m_{\text{rock}}}{\rho_{\text{rock}}h}$ with $\rho_{\text{rock}} = 3000 \text{ kg/m}^3$ and h = 1 m, the minimum estimated thickness for reacting serpentinised peridotites at depth. With these values, the maximum diffusive flux of H₂ is of $1.3 \times 10^{-15} \text{ mol H}_2/\text{day/g rock}$.

Equations S-3, S-9 and S-10 define a system of three differential equations. This system was numerically solved with an adaptive time step depending on the maximum of $n_{\rm H_2}$ variation for each process.



Supplementary Tables

Table S-1Symbols used.

Symbol	Definition	Units
$X_{\rm Fe(OH)_2}$	molar fraction of Fe(OH) ₂ in brucite	_
$X_{\rm Fe(OH)_2,eq}$	equilibrium molar fraction of $Fe(OH)_2$ in brucite at T and P	-
x	initial X _{Fe(OH)2}	_
v	parameter related to reaction progress as defined in Equation 2	_
r	hydrogen (H ₂) production rate	mol s ⁻¹
r_0	hydrogen production rate as defined in Equation S-3	mol s ⁻¹
k_0	reaction kinetic constant of ferroan brucite alteration	$mol m^{-2} s^{-1}$
kol	reaction kinetic constant of olivine serpentinisation	$mol m^{-2} s^{-1}$
k _{MD}	maximum reaction constant provided in McCollom and Donaldson (2016) for	mol kg ⁻¹ s ⁻¹
	H_2 production during olivine serpentinisation at 363 K	8
$n_{ m H_2}$	total amount of H ₂ produced in the experiment	mol
\bar{n}_{H_2}	total amount of H ₂ produced in the experiment normalised to the mass of starting	mol kg ⁻¹
-	Fe-brucite material	0
n _{H2,g}	amount of H ₂ in the gas phase	mol
n _{H2,eq}	equilibrium amount of H_2 produced at T and P	mol
n _{H2,ol}	amount of H_2 produced during Reaction S-8	mol
n _{H2,leak}	amount of H_2 leaking in the model of serpentinised dunite alteration	mol
n _{ol}	amount of olivine in the serpentinised dunite in the model of serpentinised	mol
01	dunite alteration	mor
m _{water}	mass of water in contact with the rock at each time step of the serpentinised	kg
water	dunite alteration model	1.5
m _{rock}	mass of rock in the serpentinised dunite alteration model	kg
λ	factor used to convert last measured $n_{\rm H_2}$ value into $n_{\rm H_2,eq}$	-
$n^{\circ}_{\rm Fe}, n^{\circ}_{\rm Mg}$	initial number of moles of Fe and Mg in ferroan brucite	mol
A re, n Mg	ferroan brucite specific surface area	$m^2 kg^{-1}$
SA _{MD}	surface area of the olivine used in McCollom and Donaldson (2016)	$m^2 kg^{-1}$
SA	specific surface area of olivine calculated with the relationship provided in	$m^2 kg^{-1}$
521	Brantley and Melott (2000) for a grain size of 500 μ m, typical for serpentinised	III Kg
	dunites in ophiolites (Malvoisin <i>et al.</i> , 2017)	
R	universal gas constant	J mol ⁻¹ K ⁻¹
T	temperature	K
$\frac{1}{Q_i}$	quotient of reaction <i>i</i> with $i = 1$ or 2	
$\frac{Q_i}{K_i}$	equilibrium constant of reaction <i>i</i> with $i = 1$ or 2	_
$\frac{K_l}{E_a}$	activation energy of reaction	J mol ⁻¹
$P_{\rm H_2}$	partial pressure of H_2 in the gas phase	bar
	partial pressure of H_2 in the gas phase at the equilibrium	bar
<i>P</i> _{H₂,eq} [H ₂]	concentration of H_2 in the aqueous phase	$mol m^{-3}$
$K_{\rm H}$	Henry's coefficient for H_2 gas	Pa m ³ mol ^{-1}
		m^3
V _g	volume of the gas phase	m^3
V_{aq} $\Delta_r G^{P,T}$	volume of the aqueous phase Cibbs free energy of reaction at T and P	
$\Delta_r G^{P,T}$ $\Delta_a H_i^{P,T}$	Gibbs free energy of reaction at T and P	$J \text{ mol}^{-1}$
$\frac{\Delta_{a}H_{i}}{S_{i}^{P,T}}$	apparent enthalpy of phase <i>i</i> at <i>T</i> and <i>P</i>	$J \text{ mol}^{-1}$
-	third-law entropy of phase <i>i</i> at <i>T</i> and <i>P</i>	$J \text{ mol}^{-1} \text{ K}^{-1}$
$\frac{\Delta_{\rm f}H_i^\circ}{\Delta_{\rm f}C^\circ}$	standard formation enthalpy of phase <i>i</i> (from the elements)	J mol ⁻¹
$\frac{\Delta_{\rm f}G_i^{\circ}}{\sigma^{\circ}}$	standard formation Gibbs free energy of phase <i>i</i>	$J \text{ mol}^{-1}$
S_i°	standard third-law entropy of phase <i>i</i>	$J \text{ mol}^{-1} \text{ K}^{-1}$
Cp_i	molar heat capacity function of phase <i>i</i> valid in the T_r - <i>T</i> temperature range	$J \text{ mol}^{-1} \text{K}^{-1}$
V_i^0	molar volume of phase <i>i</i> at reference P_r and T_r	$m^3 mol^{-1}$
a _i	activity of phase <i>i</i>	_
ν_i	stoichiometric coefficient of phase <i>i</i>	—



Table S-1 continued.

Symbol	Definition	Units
Tr	reference temperature (298 K)	K
$P_{\rm r}$	reference pressure (0.1 MPa)	Pa
T_j	temperature at equilibrium for experiment <i>j</i>	K
P_j	pressure at equilibrium for experiment <i>j</i>	Pa
fj	sum of parameters which can be calculated from tabulated data of Klein <i>et al.</i>	$J \text{ mol}^{-1} \text{ K}^{-1}$
	(2009) in Equation S-6	
J	constant fixing the H ₂ escape rate in the model of serpentinised dunite alteration	kg s ⁻¹
D	diffusion coefficient of H_2 in the fluid	$m^2 s^{-1}$
$ ho_{ m water}$	water density	kg m ⁻³
$ ho_{ m rock}$	rock density	kg m ⁻³
$A_{ m h}$	area of the horizontal section through which H ₂ diffuses	m ²
h	vertical thickness of the serpentinised peridotite layer reacting in the model of	m
	serpentinised dunite alteration	

Table S-2 Experimental conditions for each experiment, produced H₂ and XRD analysis results. Expected H₂ production assuming equilibrium is also calculated with PHREEQC using the thermochemical data retrieved in the present study. f-brc, ferroan brucite; mag, magnetite; pyr, pyroaurite; L/V, liquid-vapor equilibrium; n.d., not detected; n.c., not calculated; thermo., thermodynamic data used for determining thermodynamic parameters of Fe(OH)₂; and kin., kinetic data used for determining the kinetic law (predicted duration to reach measured n_{H_2} is not provided for these experiments). * 53.3 ± 16.5 wt. % Fe(III)-bearing brucite (see Carlin *et al.*, 2023) have also been detected in Run caps#t4. Errors are provided as ±3 times the standard deviation.

	Initial experimental conditions						1	XRD a	nalysis outpu	t		PHREEQC	Kinetics	
run name	duration (h)	temperature (K)	initial pressure at T (MPa)	initial f-brc mass (g)	mass water- rock ratio	measured $n_{\rm H_2}$ from GC (mol)	$X_{\rm Fe(OH)_2}$ in initial f-brc (%)	$X_{\rm Fe(OH)_2}$ in final f-brc (%)	final f-brc (wt. %)	mag (wt. %)	pyr (wt. %)	$\begin{array}{c} \text{simulated} \\ n_{\text{H}_2} \text{ at} \\ \text{equilibrium} \\ (\text{mol}) \end{array}$	predicted duration to reach measured $n_{\rm H_2}$ (h)	use
caps#1	360	378	20	0.01-0.03	3.33-10	2.87E-09	15.6±2.1	11.0±1.2	93.8±1.2	3.2±0.6	3.3±0.9	1.80E-09	146	thermo.
caps#2	624	378	20	0.032	3.13	8.49E-09	15.6±2.1	12.9±1.8	95.3±1.2	4.7±1.2	n.d.	1.80E-09	267	thermo.
caps#3	984	423	20	0.046	2.17	2.11E-07	18.5±1.8	11.6±0.6	90.8±1.2	9.2±1.2	n.d.	3.07E-08	29	thermo.
caps#4	984	473	20	0.048	2.08	1.29E-06	18.5±1.8	15.8±0.3	92.8±0.6	7.2±0.6	n.d.	3.48E-07	1	thermo.
caps#6	1344	348	20	0.049	2.06	1.91E-10	17.9±2.1	14.7±1.5	89.2±2.1	6.3±1.2	4.5±1.8	5.50E-10	214	thermo.
caps#7	1344	403	20	0.049	2.06	1.91E-09	17.9±2.1	12.9±1.5	90.3±1.5	9.7±1.5	n.d.	1.00E-08	2	thermo.
caps#9	60	378	20	0.040	2.50	1.21E-07	18.1±2.7	13.7±2.1	99.8±0.3	0.2 ± 0.3	n.d.	2.82E-09	2707	not used
caps#10	60	423	20	0.050	2.00	1.75E-07	18.1±2.7	10.2±2.4	97.4±0.9	2.6±0.9	n.d.	2.87E-08	23	thermo.
caps#11bis	60	473	20	0.050	2.00	5.88E-07	18.1±2.7	18.5±0.9	95.1±0.9	4.9±0.9	n.d.	3.27E-07	0.8	thermo.
caps#12bis	60	523	20	0.046	2.17	3.68E-06	18.1±2.7	16.2±0.9	92.6±0.9	8.4±0.9	n.d.	2.65E-06	< 0.1	thermo.
caps#13	296	523	20	0.059	1.69	1.42E-06	18.1±2.7	19.8±0.3	92.0±0.9	8.0±0.9	n.d.	2.74E-06	< 0.1	thermo.
caps#14	240	573	20	0.025	4.00	5.45E-06	18.1±2.7	12.1±0.6	83.6±1.2	16.4±1.2	n.d.	7.81E-06	< 0.1	thermo.
caps#15	48	573	20	0.053	1.89	6.08E-06	18.1±2.7	15.6±0.6	87.3±1.2	12.7±1.2	n.d.	1.14E-05	< 0.1	thermo.
SP#3	192	473	1.55 (L/V)	1.52	24.42	1.71E-04	15.6±2.1	20.1±0.3	94.0±0.9	6.0±0.9	n.d.	3.20E-04		kin./thermo.
SP#4	312	473	1.55 (L/V)	0.31	64.52	6.06E-05	18.5 ± 1.8	16.0±0.6	90.7±0.6	9.3±0.6	n.d.	1.85E-04		kin./thermo.
SP#5	408	423	0.476 (L/V)	1.52	24.41	2.45E-05	17.9±2.1	17.7±0.9	93.4±0.9	6.6±0.9	n.d.	1.16E-04		kin./thermo.
SP#6	1656	378	5	1.495	19.80	4.89E-06	18.1±2.7	19.5±1.2	94.9±1.2	3.1±0.6	2.1±0.9	2.81E-05		kin./thermo.

Table S-2 continued.

Initial experimental conditions							XRD a	nalysis outp	ut		PHREEQC	Kinetics		
run name	duration (h)	temperature (K)	initial pressure at T (MPa)	initial f-brc mass (g)	mass water- rock ratio	measured $n_{\rm H_2}$ from GC (mol)	$X_{\rm Fe(OH)_2}$ in initial f-brc (%)	$X_{\rm Fe(OH)_2}$ in final f-brc (%)	final f-brc (wt. %)	mag (wt. %)	pyr (wt. %)	simulated $n_{\rm H_2}$ at equilibrium (mol)	predicted duration to reach measured $n_{\rm H_2}$ (h)	use
caps#t1	0.5	378	0.121 (L/V)	0.0624	1.60	8.13E-10	20.5±1.8	11.5±1.5	86.6±3	1.6±0.9	11.8±3	4.19E-07		not used
caps#t2	2	378	0.121 (L/V)	0.0639	1.56	5.58E-10	20.5±1.8	14.6±1.5	95.4±1.5	1.5±0.9	3.1±0.9	4.18E-07		not used
caps#t3	143	378	0.121 (L/V)	0.0656	1.52	1.53E-08	20.5±1.8	13.5±1.2	84.8±2.4	2.8±0.9	12.4±2.4	4.17E-07		not used
caps#t4	335	378	0.121 (L/V)	0.0567	1.76	4.62E-10	20.5±1.8	n.c.	39.4±2.1*	3.1±0.9*	18.0±3.9*	4.23E-07		not used
caps#t5	0.5	423	0.476 (L/V)	0.0421	2.38	7.33E-10	20.5±1.8	13.7±1.5	90.2±2.4	1.8±0.9	8.1±2.4	2.65E-06		kin.
caps#t6	72	423	0.476 (L/V)	0.027	3.71	8.79E-09	20.5±1.8	14.6±0.9	98.2±0.6	1.8±0.6	n.d.	2.53E-06		kin.
caps#t7	163	423	0.476 (L/V)	0.0509	1.97	4.93E-08	20.5±1.8	15.4±0.9	97.9±0.6	2.1 ± 0.6	n.d.	2.67E-06		kin.
caps#t8	483	423	0.476 (L/V)	0.0519	1.93	2.02E-07	20.5±1.8	15.0±0.9	97.6±0.6	2.4±0.6	n.d.	2.67E-06		kin.
caps#t9	721	423	0.476 (L/V)	0.0527	1.90	5.57E-07	20.5±1.8	15.8±0.9	96.9±0.6	3.1±0.6	n.d.	2.68E-06		kin.
caps#t11	1872	423	0.476 (L/V)	0.0557	1.80	1.38E-06	20.5±1.8	20.8±0.3	95.6±0.6	4.4±0.6	n.d.	2.69E-06		kin.
caps#t12	3552	423	0.476 (L/V)	0.0454	2.20	2.33E-06	20.5±1.8	19.7±0.3	94.1±0.9	6.0±0.9	n.d.	2.63E-06		kin./thermo.

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Table S-3			$(n_{\rm H_2})$ measu	red with gas chromatography as a
function of time	me for SP#6 (378 K, 5 MPa).	•		
			measured	

duration (h)	measured n _{H2} from GC (mol)
0	1.12E-07
5	1.44E-07
22	1.80E-07
51	2.31E-07
77	2.53E-07
150	3.24E-07
188	3.63E-07
219	3.41E-07
312	5.17E-07
363	5.22E-07
744	2.76E-06
987	3.55E-06
1078	4.00E-06
1198	4.42E-06
1323	4.58E-06
1390	4.73E-06
1487	4.70E-06

Table S-4 Summary of available Fe(OH)₂ thermodynamic data. *a*, *b* and *c* are the coefficients of the equation of Maier and Kelley (1932) for C_p : $C_p = a + bT + cT^{-2}$.

References	Δ _f H° kJ/mol	Δ _f G° kJ/mol	S° J/mol/K	<i>a</i> J/mol/K	<i>b</i> (10 ³) J/mol/K ²	<i>c</i> (10 ⁻⁵) J K/mol
Wagman et al. (1982) - NBS	-569.0	-486.5	88.0			
Chase (1998) - NIST-JANAF	-574.045	-491.97	87.864	97.069		
Leussing and Kolthoff (1953)		-492.58				
Refait et al. (1999)		-490				
Ziemniak et al. (1995)	-583.39	-500.16	84	90		
Sverjensky and Molling (1992)		-494.97				
McCollom and Bach (2009)	-574.61*	-492.58†	88.0‡	109.035	18.192	-22.51
This study	-581.3*	-498.90	86.4	109.035**	18.192**	-22.51**

*: computed from $\Delta_{\rm f}G^{\circ} = \Delta_{\rm f}H^{\circ} - 298.15\Delta_{\rm f}S^{\circ}$ with the values of $\Delta_{\rm f}H^{\circ}$ and S° provided in the table and standard entropies of the elements taken from Helgeson *et al.* (1978).

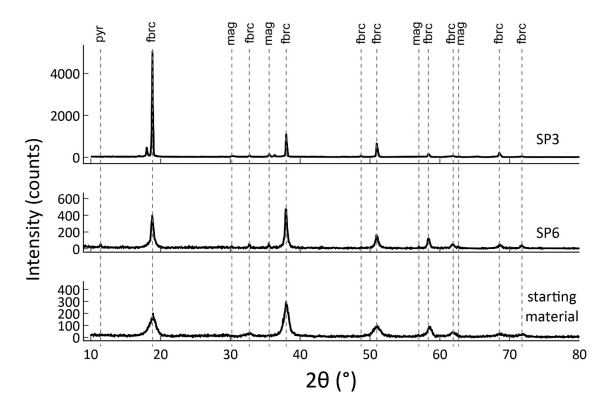
†: taken from Leussing and Kolthoff (1953).

: taken from Wagman et al. (1982).

**: taken from McCollom and Bach (2009), calculated as a linear function of brucite, greenalite and chrysotile C_p functions.



Letter



Supplementary Figures

Figure S-1 Selected XRD patterns. From bottom to top: synthetic ferroan brucite starting material, Run SP#6 (378 K, 69 days) and Run SP#3 (473 K, 8 days). fbrc, ferroan brucite; mag, magnetite; pyr, pyroaurite. CuKα radiation.

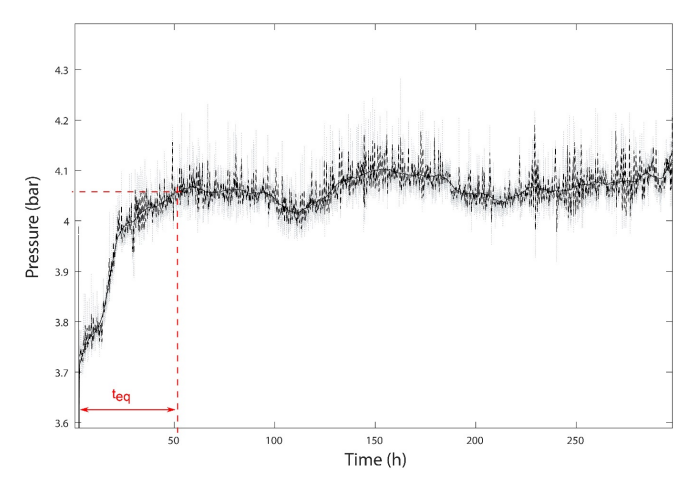


Figure S-2 In situ pressure monitoring for SP#5. The grey dotted line, the black dashed line and the solid black curve correspond, respectively, to the raw pressure data, the pressure filtered with a low pass filter, and smoothed filtered pressure. The time to reach equilibrium (t_{eq} ; red dashed lines) is graphically determined.

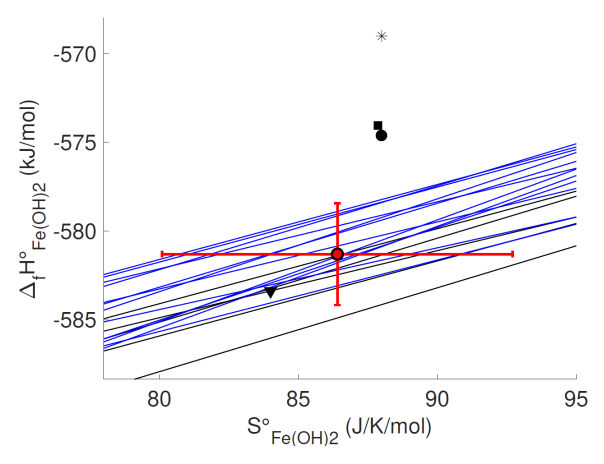


Figure S-3 $\Delta_{f}H^{\circ}_{Fe(OH)2}$ vs. $S^{\circ}_{Fe(OH)2}$ relationship determined for capsules experiments (blue lines) and titanium reactor experiments (black lines). The best-fitting $\Delta_{f}H^{\circ}_{Fe(OH)2}$ and $S^{\circ}_{Fe(OH)2}$ values \pm their standard error determined by linear regression ($R^{2} = 0.926$) are displayed with a red dot. The black circle, square, triangle and star correspond to the thermodynamic data of McCollom and Bach (2009), Chase (1998), Ziemniak *et al.* (1995) and Wagman *et al.* (1982), respectively.

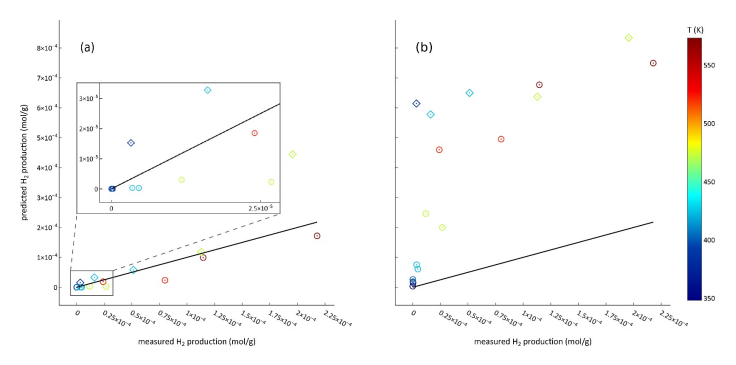


Figure S-4 Measured *vs.* predicted moles of H_2 produced per gram of ferroan brucite in experiments SP#3 to #6 and caps#t12 (diamond symbols), and caps#1 to #15 (circle symbols). The predicted amount of H_2 is equal to the measured amount of H_2 on the black solid line. The predictions were calculated with PHREEQC using the thermodynamic data for Fe(OH)₂ (a) derived here and (b) from Klein *et al.* (2009).

Supplementary Information References

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