Serpentinization-Driven H₂ Production From Continental Break-Up to Mid-Ocean Ridge Spreading: Unexpected High Rates at the West Iberia Margin

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Molecular hydrogen (H₂) released during serpentinization of mantle rocks is one of the main fuels for chemosynthetic life. Processes of H₂ production at slow-spreading mid-ocean ridges (MORs) have received much attention in the past. Less well understood is serpentinization at passive continental margins where different rock types are involved (lherzolite instead of harzburgite/dunite at MORs) and the alteration temperatures tend to be lower (<200°C vs. >200°C). To help closing this knowledge gap we investigated drill core samples from the West Iberia margin. Lherzolitic compositions and spinel geochemistry indicate that the exhumed peridotites resemble sub-continental lithospheric mantle. The rocks are strongly serpentinized, mainly consist of serpentine with little magnetite, and are generally brucite-free. Serpentine can be uncommonly Fe-rich, with \[ X_{\text{Mg}} = \frac{\text{Mg}}{(\text{Mg} + \text{Fe})} < 0.8 \], and shows distinct compositional trends toward a cronstedtite endmember. Bulk rock and silicate fraction Fe(III)/\( \sum \text{Fe} \) ratios are 0.6–0.92 and 0.58–0.8, respectively; our data show that 2/3 of the ferric Fe is accounted for by Fe(III)-serpentine. Mass balance and thermodynamic calculations suggest that the sample’s initial serpentinization produced \( \sim 120 \) to \( >300 \) mmol H₂ per kg rock. The cold, late-stage weathering of the serpentinites at the seafloor caused additional H₂ formation. These results suggest that the H₂ generation potential evolves during the transition from continental break-up to ultraslow and, eventually, slow MOR spreading. Metamorphic phase assemblages systematically vary between these settings, which has consequences for H₂ yields during serpentinization. At magma-poor rifted margins and ultraslow-spreading MORs, serpentine hosts most Fe(III). Hydrogen yields of 120 to >300 mmol and 50–150 mmol H₂ per kg rock, respectively, may be expected at temperatures of <200°C. At slow-spreading MORs, in contrast, serpentinization may produce 200–350 mmol H₂, most of which is related to magnetite formation at >200°C. Since, in comparison to slow-spreading MORs, geothermal gradients at magma-poor margins...
and ultraslow-spreading MORs are lower, larger volumes of low-temperature serpentinite should form in these settings. Serpentinization of lherzolitic rocks at magma-poor margins should produce particularly high amounts of H₂ under conditions within the habitable zone. Magma-poor margins may hence be more relevant environments for hydrogenotrophic microbial life than previously thought.

Keywords: serpentinization, hydrogen generation, magma-poor rifted margin, mid-ocean ridges, ultraslow spreading, hydrothermal alteration, chemosynthetic life, seafloor weathering

INTRODUCTION

Serpentinites in the oceanic lithosphere form when mantle peridotite interacts with seawater at temperatures below \(-450°C\). A byproduct of this reaction is molecular hydrogen (H₂), which in turn provides a source of metabolic energy for lithoautotrophic microbial communities (e.g., Sleep et al., 2004; Edwards et al., 2005; Petersen et al., 2011). Serpentinization sites have hence repeatedly been proposed as locations favorable for life to have evolved on Earth and elsewhere (e.g., Schulte et al., 2006; McCollom and Seewald, 2013; Fryer et al., 2020). The understanding of serpentinization-related H₂ production has been of great interest over the last years.

Major sites of serpentinization—and thus of H₂ production—are slow and ultraslow-spreading mid-ocean ridges (MORs) and magma-poor rifted margins, where mantle peridotite is frequently exposed to the seafloor by tectonic processes (e.g., Brun and Beslier, 1996; Cannat et al., 2010; Sauter et al., 2013). Serpentinization reactions and the associated H₂ formation at MORs have been extensively investigated in the past (e.g., Bach et al., 2006; Klein et al., 2009; Andreani et al., 2013; Tutolo et al., 2020). The hydration of mantle peridotite by seawater leads to the oxidation of ferrous Fe in primary minerals, i.e., olivine and pyroxene, to Fe(III) in secondary phases, following the generalized reaction \(2 FeO + H₂O = Fe₂O₃ + H₂(aq)\) (e.g., Sleep et al., 2004). Major hosts of ferric Fe are magnetite, \(Fe₃⁺Fe²⁺O₄\), and the Fe³⁺-serpentine varieties hisingerite, \(Fe³⁺SiO₃(OH)₄\), and cronstedite (\(MgFe²⁺Fe³⁺O₅SiO₃(OH)₄\)). Magnetite is most abundant at temperatures of \(200–350°C\), causing the production of \(>300 \text{ mmol H}_2\) per kg of dunite or harzburgite at these temperatures, but its abundance decreases toward lower temperatures at which Fe is progressively incorporated into Fe-rich brucite (Bach et al., 2006; Klein et al., 2009). At such lower temperatures, Fe(III)-serpentine is the main host for ferric Fe and the total amounts of H₂ produced are thought to be much lower (below \(\sim100 \text{ mmol per kg of rock}\); Klein et al., 2009; Andreani et al., 2013).

Serpentinization processes at magma-poor margins have received less attention. The chemical compositions of mantle rocks in these settings, however, commonly differ from peridotites at MORs. Specifically, rocks in ocean–continent transition (OCT) zones have experienced only moderate degrees of melt depletion and are hence compositionally closer to sub-continental lithospheric mantle (SCLM). Such rocks contain higher modal amounts of pyroxene than peridotite at MORs and thus more Si, Al, Ca, and Na (e.g., Hébert et al., 2001; Bodinier and Godard, 2014). Owing to these differences in bulk composition, serpentinization pathways likely differ from those at MORs where generally more depleted peridotites are serpentinized. For instance, higher Si in the bulk system potentially reduces the amount of brucite being formed (cf. Frost and Beard, 2007). Instead, more serpentine may form, some of which may hold Fe(III), which itself would increase the rates of H₂ production (see, e.g., reviews by Evans et al., 2013; Mayhew and Ellison, 2020).

The most prominent example of a magma-poor margin is the one at West Iberia, which formed during the break-up of Pangaea. Here, continental rifting caused thinning of the crust and eventually the widespread exposure of lithospheric mantle peridotite to the young North Atlantic Ocean seafloor (e.g., Boillot et al., 1988; Pickup et al., 1996; Pérez-Gussinyé, 2012). These exposures include the OCT, i.e., exhumed mantle that ranges in composition from weakly depleted sub-continental lherzolite to more depleted harzburgitic/dunitic rocks in the oceanic lithosphere (e.g., Hébert et al., 2001; Chazot et al., 2005). Studies of serpentinization at the West Iberia margin have mostly focused on peridotites from the depleted oceanic-type mantle (Sites 897 and 899 drilled during Ocean Drilling Program (ODP) Leg 149, and Sites 1067, 1068, and 1070 of Leg 173; e.g., Beard and Hopkinson, 2000; Skelton and Valley, 2000; Hopkinson et al., 2004; Schwarzenbach et al., 2013). To our knowledge, there are no detailed investigations on the serpentinization of the more continental-type mantle exposures—these reactions hence remain incompletely understood.

In this work we investigate serpentinization processes and the associated production of H₂ in pyroxene-rich mantle peridotite from the OCT at the West Iberia margin. By discussing the effects of bulk rock composition, spreading velocities, and temperature conditions during alteration, this study outlines the systematic changes in serpentinization reaction pathways and H₂ yields during the evolution from continental break-up to mid-ocean ridge divergence at ultraslow and eventually slow spreading rates.

Geological Setting and Samples

The West Iberia margin is a hyperextended magma-poor margin that documents the break-up of Pangaea and the early stages of the opening of the North Atlantic. Combined seismic studies and oceanic research drilling (ODP Legs 103, 149, and 173) have revealed the evolution from thinned continental crust over the OCT to purely oceanic crust (e.g., Boillot et al., 1987; Manatschal et al., 2001; Pérez-Gussinyé and Reston, 2001; Whitmarsh et al., 2001; Reston and McDermott, 2011; Dean et al., 2015). The OCT
is about 40–170 km wide and consists of several north–south-trending "peridotite ridges" at which lithospheric mantle has been exhumed to the seafloor along a series of extensional faults (Figure 1; see references above). The extension that lead to this mantle exhumation took place at rates of ~10 mm/yr (Whitmarsh and Miles, 1995). The peridotites are generally characterized by variable degrees of melt depletion, in concert with the evolution from sub-continental to oceanic lithospheric mantle (e.g., Abe, 2001; Hébert et al., 2001; Chazot et al., 2005). Seawater influx along numerous extensional faults led to extensive serpentinization of the ultramafic rocks. Up to 100% of the peridotite has been transformed into serpentinite to a depth of ~4 km below the seafloor, as evidenced by drill cores and suggested by seismic imaging (e.g., Boillot et al., 1988; Dean et al., 2000; Beslier et al., 2001; Bayrakci et al., 2016). At even greater depths, serpentinites may account for up to 25% of the basement, in particular where deep faults are present (e.g., Dean et al., 2000). The timescales for serpentinization were estimated to be ~10^5–10^6 years (Skelton et al., 2005) even though some hydration of the uppermost mantle may have taken place before its exhumation (Skelton and Valley, 2000; Pérez-Gussinyé and Reston, 2001; Pérez-Gussinyé et al., 2006). The bulk of the serpentinization took place at temperatures of 120–200°C (Agrinier et al., 1988; Agrinier et al., 1996; Schwarzenbach et al., 2013; Klein et al., 2014).

The rock samples investigated here originate from ODP Leg 103, Hole 637A—a drill site within the transition from SCLM to oceanic mantle at the Galicia Bank (Figure 1). Coring extended ~75 m into the basement and recovered dominantly clinopyroxene-rich spinel harzburgites and lherzolites (Boillot et al., 1988). The rocks are rich in Na, Al, and Ti, and spinel has particularly low Cr/Al and high Mg/Fe ratios, suggesting low degrees of depletion by melt extraction (<10%; Evans and Girardeau, 1988). The peridotites were likely exposed to the seafloor prior to the establishment of seafloor spreading (Evans and Girardeau, 1988). They are strongly serpentinized with dominantly pervasive replacement textures (Boillot et al., 1988). Shallow within the drill hole, the serpentinites have a leached appearance and replacement by calcite is common; further downcore, calcite is rather present in veins (Evans and Baltuck, 1988). Our samples originate from ~230 to 267 m below the seafloor.

**METHODOLOGY**

Polished thin sections were examined with a standard petrographic microscope. Primary and secondary fabrics and mineral phase assemblages and compositions were further evaluated on the basis of backscattered electron (BSE) imaging, energy-dispersive X-ray spectroscopy, and electron microprobe analyses. For bulk rock investigations, 10–15 g of sample were homogenized and aliquots of these were used for the different analyses. Sample preparation included cutting with a rock saw, cleaning with deionized water in an ultrasonic bath, and drying of the samples at 50°C. The samples were pulverized using a planetary agate ball mill.
Bulk Rock Elemental Analyses
Bulk rock compositions were analyzed on fused glass beads at the Institute for Chemistry and Biology of the Marine Environment at the University of Oldenburg. Preparation included the preoxidation and melting of weighed portions of sample, Li2B4O7 and NH4NO3 at 500 and 1,450°C, respectively, following Eckert et al. (2013). Concentrations of major and trace element concentrations were measured with a wavelength dispersive PANalytical Axios plus X-ray fluorescence spectrometer. Analyses of in-house reference materials were used to monitor the data quality. Analytical accuracies are better than 2.6% for major elements, except for Na2O.

Carbon contents were measured on a LECO CS744 combustion analyzer at the Department of Geosciences, University of Bremen. Contents of total carbon were determined on ~0.1 g dried bulk rock powder, and contents of total organic carbon were analyzed on ~0.1 g aliquots of rock powder that were previously treated with HCl (12.5%) to remove all inorganic carbon. Amounts of total inorganic carbon were calculated by subtracting total organic carbon from total carbon contents.

Magnetometry and Saturation Magnetization
For measurement of magnetic hysteresis, between 20 and 30 mg of sample powder were filled into a plastic straw and then hardened by applying a small amount of glue (cyanoacrylate). Measurements were performed at the Department of Geosciences, University of Bremen, on a Princeton Measurement Corporation Micromag 2,900 Alternating Gradient Magnetometer (Flanders, 1988). The accuracy of the instrument is 2% vs. the calibration sample (77.67 × 10−6 Am2, standard reference material 2,853, Yttrium Iron Garnet); its sensitivity is 10 × 10−12 Am2 standard deviation. The maximum applied fields were 0.3 and 1 T, respectively, which are sufficient to saturate magnetite; hematite and goethite need much larger field strengths to become magnetically saturated. The saturation magnetization of magnetite, hematite, and goethite are 480, ~2.5 and ~2 kA m−1, respectively (Dunlop and Özdemir, 1997). These numbers result in specific magnetizations of 92.36, 0.47, and 0.47 Am2 kg−1 when applying densities of 5.197, 5.271 and 4.264 g cm−3, respectively (Dunlop and Özdemir, 1997). After conducting the magnetic hysteresis measurements, we subtracted the linear slope from the data calculated for the field interval larger than 70% of the peak field, which is due to para- and/or diamagnetic constituents.

We estimated magnetite concentrations by assuming that it contributes the largest part to the saturation magnetization of the entire sample, also in the presence of other Fe oxides or hydroxides, due to its ~200 times higher specific magnetization compared to hematite and goethite. (Partial) oxidation of magnetite, although decreasing its mass magnetization, would not dramatically affect the computed magnetite concentrations, because the mass magnetization of fully oxidized maghemite (γ-Fe2O3) is still 74.89 Am2 kg−1 (Dunlop and Özdemir, 1997).

Fe(III)/ΣFe Analyses
In order to further examine possible oxidation of the magnetic mineral phases we exemplarily performed temperature-dependent remanence measurements and additional magnetic hysteresis measurements in peak fields of 1 T on two samples (25R-4 and 27R-2, 126–130 cm). The relative sensitivity of the used Quantum Design MPMS XL-7 superconducting quantum interference device (SQUID) magnetometer operated at the University of Bremen is <1 × 10−11 Am². An isothermal remanent magnetization was imparted to the samples in a steady magnetic field of 5 T at 300K. Afterward, the sample was cooled in zero-field to 5K and subsequently re-warmed to 300K (cf. Özdemir and Dunlop, 2010).

Fe(III)/ΣFe was determined for all samples using Mössbauer spectroscopy at Bayerisches Geoinstitut, University of Bayreuth. Roughly 200 mg of each sample were mounted in plastic holders, which corresponds to a Mössbauer thickness of 13 mg Fe cm−2. Mössbauer spectra were recorded at room temperature in transmission mode on a constant acceleration Mössbauer spectrometer with a nominal 1.85 GBq 57Co source in Rh. The velocity scale was calibrated relative to metallic iron. Spectra were collected for between one and five days and fit using MoSSA software (Prescher et al., 2012). Iron-containing phases could be identified based on hyperfine parameter values, and the distribution of Fe between phases was calculated from relative areas.

Electron Microprobe Analyses
Mineral chemical compositions were obtained with a CAMECA SX100 microprobe at the Department of Geosciences, University of Bremen. The device is equipped with four wavelength dispersive spectrometers. Analytical conditions included accelerating voltages of 15 kV and beam currents of 10–20 nA. Peak counting times varied between 20 and 40 s per element; analyses of primary phases were carried out with a focused beam whereas alteration phases were measured with a beam of 5 µm diameter. The built-in PAP correction was applied for data reduction and correction. Natural silicate and oxide standards (olivine, USNM 143965, number of analyses, n = 25; hornblende, USNM 143965, n = 20; chromite, USNM 117075, n = 22; Jarosewich et al., 1980) were repeatedly analyzed to check data quality. For elements with concentrations >1 wt%, the external reproducibility is better than 2.5%, except for Na2O (3.2%), and the averaged relative error as a measure for accuracy is below 2.5%, except for TiO2 (3.2%) and Na2O (6.5%).

Thermodynamic Calculations
Modeling was conducted using EQ3/6 version 8.0 (Wolery and Jarek, 2003) and the customized database from Klein et al. (2013), which has been assembled using SUPCRT92 (Johnson et al., 1992). The database contains log K values for temperatures from 0 to 400°C in 25°C increments at a constant pressure of 50 MPa. Details of thermodynamic data and on the calculation of activity coefficients in the database can be found in Klein et al. (2013). The models include solid solutions for olivine (forsterite, fayalite), orthopyroxene (enstatite, ferrosilite), clinopyroxene (diopside, hedenbergite), serpentinite (containing the endmembers
The calculations were done in three steps that mimic the inflow of seawater into peridotite basement, changes in fluid chemistry during the reaction with the basement, and the eventual serpentinization of peridotite by the seawater-derived fluid. We first speciated 1 kg of sulfate-free, seawater-like fluid (composition as in Table 3 of Klein et al. (2009)), except that we excluded HCO$_3^-$ and SO$_4^{2-}$, at 25°C and 50 MPa using EQ3NR. In the second step, 1 kg of solid reactant was added in increments to the seawater and equilibrated for the same conditions using EQ6. We chose two rock compositions as solid reactants: 1) sample 25R-2 as an example for rocks from the OCT; 2) an undepleted hypothetical endmember composition (Salters and Stracke, 2004). In the last modeling step, the system was heated to 400°C in small increments and equilibrated at each step, also using EQ6. All calculations were for a water-to-rock (w/r) mass ratio of unity, and the mineral and fluid compositions as a function of temperature. Such a w/r ratio is thought to prevail during pervasive serpentinization (cf. Barnes and Sharp, 2006) and is likely representative of the hydration of the investigated samples. Further, the calculated equilibrium assemblages/conditions are well representative of the West Iberia margin samples since serpentinization reactions proceed relatively fast and local equilibria have likely been reached throughout most of the samples. In addition, a set of calculations for a w/r ratio of 10 was performed to investigate the higher fluxes that may have prevailed active fault zones.

### RESULTS

#### Petrology

The fifteen samples are strongly serpentinized lherzolites and harzburgites, and one dunite. An overview of the samples, the presumed protolith types, and the degree of alteration is presented in Table 1.

Protolith fabrics were mostly overprinted during serpentinization. Where preserved, they can differ on a centimeter scale (Figures 2A,B). In some samples, deformation of peridotite is illustrated by elongated pyroxene and Cr spinel grains (e.g., samples 25R-3 and 29R-1) and in some cases by foliation or brittle deformation textures (e.g., sample 27R-1, 147–150 cm). Other samples lack such deformation fabrics and instead show non-elongated pyroxene and/or Cr spinel (e.g., samples 28R-3 and 27R-4). Unaltered olivine was not observed but primary clinopyroxene, orthopyroxene, and spinel are locally present (Figures 2A–C). Pyroxene frequently shows undulatory extinction and cleavage planes. Clinopyroxene grains are usually up to ~2 mm long, but may reach lengths of >4 mm (e.g., sample 25R-5), and often occurs close to
FIGURE 2 | Petrographic and textural features of the studied samples (A) Thin section scan showing a typical sample with domains of magnetite-poor serpentine mesh textures and darker domains enriched in magnetite. Pyroxene is partly and spinel is mostly unaltered. Light-colored serpentine veins crosscut previous textures. Carbonate fills voids, many of which may have been former mesh centers. Sample 27R-4, 75–79 cm, plane polarized light (PPL) (B) Thin section scan showing a domain rich in mostly unaltered pyroxene. Magnetite appears concentrated in elongated domains (dark areas). Carbonate veins crosscut the sample. Sample 25R-3, 10–13 cm, PPL. (C) Photomicrograph depicting mostly unaltered clinopyroxene with some serpentine present along intragranular fractures. Sample 25R-5, 119–122 cm, PPL. (D) Photomicrograph showing typical serpentinite mesh texture. Carbonate occurs in some mesh centers. Sample 27R-2, 126–130 cm, PPL (left) (Continued)
Orthopyroxene or in its exsolution lamellae (e.g., sample 25R-4). Orthopyroxene is more frequent than clinopyroxene and is mostly ~3–4 mm in size (e.g., sample 27R-2, 126–130 cm). Typically, pyroxene is serpentinized along cleavage planes and microcracks; only minor parts are unaltered (Figures 2A–C). Spinel occurs in anhedral or holly-leaf shapes (e.g., samples 27R-1, 147–150 cm, and 27R-4, respectively) with grain sizes of up to ~2.5 mm. Some grains are rimmed by or locally altered to magnetite.

The samples are extensively altered, with degrees of serpentinization generally >90% (Table 1). Whereas shallow samples appear yellowish–brownish, dark brown–greyish colors dominate further downhole. Secondary mineral assemblages include serpentine, carbonate, oxides, and minor chlorite and amphibole (Figures 2A–I). Serpentinite, the dominant phase, has in most cases pervasively replaced primary minerals but also can be found as precipitates in veins crosscutting the rocks (Figures 2A,D–F). Despite the pervasive replacement of olivine, its pseudomorphs can usually only be identified using polarized light microscopy (Figure 2F). Bastite serpentines textures, by contrast, are commonly well discernible using polarized light microscopy (e.g., sample 28R-3). Highly porous serpentinite that appears to have grown in void space—which we interpret to represent dissolved centers of former olivine and/or serpentine—is frequent throughout all samples. Micrometer-sized Fe oxides, presumably magnetite, are distributed in serpentine that pervasively replaced olivine as well as in veins. In some cases, magnetite forms bands crosscutting serpentine (Figure 2E). Accumulations of magnetite cause a local dark-colored appearance of serpentinite (Figure 2B). In veins, grains can be up to 50 µm large (Figure 2G). Magnetite is less common within altered pyroxene. Up to 20 µm large, euhedral Fe-oxide grains with stoichiometries matching that of magnetite can further occur within void space (Figure 2H) or within carbonate veins (Figure 2I; see below). In a few samples we observed chlorite, mostly in the direct vicinity of spinel (e.g., sample 29R-1). Minor amounts of amphibole are associated with fresh or altered pyroxene in the lherzolitic samples; amphibole is usually anhedral but shows needle-like shapes in a few instances (e.g., samples 27R-1, 45–48 cm and 27R-2, 126–130 cm). The majority of the samples experienced late-stage carbonation, with frequent calcite occurrences in millimetric to centimeter-wide veins (e.g., samples 25R-4 and 26R-3) and in the centers of former olivine grains (Figure 2D). In some samples, euhedral Fe and/or Mn oxide grains are enclosed by calcite or occur in between the carbonate and the host mineral (e.g., samples 27R-1, 45–48 cm, and 28R-3).

**Bulk Rock Geochemistry**

The serpentinites are characterized by variable major elemental concentrations, in particular SiO$_2$ (23.7–41.2 wt%), MgO (21.1–35.7 wt%), Fe$_2$O$_3$ (6.2–9.5 wt%), and CaO (0.7–21.5 wt%; Supplementary Table S1). Total inorganic carbon (TIC) ranges from 0.4 to 17.1 wt% and the loss on ignition (LOI) from 13.1 to 25.1 wt%. TIC and CaO strongly correlate with LOI and SiO$_2$ whereas MgO anticorrelates with CaO, TIC, and LOI, respectively. The data imply that CaCO$_3$ was added to the peridotites, resulting in a strong dilution of some samples. To assess the peridotite geochemistry before the addition of CaCO$_3$ we back-calculated the bulk rock compositions assuming that all TIC is present in the form of CaCO$_3$. The corrected values are presented in Table 2 and Figure 3 (dry rock). Ratios of Al$_2$O$_3$/SiO$_2$ and MgO/SiO$_2$ average at 0.041 ± 0.013 (1 SD) and 0.89 ± 0.033, respectively.

Trace elemental compositions are similar to previously published data from the Iberia margin (e.g., Seifert and Brunotte, 1996; Hébert et al., 2001).

**Iron Oxidation State and Magnetite Contents**

Bulk rock Mössbauer analyses revealed considerable distinctions between the samples and provide a means to identify different Fe oxides. These results allow a subdivision of the samples into three groups (Table 1 and Supplementary Table S3). In the first two groups, about 3/4 of all Fe is present within silicate phases, which we will refer to as “least modified” and “oxidized” from here on. The bulk of the remaining Fe is incorporated in magnetite in the least modified samples and in goethite and/or hematite in the oxidized samples. Within the third group, the “metasomatised” samples, silicates are markedly less dominant and instead, goethite and/or hematite can contain up to 4/5 and hence most of the Fe (on average 56% vs. 6% in the least modified and 20% in the oxidized samples). Mössbauer spectra are available in Supplementary Figure S1 and hyperfine parameter values in Supplementary Figure S2.

Bulk rock oxidation states, i.e., Fe(III)/ΣFe, calculated on the basis of these data are about 0.6–0.65 for the least modified samples, around 0.8 in the oxidized samples, and up to 0.93 in the metasomatized ones. In line with these results, bulk magnetization analyses suggest magnetite as the dominant magnetic mineral in the least modified samples and significant partial oxidation in the oxidized group. In greater detail, magnetic hysteresis loops of samples from both groups (25R-4 and 27R-2, 126–130 cm, respectively) indicate the dominance of low-coercivity minerals such as magnetite.
TABLE 2 | Corrected bulk rock geochemistry, Fe oxidation state, and magnetite contents of the Hole 637A samples.

| Sample | SiO₂ wt% | Al₂O₃ | MgO | CaO | Na₂O | LOI | MgO/SiO₂ | Fe∑ | Fe(III)/Fe∑ | Ba | As | Mo | Nb | V | Zn | Y | Pb | Sr | Th | U | V | Cr | Cu | Ga | Mo | Nb | Ni | Pb | Rb | Sr | Th | U | V | Cr | Cu | Ga | Mo | Nb | Ni | Pb | Rb |
|--------|---------|-------|-----|-----|------|-----|-----------|------|------------|----|----|----|----|---|----|---|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|
| 637A-25R-2, 62-65 cm | 45.14 | 44.05 | 44.72 | 46.23 | 41.62 | 45.29 | 45.97 | 46.58 | 46.35 | 47.55 | 45.25 | 42.36 | 44.78 | 38.62 | 36.64 |
| 637A-29R-3, 10-13 cm | 2.62 | 1.66 | 2.55 | 2.42 | 0.84 | 1.05 | 1.73 | 1.23 | 2.18 | 1.41 | 2.33 | 1.42 | 2.06 | 1.61 | 2.47 |
| 637A-25R-4, 28-32 cm | 8.34 | 8.67 | 8.53 | 7.83 | 7.77 | 8.35 | 9.08 | 8.52 | 8.63 | 8.09 | 9.01 | 9.21 | 10.77 | 10.82 |
| 637A-25R-5, 119-122 cm | 0.09 | 0.08 | 0.08 | 0.10 | 0.32 | 0.17 | 0.08 | 0.08 | 0.08 | 0.07 | 0.08 | 0.13 | 0.10 | 0.24 | 0.10 |
| 637A-25R-6, 66-69 cm | 38.85 | 37.70 | 39.92 | 39.34 | 38.99 | 40.79 | 40.20 | 41.00 | 40.63 | 40.74 | 41.06 | 36.90 | 40.20 | 34.35 | 42.50 |
| 637A-27R-1, 45-48 cm | 2.65 | 5.12 | 1.81 | 1.19 | 3.00 | 2.73 | 1.47 | 0.61 | 0.85 | 0.84 | 0.64 | 4.61 | 10.03 | 7.14 | 0.03 |
| 637A-27R-2, 107-110 cm | 0.32 | 0.35 | 0.31 | 0.28 | 0.24 | 0.23 | 0.26 | 0.13 | 0.18 | 0.18 | 0.19 | 0.29 | 0.36 | 0.34 | 0.10 |
| 637A-27R-3, 147-150 cm | 0.09 | 0.12 | 0.05 | 0.04 | 0.01 | 0.03 | 0.04 | 0.01 | 0.02 | 0.01 | 0.03 | 0.03 | 0.03 | 0.05 | 0.01 |
| 637A-27R-4, 65-68 cm | 83.70 | 81.19 | 84.48 | 85.22 | 82.03 | 83.53 | 84.21 | 86.53 | 86.20 | 87.34 | 83.33 | 79.01 | 77.82 | 73.73 | 84.84 |
| 637A-28R-1, 126-130 cm | 15.17 | 17.20 | 14.16 | 13.77 | 16.36 | 15.12 | 14.45 | 13.01 | 13.32 | 12.96 | 15.09 | 17.58 | 18.26 | 20.43 | 14.76 |
| 637A-28R-2, 138-141 cm | 0.058 | 0.038 | 0.057 | 0.052 | 0.020 | 0.023 | 0.038 | 0.027 | 0.047 | 0.030 | 0.051 | 0.034 | 0.046 | 0.042 | 0.057 |
| 637A-28R-3, 68-70 cm | 0.961 | 0.856 | 0.893 | 0.851 | 0.937 | 0.901 | 0.874 | 0.880 | 0.877 | 0.857 | 0.907 | 0.871 | 0.898 | 0.890 | 0.974 |
| 637A-28R-4, 36-40 cm | 0.787 | 0.792 | 0.797 | 0.803 | 0.931 | 0.827 | 0.805 | 0.648 | 0.597 | 0.607 | 0.611 | 0.859 | 0.922 | 0.906 | 0.639 |

- From Mössbauer spectroscopy.
- From magnetometry.
Temperature cycling of the latter sample points to a chemical composition close to stoichiometric magnetite and hence strengthens the above interpretation. Results of sample 25R-4, by contrast, suggest significant partial oxidation of magnetite. Further details can be found in the caption of Supplementary Figure S3.

Bulk magnetization analyses revealed magnetite contents from 0.29 to 1.73 wt%, with a median of 0.53 wt% (Table 2). The least modified samples hold a median of 0.83 wt% of magnetite whereas contents in oxidized and metasomatized samples are 0.53 and 0.41 wt%, respectively. No clear relation between magnetite content and coring depth has been observed.

Mineral Chemistry

Clinopyroxene has $X_{\text{Mg}}$ values typically in the range of 0.92–0.94, $\text{Al}_2\text{O}_3$ contents ranging from 3.3–6.2 wt%, TiO$_2$ between 0.1 and 0.4 wt%, and Na$_2$O contents between 0.58 and 1.09 wt%. Orthopyroxene has slightly lower $X_{\text{Mg}}$ values (0.91–0.92), and typically contains 3.5–5.2 wt% Al$_2$O$_3$, <1 wt% CaO (except where close to clinopyroxene exsolution lamellae), and <0.1 wt% Na$_2$O. Primary spinel has $X_{\text{Cr}} = \text{Cr}/(\text{Cr} + \text{Al})$...
values usually within 0.13 and 0.45 and $X_{\text{Mg}}$ of 0.52–0.76 (Figure 4A). The cores of large grains are usually unzoned; in a few instances, for example when chlorite occurs close to spinel, slight increases in Cr and Fe toward the rims were observed. Contents of TiO$_2$ in spinel are mostly <0.15 wt% but can reach up to 0.25 wt% (Figure 4B).

Secondary phases show variable compositions. The bulk of these phases can be identified as serpentine, with median $X_{\text{Mg}}$ of 0.93 ± 0.05 (1 SD; n = 256; Figure 5A). Elevated Fe contents in a number of analyses ($X_{\text{Mg}}$ as low as 0.43) point toward a cronstedtite component in serpentine. The Fe-rich serpentine occurs either in the centers of mesh textures (Figure 2F and Supplementary Figures S4B–D) or as porous, void-filling precipitates (Figure 2H). We could, however, not record clear cronstedite spectra during reconnaissance micro-Raman analysis, presumably due to an overprint by lizardite (Supplementary Figure S5).

Mineral compositional data are reported in Supplementary Table S2.

**Geochemical Modeling Results**

We applied thermodynamic calculations to track the amounts of H$_2$ that are being produced during serpentinization at magma-poor margins such as the West Iberia margin. Figures 6A–D summarize results for seawater–basement reactions at an OCT, based on the composition of sample 25R-2, 62–65 cm, demonstrating intermediate composition between those of SCLM and strongly depleted harzburgitic/dunitic rocks commonly found at MORs (cf. Figure 2; see also Degree of Melt Depletion at Hole 637A). Over most of the investigated...
temperature range, the equilibrium mineral assemblage (Figure 6A) consists of serpentine, amesite, and clinopyroxene. Olivine and magnetite are predicted to be stable only at/above 400°C. Below ∼150°C, trace amounts of brucite are predicted. The $X_{\text{Mg}}$ of serpentine plots at ∼0.89 independent of temperature, with a fraction of ferric Fe in serpentine of ∼0.2 at 380–150°C that decreases toward ∼0.1 at 25°C once brucite has joined the assemblage (Figure 6B). As a consequence, the predicted amounts of H$_2$(aq) per kg rock are ∼150 mmol at ∼380–150°C but <100 mmol toward seawater temperatures (Figure 6D).

Figures 6E–H demonstrate the consequences of fluid–SCLM interactions, based on the undepleted mantle composition by Salters and Stracke (2004). We chose this composition as a counterpart to the strongly depleted compositions found at MORs to demonstrate the consequences of high modal pyroxene contents for serpentinization. The predicted equilibrium assemblage yields slightly higher amounts of magnetite and brucite, which are stable over larger temperature ranges (Figure 6E). The $X_{\text{Mg}}$ of brucite drops from 0.83 at ∼220°C to 0.59 at 25°C (Figure 6F). The fraction of ferric Fe is highest at ∼380–220°C where both serpentine and magnetite are stable, with Fe(III)/∑Fe of 0.32, but decrease as brucite becomes stable and progressively richer in Fe (Figure 6F). Up to 0.37 moles Fe(III) are produced by the co-occurrence of serpentine and magnetite; once magnetite is no longer part of the equilibrium assemblage, this number drops to <0.3 moles and further decreases with decreasing temperature (Figure 6G). The predicted amount of H$_2$(aq) follows that of ferric Fe (Figure 6H).

Results from model runs at w/r ratios of 10 as well as those in which the formation of garnet is allowed are shown in Supplementary Figures S6, S7.
DISCUSSION

Hydrothermal Alteration at Hole 637A
Initial Pervasive Serpentinization

The serpentinites investigated here show evidence for a high-temperature strain and metamorphism history, as evidenced by microtextures indicative of ductile deformation and the occasional presence of amphibole, respectively (e.g., Figure 2B; cf. Agrinier et al., 1988; Kimball and Evans, 1988). Subsequent to these, extensive fluid–rock interactions took place at lower temperatures. Early serpentinization of the samples is distinguishable from prolonged interaction with seawater on the basis of petrographic observations and bulk rock and mineral chemical data.

The initial stage of serpentinization was mostly pervasive. Whereas primary spinel is preserved in all samples, olivine is completely altered to serpentine minerals and trace amounts of Fe oxides, often in pseudomorphic mesh textures. Microtextures
indicative of deformation at higher temperatures are statically replaced by serpentine. Higher $X_{Mg}$ in serpentine (median = 0.93) relative to that of its precursor olivine (ranging from 0.894 to 0.906; Kimball and Evans, 1988) implies that Fe has been redistributed during alteration. We thereby exclude that large amounts of Fe have been incorporated into brucite, since the breakdown of pyroxene likely caused high SiO$_2$ activities ($a$SiO$_2$) during serpentinization that prevented brucite formation in the first place (cf. Frost and Beard, 2007). Instead, it is likely that most of the serpentinization occurred via the reaction $Mg_2SiO_4 + 2SiO_2 + 4 H_2O = 2 Mg_3Si_2O_5(OH)_4$ with SiO$_2$ derived from the dissolution of pyroxene through the reaction $3 MgSiO_3 + 2H_2O = Mg_3Si_2O_5(OH)_4 + SiO_2$. In line with this conclusion are pyroxene grains that are in most cases strongly reacted (cf. Figure 2), the lack of brucite in any of our samples (Figure 5), and thermodynamic calculations predicting only trace amounts of brucite for the reaction of a representative sample with seawater-derived fluid (Figure 6). Hence we do not interpret the putative loss of bulk MgO in some of our samples (Figure 3C) as the dissolution of brucite during prolonged seawater interaction (e.g., Snow and Dick, 1995; Klein et al., 2020), but as a metasomatism-related dilutional effect (it is the samples with high CaO that appear particularly MgO-poor). Thus, instead of being partitioned into brucite, we infer that most Fe has likely been partitioned into serpentine and/or scarce magnetite, the latter of which occurs evenly distributed in some but very heterogeneously in other samples.

Initial serpentinization resulted in $Fe(III)/\Sigma Fe$ of $\sim$0.6 (least modified samples in Figure 7A), very similar to the Fe oxidation state in serpentinites from the Mid-Atlantic Ridge (MAR; Andreani et al., 2013; Klein et al., 2014). Serpentine thereby contains more than 3/4 and thus the bulk of the ferric Fe and magnetite holds another $\sim$15–20% (Figure 7B). The Fe-rich serpentine mesh centers presumably host a large portion of the Fe(III) but in samples without such mesh centers, Fe(III) must be present elsewhere—bulk magnetization analyses suggest a median of 0.83 wt% magnetite for the least modified samples (Table 2). However, this number may serve as a minimum estimate for magnetite production during initial serpentinization since minor amounts of goethite and hematite may have formed after magnetite and hence, initial magnetite contents may have been slightly higher. This alteration likely also lowered the measured relative fraction of Fe(III) in serpentine (Figure 7B). Nevertheless, conclude that generally little magnetite has been produced during initial serpentinization.

Our reaction path models match this assumption as they do not predict any magnetite in the equilibrium assemblage over most of the temperature range (Figure 6A). This is at least true for the case of a composition representing low to intermediate stages of melt depletion, as was used in our models. But the peridotites recovered from Hole 637A are compositionally very diverse, ranging from lherzolitic to dunitic (Boillot et al., 1987) and it is hence likely that other lithologies have influenced the fluid–rock interactions. For instance, $\sim$1 wt% of magnetite would
be produced during the hydration of undepleted lithologies (Figure 6E), and even higher portions for strongly depleted rocks such as harzburgites or dunites (Klein et al., 2009; McCollom and Bach, 2009). Furthermore, aside from the composition of the protolith, the alteration temperature and the extent of serpentinization are factors that to a major degree influence the amounts of magnetite being produced. Andreani et al. (2013) and Klein et al. (2014) demonstrated that magnetite contents non-linearly correlate with increases in temperature and the serpentinization extent. All peridotites recovered from Hole 637A as well as from nearby holes drilled during ODP Legs 149 and 173 are extensively serpentinized and yet hold little magnetite (Boillot et al., 1987; Sawyer et al., 1994; Whittmarsh et al., 1998), indicating low hydration temperatures. This general interpretation has been validated by results from O stable isotopes, which consistently imply serpentinization temperatures at the West Iberia margin of below 200°C (Agrinier et al., 1988; Agrinier et al., 1996; Schwarzenbach et al., 2013; Klein et al., 2014). Altogether, the median magnetite content of our least modified samples (0.83%) falls markedly below contents typical for MORs (4–6%; e.g., Klein et al., 2014).

Instead of having been partitioned into brucite and/or magnetite, Fe was incorported into serpentine as indicated by our mineral chemical data (Figure 5). About 16% of the total analyses (n = 256) exhibit $X_{Mg} < 0.9$ and the median $X_{Mg}$ of the lowest 10% is 0.83. In particular, the centers of serpentine mesh textures can be enriched in Fe, whereas mesh rims and intergranular serpentine usually represent “typical” Mg-rich serpentine (Figure 2F and Supplementary Figures S4B–D). Such serpentinite with higher Fe contents in mesh centers has previously been described by Kimball and Evans (1988). Our data imply a high cronstedtite component, which is further discussed in Formation of Cronstedtite and $H_2$ During Initial Serpentinization. Ortho- and clinopyroxene are mostly altered to bastite serpentine but domains of un-serpentinized pyroxene are locally preserved in most samples. This can either suggest that the alteration conditions were favorable for the (meta-)stable preservation of pyroxene or that pyroxene reacted at considerably slower rates in comparison to olivine. The relative rates of pyroxene and olivine reaction are a matter of recent debate (see, e.g., discussion in McCollom et al., 2020). Thermodynamic modeling results suggest that clinopyroxene is stable throughout the considered temperature range at w/r ratios close to unity (Figure 6). Somewhat contrasting, petrographic observations in some cases revealed clinopyroxene that is partially serpentinized (Figure 2C). These observations are more in agreement with model runs at higher w/r ratios, at which pyroxene is no longer part of the equilibrium assemblage (Supplementary Figure S6A). Prolonged late-stage interaction with seawater (see below) is also not accounted for by the models but could explain local clinopyroxene serpentinization. Orthopyroxene is not predicted to be stable in any model run, which fits with petrographic observations revealing that orthopyroxene is in most cases strongly reacted.

**Oxidation and Mass Transfer During Prolonged Fluid–Rock Interactions**

The interaction with seawater did not cease subsequent to the initial serpentinization but instead continued and led to further mineralogical and geochemical modifications. Calcite is very common and occurs in veins and void space (Figures 2B,D,I), the latter of which probably represent the centers of former olivine grains. Calcite abundance is reflected in the bulk geochemistry, which can exhibit up to 21.5 wt% CaO and 17.1 wt% CO$_2$ (uncorrected data, Supplementary Table S1). Both CaO and CO$_2$ were added to the rocks by cold, circulating seawater as was proposed earlier for West Iberia margin drill sites (Evans and Baltuck, 1988; Gibson et al., 1996). Euhedral Fe oxides that match the stoichiometry of magnetite are present within many calcite veins (Figure 2I; see also Evans and Baltuck, 1988), suggesting that Fe was added to the seafloor together with the calcite or that Fe was released during the dissolution of Fe-containing minerals and was subsequently re-precipitated in the present form. The former may be true for samples in which bulk Fe$_2$O$_3$ is increased relative to “typical” serpentine (Figures 3, 7). Bulk Fe$_2$O$_3$ concentrations of most samples do not, however, display noticeable enrichments and thus support the latter possibility. The common observation of dissolved mesh center argues for the dissolution of olivine and/or serpentine during late fluid–rock interactions. Calcite that occurs in mesh centers which are comparable in size and location to the olivine or serpentine cores found in other samples has previously been described from Hole 637A (Kimball and Evans, 1988). Similar conclusions have been drawn based on observations in weathered peridotites from the MAR, in which open voids in the centers of mesh textures host Fe hydroxides that apparently precipitated from void-filling fluids after the dissolution of brucite and/or olivine (Jöns et al., 2017). These rocks also contain abundant carbonate that, besides being present in veins, fills the voids alongside Fe hydroxide.

The Site 637 samples further show evidence for the formation of a late serpentine generation (Figure 2H) with highly porous serpentine that frequently exhibits $X_{Mg} <$ 0.9 (Supplementary Figure S4B) and elevated Na contents. Microtextures suggest precipitation in void space and the mineral chemistry implies intergrowth of serpentine with a strong cronstedtite-component, stevensite, and talc (Figure 5). It presumably is this cronstedtite-rich serpentine that accounts for the shift of the fraction of Fe(III) in serpentine from ~0.6 in the least modified to ~0.75 in oxidized samples (Figure 7A). This shift can be observed in 1/3 of our samples and we suggest that this is related to heterogeneities in porosity and permeability between the individual portions of rock that led to different degrees of seawater influx and reaction. Indeed, the precipitation of carbonate may even clog further inflow of seawater and hence reduce the permeability (cf. Jöns et al., 2017). The bulk Fe(III)/ΣFe of these samples has increased to ~0.8 (Figure 7A) and highlights the general oxidation of the seafloor caused by the prolonged interaction with seawater. It is within these samples from the shallowest core sections that goethite and hematite appear to have replaced most of the magnetite, similar to what has been described for Site 1068 (Beard and Hopkinson, 2000).
average, more than 20% of the total Fe(III). The presence of maghemite is also likely (cf. Zhao, 2001; Ouf et al., 2002) but it cannot be confirmed with the analytical techniques used in this study.

Overall, these observations document the prolonged low-temperature interaction with fluids at/close to the seafloor. This interpretation is supported by the presence of calcite that precipitated at or close to ambient deep-sea temperatures (Agrinier et al., 1988; Evans and Baltuck, 1988). Similarly, serpentine may at least in parts have formed at temperatures of pure seawater (cf. Barnes and Sharp, 2006). Intergrowth of stevensite and talc, however, may also suggest that highest water/rock ratios during alteration. The generation, most of which we observed in samples from the shallow core sections. For these sections, Evans and Baltuck (1988) suggest highest water/rock ratios during alteration. The intergrowth of stevensite and talc, however, may also suggest that the serpentinized interacted with sedimentary pore waters instead of pure seawater (cf. Barnes and Sharp, 2006).

In the metasomatized samples, high abundances of goethite and hematite are remarkable (Figure 7B). These samples contain high amounts of calcite (whole rock data show consistent enrichments in CaCO3), host Fe oxides and oxyhydroxides, and in some cases serpentine with variable X_{Mg}. Microtextures are in many cases indicative of mineral growth in equilibrium with a fluid. Oxides rich in Mn, Co, and/or Ti are present in trace amounts. It appears likely that the former fractures—now filled with vein precipitates—provided pathways for enhanced seawater flow through these portions of the seafloor. This seawater interaction also led to the addition of Fe in some of the metasomatized samples, as previously proposed by Gibson et al. (1996), and in all samples to the oxidation of most of the Fe that is not contained in serpentine. The portions of ferric Fe in serpentine, however, appear unchanged as they are within or close to the range of the least modified or oxidized samples.

**H₂ Production During Seawater–Basement Interactions at Hole 637A**

**Formation of Cronstedtite and H₂ During Initial Serpentinization**

During serpentinization, H₂ is produced during the oxidation of ferrous to ferric Fe. Ferric Fe is incorporated in magnetite, Fe₂Fe₃+O₄, and/or the serpentine-group endmember cronstedtite, (Mg,Fe²⁺,³⁺)₂Fe₃+SiO₃(OH)₄. In the samples investigated here, the amount of ferric Fe in serpentine and hence H₂ production associated with the formation of cronstedtite outweighs that in/related to magnetite (Figure 7B).

Cronstedtite substitution in serpentine describes the partial exchange of Fe²⁺ and/or Mg with Fe³⁺, which is accompanied by a Fe²⁺(Fe³⁺Si)₁ exchange to maintain charge balance (e.g., O’Hanley and Dyar, 1993). If it went to completion, the substitution would result in the endmembers Fe-cronstedtite, Fe₂⁺Fe₃+SiO₃(OH)₄, or Mg-cronstedtite, Mg₂Fe₂⁺SiO₃(OH)₄, or mixtures thereof.

The presence of cronstedtite by itself is not unusual for marine serpentinites. It can be approximated by analyzing serpentinite Fe(III)/∑Fe; generally, between 50 and 60% of the Fe in MOR and passive margin serpentinites are ferric, even in ones containing little magnetite (Mayhew and Ellison, 2020). But X_{Mg} values for serpentine in these rocks rarely fall below 0.9 and the total contents of Fe₂O₃ in serpentinites are usually below 7–8 wt% (cf. Mayhew and Ellison, 2020).

Most of the serpentine analyzed here plot in the typical X_{Mg} range of 0.9–0.95 (median = 0.93) but several samples also exhibit serpentine with X_{Mg} values well below 0.9 in mesh centers and strongly suggest a pronounced cronstedtite component (Figures 2F, 5A). The stability of cronstedtite is primarily dependent on aSiO₂ and H₂ fugacity during serpentinization (e.g., Zolotov, 2014; Tutolo et al., 2020). Serpentine with minor cronstedtite fractions can be formed under rather variable conditions, but higher cronstedtite components require increasing aSiO₂ (Figure 8). As discussed above, such elevated-aSiO₂ conditions likely prevailed during the initial serpentinization of the lherzolitic rocks as well as during their late interaction with (sediment-derived) fluids.

Based on our Mössbauer data, 60–65% of the Fe in serpentine of the least modified samples is ferric (Figure 7A). Serpentine of such composition can be explained by mixing of serpentine–greenalite with Mg-cronstedtite (Figure 9). One kg of pure serpentine with Fe(III)/∑Fe of 0.65 and X_{Mg} of 0.95 would produce about 150–170 mmol H₂, assuming that H₂O was the only oxidant. This hypothetical value is in the same order of magnitude as our modeling results (~120–160 mmol H₂ per kg rock at 120–200°C; Figures 6D,H) and previous thermodynamic models for lherzolitic compositions (Klein et al., 2013). By comparison, ~430 mmol H₂ would be produced per kg serpentine with X_{Mg} = 0.88, as observed in mesh centers (Figure 9). Assuming that such serpentine accounts for 1/5 to 1/4 of the entire serpentine at the West Iberia margin, H₂ yields of ~215–230 mmol H₂ appear realistic per kg serpentine—which is almost twice as much as if such Fe-rich serpentine were not present. Hydrogen fluxes calculated based on our Mössbauer data are even higher (up to 260 mmol H₂ per kg serpentinite; Figure 7C), but these likely are overestimations since some of the oxidation may have been caused by oxidizing agents other than H₂O (e.g., SO₄²⁻).

The scarcity or absence of magnetite in the least modified samples (median = 0.83 wt%; Table 2) hence does not signify a lack of oxidation or that no hydrolysis has taken place. However, magnetite formation in these rocks has even increased the fluxes of H₂ up to >300 mmol per kg bulk rock (Figures 7B,C).

The bulk of the pervasive serpentinization likely took place during seawater seepage into the seafloor at w/r ratios close to unity. In addition, however, tectonic activity likely provided fluid pathways at/close to which each rock unit reacted with larger amounts of fluid. Our thermodynamic estimates revealed mineral assemblages dominated by serpentine for both undepleted to weakly-depleted compositions at w/r ratios of 10 (Supplementary Figure S6). Amounts of Fe(III) are <0.4 mole per kg rock in both cases, translating to 50–120 mmol H₂(aq) produced at >250°C but ≤50 mmol at lower temperatures.
Late-Stage Serpentinite Oxidation and H₂ Production

Seawater–serpentinite reaction after their exposure to the seafloor caused additional H₂ fluxes, related to the formation of Fe(III)-serpentine–stevensite–talc mixtures and the formation and transformation of Fe oxides and/or hydroxides. The former was favored by increased aSiO₂ (cf. Figure 8) and resulted in the release of another ~50 mmol H₂ per kg rock (Figure 7C). The latter was particularly pronounced in the shallow, strongly oxidized core sections of Hole 637A, in which much of the magnetite has turned into hematite and goethite (Figures 7A,B). This modification likely took place in two steps, namely the oxidation of magnetite to hematite, 4 Fe₃O₄ + H₂O = 6 Fe₂O₃ + H₂(aq), and the formation of goethite at the expense of hematite, Fe₂O₃ + H₂O = 2 FeO(OH) (cf. Klein et al., 2017). The decisive factor for the stability of hematite and magnetite is the H₂ fugacity of the system (Frost, 1985). The abundance of hematite in the oxidized samples and even more so in some of the metasomatized samples, in which hematite holds
between 2/5 and 3/5 of all Fe(III), suggests that the H$_2$ fugacities did not cross the critical level at which hematite destabilizes and magnetite is predicted—despite the fact that H$_2$ is produced during the above reaction. In fact, the transformation of magnetite into hematite and/or goethite accounts for the estimated shift in H$_2$ production per kg rock from roughly 300–350 mmol to >400 mmol per kg rock (Figure 7C). It appears counterintuitive that a more oxidized assemblage comes with a high H$_2$ yield. But this reaction likely occurred over long timescales, presumably tens of millions of years before the seawater circulation through the basement ceased owing to the burial of the rocks by sediments (e.g., Tucholke and Sibuet, 2007). Thus, despite the fact that overall ~100 mmol H$_2$ per kg rock were produced during this late-stage alteration, the flux of H$_2$ per increment of time as well as the H$_2$ fugacities were presumably rather low.

We assume that the oxidation to ferric Fe would have continued and likely would have proceeded to completion if the thick layer of sediment had not restricted seawater–basement interaction. The Fe in extensively weathered serpentinites from other marine settings has been shown to be almost entirely ferric (e.g., Klein et al., 2017; Dessimouilie et al., 2020) since much of the magnetite and spinel hosting Fe(II) turned into Fe(III)-bearing hematite, goethite, and ferrichromite. The dissolution of unserpentinized olivine, serpentine, and brucite (if present) also releases ferrous Fe that is subsequently precipitated as Fe hydroxides (e.g., Jöns et al., 2017). Under the prerequisite that H$_2$O is the oxidizing agent, additional H$_2$ would be formed during each of these reactions.

**H$_2$ Production From Continental Break-Up to Mid-Ocean Ridge Spreading**

Degree of Melt Depletion at Hole 637A

Across OCTs, seawater should principally interact with SCLM at proximal and with oceanic lithospheric mantle at more distal settings. To put the findings from this communication into the context of such a transition from continental rifting and break-up to MOR spreading, we next elaborate to what extent our samples depict serpentinized SCLM. Mantle rocks consist mainly of olivine + orthopyroxene + clinopyroxene ± garnet ± spinel, the proportions of which vary with the progressive removal of basaltic components during partial melting (e.g., Arai, 1994). Lherzolites are considered to be best representative of SCLM, and persistent partial melting in response to the above-mentioned transition causes the transformation from lherzolitic (at passive margins) to harzburgitic and/or dunitic compositions (at MORs). This general change in composition occurs across the Iberia margin OCT (e.g., Hebert et al., 2001).

Geochemically, undepleted lherzolite should plot close to the primitive mantle estimates (Figure 3) and the rocks from the peridotite ridges at the West Iberia margin—generally thought of as variably depleted SCLM—should ideally fall along the melt depletion trend (i.e., the “terrestrial array” in Figure 2A in Niu, 2004). This is indeed the case for our as well as other Iberia margin samples. However, our samples display very heterogeneous compositions, some of which suggest little whereas others imply advanced stages of depletion. These compositions well mirror the lithological variations throughout all drill cores at the West Iberia margin, in which lherzolitic core sections can alternate with harzburgitic and dunitic ones on centimeter scales (e.g., Boillot et al., 1987; Sawyer et al., 1994; Whitmarsh et al., 1998). In addition, further heterogeneity has potentially been caused by melt infiltration (e.g., Chazot et al., 2005; Müntener and Manatschal, 2006) and/or extensive metasomatism (e.g., this study; Seifert and Brunotte, 1996). Pinning down the degree of melt depletion on the basis of our bulk compositions would hence be speculative.

A more accurate way is offered by the compositions of spinel. Prolonged melt extraction turns initially Mg- and Al-rich spinel into spinel with a higher chrome fraction, i.e., low $X_{Mg}$ and high $X_{Cr}$ (Arai, 1994). Spinel in our samples is particularly rich in Al and Mg and plots among the least-depleted spinels of rifted margin and MOR samples (Figure 4A). Further, TiO$_2$ contents below 0.15 wt% imply that most of our samples were not affected by significant melt impregnation processes (Figure 4B; cf. Müntener and Manatschal, 2006). Based on these data we infer that the investigated rocks can, overall, not have undergone large degrees of melt extraction. This interpretation is also consistent with up to 5.2 wt% Al$_2$O$_3$ in orthopyroxene and with high Al$_2$O$_3$, Na$_2$O, and TiO$_2$ contents and a high average value of 6.2 for 100*Na/(Na + Ca) for clinopyroxene (see also Evans and Girardeau, 1988). In fact, Evans and Girardeau estimated that the Site 637 peridotites have experienced less than 10% of partial melting. Based on these data and on the (reconstructed) lherzolitic protolith compositions, we are confident that the investigated rocks well resembled SCLM prior to initial serpentinization. The samples from Hole 637A hence depict a compositional endmember opposite to the dunitic and harzburgitic lithologies at MORs.

**Serpentinization From Rift to Ridge**

Serpentinization at Slow- and Ultraslow-Spreading Ridges

In this work we have analyzed the serpentinization and derived H$_2$ fluxes associated with SCLM at the most landward domain of the Iberia margin OCT. In contrast, sustained divergence led to the exposure of oceanic peridotite rather than SCLM in more oceanward domains and all along to the modern MAR. Similar transitions have been described from many settings worldwide and throughout Earth’s history (e.g., Louden and Chian, 1999; Pérez-Gussinyé, 2001; Manatschal, 2004; Pérez-Gussinyé, 2006). Both rock types undergo hydration by seawater. To obtain an overview of H$_2$ yields across such transitions, we next review serpentinization reactions affecting oceanic mantle. We thereby explicitly focus on “background” serpentinization, i.e., pervasive hydration of the mantle that is induced by the percolation of seawater into the seafloor (contrary to serpentization that can occur along deep faults).

We also need to take account of changing spreading rates over time. During the evolution to its present state, the MAR developed from an ultraslow- (at the OCT, about 130 Ma ago; e.g., Whitmarsh and Miles, 1995) into a slow-spreading ridge (100 s of km to the west of the OCT). Aside from the Newfoundland–Iberia conjugate margin, such increases in...
extension rate have been suggested for several other break-up/spreading settings (e.g., Bullock and Minshull, 2005; Vissers et al., 2013; Williams et al., 2019). Whereas serpentinization at slow-spreading ridges is generally well understood, much less work has been conducted at their ultraslow counterparts. But previous investigations indicate that the styles of serpentinization differ between such settings.

In both cases, the extent and nature of serpentinization strongly depend on the crustal architecture, which itself is a function of variable melt supply and tectonic extension. The ultraslow-spreading Gakkel and Southwest Indian Ridges are considered to be modern counterparts of the Cretaceous MAR (cf. Whitmarsh et al., 1996; Hopper et al., 2004; Cannat et al., 2009). They are characterized by generally little magmatic activity and long stretches of sparsely non-magmatic ridge sections separated by discontinuous volcanic centers that formed by melt focusing. Mantle blocks emplaced at the seafloor via detachment faulting dominate the melt-starved sections (e.g., Cannat et al., 2006; Escartín et al., 2008). Overall, the along-ridge variation in crustal thickness is immense, with little to no volcanic crust at the magmatic sections and several kilometer thick mafic layers at volcanic centers (e.g., Jokat et al., 2003; Schmid and Schlindwein, 2016). Similarly, slow-spreading ridges are strongly segmented and melt supply at segment ends is usually low (e.g., Tolstoy et al., 1993; Detrick et al., 1995), which as well commonly results in the uplift of ultramafic seafloor to shallow depths. However, it appears that these mantle sections in many cases comprise variable volumes of gabbroic intrusions representing magma that crystallized at depth. The intrusion of such magmatic bodies results in higher pressure–temperature gradients than that expected for exhumed mantle without magmatic intrusions. Estimates for axial geotherms range from 45°C km⁻¹ (Cannat et al., 2019) to 100–150°C km⁻¹ (Bach et al., 2011) for magma-poor sections at slow-spreading ridges, which would place the 400°C isotherm and hence the window in which serpentinization can occur to a maximum depth of 8–9 km. Gradients at ultraslow-spreading ridges are lower, varying between about 25 and 50°C km⁻¹, depending on the magmatic activity in the respective section (Cannat et al., 2019). The depth at which serpentinization could take place would thus be extended to about 15 km. Indeed, results from microseismicity studies at the ultraslow-spreading Mid-Cayman Rise and Southwest Indian Ridge indicate extensive hydration of mantle rocks until depths of 15 km and more (Schlindwein and Schmid, 2016; Greveneyer et al., 2019). Given that the overall amount of hydration decreases with increasing depth below seafloor (Cannat et al., 2010, proposed that significant background serpentinization does generally not extend to depths greater than 3–4 km below seafloor), these differences in the lithosphere’s thermal state suggest generally lower serpentinization temperatures at ultraslow- in comparison to slow-spreading ridges. In other words, larger volumes of low-temperature serpentine should form at ultraslow-spreading ridges—but also at rifted margins that possess similar geotherms.

This reasoning is in line with serpentinization temperature estimates from O isotope thermometry that point at much higher values for spreading systems that are magmatically active. At the MAR and Hess Deep, seawater–peridotite interaction ranges from about 200 to >400°C; those at the nonvolcanic Iberia margin are considered to have occurred at <200°C (e.g., Früh-Green et al., 2004; Barnes, et al., 2009; Klein et al., 2014).

Such distinctions presumably apply for serpentinization evoked by seawater that pervasively infiltrates the permeable basement or flows in through the ubiquitous minor faults and fractures existing in spreading systems. However, this does not exclude that higher-temperature serpentinization may also take place where geotherms are low. Detachment fault surfaces that act as fluid pathways can facilitate seawater penetration into the seafloor to depths of >10 km (e.g., deMartin et al., 2007). Such inflow likely evokes serpentinization of the detachment wall rock to proceed at great depth, i.e., at the upper end of the serpentinization temperature window (e.g., Cannat et al., 2019). Continued spreading along these detachments may cause the exposure of such high-temperature serpentine, even at ultraslow-spreading ridges. Samples collected with seafloor dredges may undoubtedly represent such former detachment surfaces and may not be representative for background serpentinization (e.g., Rouméjon et al., 2018; Tao et al., 2020).

**Dependency of H₂ Yields on Rock Type and Serpentinization Temperature**

The quantity of H₂ generated during serpentinization primarily depends on rock type, alteration temperature, w/r ratio, and oxidation potential of the fluid. Water/rock ratios and the nature of the fluid (usually seawater-derived) are likely similar at slow- and ultraslow-spreading ridges as well as at rifted margins. But rock types and serpentinization temperatures systematically vary between the different settings, as discussed above. In Figure 10 we conceptualized the H₂ production potential during the evolution from continental break-up to MOR spreading.

At slow-spreading ridges, more than 80% of the H₂ generated during the serpentinization of harzburgites and dunites at temperatures in excess of 200°C originates from the formation of magnetite (Figure 5 in Klein et al., 2009). Hydrogen fluxes increase with temperature from about 200 to 350 mmol per kg of rock (dotted lines in Figures 6D, H); highest concentrations are expected where the maximum amounts of magnetite are formed (toward ~320°C). However, a small portion of such H₂ is generated by the oxidation of Fe that is subsequently incorporated in serpentine. Thermodynamic modeling predicts 20–35% of the Fe in serpentine to be ferrous within the temperature range of 200–280°C (Klein et al., 2009). Such serpentine compositions can be explained by mixing of serpentine–greenalite and Mg-cronstedtite (Figure 9); serpentine with X_Mg of 0.95 (as predicted by Klein et al. models) may produce about 50 mmol H₂ per kg rock. More generally, recorded serpentinization compositions from MOR settings worldwide usually scatter around X_Mg of 0.9–0.95; no unambiguous trends toward any endmember containing ferric Fe are discernable. Even so, analyses of natural rocks from MORs (e.g., Figure 5F in Mayhew and Ellison, 2020) suggest that up to 2/3 of the Fe in serpentine can be ferric. Such increase in Fe(III)/ΣFe would raise the amounts of H₂ generated solely by serpentine
to levels of ∼180 mmol per kg rock (Figure 9). Microprobe data plotted on the frequently-used SiO₂–FeO–MgO plane would not reveal such high Fe(III) contents.

In contrast, H₂ generation at ultraslow-spreading ridges should be mainly driven by Fe(III)-serpentine because of considerably lower serpentinization temperatures. In fact, thermodynamic modeling predicts magnetite-free mineral assemblages at temperatures below ∼160°C (Klein et al., 2009). If a little more than 20% of the Fe in serpentine were ferric (Figure 5B in Klein et al., 2009), such rocks would produce between about 50 and 100 mmol H₂ for Xₘ₉ values of 0.95 and 0.9, respectively (Figure 9). Higher Fe(III)/ΣFe in serpentine would considerably increase the levels of H₂ (see above). The serpentinized harzburgites and dunites recovered from the distal parts of the Iberia margin OCT can be considered as examples for ultraslow-spreading MORs; these rocks hold very little magnetite owing to rather low serpentinization temperatures and, instead, the Fe partitioned into brucite (Agrinier et al., 1996; Beard and Hopkinson, 2000; Klein et al., 2014).

In the proximal parts of OCTs, where the exposed mantle is rather lherzolitic in nature, background serpentinization also occurs at low temperatures and the main driver for H₂ production is Fe(III)-rich serpentine. The presence of such cronstedtite-rich serpentine is promoted by rather high aSiO₂ during the serpentinization of pyroxene-rich peridotite (in comparison to the alteration of harzburgites and dunites at MORs; Figure 8). Our results imply that considerable amounts of H₂ can be produced. With about 2/3 of the Fe being ferric in our least-modified samples, >300 mmol H₂ per kg serpentine may have been generated (Figure 7C). This value is much higher in comparison to ultraslow-spreading ridges and similar to magnetite-dominated high-temperature serpentinization at slow-spreading ridges. Considering that not all of the oxidation of these samples has been caused by seawater, the true H₂ yields may have been somewhat lower. But even with H₂O as the only oxidant, ∼120–160 mmol H₂ per kg rock are thermodynamically realistic (Figures 6D,H) and can be considered as an absolute minimum estimate.

Magnetic properties of drilled serpentinites support our conjecture that H₂ yields can predominantly be ascribed to magnetite at slow-spreading ridges but not at magma-poor systems. Comparing serpentinite magnetic susceptibilities reveals the abundance of magnetite at the slow-spreading MAR and thus highlights the distinction in the alteration mineral assemblages in comparison to the West Iberia margin (Figure 11). Serpentinization at the MAR can produce large amounts of magnetite and consequently forms rocks with high albeit variable magnetic susceptibilities. Since the production of magnetite enhances with the degree of serpentinization, magnetic susceptibilities increase rapidly when the degree of alteration exceeds 75% (Toft et al., 1990; Ouﬁ et al., 2002). In this regard, Ouﬁ et al. suggested a threshold at about 0.05 SI that separates weakly from extensively serpentinized samples (dashed gray bar in Figure 11; see also Figure 5C in Maffione et al., 2014). By contrast, recovery from the Iberia and Newfoundland margins
exhibits low magnetic susceptibilities, even though it is extensively serpentinized. Long-lasting weathering of these rocks at the seafloor may have obscured the initial magnetite contents to a certain degree but our results demonstrate that at least the least modified samples well reflect initial serpentinization products. The magnetite contents of these samples are, however, also low.

In addition to H₂ produced by serpentine (and magnetite), the formation of garnet during serpentinization may even increase H₂ yields at sites of continental break-up and at OCTs. Our thermodynamic calculations predict up to 8 vol% garnet dominated by an andradite component for lherzolitic rocks (Supplementary Figure S7). Its formation would raise the levels of H₂(aq) to ∼100–170 mmol at temperatures below ∼250°C, which is an increase by a factor of two to three in comparison to serpentinization of dunites and/or harzburgites (Supplementary Figures S7D,H). At the Iberia margin, andradite and hydroandradite have indeed been observed at Site 1068 (e.g., Beard and Hopkinson, 2000).

At both rifts and ridges, late-stage alteration of serpentinites produces further H₂ flux to the ocean. Since serpentinized dunites and harzburgites possess generally larger pools of Fe(II) in comparison to lherzolites (Fe(III)/∑Fe = 0.57 and 0.58 vs. 0.65, respectively; Mayhew and Ellison, 2020) their potential for H₂ production during weathering is slightly larger. Low-temperature weathering and the related mineralogical and geochemical changes are presumably long-lasting, widespread, and hence even more important at MORs (over OCTs) because sedimentation rates at these settings are usually lower and inefficiently inhibit seawater–serpentine interactions.

**CONCLUSION**

Hydrogen yields during seawater–peridotite interactions primarily rely on the serpentinization mineral assemblages, which in turn strongly depend on protolith type and alteration temperature. During the evolution from continental rifting and break-up affecting Pangea to MOR spreading in the North Atlantic, a succession of lherzolitic to harzburgitic and dunitic rocks underwent serpentinization—at low temperatures (<200°C) during rifting and early, ultraslow spreading and at higher temperatures (200–280°C) during later spreading at slow rates. During low-temperature serpentinization, the incorporation of Fe(III) in serpentine was mainly responsible for H₂ production. Hydrogen yields were likely higher by a factor of two or more during the seawater–lherzolite reactions at the OCT in comparison to those resulting from the hydration of harzburgites/dunites at the early, ultraslow-spreading MAR, because much ferrous Fe was partitioned into brucite in the latter. Once higher geotherms and thus higher serpentinization temperatures were established, the formation of larger amounts of magnetite again increased the H₂ formation. Continued weathering during cold, late-stage weathering of the serpentinites at the seafloor caused additional H₂ fluxes to the ocean.

These concepts and general changes in H₂ production likely apply to the majority of magma-poor margins as well as ultraslow- and slow-spreading MORs. We speculate that sufficient catabolic energy (in the form of H₂) is available at magma-poor rifted margins and ultraslow-spreading ridges to enable lithoautotrophic life (see discussion by Lang and Brazelton, 2020). Indeed, particularly large volumes of serpentinite should form at temperatures close to those at/below the limit of life (Figure 10), which likely favors geological–biological interactions and the presence of a hydrogenotrophy-based biosphere.

**DATA AVAILABILITY STATEMENT**

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author.
AUTHOR CONTRIBUTIONS
EA and MP-G designed the study. EA re-sampled the drill cores, investigated the samples, and conducted the analyses, except for magnetometry (TF) and Mössbauer spectroscopy (CM). EA and WB performed the modeling. EA wrote the manuscript with contributions from all co-authors.

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SUPPLEMENTARY MATERIAL
The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/feart.2021.673063/full#supplementary-material

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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