The effect of pH on rates of reaction and hydrogen generation during serpentinization

Thomas M. McCollom¹, Frieder Klein², Peter Solheid³,⁴ and Bruce Moskowitz³,⁴

¹Laboratory for Atmospheric and Space Physics, University of Colorado, Boulder, CO 80309, USA
²Marine Chemistry and Geochemistry Department, Woods Hole Oceanographic Institution, Woods Hole, MA 02543, USA
³Department of Earth Sciences, and ⁴Institute for Rock Magnetism, University of Minnesota, Minneapolis, MN 55455, USA

A series of three laboratory experiments were conducted to investigate how pH affects reaction pathways and rates during serpentinization. Two experiments were conducted under strongly alkaline conditions using olivine as reactant at 200 and 230°C, and the results were compared with previous studies performed using the same reactants and methods at more neutral pH. For both experiments, higher pH resulted in more rapid serpentinization of the olivine and generation of larger amounts of H₂ for comparable reaction times. Proportionally greater amounts of Fe were partitioned into brucite and chrysotile and less into magnetite in the experiments conducted at higher pH. In a third experiment, alkaline fluids were injected into an ongoing experiment containing olivine and orthopyroxene to raise the pH from circumneutral to strongly alkaline conditions. Increasing the pH of the olivine-orthopyroxene experiment resulted in an immediate and steep increase in H₂ production, and led to far more extensive reaction of the primary minerals compared to a similar experiment conducted under more neutral conditions. The results suggest that the development of strongly alkaline conditions in
actively serpentinizing systems promotes increased rates of reaction and \( H_2 \) production, enhancing the flux of \( H_2 \) available to support biological activity in these environments.

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1. Introduction

Two of the defining characteristics of fluids discharged from low temperature (less than approx. 250°C), actively serpentinizing systems are their strongly alkaline \( pH \) and highly elevated \( H_2 \) concentrations (e.g. [1–6]). The \( H_2 \) in these fluids provides a source of metabolic energy to support lithoautotrophic microbial communities in alkaline springs on land and in serpentinite-hosted hydrothermal vents on the ocean floor [5–8]. Because the ultramafic rocks that undergo serpentinization are widespread in the solar system, similar processes may also generate \( H_2 \) and support biological activity on other rocky planetary bodies [9–11]. Understanding the capacity of serpentinizing systems to support biological communities on Earth and beyond, therefore, requires knowledge of the flux of \( H_2 \) that is generated as the reaction proceeds.

For this and other reasons, the serpentinization of ultramafic rocks has been the subject of progressively increasing numbers of laboratory experimental studies over the last few decades. Many of these studies have focused on understanding the impact that environmental variables such as temperature, rock composition, salinity and the activities of dissolved \( SiO_2 \) and Al have on reaction rates and on the composition of products during serpentinization [12–28]. To date, however, there has been no systematic effort to study how \( pH \) affects serpentinization reactions. Nevertheless, the strongly alkaline conditions that prevail in many low-temperature serpentinizing environments suggests that the potential contribution of \( pH \) to reaction progress needs to be evaluated.

Previous experimental studies provide only limited information about the effect of strongly alkaline \( pH \) on serpentinization reactions. Lafay et al. [24] reacted olivine with extremely alkaline solutions (\( pH 13.5 \)) and observed overall reaction rates that were much higher than observed in other recent studies of olivine serpentinization that were conducted at comparable temperatures and lower \( pH \) [12,13]. On the other hand, the rates observed by Lafay et al. were similar to those reported in other experimental studies [27,28], although a lack of information on the \( pH \) of the latter studies precludes evaluation of its possible role in higher reaction rates. In some other experiments, a transition from mildly alkaline to strongly alkaline \( pH \) during serpentinization was observed to be accompanied by a steep increase in \( H_2 \) generation [14,19,25,26]. Taken together, these results suggest that higher \( pH \) may increase overall reaction rates and alter reaction pathway to enhance \( H_2 \) production, but the role of \( pH \) during these experiments remains undetermined. By contrast to the results of serpentinization studies, laboratory experiments designed to monitor the dissolution of olivine or pyroxene have observed monotonically decreasing rates with increasing \( pH \) [29–32]. If the dissolution of the primary minerals is the rate-limiting step during serpentinization as suggested by some experimental studies [12,16], these observations would imply that the process should become slower with increasing \( pH \).

In an effort to better understand the effect of \( pH \) on reaction pathways and rates during serpentinization, a series of three hydrothermal laboratory experiments were conducted for this study. The results reveal modified reaction pathways and increased rates of serpentinization and of \( H_2 \) formation at high \( pH \). We discuss possible mechanisms to account for observed \( pH \) effects and assess implications for natural systems undergoing serpentinization at moderate to low temperatures.

2. Methods

Three separate experiments were performed for this study, with initial ingredients and reaction parameters summarized in table 1. Two experiments were conducted by reacting olivine alone
reaction cell with titanium fittings [33]. The reaction cell was contained within a stainless steel
experiment. All titanium fittings that were exposed to reactants were combusted in air for at least
samples were taken during the experiments in order to monitor reaction progress over time. The
phase. The cell was equipped with a capillary tube attached to a valve through which fluid
eliminated the presence of a vapour headspace so that reactions were confined to the aqueous
reaction cell allowed fluids to be sampled from the reaction cell without loss of pressure and
pressure housing, with water used as the external pressurizing fluid. The flexibility of the gold
previous experiments are also listed in table 1). This parallel design allows for direct evaluation
a less strongly alkaline pH [13] (to facilitate comparison, the starting ingredients for the
were designed to closely parallel previous olivine serpentinization experiments that attained
of the impact of pH on the reactions, since pH is essentially the only reaction parameter
that varied between the experiments. The third experiment reacted a mixture of olivine and
orthopyroxene (Opx) at 230°C (OlivOpx230pH). This experiment was allowed to react at 230°C
and circumneutral pH for an extended period of time (approx. 4500 h). Additional fluid was
then injected to raise the pH, and the experiment was allowed to continue to react for another
3200 h. Again, the reactants and methodology of this experiment were designed to closely parallel
previous experiments with olivine–Opx mixtures as starting materials that developed a less
alkaline pH as serpentinization proceeded [14].

The experiments were conducted by heating powdered mineral substrates in the presence
of aqueous solutions. The fluid composition was monitored during the experiments for the
production of H₂ and other dissolved compounds as the reactions proceeded. Solids were
recovered at the end of the experiments and analysed for their mineral and chemical composition.
Methods used to analyse the fluids and solid reaction products are summarized briefly below,
with additional details provided as electronic supplementary material. All aspects of the
experimental and analytical procedures are the same as those used in other recent studies [13,14].

The experiments were conducted in a flexible-cell hydrothermal apparatus using a gold
reaction cell with titanium fittings [33]. The reaction cell was contained within a stainless steel
pressure housing, with water used as the external pressurizing fluid. The flexibility of the gold
reaction cell allowed fluids to be sampled from the reaction cell without loss of pressure and
eliminated the presence of a vapour headspace so that reactions were confined to the aqueous
phase. The cell was equipped with a capillary tube attached to a valve through which fluid
samples were taken during the experiments in order to monitor reaction progress over time. The
valve also allowed additional fluid to be injected into the reaction cell without disturbing the
experiment. All titanium fittings that were exposed to reactants were combusted in air for at least

| experiment      | source         | temperature (°C) | mineral reactants          | initial fluid (g) | fluid composition          |
|-----------------|----------------|------------------|---------------------------|------------------|---------------------------|
| Oliv230pH       | this study     | 230              | 15.0 g olivine, <53 μm    | 45.8             | 480 m NaCl, 3 m Na₂CO₃, 33.8 m NaOH; pH 12.5 |
| Oliv230fine     | (a)            | 230              | 16.0 g olivine, <53 μm    | 39.5             | 485 m NaCl, 20 m NaHCO₃; pH 7.8 |
| Oliv200pH       | this study     | 200              | 14.1 g olivine, 38–53 μm  | 51.2             | 483 m NaCl, 2 m Na₂CO₃, 32.9 m NaOH; pH 12.4 |
| Oliv200fine     | (a)            | 200              | 17.0 g olivine, 38–53 μm  | 46.2             | 486 m NaCl, 20 m NaHCO₃; pH 6.4 |
| OlivOpx230pH    | this study     | 230              | 16.7 g olivine + 2.8 g Opx, 53–212 μm | 35.8         | 486 m NaCl, 20 m NaHCO₃; pH 7.8 |
| OlivOpx230med   | (b)            | 230              | 18.0 g olivine + 3.2 g Opx, 53–212 μm | 45.4         | 485 m NaCl, 20 m NaHCO₃; pH 7.8 |
| OlivOpx230      | (b)            | 230              | 21.0 g olivine + 3.5 g Opx, 53–212 μm | 44.0         | 485 m NaCl, 20 m NaHCO₃; pH 7.8 |

Table 1. Summary of experimental parameters, with parameters used in comparable lower pH studies discussed in the text shown for reference. m = mmolal. Sources: (a) McCollom et al. [13] and (b) McCollom et al. [14].
48 h at 400°C prior to use in the experiments in order to form a relatively inert TiO₂ surface layer. Experiments were conducted at a pressure of 35 MPa.

The initial reactant fluid contained 485 mmol NaCl kg⁻¹ to approximate seawater salinity, with NaOH added to increase the pH as appropriate. In order to monitor for the possible formation of hydrocarbons from the reduction of inorganic carbon [34], the experiments also included a source of dissolved inorganic carbon (DIC) in the form of either NaHCO₃ or Na₂CO₃ at concentrations ranging from 2 to 20 mmol kg⁻¹. The initial water : rock mass ratio ranged from 1.8 to 3.6, although the ratio within the reaction vessel decreased somewhat during the experiments as fluid aliquots were removed for chemical analysis and as water was incorporated into secondary phases.

The reactant minerals were pulverized by hand using a ceramic mortar and pestle, always avoiding the use of metal tools that could contaminate the reactants with metal particles that might generate H₂ during heating. The resulting powders were sieved to achieve a defined range of grain sizes. For experiments using grains in the 38–53 µm or 53–212 µm size range, the sieved materials were washed repeatedly with deionized water (DI) to remove small particulates. The minerals used in the experiments were purchased from Excalibur Minerals (Charlottesville, VA, USA). The source of the olivine was mantle xenoliths from San Carlos, Arizona. Only olivine crystals larger than approximately 4 mm in diameter were used to prepare the mineral reactants, and partially crushed materials were hand-picked under a binocular microscope to exclude grains with obvious signs of weathering or inclusions of other minerals. Despite efforts to purify the reactant minerals, the olivine used in the experiments contained trace amounts (≪1 vol%) of chromian spinel, Opx and clinopyroxene. The Opx used in the experiments was from Bamble, Norway [35]. The prepared Opx reactant included less than 1% of several other contaminants including quartz, Ca-phosphate (probably chlorapatite), talc and albite. Magnetization analysis indicated the olivine and Opx prepared for the experiments contained 0.0001% and approximately 0.14 wt% magnetite, respectively. Chemical compositions of the reactant minerals are summarized in table 2.

At intervals throughout the experiments, several fluid aliquots (0.3–1 g each) were taken directly into glass gas-tight syringes and analysed for dissolved volatile species including H₂, total dissolved CO₂ (\[\sum CO₂ = CO₂(\text{aq}) + HCO₃⁻ + CO₃²⁻\]), CH₄ and C₂–C₆ hydrocarbons. These compounds were measured by gas chromatography (GC) with thermoconductivity detection (TCD) or flame ionization detection (FID) following a headspace extraction. In most cases, the reported H₂ concentrations are averages of three to four separate fluid aliquots, while other dissolved gases were typically only measured for a single aliquot to conserve fluid. For comparative purposes, the measured H₂ concentrations were normalized to the total number of moles generated per gram of reactant minerals to account for variations in the amounts of solids used in different experiments and for the changing mass of fluid present in the reaction vessel as aliquots are removed during sampling. Only trace amounts (less than 1 µmol kg⁻¹) of C₂–C₆ hydrocarbons were observed in any of the experiments, so individual results are not reported. Estimated errors for volatile analyses are ±5%. Fluid aliquots remaining after extraction of volatiles were used for measurements of pH, total dissolved SiO₂ (\[\sum SiO₂\]) and major elements. Detailed methods for fluid analyses are provided in the electronic supplemental material.

At the termination of the experiments, the solids were removed, filtered and rinsed with DI water. The solids were characterized by a variety of methods including scanning electron microscopy (SEM) coupled with energy-dispersive X-ray spectroscopy (EDS), X-ray diffraction (XRD), electron microprobe analysis (EMPA), confocal Raman spectroscopy, thermogravimetric analysis (TGA) and magnetic methods (see electronic supplementary material for details). To obtain additional information on the distribution of Fe among secondary phases, Mössbauer spectroscopy (MB) was also performed on the reaction products.

Thermodynamic calculations were conducted to determine fluid speciation, estimate in situ pH at experimental conditions, and construct mineral stability diagrams. The calculations used the 35 MPa database described by McCollom & Bach [36], which is based on thermodynamic data from Helgeson et al. [37], Shock & Helgeson [38] and Shock et al. [39,40]. Thermodynamic properties for solid solutions were calculated from end-member data assuming ideal mixing.
Fluid speciation calculations to estimate in situ pH and silica activity were performed in two steps using the computer program EQ3 [41]. First, the fluid was speciated at 25°C using the measured fluid compositions, including the room temperature pH, and adjusted for charge balance with Na. The total dissolved Na calculated at 25°C was then used with other measured concentrations to respeciate the fluid at the reaction temperature, with charge balance determining the in situ pH. In those cases where Mg, Ca and Fe were below detection limits, the total dissolved abundance of these elements were assumed to be 1 µmol kg⁻¹, 1 µmol kg⁻¹, 0.1 µmol kg⁻¹, respectively. Mineral stability diagrams were constructed using the computer program Geochemist’s Workbench (Aqueous Solutions LLC, Champaign, IL, USA).

3. Results

The results of individual experiments are summarized in the following subsections. Key observations from the experiments are summarized in Table 3. Selected components of the fluids are shown in the accompanying figures, along with fluid compositions from analogous experiments from previous studies performed under circumneutral to mildly alkaline conditions. A table providing more complete measured fluid compositions for experiments from the present study is included in the electronic supplementary material, Table S1. Analyses of solid products by XRD are shown in electronic supplementary material, figure S1, and are incorporated into the summaries below.

(a) Reaction of olivine at 230°C

Experiment Oliv230pH was conducted to evaluate the reaction of olivine with a strongly alkaline pH solution at 230°C (Table 1). The temperature and initial reactants for this experiment were
Table 3. Summary of experimental results from this and previous studies. Numbers in parentheses are standard deviations. The *in situ* pH is calculated from the measured room temperature pH as described in the methods section. ‘—’, not determined or not present; b.d., below detection; Srp, serpentine; Brc, brucite; Mgt, magnetite.

| experiment       | temperature (°C) | duration (h) | final pH₂5°C | final pH₂,situ | final H₂ conc. (mmol kg⁻¹) | total H₂ generated (μmol g⁻¹ minerals) | Mg#   | srp. | brc. | Wt% mgt. | estimated wt% reaction |
|------------------|------------------|--------------|---------------|----------------|-----------------------------|------------------------------------------|-------|------|------|----------|------------------------|
| Oliv230pH        | 230              | 2972         | 12.3          | 9.3            | 47                          | 105                                      | 93 (0.9) | 78 (1) |      | 1.6      | 70                     |
| Oliv230finea     | 230              | 4293         | 9.1           | 7.8            | 25                          | 47                                       | 97 (0.4) | 92 (1.2) |      | 1.75     | 25                     |
| Oliv200pH        | 200              | 7677         | 11.3          | 8.5            | 8.4                         | 20.9                                     | 95 (0.7) | 84 (1.8) |      | 1.0      | 41                     |
| Oliv200finea     | 200              | 9219         | 9.2           | 8.2            | 2.9                         | 5.9                                      | 95 (0.8) | 91 (1.1) |      | 0.25     | 5.9                    |
| OlivOpx230pH     | 230              | 7715         | 10.3          | 8.7            | 7.4                         | 13.5                                     | 88 (1.4) |      |      | 0.02     | 17                     |
| OlivOpx230medb   | 230              | 2709         | 7.2           | 7.8            | 0.008                       | 0.016                                    | —      |      |      | b.d.      | 1.9                    |
| OlivOpx230b      | 230              | 9287         | 11.7          | 8.8            | 61.0                        | 90.7                                     | 91 (0.9) | 87 (1.7) |      | 1.52     | 53                     |

*a* McCollom et al. [13]; note that values for estimated wt% reaction are slightly lower than originally reported owing to an error discovered in the calculations.  
*b* McCollom et al. [14].
Figure 1. Fluid compositions during olivine experiment Oliv230pH (squares), with results for Oliv230fine (diamonds) from McCollom et al. [13] shown for comparison. (a) Total amount of H₂ generated per gram mineral reactant. (b) Measured room temperature pH (pH₂₅°C) and calculated in situ pH (pH_in_situ). (c) Total dissolved SiO₂ concentration. (Online version in colour.)

essentially the same as those used in experiment Oliv230fine reported in McCollom et al. [13] (table 1), except that the initial pH of the reactant fluid for Oliv230pH was increased to 12.4 by the addition of NaOH (versus an initial pH of 7.8 in Oliv230fine). The amount of DIC added was also reduced (from 20 mmol kg⁻¹ to 3 mmol kg⁻¹) to limit the pH buffering capacity of the fluid and the precipitation of carbonate minerals. Measured compositions of selected components in the fluid during Oliv230pH are shown in figure 1, with fluid compositions during Oliv230fine shown for comparison.

The room temperature pH (pH₂₅°C) of Oliv230pH was 11.8 after 2 days of heating, but rose to 12.3 by the time of the second sample at 333 h and then remained near this level for the remainder of the experiment (figure 1b). The calculated pH at experimental conditions (pH_in_situ) reached 9.3 after 333 h and remained at this level throughout the experiment, reflecting strongly alkaline conditions (figure 1b; for reference, neutral pH at the temperature of the experiment is about 5.5). In experiment Oliv230fine without the added NaOH, the pH₂₅°C was near 8.9 at the start of the experiment, and levelled off at around 9 for the remainder of the experiment. The calculated pH_in_situ for Olive230fine rose to 8.2 after 262 h, and then slowly declined to a final value of 7.8 at the termination of the experiment (figure 1b).
Generation of H₂ in Oliv230pH began immediately upon heating and continued throughout the experiment, although the rate appeared to taper off slightly as the reaction progressed (figure 1a). By the end of the experiment, a total of 105 µmol (g olivine)^−1 of H₂ had been produced (note that here and elsewhere the amount of H₂ generated is indexed to the mass of the original amount of mineral reactant, not the proportion of minerals that had reacted). The generation of H₂ also began immediately in Oliv230fine and increased steadily during the experiment. However, rates of H₂ generation in Oliv230pH were about three times higher than was observed for Oliv230fine (figure 1a).

The concentration of total dissolved silica (ΣSiO₂) was relatively high (480 µmol kg^−1) at the start of Oliv230pH, and decreased to lower levels as the experiment progressed to attain a final concentration of 240 µmol kg^−1 (figure 1c). The ΣSiO₂ concentration at 333 h was significantly lower than other measurements during this experiment; however, it is not clear whether this represents an actual dip in concentration or a spurious measurement. In any case, ΣSiO₂ concentrations throughout Oliv230pH remained well above those observed in the lower pH Oliv230fine experiment (less than or equal to 43 µmol kg^−1; figure 1c). The concentration of Na in the fluid from Oliv230pH remained constant during the experiment, and most other cations (Mg, Fe, Ca, Al) were present at levels below a few micromoles per kilogram (electronic supplementary material, table S1). However, significant amounts of dissolved K at approximately 7 mmol kg^−1 were observed in all samples. The source of K for the experiments is uncertain, but is likely an unidentified trace mineral present in the initial reactants. Similar concentrations of cations were observed during Oliv230fine, although K was much lower at approximately 0.18 mmol kg^−1.
Secondary mineral products of Oliv230pH were dominated by chrysotile and brucite, along with minor magnetite and trace amounts of calcite (figure 2; electronic supplementary material, figure S1). Also present scattered among the products were very fine-grained particles with elevated Ni and Fe contents as indicated by EDS analysis (electronic supplementary material, figure S2). The lack of elevated amounts of S or O associated with these particles suggests that they likely consist of awaruite (nominally Ni$_3$Fe). In addition to these secondary products, a small fraction of the original olivine remained at the termination of the experiment (table 3).

The chrysotile formed thick fibrous accumulations, while the brucite was generally present as platy hexagonal crystals embedded in the chrysotile (figure 2a,c). Although the original reactants were powders, the chrysotile fibres cemented the products into a solid mass with limited porosity (figure 2b). In some cases, the brucite was found in close contact with remnant olivine (figure 2b). The chrysotile has a Mg# [ = 100 \times \text{Mg}/(\text{Mg} + \text{Fe}), molar basis] equal to 93, while brucite has a Mg# of 78 (table 2). No spatial variation in the chemical composition of these minerals was observed. Analyses of brucite crystals by EMPA and EDS showed that it contains up to 2 wt% Cl, indicating the brucite is intergrown with iowaite as observed in previous studies [13,14]. For both chrysotile and brucite, the minerals are more enriched in Fe (i.e. lower Mg#) than was observed for Oliv230fine (table 3), which reflects favoured partitioning of Fe into chrysotile and brucite in the higher pH experiment. The products of Oliv230pH contained 1.60 wt% magnetite versus 1.75 wt% in Oliv230fine.

Mössbauer analysis of the secondary products from Oliv230pH indicated the presence of four doublets, with two peaks assigned each to Fe$^{II}$ and Fe$^{III}$ (table 4). The three main doublets (Fe$^{II}$ and Fe$^{III}$) had hyperfine parameters consistent with chrysotile, with the occupation of both octahedral and tetrahedral sites by the Fe$^{III}$ [42]. Adding a second ferrous doublet (assigned to...
Figure 3. Results of thermogravimetric analysis of reacted solids for laboratory experiments, with results from previous studies shown for comparison. (a) Olivine experiments at 230°C, (b) olivine experiments at 200°C, and (c) mixed olivine-Opx experiments at 230°C. Mass loss between 250 and 450°C is assigned to brucite and that for 500–750°C is assigned to serpentine (chrysotile). (Online version in colour.)

brucite) slightly improved the fit, but only accounts for 8% of the total spectral area. The FeII doublets for brucite and chrysotile overlap making it difficult to resolve these two phases in the MB spectra, particularly when brucite is subordinate. However, because brucite was identified in this sample by other methods, fitting with two ferrous doublets was justified. Note that the MB analyses for experiments in this study were performed on secondary products that had been physically separated from remnant reactant minerals by sonication and suspension in ethanol, so there is no doublet for remnant olivine (see electronic supplementary material).

Thermogravimetry of Oliv230pH resulted in mass losses from the decomposition of chrysotile and brucite (figure 3). About 8.1 wt% mass loss is attributable to dehydration of chrysotile while 2.3 wt% is attributable to brucite, which translates to a molar chrysotile:brucite ratio of
approximately 1.8 (table 5). Based on the TGA results, approximately 70 wt% of the olivine had reacted by the time the experiment was terminated (table 5). This is a substantially greater amount of reaction progress than was observed for Oliv230fine (approx. 25 wt% olivine reacted) despite the considerably longer reaction time for that experiment (4993 h for Oliv230fine versus 2972 h for Oliv230pH).

(b) Reaction of olivine at 200°C

Experiment Oliv200pH was conducted to evaluate the reaction of olivine in the presence of strongly alkaline pH at 200°C (table 1). Initial conditions for this experiment were designed to parallel those of Oliv200fine from McCollom et al. [13] except for the addition of NaOH to raise the initial pH and inclusion of a smaller amount of DIC. Measured concentrations of selected dissolved components in the fluid during this experiment are shown in figure 4, with measurements of the same components from Oliv200fine shown for comparison.

The pH25°C in Oliv200pH was initially 12.4 and remained at this level for around 1300 h before gradually decreasing to a final value of 11.3 (figure 4b). The calculated pHin situ was 9.6 at the first measurement and decreased over time to a final value of 8.5. Generation of H2 in Oliv200pH began soon after heating and the amount produced had risen to 2.3 µmol (g olivine)-1 by the time of the second measurement at 502 h (figure 4a). However, the H2 abundance in the fluid then decreased slightly over the next approximately 2000 h before increasing once again to reach a final amount of 20.9 µmol (g olivine)-1 at 7677 h when the experiment was terminated (figure 4a). For comparison, the measured pH25°C in Oliv200fine at the first measurement was 6.4, but then increased to 8.8 by the second sample at 2018 h and remained near this value for the remainder of the experiment (figure 4b). The corresponding pHin situ was initially 7.1 but then rose to level off at values around 8.3, similar to the pHin situ in the latter stages of Oliv200pH. The amount of H2 generated in Oliv200fine increased steadily over time to a final value of 5.9 µmol (g olivine)-1 at 9219 h, less than 30% of the amount produced in Oliv200pH (figure 4a).

The dissolved silica concentration was relatively high (430 µmol kg−1) at the beginning of Oliv200pH, but decreased to 200 µmol kg−1 in the first 500 h and then continued to decrease steadily as the experiment progressed to attain a final concentration of 60 µmol kg−1 (figure 4c).

Table 5. Results of thermogravimetric analyses for experiments from this study and for comparable experiments at lower pH from previous studies.

|                      | Oliv230pH | Oliv230fine | Oliv200pH | Oliv200fine | OlivOpx230pH | OlivOpx230med |
|----------------------|-----------|-------------|-----------|-------------|--------------|--------------|
| TGA weight loss (wt%)|           |             |           |             |              |              |
| serpentine           | 8.1       | 3.0         | 4.7       | 0.75        | 2.3          | 0.27         |
| brucite              | 2.3       | 0.9         | 1.9       | 0.15        | 0            | 0            |
| secondary minerals (wt%)|     |             |           |             |              |              |
| serpentine           | 64.0      | 24.0        | 37.0      | 6.0         | 18.6         | 2.2          |
| brucite              | 8.0       | 3.1         | 6.7       | 0.51        | 0            | 0            |
| magnetite            | 1.6       | 1.75        | 1.0       | 0.25        | 0.02         | 0            |
| extent of reaction (%)d | 70       | 25          | 41        | 5.9         | 19           | 1.9          |
| serpentine : brucite | 1.76      | 1.65        | 1.24      | 2.5         | —            | —            |

aMcCollom et al. [13].
bMcCollom et al. [14].
cFrom magnetization measurements.
dEstimated percent of total primary minerals reacted, by weight.
Concentrations of $\Sigma$SiO$_2$ during the latter stages of Oliv200pH converged on those in Oliv200fine (10–30 µmol kg$^{-1}$), but remained slightly higher. Measured concentrations of K in Olvi200pH ranged from 500 to 1000 µmol kg$^{-1}$ during the experiments, while concentrations of other cations (Mg, Ca, Fe, Al) remained at very low levels (70 µmol kg$^{-1}$ or less; electronic supplementary material, table S1). Similar cation abundances were observed in Oliv200fine.
Figure 5. Fluid compositions during olivine experiment OlivOpx230pH (circles), with results for OlivOpx230med (triangles) from McCollom et al. [14] shown for comparison. The vertical dashed lines denote the timepoint of pH increase induced by the injection of a strongly alkaline solution. (a) Total amount of H₂ generated per gram mineral reactant. (b) Measured room temperature pH (pH₂5°C) and calculated in situ pH (pH løs). (c) Total dissolved SiO₂ concentration. (Online version in colour.)

Chrysotile and brucite were the predominant secondary mineral products of Oliv200pH, along with magnetite and trace calcite (figure 2; electronic supplementary material, figure S1). A substantial fraction of the original olivine also remained at the end of the experiment. The morphology of chrysotile and brucite were very similar to the products of Oliv230pH (figure 2d–f). The chrysotile in Oliv200pH had the same chemical composition (Mg# = 95) as in Oliv200fine, but brucite had higher Fe contents in Oliv200pH (Mg# = 84) than Oliv200fine (Mg# = 91; table 3). As in other similar experiments, the brucite in Oliv200pH contained substantial amounts of Cl indicating it was intergrown with iowaite [13,14]. Magnetization measurements indicated that 1.0 wt% magnetite was produced during Oliv200pH, substantially more than in Oliv200fine (0.25 wt%).

Mössbauer analysis of the products from Oliv200pH identified three main doublets, with one peak assigned to FeII and two peaks to FeIII (table 4). The hyperfine parameters for all three peaks are consistent with chrysotile, with the occupation of both octahedral and tetrahedral sites by the FeIII. Similar to the Oliv230pH results, adding another FeII doublet assigned to brucite (12 mol% of total Fe) slightly improved the final fit. Approximately equal amounts of FeII and
Fe$^{III}$ are present in the chrysotile, with about two-thirds of the Fe$^{III}$ occupying the octahedral site. Thermogravimetry of Oliv200pH showed significant weight loss attributable to dehydration of both chrysotile (4.7 wt%) and brucite (1.9 wt%) corresponding to a chrysotile:brucite mass ratio of 1.2, with substantially greater amounts of both products than in Oliv200fine (figure 3; table 3). The proportion of olivine reacted in Oliv200pH was about 41 wt%, much greater than in Oliv200fine (5.9 wt%).

(c) Reaction of olivine–Opx mixture at 230°C

Experiment OlivOpx230pH was performed with a mixture of olivine and Opx as reactants at 230°C, and involved increasing the pH during the experiment from circumneutral to strongly alkaline conditions (table 1). The reactants for this experiment were designed to parallel those used in a previous series of experiments with olivine–Opx mixtures performed under the same conditions but without externally adjusting the pH [14]. Fluid compositions during the experiment are shown in figure 5, along with the fluid composition from a previous experiment using essentially the same initial ingredients but performed for a somewhat shorter period of time at circumneutral pH (OlivOpx230med; table 1).

The initial reactant fluid for OlivOpx230pH had a circumneutral pH, and the measured pH$_{25°C}$ in the experiment after 52 h of heating was 7.4 (figure 5b). A first attempt to increase the pH was made after 1530 h, when a fluid with pH$_{25°C}$ = 10.6 was injected into the reaction cell (electronic supplementary material, table S1). However, analysis of the fluid approximately 200 h after the injection indicated that the pH$_{25°C}$ had risen only slightly to 7.6, indicating the pH of the fluid had been buffered by reaction with the solids in the reaction cell. The experiment was then allowed to continue to react at circumneutral pH for another 2800 h. Throughout this initial stage of the experiment, the pH$_{25°C}$ of OlivOpx230pH remained at levels very similar to OlivOpx230med (figure 5b). A second attempt to increase pH was made at 4526 h, when a fluid with pH$_{25°C}$ = 12.3 was injected (electronic supplementary material, table S1). This injection successfully raised the pH$_{25°C}$ of the fluid in the reaction cell to 10.6, and it remained above 10 for the remainder of the experiment (figure 5b).

Very little production of H$_2$ was observed in OlivOpx230pH during the first 4503 h of the experiment prior to the increase of pH, although there was a very slight increase in the rate of H$_2$ accumulation following the first injection at 1530 h (figure 5a). After the increase in pH, however, there was a steep increase in H$_2$ production that continued at a steady rate for the remainder of the experiment (figure 5a). By the end of the experiment, 13.5 µmol of H$_2$ had been generated per gram of reactant minerals (figure 5a). Virtually no H$_2$ production was observed in the previous OlivOpx230med experiment at a lower pH.

The $\Sigma$SiO$_2$ concentration in OlivOpx230pH remained fairly steady at around 400 µmol kg$^{-1}$ up until the time of the pH increase, with concentrations slightly lower than those observed in OlivOpx230med (figure 5c). Shortly after the pH increase, the $\Sigma$SiO$_2$ concentration in OlivOpx230pH began to steadily decrease and reached 100 µmol kg$^{-1}$ at the termination of the experiment. Dissolved Ca, Mg and Al were observed in the early stages of OlivOpx230pH at concentrations of 50–100 µmol kg$^{-1}$, but only Ca persisted at levels above the detection limit after the pH increase (electronic supplementary material, table S1). Dissolved K was present at around 200 µmol kg$^{-1}$ before the pH increase but fell below detection afterwards, while Fe was below detection throughout the experiment. Concentrations of Na and Cl remained constant within analytical errors throughout the experiment. Calcium was the only cation detected other than Na in OlivOpx230med, with a final concentration of 60–100 µmol kg$^{-1}$.

The secondary reaction products of OlivOpx230pH were dominated by chrysotile (figure 2g–i). The chrysotile fibres formed thin mats that covered the surfaces of olivine and Opx crystals, which curled up into the void spaces between the reactant minerals (figure 2i). The solid reaction products also included trace amounts of magnetite and calcite. Both relict olivine and Opx remained at the end of the experiment; however, analysis of the reacted solids by XRD (electronic supplementary material, figure S1) showed that the olivine:Opx ratio was far
higher in the products than in the unaltered reactants, and examination of the products by SEM detected very few remnants of unreacted Opx. Therefore, Opx evidently reacted more extensively than olivine during the experiment. In the experiment OlivOpx230med at lower pH, only a very limited amount of secondary products was observed, forming a very thin rind on the surfaces of olivine and Opx crystals [14]. Alteration products in the rind from OlivOPx230med were poorly crystalline, but were composed of a mixture of serpentine (most likely lizardite) and minor amounts of talc.

The chrysotile that formed in OlivOpx230pH had an Mg\# = 87, intermediate between the Mg\#s of the original olivine and Opx (table 2). Mössbauer analysis indicates that approximately 29 mol% of the total Fe in the serpentine was present as Fe\textsuperscript{III}, with nearly equal amounts occurring in octahedral and tetrahedral coordination (table 4). The reacted solids included only 0.02 wt% magnetite. Since this is about the same amount that would have been present in the reactants from Opx, it is not clear if any of this magnetite was produced during the experiment. Weight loss during TGA was entirely attributable to dehydration of serpentine (2.3 wt%), corresponding to reaction of about 19 wt% of the original mineral reactants (figure 4; table 3). Only 1.9 wt% of the original minerals had reacted in OlivOpx230med, which lasted only approximately 35% as long as OlivOpx230pH.

4. Discussion

(a) Effect of pH on reaction of olivine

Both of the olivine serpentinization experiments performed for this study show evidence for steep increases in overall reaction rates and the rate of H\textsubscript{2} production at higher pH. At 230°C, an increase in pH\textsubscript{in situ} of approximately 1.2 units between Oliv230pH and Oliv230fine resulted in a nearly threefold increase in the extent of olivine reaction and greater than twofold increase in the amount of H\textsubscript{2} generated, despite the significantly shorter reaction time for Oliv230pH (table 3). At 200°C, the pH\textsubscript{in situ} of Oliv200pH and Oliv200fine converged as the reactions progressed, yet the extent of olivine reaction was still about seven times greater in Oliv200pH and more than three times as much H\textsubscript{2} was generated.

To a large extent, the greater rate of H\textsubscript{2} production in the higher pH experiments can be attributed to the increased overall rate of olivine reaction. However, in both of the high pH olivine experiments, the increase in the amount of H\textsubscript{2} generated is less than the increase in the extent of reaction (table 3). Since generation of H\textsubscript{2} during serpentinization is linked to the conversion (oxidation) of ferrous Fe (Fe\textsuperscript{II}) from the primary minerals to ferric Fe (Fe\textsuperscript{III}) in the secondary mineral products [36,43], the lower H\textsubscript{2} yields as a function of reaction progress in the higher pH experiments must reflect changes in the distribution of Fe among the solid products.

Analyses of mineral compositions show significant differences in how Fe is distributed among the reaction products in the high pH experiments relative to their lower pH counterparts (table 3). Despite a much greater overall extent of reaction, less magnetite was produced during Oliv230pH (1.6 wt%) than in Oliv230fine (1.75 wt%). While more magnetite was produced during Oliv200pH (1.0 wt%) than in Oliv200fine (0.25 wt%), the amount produced relative to the proportion of olivine reacted was substantially less in Oliv200pH. The Fe content of brucite was significantly greater in both of the high pH experiments relative to their lower pH counterparts, and the Fe content of chrysotile was also greater in Oliv230pH (table 3). Furthermore, the Fe\textsuperscript{III}:Fe\textsubscript{total} ratio in chrysotile was substantially higher in Oliv230pH and Oliv200pH (0.53 and 0.51, respectively) than in lower pH olivine experiments, where the ratio for chrysotile is consistently in the 0.14–0.24 range (table 4; [13]).

These observations indicate that the higher pH experiments resulted in more Fe being partitioned into brucite and chrysotile and less into magnetite from each increment of olivine reacted. At the same time, a greater proportion of Fe\textsuperscript{III} was partitioned into chrysotile rather than into magnetite. The ultimate outcome was that each increment of olivine that underwent
serpentinization generated less H$_2$ in the higher pH experiments, with proportionally less of the 
H$_2$ derived from Fe$^{III}$ partitioned into magnetite and more from Fe$^{III}$ precipitated as chrysotile.

Further study is required to determine the mechanisms that caused the changes in the 
partitioning of Fe among the reaction products at higher pH. One possibility, however, is that 
higher pH may alter the speciation of dissolved components in ways that effect the relative 
rates of mineral precipitation and uptake of Fe species. For instance, higher $\Sigma$SiO$_2$ and OH$^-$ 
concentrations may have allowed more rapid precipitation and uptake of Fe by chrysotile and 
brucite, and slowed precipitation of magnetite. Another possibility is that the higher pH altered 
the surface charge of precipitating minerals and allowed Fe to be more efficiently taken up by 
chrysotile and brucite. Additionally, at higher pH, a greater proportion of the dissolved Fe would 
be present as hydroxide complexes such as FeOH$^+$ (e.g. \[26\]), and such positively charged species 
may more readily interact with the surfaces of precipitating brucite and lead to more rapid uptake.

Other investigators have suggested that the greater proportion of FeOH$^+$ at higher pH may 
also lead to increased H$_2$ generation and magnetite precipitation owing to faster rates of oxidation 
of FeOH$^+$ relative to other dissolved Fe species \[26,44\]. For instance, Syverson et al. \[26\] observed 
a substantial increase in the rate of H$_2$ generation following an increase in pH during the reaction 
of an olivine–talc mixture at 300°C, and attributed the higher rate of H$_2$ generation to a faster rate 
of Fe oxidation and magnetite precipitation following the change in pH. However, the results of 
the present study suggest that much of the increase in H$_2$ generation in their experiment might 
instead be attributable to an increase in the overall rate of serpentinization of olivine following 
the increase in pH rather than to a change in Fe speciation (as in this study, Syverson et al. only 
examined the mineral products at the termination of the experiments, so the possibility of changes 
in reaction rates during their experiment cannot be directly evaluated). Furthermore, rather than 
enhancing magnetite precipitation as inferred by Syverson et al. \[26\], the results of the present 
study suggest that higher pH may actually favour precipitation of chrysotile and brucite relative 
to magnetite. Nevertheless, it is possible that changes in Fe speciation may have affected to some 
degree the rate of oxidation and resulting distribution of Fe$^{III}$ between magnetite and chrysotile 
in the present set of experiments.

The distribution of Fe among product minerals may also have been influenced by 
thermodynamic factors. For example, thermodynamic equilibrium favours greater partitioning 
of Fe into brucite in coexistence with magnetite as H$_2$ activities increase, as reflected in the 
generalized reaction

\[
3[\text{Fe(OH)}_2]_{\text{brucite}} \leftrightarrow \text{Fe}_3\text{O}_4[\text{magnetite}] + 2\text{H}_2\text{O} + \text{H}_2, \quad (4.1)
\]

where $[\text{Fe(OH)}_2]_{\text{brucite}}$ represents the Fe component of brucite. As the serpentinization reaction 
progresses and H$_2$ accumulates in solution, equilibration of this reaction should, therefore, 
favour higher Fe contents in brucite and limit the amount of Fe available to precipitate as 
magnetite. This may help explain, in part, the greater Fe contents of brucite and relatively smaller 
amounts of magnetite in the higher pH experiments, since these experiments accumulated higher 
concentrations of H$_2$ (\textbf{table 3}). However, Rxn. 1 would also predict that magnetite production 
should decrease with increasing reaction progress as H$_2$ accumulates, but other experimental 
studies of olivine serpentinization have found that the amount of magnetite produced increased 
at a steady rate as the reactions progressed \[12,13\]. Consequently, the role of thermodynamic 
factors in regulating the distribution of Fe between magnetite and brucite remains uncertain.

Marcaillou et al. \[45\] observed that the Fe$^{III}$:Fe$_{\text{total}}$ ratio of serpentine (lizardite) increased 
steadily with increasing reaction progress during the serpentinization of lherzolite at 300°C. The 
greater partitioning of Fe$^{III}$ into serpentine with increasing extent of reaction in their experiment 
was unrelated to alkaline pH, since the fluid was mildly acidic throughout the reaction (pH$_{25^\circ\text{C}}$ 
5.1–5.8). Nevertheless, the olivine experiments in the present study appear to display a similar 
correspondence between higher Fe$^{III}$:Fe$_{\text{total}}$ ratio with a greater extent of reaction, suggesting 
that increased partitioning of Fe$^{III}$ into serpentine with increasing reaction progress may occur 
over a broad range of conditions.
(b) Effect of pH on reaction of olivine–pyroxene mixtures

The increase in pH during the experiment OlivOpx230pH also led to substantially increased rates of reaction and H₂ generation. The OlivOpx230med experiment provides some insight into the state of OlivOpx230pH prior to the pH increase. After 2709 h of heating at 230°C, the reaction of the olivine–Opx mixture in OlivOpx230med resulted in only a very limited extent of reaction (approx. 1.9 wt% of reactants) and virtually no H₂ production [14] (table 3). Reaction products in that experiment were limited to a thin alteration rind only a few micrometre thick deposited on the surfaces of the original reactant minerals and composed of an intimate mixture of poorly crystalline lizardite and talc. Similar reaction products were likely present in OlivOpx230pH at the time that the fluid was injected to increase pH at 4526 h, although the extent of reaction may have been somewhat greater owing to the longer reaction time.

Comparison with OlivOpx230med suggests that about 4 wt% of the primary minerals in OlivOpx230pH had reacted prior to raising the pH while the extent of reaction had increased to 19 wt% by the end of the experiment at 7715 h, indicating a greater than fivefold increase in overall reaction rate following the pH increase. In both OlivOpx230pH and OlivOpx230med, the olivine : Opx ratio of the reactants that remained at the end of the experiments was much higher than in the starting materials (electronic supplementary material, figure S1), indicating reaction of Opx proceeded more rapidly than olivine. At present, however, there is not sufficient data to determine whether the increase in pH accelerated the reaction of both minerals equally or not.

The results of OlivOpx230pH can also be compared with those of OlivOpx230 reported by McCollom et al. [14]. The composition of reactant minerals and experimental conditions for OlivOpx230 were very similar to both OlivOpx230pH and OlivOpx230med (table 1), and OlivOpx230 was allowed to react for a period comparable to OlivOpx230pH. Unlike the other experiments, however, OlivOpx230 spontaneously transitioned from circumneutral to strongly alkaline conditions after approximately 4000 h of reaction without any external intervention (figure 6b). The reason why the pH spontaneously increased during experiment OlivOpx230 but not in the other experiments is undetermined at this time, but it appears to be related to a small amount of clinopyroxene present as a contaminant in the starting minerals for that experiment [14].

As in the experiment OlivOpx230pH, the increase in pH in OlivOpx230 was accompanied by a steep increase in H₂ production (figure 6a). The overall extent of reaction and rate of H₂ generation was significantly greater in OlivOpx230 than in OlivOpx230pH, but are comparable to the extent of reaction and H₂ generation observed in the olivine-only experiment Oliv230pH at high pH (table 3). Reaction products of OlivOpx230 included substantial amounts of brucite and magnetite in addition to chrysotile, which likely reflects the domination of the products by the dissolution of olivine during the latter stages of the experiment [14]. No Opx remained at the end of the experiment OlivOpx230.

The spontaneous increase in pH in experiment OlivOpx230 clearly caused similar increases in rates of overall reaction and H₂ generation to those observed following the induced increase in pH during OlivOpx230pH (figure 6a; table 3). More rapid reaction rates were attained in OlivOpx230 than OlivOpx230pH, which might be attributable to somewhat more strongly alkaline conditions in the former experiment (figure 6b) or to lower concentrations of ΣSiO₂ (figure 6c). Experiment OlivOpx230 also apparently transitioned to a state where olivine dominated the overall reaction leading to precipitation of brucite and magnetite in addition to chrysotile [14], which may have contributed to greater rates of H₂ generation.

(c) Possible mechanism for increased rates

The observed increased rates of serpentinization and H₂ generation at higher pH contrast with laboratory studies of olivine and pyroxene dissolution where rates have been found to decrease monotonically with increasing pH, at least up to 150°C and pH 12 [29–32]. Moreover, dissolution rates of olivine and pyroxene observed in laboratory experiments are much faster than the...
Figure 6. Comparison of (a) $H_2$ production, (b) room temperature pH and (c) total dissolved SiO$_2$ concentration during experiments with olivine–Opx mixtures at 230°C. The dashed vertical line denotes the timing of the injection of alkaline fluids to increase pH in the experiment OlivOpx230pH. Data for OlivOpx230 and OlivOpx230med from McCollom et al. [14]. (Online version in colour.)

rates of serpentinization observed in the present study. For example, using the equation for the dissolution of olivine as a function of pH and temperature from Rimstidt et al. [31] based on laboratory experiments (their eqn (6)), only about 3 days should have been required to dissolve 90 wt% of the original olivine in Oliv200pH and less than 8 h for Oliv230pH, far shorter than the duration of either experiment (see electronic supplemental material for calculations). Although the rate equation of Rimstidt et al. is only calibrated to 150°C and ambient pressure, there is no reason to believe that rates should be slower at the higher temperatures and pressures of the present serpentinization experiments. Extrapolation of enstatite dissolution rates from laboratory studies [31] to the temperature of OlivOpx230pH would also predict that Opx should have completely dissolved on a much shorter timescale than the experiment. Evidently, feedbacks from the solution prevented the olivine and Opx in the serpentinization experiments from dissolving at the rates observed in the far-from-equilibrium laboratory dissolution studies.

While the reaction rates of olivine in the serpentinization experiments are substantially slower than dissolution studies would imply, the mechanism of dissolution determined from those studies may provide some insight into the cause for the increased rates of serpentinization at higher pH relative to those observed at circumneutral conditions. During the steady-state dissolution of olivine at acidic to slightly alkaline pH, a very thin layer (~a few unit-cells thick) forms at the olivine surface that is enriched in Si and depleted in Mg [29,30]. At pH
greater than approximately 9, however, the dissolution mechanism shifts, and a Si-depleted, Mg-enriched surface layer develops. The structure of this thin Mg-rich layer apparently resembles the individual sheets of brucite [29]. The rate-limiting step for dissolution at higher pH is thought to involve hydrolysis and detachment of Mg ions from this Si-depleted surface layer [29,30].

A similar Mg-enriched, Si-depleted layer may develop on the surface of olivine during the reaction at the higher temperatures of the serpentinization experiments. Higher pH may then increase the rate of the detachment of Mg from the surface of the Si-depleted layer, leading to faster overall reaction rates of the olivine. When brucite is present, it may enhance the dissolution rate at higher pH by lowering the concentration of Mg and continuously removing it from the solution. That is, for precipitation of brucite from solution according to the reaction (written for the pure Mg endmember)

\[
Mg^{2+} + 2OH^- \leftrightarrow Mg(OH)_2 \text{ [brucite]},
\]

the increased concentration of OH\(^-\) at higher pH results in a decreased concentration of Mg in the fluid in equilibrium with brucite. Although Mg\(^{2+}\) and OH\(^-\) would be rapidly removed from solution by brucite precipitation, they would be continuously replenished by olivine dissolution

\[
Mg_2SiO_4 \text{ [olivine]} + 2H_2O \rightarrow SiO_2(aq) + 2Mg^{2+} + 4OH^-. \]

Notably, the Si-depleted layer forms on dissolving olivine even when the solution is undersaturated with respect to brucite [29,30], so this layer may also be present at the olivine surface in the mixed olivine–Opx experiments at higher pH [14].

The concentrations of ΣSiO\(_2\) in the fluids provides support for the formation of an Mg-rich layer on the olivine surface during the experiments. Figure 7a shows the calculated activities of dissolved silica \([a_{SiO_2(aq)}]\) and pH\(_{\text{in situ}}\) during the 230°C experiments projected onto an equilibrium mineral stability diagram for the experimental conditions. It can be seen from the figure that the fluid compositions for the olivine-only experiments plot within the chrysotile stability field and well above the chrysotile–brucite boundary, despite the presence of abundant

**Figure 7.** Calculated *in situ* fluid compositions for laboratory experiments projected onto mineral stability diagrams for Mg-bearing minerals in the Mg–Fe–Si–O–C–H system as a function of pH and activity of dissolved silica. (a) shows stable equilibrium mineral assemblages at 230°C, while (b) shows metastable assemblages calculated by excluding equilibration with chrysotile. Numbers in (a) identify sample numbers for OlivOpx230P and in (b) the first and last samples for OlivOpx230 are identified. Compositions of chrysotile and brucite are constrained to approximate those from the experiments. For the construction of the diagrams, the activity of dissolved Mg was set to \(10^{-5}\), H\(_2\) to \(10^{-2}\) and HCO\(_3^-\) to \(10^{-2.5}\), while dissolved Fe was assumed to be in equilibrium with magnetite. Dashed vertical lines separate predominance fields for CO\(_2\) and HCO\(_3^-\) for dissolved carbon species. (Online version in colour.)
brucite in both experiments. Under these conditions, brucite should react with dissolved SiO$_2$ to form chrysotile, lowering the $a_{\text{SiO}_2(aq)}$ to the chrysotile–brucite equilibrium boundary. The persistence of $a_{\text{SiO}_2(aq)}$ well above this level probably indicates kinetic inhibition of the reaction between brucite and dissolved SiO$_2$ [23], precluding equilibrium between brucite and chrysotile.

Instead, it appears that the activity of dissolved silica in the experiments may be controlled by steady-state reactions near the brucite–olivine equilibrium boundary. Figure 7b shows a mineral stability diagram where equilibrium with chrysotile has been suppressed. As seen in the diagram, the fluid compositions during the olivine experiments converge on the olivine–brucite boundary, suggesting that $a_{\text{SiO}_2(aq)}$ levels are being regulated to values near equilibrium between olivine and brucite (or a brucite-like phase). In this case, this would represent a metastable equilibrium, since olivine is thermodynamically unstable and would be replaced by chrysotile at full equilibrium. Prior to the increase in pH in the experiment OlivOpx230pH, the $a_{\text{SiO}_2(aq)}$ was apparently buffered at relatively high levels by chrysotile–talc equilibrium (figure 7a).

However, following the pH increase the $a_{\text{SiO}_2(aq)}$ in this experiment also converged on the olivine–brucite equilibrium boundary towards the end of the experiment (figure 7b). Silica activities consistent with metastable olivine–brucite equilibrium have also been observed in several other serpentinization experiments [13,14], as illustrated by the results for Oliv230fine and OlivOpx230 shown in figure 7b.

It may be that the dissolved silica activities in these experiments are being regulated by steady-state dissolution and interactions between the Mg-enriched surface layer and the underlying olivine [14]. At the interface between the surface layer and olivine, the exchange of SiO$_2$ with the fluid may be rapid and maintain a local metastable equilibrium between the mineral surface and dissolved silica in the fluid. Given the apparent brucite-like structure of the Mg-enriched layer, the resulting metastable equilibrium between the surface layer and underlying olivine would be expected to occur at a silica activity close to brucite–olivine equilibrium.

It should be noted that although the silica activities converge on similar values in these experiments, that does not necessarily mean that the experiments will have similar concentrations of total dissolved silica since the speciation varies substantially as a function of pH. As shown in electronic supplementary material, figure S4, at pH$_{\text{in situ}}$ less than 8.4 the speciation of dissolved silica is dominated by SiO$_2(aq)$, but as the pH$_{\text{in situ}}$ rises above 8.4 the speciation is increasingly dominated by HSiO$_3^-$ and possibly NaHSiO$_3(aq)$, although the stability of the latter under hydrothermal conditions is currently uncertain [46]. As a consequence, at constant $a_{\text{SiO}_2(aq)}$ the total dissolved silica concentration (i.e. $\Sigma\text{SiO}_2$) increases steeply with increasing pH$_{\text{in situ}}$ above 8.4 (electronic supplementary material, figure S4). This may help explain why the $\Sigma\text{SiO}_2$ concentrations in the higher pH experiments are substantially higher than those in the lower pH experiments (figures 1c and 4c).

The elevated $\Sigma\text{SiO}_2$ concentrations in the higher pH experiments may also contribute to the more rapid reaction rates observed for those experiments. The elevated $\Sigma\text{SiO}_2$ concentrations may promote more rapid precipitation of chrysotile and therefore the dissolution of olivine, leading to more rapid overall reaction relative to less alkaline pH conditions where $\Sigma\text{SiO}_2$ concentrations are lower. In addition, the steep increase in $\Sigma\text{SiO}_2$ concentrations at increasingly alkaline conditions may help regulate the pH in the experiments, since raising the pH further would require increasingly higher concentrations of $\Sigma\text{SiO}_2$ that would be counteracted by precipitation of chrysotile.

The higher pH might also induce an increase in the precipitation rate of brucite, and thereby lead to an increase in overall reaction rate. Specifically, removal of Mg and Fe from solution owing to faster precipitation of brucite may cause these elements to be removed at a greater rate from the surface of olivine, leading to an increase in olivine dissolution rate. A detachment of Mg ions from the surface is thought to be the rate-limiting step for the dissolution of olivine at alkaline pH in laboratory dissolution studies [29,30], and the same may hold true during serpentinization.

Testing of each of these possibilities will require additional experimental studies that continue to probe the effect of different fluid parameters on rates of individual reactions taking place during serpentinization (e.g. [23]). However, regardless of the underlying cause, the experiments
in this study indicate that the strongly alkaline pH that develops in many serpentinizing systems will result in a steep increase in overall reaction rates. Because this outcome is contrary to expectations based on lower temperature dissolution studies, further study and elucidation of the underlying mechanisms is warranted in order to better understand serpentinization in natural systems.

(d) Implications for natural systems

The results of this study suggest that higher pH fluids circulating through ultramafic rocks are likely to induce more rapid serpentinization and higher H₂ fluxes than lower pH fluids. In subseafloor settings, for example, this may result in a greater extent of serpentinization of the rocks and higher concentrations of H₂ discharged at the seafloor in hydrothermal fluids. As seawater begins to circulate through subseafloor hydrothermal systems, it has a slightly alkaline pH (≈8) and any interaction of the fluid with ultramafic rocks may initially result in modest rates of reaction and H₂ generation. However, as the circulating fluid evolves through water–rock interactions and transitions to strongly alkaline pH (as in OlivOpx230; figure 6), rates of reaction and H₂ generation would be expected to accelerate. Other factors such as the rate of fluid circulation, bulk rock composition and temperature will undoubtedly influence the amount of H₂ that is generated as well, but the experimental results indicate that the evolving pH along the flow path of circulating fluids may be a critical factor in regulating the flux of H₂ discharged from actively serpentinizing systems.

The experimental results also indicate that pH affects Fe partitioning among the mineral products during serpentinization. Serpentinized rocks recovered from the ocean floor and from ophiolites exhibit substantial variation in the Fe contents of chrysotile and brucite and in the amount of magnetite present (e.g. [47–55]), but the underlying factors that regulate these variations remain incompletely understood. The results of this study indicate that pH needs to be considered along with other factors such as temperature, rock composition and silica activity in interpreting the distribution of Fe in serpentinized rocks. If the experiments from this study are representative, the results suggest that higher pH will lead to increased partitioning of Fe into serpentine and brucite rather than magnetite, as well as greater incorporation of FeIII in serpentine. However, further study is required to confirm these trends and more confidently determine the role of pH in the distribution of Fe during serpentinization.

Data accessibility. Supporting data for the figures are included in the electronic supplementary material. Authors’ contributions. T.M.M. conceived of and conducted the experiments, interpreted data and led the authorship of the manuscript. F.K. contributed to the interpretation of data, provided thermogravimetric analysis and helped write the manuscript. B.M. and P.S. provided Mössbauer and magnetic analyses, and contributed to the interpretation of data and writing of the manuscript. Competing interests. We declare we have no competing interests. Funding. This research was supported by the NASA Astrobiology Institute through Cooperative Agreement no. NNA15BB02A and NASA Solar Systems Workings program grant no. NNX16AL74G. The IRM is supported by the Instruments and Facilities Program of the NSF Division of Earth Science. Acknowledgements. Christopher Donaldson is thanked for his help with many of the experimental measurements. This is IRM contribution 1902. We thank Benjamin Tutolo, Andrew McCaig and an anonymous reviewer for helpful comments that resulted in significant improvements to this communication.

References

1. Barnes I, O’Neil JR. 1969 The relationship between fluids in some fresh Alpine-type ultramafics and possible modern serpentinization, western United States. Geol. Soc. Am. Bull. 80, 1947–1960. (doi:10.1130/0016-7606(1969)80[1947:TRBFIS]2.0.CO;2)
2. Barnes I, O’Neil JR, Trescases JJ. 1978 Present day serpentinization in New Caledonia, Oman, and Yugoslavia. Geochim. Cosmochim. Acta 42, 144–145. (doi:10.1016/0016-7037(78)90225-9)
3. Neal C, Stanger G. 1983 Hydrogen generation from mantle source rocks in Oman. Earth Planet. Sci. Lett. 66, 315–320. (doi:10.1016/0012-821X(83)90144-9)
4. Abrajano TA, Sturchio NC, Kennedy BM, Lyon GL, Muehlenbachs K, Bohlke JK. 1990 Geochemistry of reduced gas related to serpentinization of the Zambales ophiolite, Philippines. *Appl. Geochem.* **5**, 625–630. (doi:10.1016/0883-2927(90)90060-I)

5. Kelley DS et al. 2005 A serpentinite-hosted ecosystem: The Lost City hydrothermal field. *Science* **307**, 1428–1434. (doi:10.1126/science.1102556)

6. Morrill PL, Kuenen JG, Johnson OJ, Suzuki S, Rietze A, Sessions AL, Fogel ML, Nealon KH. 2013 Geochemistry and geobiology of a present-day serpentinization site in California: The Cedars. *Geochim. Cosmochim. Acta* **109**, 222–240. (doi:10.1016/j.gca.2013.01.043)

7. Schrenk MO, Brazelton WJ, Lang SQ. 2013 Serpentinization, carbon, and deep life. In *Carbon in Earth, reviews in mineralogy and geochemistry*, vol. **75** (eds RM Hazen, AP Jones, J Baross), pp. 575–606. Chantilly, VA: Mineralogical Society of America (doi:10.2138/rmg.2013.75.18)

8. McCollom TM, Seewald JS. 2013 Serpentinites, hydrogen, and life. *Elements* **9**, 129–134. (doi:10.2113/gselements.9.2.129)

9. Schulte M, Blake D, Hoehler T, McCollom T. 2006 Serpentinization and its implications for life on the early Earth and Mars. *Astrobiology* **6**, 364–376 (doi:10.1089/ast.2006.6.364)

10. Waite JH et al. 2017 Cassini finds molecular hydrogen in the Enceladus plume: evidence for hydrothermal processes. *Science* **356**, 155–159. (doi:10.1126/science.aai8703)

11. Vance SD, Hand KP, Pappalardo RT. 2016 Geophysical controls of chemical disequilibria in Europa. *Geophys. Res. Lett.* **43**, 4871–4879. (doi:10.1002/2016GL068547)

12. Malvoisin B, Brunet F, Carlut J, Roumejon S, Cannat M. 2012 Serpentinization of oceanic peridotites: 2. Kinetics and processes of San Carlos olivine hydrothermal alteration. *J. Geophys. Res.* **117**, B04102. (doi:10.1029/2011JB008842)

13. McCollom TM, Klein F, Robbins M, Moskowitz B, Berqué TS, Jöns N, Bach W, Templeton A. 2016 Temperature trends for reaction rates, hydrogen generation, and partitioning of iron during experimental serpentinization of olivine. *Geochim. Cosmochim. Acta* **181**, 175–200. (doi:10.1016/j.gca.2016.03.002)

14. McCollom TM, Klein F, Robbins M, Moskowitz B, Berqué TS, Bach W, Templeton A. Submitted. Hydrogen generation and iron partitioning during experimental serpentinization of olivine–pyroxene mixtures. *Geochim. Cosmochim. Acta.*

15. Ogasawara Y, Okamoto A, Hirano N, Tsuchiya N. 2013 Coupled reactions and silica diffusion during serpentinization. *Geochim. Cosmochim. Acta* **119**, 212–230. (doi:10.1016/j.gca.2013.06.001)

16. Okamoto A., Ogasawara Y., Ogawa Y., Tsuchiya N. 2011 Progress of hydration reactions in olivine-H2O and orthopyroxenite-H2O systems at 250°C and vapour-saturated pressure. *Chem. Geol.* **289**, 245–255. (doi:10.1016/j.chemgeo.2011.08.007)

17. Andreani M, Daniel I, Pollet-Villard M. 2013 Aluminum speeds up the hydrothermal alteration of olivine. *Am. Mineral.* **98**, 1738–1744. (doi:10.2138/am.2013.4469)

18. Pens M, Andreani M, Daniel I, Perrillat J-P, Cardon H. 2016 Contrasted effect of aluminum on the serpentinzation rate of olivine and orthopyroxene under hydrothermal conditions. *Chem. Geol.* **441**, 256–264. (doi:10.1016/j.chemgeo.2016.08.007)

19. Seyfried Jr WE, Foustoukos DI, Fu Q. 2007 Redox evolution and mass transfer during serpentinization: an experimental and theoretical study at 200°C, 500 bar with implications for ultramafic-hosted hydrothermal systems at Mid-Ocean Ridges. *Geochim. Cosmochim. Acta* **71**, 3872–3886. (doi:10.1016/j.gca.2007.05.015)

20. Lamadrid H. M., Rimstidt J. D., Schwarzenbach E. M., Klein F., Ulrich S., Dolocan A. and Bodnar R. J. 2017 Effect of water activity on rates of serpentinization of olivine. *Nat. Commun.* **8**, 16107. (doi:10.1038/ncomms16107)

21. Shibuya T, Yoshizaki M, Sato M, Shimizu K, Nakamura K, Omori S, Suzuki K, Takai K, Tsunakawa H, Maruyama S. 2015 Hydrogen-rich hydrothermal environments in the Hadean ocean inferred from serpentinization of komatiites at 300°C and 500 bar. *Prog. Earth Planet. Sci.* **2**, 46. (doi:10.1186/s40645-015-0076-z)

22. Huang R, Song M, Ding X, Zhu S, Zhan W, Sun W. 2012 Influence of pyroxene and spinel on the kinetics of peridotite serpentinization. *J. Geophys. Res.* **122**, 7111–7126. (doi:10.1002/2017JB014231)

23. Tutolo BM, Luhmann AJ, Tosca NJ, Seyfried Jr WE. 2018 Serpentinization as a reactive transport process: the brucite silicification reaction. *Earth Planet. Sci. Lett.* **484**, 385–395. (doi:10.1016/j.epsl.2017.12.029)
24. Lafay R, Montes-Hernandez G, Janots E, Chiriac R, Findling N, Toche F. 2012 Mineral replacement rate of olivine by chrysotile and brucite under high alkaline conditions. J. Cryst. Growth 347, 62–72. (doi:10.1016/j.jcrysgro.2012.02.040)

25. Grozeva NG, Klein F, Seewald JS, Sylva SP. 2017 Experimental study of carbonate formation in oceanic peridotite. Geochim. Cosmochim. Acta 199, 264–286. (doi:10.1016/j.gca.2016.10.052)

26. Syverson DD, Tutolo BM, Borrok DM, Seyfried Jr WE. 2017 Serpentinization of olivine at 300°C and 500 bars: an experimental study examining the role of silica on the reaction path and oxidation state of iron. Chem. Geol. 475, 122–134. (doi:10.1016/j.chemgeo.2017.11.006)

27. Martin B, Fyte WS. 1970. Some experimental and theoretical observations on the kinetic of hydration reactions with particular reference to serpentinization. Chem. Geol. 6, 185–202. (doi:10.1016/0009-2541(70)90018-5)

28. Wegner WW, Ernst WG. 1983 Experimentally determined hydration and dehydration reaction rates in the system MgO–SiO2–H2O. Am. J. Sci. 283–A, 151–180.

29. Pokrovsky OS, Schott J. 2000 Kinetics and mechanism of forsterite dissolution at 25°C and pH from 1 to 12. Geochim. Cosmochim. Acta 64, 3313–3325. (doi:10.1016/S0016-7037(00)00434-8)

30. Oelkers EH, Declerq J, Saldi GD, Gislason SR, Schott J. 2018 Olivine dissolution rates: a critical review. Chem. Geol. 500, 1–19. (doi:10.1016/j.chemgeo.2018.10.008)

31. Rimsdidt JD, Brantley SL, Olsen AA. 2012 Systematic review of forsterite dissolution rate data. Geochim. Cosmochim. Acta 99, 159–178. (doi:10.1016/j.gca.2012.09.019)

32. Oelkers EH, Schott J. 2001 An experimental study of enstatite dissolution rates as a function of pH, temperatures, and aqueous Mg and Si concentration, and the mechanism of pyroxene/pyroxenoid dissolution. Geochim. Cosmochim. Acta 65, 1219–1231. (doi:10.1016/S0016-7037(00)00564-0)

33. Seyfried Jr WE, Janecky DR, Berndt ME. 1987 Rocking autoclaves for hydrothermal experiments, II. The flexible reaction-cell system. In Hydrothermal experimental techniques (eds GC Ulmer, HL Barnes), pp. 216–239. New York, NY: John Wiley and Sons.

34. McCollom TM. 2016 Abiotic methane formation during experimental serpentinization of olivine. Proc. Natl Acad. Sci. USA 113, 13 965-13 970. (doi:10.1073/pnas.1611843113)

35. Closmann C, Knittle E, Bridges F. 1996 An XAFS study of the crystal chemistry of Fe in orthopyroxene. Am. Mineral. 81, 1321–1331. (doi:10.2138/am-1996-11-1203)

36. McCollom TM, Bach W. 2009 Thermodynamic constraints on hydrogen generation during serpentinization of ultramafic rocks. Geochim. Cosmochim. Acta 73, 856–875. (doi:10.1016/j.gca.2008.10.032)

37. Helgeson HC, Delany JM, Nesbitt HW, Bird DK. 1978 Summary and critique of the thermodynamic properties of rock-forming minerals. Am. J. Sci. 278-A, 1–229.

38. Shock EL, Helgeson HC. 1988 Calculation of the thermodynamic and transport properties of aqueous species at high pressure and temperatures: Correlation algorithms for ionic species and equation of state predictions to 5 kb and 1000°C. Geochim. Cosmochim. Acta 52, 2009–2036. (doi:10.1016/0016-7037(88)90181-0)

39. Shock EL, Helgeson HC, Sverjensky DA. 1989 Calculation of the thermodynamic and transport properties of aqueous species at high pressures and temperatures: Standard partial molal properties of inorganic neutral species. Geochim. Cosmochim. Acta 53, 2157–2183. (doi:10.1016/0016-7037(89)90341-4)

40. Shock EL, Sassani DC, Willis M, Sverjensky DA. 1997 Inorganic species in geologic fluids: correlations among standard molal thermodynamic properties of aqueous ions and hydroxide complexes. Geochim. Cosmochim. Acta 61, 907–950. (doi:10.1016/S0016-7037(96)00339-0)

41. Wolery TJ, Jarek RL. 2003 Software user’s manual: EQ3/6, version 8.0. Livermore, CA: Lawrence Livermore National Laboratory.

42. O’Hanley DS, Dyar MD. 1998 The composition of chrysotile and its relationship with lizardite. Can. Mineral. 36, 727–739.

43. Klein F, Bach W, McCollom TM. 2013 Compositional controls on hydrogen generation during serpentinization of ultramafic rocks. Lithos 178, 55–69. (doi:10.1016/j.lithos.2013.03.008)

44. Janecky DR, Seyfried Jr WE. 1986 Hydrothermal serpentinization of peridotite within the ocean crust: experimental investigation of mineralogy and major element chemistry. Geochim. Cosmochim. Acta 50, 1357–1378. (doi:10.1016/0016-7037(86)90311-X)

45. Marcaillou C, Muñoz M, Vidal O, Parra T, Harfouche M. 2011 Mineralogical evidence for H2 degassing during serpentinization at 300°C/300 bar. Earth Planet. Sci. Lett. 303, 281–290. (doi:10.1016/j.epsl.2011.01.006)
46. Tutolo BM, Tosca NJ. 2018 Experimental examination of the Mg-silicate-carbonate system at ambient temperature: implications for alkaline chemical sedimentation and lacustrine carbonate formation. *Geochim. Cosmochim. Acta* **225**, 80–101. (doi:10.1016/j.gca.2018.01.019)

47. Bach W, Paulick H, Garrido CJ, Ildefonse B. 2006 Unraveling the sequence of serpentinization reactions: petrography, mineral chemistry, and petrophysics of serpentinites from MAR 15°N (ODP Leg 209, Site 1274). *Geophys. Res. Lett.* **33**, L13306. (doi:10.1029/2006GL025681)

48. Klein F, Bach W, Humphris SE, Kahl W-A, Jöns N, Moskowitz B, Berquó TS. 2014 Magnetite in seafloor serpentinite – some like it hot. *Geology* **42**, 135–138. (doi:10.1130/G35068.1)

49. Klein F, Bach W, Jöns N, McCollom T, Moskowitz B, Berquó TS. 2009 Iron partitioning and hydrogen generation during serpentinization of abyssal peridotites from 15°N on the Mid-Atlantic Ridge. *Geochim. Cosmochim. Acta* **73**, 6868–6893. (doi:10.1016/j.gca.2009.08.021)

50. Toft PB, Arkani-Hamed J, Haggerty SE. 1990 The effects of serpentinization on density and magnetic susceptibility: a petrophysical model. *Phys. Earth Planet. Inter.* **65**, 137–157. (doi:10.1016/0031-9201(90)90082-9)

51. Oufi O, Cannat M, Horen H. 2002 Magnetic properties of variably serpentinized abyssal peridotite. *J. Geophys. Res.* **107**, B5. (doi.org/10.1029/2001JB000549)

52. Evans BW. 2008 Control of the products of serpentinization by the Fe$^{2+}$Mg$^{-1}$ exchange potential of olivine and orthopyroxene. *J. Petrol.* **49**, 1873–1887. (doi:10.1093/petrology/egn050)

53. Andreani M, Muñoz M, Marcaillou C, Delacour A. 2013 µXANES study of iron redox state in serpentine during oceanic serpentinization. *Lithos* **178**, 70–83. (doi:10.1016/j.lithos.2013.04.008)

54. Frost BR, Beard JS. 2007 On silica activity and serpentinization. *J. Petrol.* **48**, 1351–1368. (doi:10.1093/petrology/egm021)

55. Mayhew LE, Ellison ET. 2020 A synthesis and meta-analysis of the Fe chemistry of serpentinites and serpentine minerals. *Phil. Trans. R. Soc. A* **378**, 20180420. (doi:10.1098/rsta.2018.0420)