Theoretical predictions vs environmental observations on serpentinization fluids: Lessons from the Samail ophiolite in Oman

James A. M. Leong¹, Alta E Howells¹, Kirtland J Robinson¹, Alysia Cox², Randall Vincent Debes¹, Kristopher Fecteau¹, Panjai Prapaipong¹, and Everett Shock¹

¹Arizona State University ²Montana Technological University

November 21, 2022

Abstract

Thermodynamic calculations provide valuable insights into the reactions that drive the profound fluid transformations during serpentinization, where surface fluids are transformed into some of the most reduced and alkaline fluids on Earth. However, environmental observations usually deviate from thermodynamic predictions, especially those occurring at low temperatures where equilibrium is slowly reached. In this work, we sampled and analyzed >100 low-temperature ($<40^{\circ}$ C) fluids from the Samail ophiolite in Oman to test thermodynamic predictions with environmental observations. Additional simulations (*e.g.*, fluid mixing, mineral leaching) were also conducted to account for deviations from equilibrium expectations. Type 1 circumneutral (pH 7 to 9) fluids result from fluid interactions with completely serpentinized rocks common in the shallow subsurface. Type 2 hyperalkaline (pH >11) fluids approach equilibrium with diopside, and serpentine and brucite actively forming during advanced stages of serpentinization. We also investigated fluids with pH values of 9 to 11 to test whether these fluids are indicative of intermediate stages of serpentinization or mixing between the above end-member fluids. Fluids at intermediate stages of serpentinization and fluids derived from mixing can have the same pH, but the former have considerably lower dissolved Si that can be attributed to concomitant subsurface serpentinization and mineral carbonation processes. Overall, this work demonstrates that predicted and measured compositions of serpentinization-derived fluids can be successfully reconciled using a combination of equilibrium and fluid-transport simulations. This work substantiates these calculations as useful tools in exploring serpentinization reactions in deep subsurface aquifers on Earth as well as those beyond our own planet.

1	Theoretical Predictions vs Environmental Observations on Serpentinization
2	Fluids: Lessons from the Samail Ophiolite in Oman
3	
4	J. A. M. Leong ^{1,2} , A. E. Howells ^{1,3} , K. J. Robinson ^{1,4} , A. Cox ⁵ , R. V. Debes II ^{1,2} , K. Fecteau ^{1,4} , P.
5	Prapaipong ^{1,2} , and E. L. Shock ^{1,2,4}
6	
7	¹ Group Exploring Organic Processes In Geochemistry (GEOPIG), Arizona State University,
8	Tempe, AZ 85287, USA
9	² School of Earth & Space Exploration, Arizona State University, Tempe, AZ 85287, USA
10	³ School of Life Sciences, Arizona State University, Tempe, AZ 85287, USA
11	⁴ School of Molecular Sciences, Arizona State University, Tempe, AZ 85287, USA
12	⁵ Laboratory Exploring Geobiochemical Engineering and Natural Dynamics (LEGEND),
13	Montana Technological University, Butte, MT 59701, USA
14	
15	Corresponding author: J. A. M. Leong (jmleong@asu.edu)
16	
17	Key Points:
18	1. Serpentinizing fluids from the Oman ophiolite were sampled and analyzed to test
19	thermodynamic predictions with environmental observations
20	2. Compositions of a few fluid samples can be accounted for by equilibrium thermodynamic
21	predictions
22	3. Most samples deviate from equilibrium expectations and can be accounted for by simulations
23	of additional processes (e.g., fluid mixing)

24 Abstract

25 Thermodynamic calculations provide valuable insights into the reactions that drive the profound fluid transformations during serpentinization, where surface fluids are transformed into some of 26 the most reduced and alkaline fluids on Earth. However, environmental observations usually 27 deviate from thermodynamic predictions, especially those occurring at low temperatures where 28 29 equilibrium is slowly reached. In this work, we sampled and analyzed >100 low-temperature (<40°C) fluids from the Samail ophiolite in Oman to test thermodynamic predictions with 30 31 environmental observations. Additional simulations (e.g., fluid mixing, mineral leaching) were 32 also conducted to account for deviations from equilibrium expectations. Type 1 circumneutral (pH 7 to 9) fluids result from fluid interactions with completely serpentinized rocks common in 33 the shallow subsurface. Type 2 hyperalkaline (pH >11) fluids approach equilibrium with 34 diopside, and serpentine and brucite actively forming during advanced stages of serpentinization. 35 We also investigated fluids with pH values of 9 to 11 to test whether these fluids are indicative of 36 37 intermediate stages of serpentinization or mixing between the above end-member fluids. Fluids at intermediate stages of serpentinization and fluids derived from mixing can have the same pH, 38 but the former have considerably lower dissolved Si that can be attributed to concomitant 39 40 subsurface serpentinization and mineral carbonation processes. Overall, this work demonstrates that predicted and measured compositions of serpentinization-derived fluids can be successfully 41 42 reconciled using a combination of equilibrium and fluid-transport simulations. This work 43 substantiates these calculations as useful tools in exploring serpentinization reactions in deep 44 subsurface aquifers on Earth as well as those beyond our own planet.

45

47 Plain Language Summary

The lithosphere is directly involved in the habitability of a planet. Interaction between water and 48 rocks mobilizes nutrients and facilitates the transfer of energy from the lithosphere to the 49 biosphere. Of all water and rock reactions on Earth, perhaps one of the most profound is 50 serpentinization as it produces some of the most reduced and alkaline fluids on the planet. Fluids 51 52 generated through serpentinization support microbial communities and thus are attractive for their potential to support life in the deep subsurface as well as in rocky bodies outside our own 53 planet. Thermodynamic simulations allow predictions of the compositions of fluids and 54 55 microbial energy supplies in these less accessible serpentinizing environments. However, deviations from predictions will likely occur, especially at low temperatures where equilibrium is 56 slowly reached. We sampled and analyzed >100 fluids from the Samail ophiolite in Oman to test 57 equilibrium expectations with environmental observations. Further simulations that can reconcile 58 predicted fluid compositions with actual measurements were also conducted. By demonstrating 59 that simulations of low-temperature serpentinization can be successfully applied to analogous 60 processes here on Earth, this work substantiates these calculations as useful tools in exploring 61 serpentinization reactions occurring beyond our own planet. 62 63

05

- 65
- 66
- 67

68 1. Introduction

When surface water meets ultramafic rocks, reactions occur that ultimately generate some 69 of the most alkaline and reduced fluids on Earth. This process, known as *serpentinization* 70 involves the hydrous mineral serpentine mainly replacing the anhydrous minerals olivine and 71 pyroxene that originally composed ultramafic rocks. During serpentinization, fluids become H₂-72 73 rich through the coupled oxidation of ferrous iron to ferric iron in minerals and reduction of water to hydrogen. Ultimately, the enormous redox disequilibria generated when these reduced 74 fluids mix with oxic surface fluids provide energy to drive many metabolic reactions 75 76 (McCollom, 2007; Amend et al., 2011; Schrenk et al., 2013; Canovas et al., 2017) and the synthesis of organic compounds (Shock and Canovas, 2010; Lang et al., 2010). Geochemical 77 energy supplies for microbial life make serpentinizing systems attractive model ecosystems for 78 assessing the link between the geosphere and the biosphere, including those occurring in the 79 Earth's subsurface (Schrenk et al., 2013; Klein et al., 2015; Rempfert et al., 2017; Fones et al., 80 81 2019) and possibly, in other rocky bodies in our solar system (Schulte et al., 2006; Vance et al., 2007; Holms et al., 2015; Glein and Zolotov, 2020). The alkaline and reduced environments 82 resulting from serpentinization are also proposed to have facilitated the emergence of life on 83 84 Earth (e.g., Russell et al., 1994, 2010; Sleep et al., 2004; Holm et al., 2006; Martin and Russell, 2007; Martin et al., 2008; Russell, 2018). 85

Submarine vents releasing >100°C fluids to the ocean, such as those in the Lost City
(~120°C, Kelley et al., 2001; Seyfried et al., 2015) and Rainbow vent fields (~360°C, Charlou et
al., 2002), are famous examples of serpentinizing environments, known for their towering
chimneys and vigorous vents surrounded by diverse deep-sea biological communities. Less
conspicuous, but likely far more widespread, are fluids seeping diffusively from ultramafic rocks

91	at temperatures <100°C, such as those found near spreading ridges (Früh-Green et al., 2018),
92	serpentine mud volcanoes near subduction zones (Mottl et al., 2003, 2004; Wheat et al., 2008;
93	2020; Mottl, 2009), and ultramafic bodies in continents, where serpentinization-generated fluids
94	are documented in ophiolites, ultramafic massifs, metamorphic belts, and kimberlite fields in ~20
95	countries (for a list of these studies, see Etiope and Whiticar, 2019 and Leong and Shock, 2020).
96	The common occurrence of serpentinized fluids in many continental settings where ultramafic
97	rocks and water can interact illustrates that serpentinization can actively proceed even at ambient
98	conditions (25°C, 1 bar).
99	Mass-transfer calculations based on thermodynamic equilibrium provide insights into
100	reactions that can drive the transformation of surface fluids such as rain and seawater into
101	reduced, high-pH fluids. When applied, past studies (Pfeifer, 1977; Bruni et al., 2002; Palandri
102	and Reed, 2004; Cipolli et al., 2004; Sader et al., 2007; Boschetti and Toscani, 2008; Marques et
103	al., 2008; Okland et al., 2012; Paukert et al., 2012) show that mass-transfer calculations can
104	account for the overall compositional trends of ultramafic-hosted fluids even at low-temperature
105	conditions. However, at low temperatures where equilibrium is slowly reached, deviations from
106	predictions can occur. Fluid compositions that deviate from equilibrium expectations have not
107	been a focus of previous theoretical studies but can inform us of the various fluid transport
108	processes that may contribute to assessing the habitability of the serpentinizing subsurface.
109	Serpentinization is not unique to Earth and can occur in other rocky bodies in the solar system
110	such as Mars (Schulte et al., 2006; Ehlmann et al., 2010) and ice-covered ocean worlds in the
111	outer solar system like Europa and Enceladus (Vance et al., 2007; Glein et al., 2015; Waite et al.,
112	2017; Glein and Zolotov, 2020). With limited existing data, and in preparation for future
113	sampling, expectations for the compositional variabilities of fluids and microbial communities

114	possible in these less accessible environments can be informed by thermodynamic simulations. It
115	is thus imperative that we test thermodynamic simulations in accessible and analogous low-
116	temperature serpentinizing systems here on Earth to assess their utility in predicting
117	consequences of serpentinization in other planetary bodies. In this study, we test predictions on
118	fluid compositions by recent calculations of Leong and Shock (2020) with fluids sampled from
119	the Samail ophiolite in the Sultanate of Oman.
120	The Samail ophiolite hosts some of the largest and best preserved continental exposures
121	of ultramafic rocks (Nicolas et al., 2001) as well as several low-temperature (<40°C),
122	hyperalkaline fluids that were documented by numerous studies in the past (Barnes et al., 1978;
123	Neal and Stanger, 1983; 1984; 1985; Bath et al., 1987; Taylor et al., 1991; Sano et al., 1993;
124	Dewandel et al., 2005; Matter et al., 2006; Paukert et al., 2012; 2019; Chavagnac et al., 2013a;
125	2013b; Boulart et al., 2013; Ollson et al., 2014; Miller et al., 2016; Rempfert et al., 2017;
126	Canovas et al., 2017; Vacquand et al., 2018; Zgonnik et al., 2019; Fones et al., 2019;
127	Giampouras et al., 2020; Boyd et al., 2020). To test thermodynamic predictions, we sampled and
128	analyzed 138 compositionally diverse fluids from 17 locations in the ophiolite with pH values
129	ranging from near-neutral to hyperalkaline.
130	We investigated reactions that account for fluids with end-member compositions: (1)
131	near-neutral and (2) hyperalkaline pH. The former have pH values ranging from 7 to 9, are
132	enriched in Mg ⁺² and HCO ₃ ⁻ , and typically occur in shallow aquifers and surface streams. In

133 contrast, the latter are highly alkaline (pH >11) and enriched in Ca^{+2} , OH⁻, and gases such as H₂

and/or CH₄. Hyperalkaline fluids occur mainly in deeper aquifers (>500 m) and manifest at the

surface when deep-seated fluids discharge as springs (Dewandel et al., 2005). We also examined

136 fluids with intermediate pH (9 to 11), which have been less of a focus in previous studies, to

137	assess if they are consequences of the serpentinization pathway where meteoric water is
138	transformed into hyperalkaline fluids, or if they result from mixing between the end-member
139	fluids. A field-based method to identify these fluid types is also proposed that can help inform
140	decisions involving exploration, sampling, and experiments while in the field. Ultimately,
141	through the aid of predictions, we can quantify the contributions of various stages of
142	serpentinization to observed fluid compositions. Eventually, these calculations, tested with
143	analogous processes at Earth's surface, can provide constraints for the exploration of low-
144	temperature serpentinizing environments such as those construed for the deep and early Earth, as
145	well as in other rocky bodies in our Solar System.
146	2. Methods
147	2.1. Computational Model
148	Thermodynamic reaction-path calculations simulate reactions of minerals with a fluid and
149	determine the compositions of coexisting solid phases and fluid constituents attained at
150	equilibrium at various extents of the overall reaction progress. Details on the model setup are
151	described in Leong and Shock (2020) and are summarized in the supporting materials
152	accompanying this work. In addition to reaction-path calculations, dissolved concentrations of
153	solutes as constrained by mineral solubilities were also predicted, as described in detail in the
154	supplementary materials.
155	2.2. Field and Analytical Methods
156	2.2.1. Field Sites
157	A total of 138 fluid samples were taken from nine surface (springs) and eight subsurface

sites in the Samail ophiolite are shown in Figure 1. All sites are hosted in ultramafic rocks except

158

(wells) sites in the Samail ophiolite in 2009, 2010, 2012, and 2014. Locations of these sampling



160

Figure 1. Locations of sample sites overlain on a satellite map of the Oman ophiolite (Google
Earth). Filled black circles represent sites where fluids were sampled from the surface (springs,
streams) while open circles represent sites where fluids were sampled from wells. Names of the
various massifs that comprise the ophiolite are indicated in red texts.

165

166 for WDA-16 and Masibt, which are located within a gabbroic body. The occurrence of hyperalkaline fluids at most sites can be associated with geologic structures (basal thrust and 167 lithological boundaries) that can promote flow of deep-seated fluids to the surface through 168 169 fractures. Three sites (Al Hilayw, Falaij, and Al Bana) are in outcrops close to the base of the ophiolite (basal thrust) corresponding to where the ophiolite is thrust over the Arabian Peninsula. 170 The other sites are located within the ophiolite and many are hosted in ultramafic rocks near the 171 ultramafic-gabbro transition. The 17 sites are distributed across several of the massifs that 172 comprise the Samail ophiolite. Sites at Dima, Qafifah, and Falaij, together with the wells NSHQ-173





186

187 Figure 2. Hyperalkaline pools covered in white calcite films flowing into surface streams at (a)

188 Qafifah and (b) Dima. (c) Hyperalkaline fluids comprising a major component of the stream flow

at Shumayt.

190 2.2.2. Field and Analytical Methods

Portable electrodes and meters were used for *in situ* measurements of pH, temperature, 191 and conductivity. pH electrodes (WTWTM SenTixTM 41 gel electrode) attached to meters 192 (WTWTM 3300*i* and 3110) were calibrated daily using 7.0, 10.01 and 12.46 pH buffers. 193 Temperature and conductivity were measured using a YSI 30 handheld meter. Samples were 194 filtered in the field (Pall Acrodisc[®] 32 mm PF syringe filter with 1.2 and 0.8/0.2 µm Supor[®] 195 membrane) and stored in containers for further laboratory analyses. Measurements of redox-196 sensitive species (dissolved ferrous iron, O₂, sulfide, nitrate, and ammonia) were performed on-197 site using field spectrophotometers (Hach[®] 2400) with commercially supplied reagents from 198 Hach[®]. All analyses, except dissolved O₂ and sulfide, were measured using filtered water 199 samples. Dissolved ferrous iron concentrations were measured using the 1,10 phenanthroline 200 method. Total dissolved nitrate and ammonia concentrations were determined using the cadmium 201 202 reduction and salicylate methods, respectively, during 2009, 2010, and 2012 fieldwork. For samples collected in 2014, nitrate and ammonia were not measured in the field and 203 concentrations were instead determined using ion chromatography (see below). Dissolved O₂ or 204 DO values were determined using Hach[®] AcuVac Ampuls[®] with the indigo carmine and the 205 206 HRDO methods for low and high DO values, respectively, while dissolved sulfide values were measured using the methylene blue method. 207

For fluid samples taken in 2014, concentrations of major anions (F⁻, Br⁻, Cl⁻, SO₄⁻², NO₃⁻, NO₂⁻) and cations (Li⁺, Na⁺, K⁺, Ca⁺², Mg⁺², NH₄⁺) were determined on separate Dionex DX-600 ion chromatography (IC) systems using suppressed conductivity detection and operated by Chromeleon software (version 6.8). Samples were stored in high-density polyethylene (HDPE) bottles rinsed and soaked with deionized water. Samples for cations were acidified with 6 N

213	methanesulfonic acid (MSA) to approximately 19 mM final concentration to suppress carbonate
214	precipitation and ammonia degassing. The anion system employed a potassium hydroxide eluent
215	generator, a carbonate removal device, and AS11-HC/AG11-HC columns. The cation system
216	was equipped with CS-16 and CG-16 columns and cations were eluted isocratically with 19 mM
217	MSA at 0.5 mL/minute. Both systems were plumbed with an external source of deionized water
218	for suppressor regeneration to improve the signal-to-noise ratio of the analyses. Further details
219	on the above method can be found in Fecteau (2016) and Robinson (2018). For 2012 samples,
220	anions and cations were measured using similar methods described in Lindsay et al. (2018). For
221	fluids sampled in 2009 and 2010, anion and cation concentrations were determined using
222	methods described in Canovas et al. (2017). On average, analytical precision for cation and anion
223	analysis was better than 2% and 5% relative standard deviation (RSD), respectively.
224	An inductively coupled plasma-mass spectrometer (ICP-MS, Thermo Finnigan Element
225	2 TM High Resolution ICP-MS) was used to determine dissolved Al concentrations, and for
226	samples taken in 2014, dissolved Si values, both with precisions mostly within 3% RSD. For
227	samples taken in 2009, 2010, and 2012, dissolved Ca values were determined using both the IC
228	and the ICP-MS (precision mostly within 5% RSD). Generally, both instruments yielded similar
229	values (see Figure S1a) except for a few samples where the IC returned lower concentrations
230	than the ICP-MS. These discrepancies are mostly attributed to fluids sampled during the 2009,
231	2010, and 2012 fieldwork where IC cation samples were not acidified. Hence, Ca values for all
232	fluids sampled before 2014 are reported using the ICP-MS results. For fluids sampled in 2009,
233	2010, and 2012, dissolved Si and Mg concentrations were measured using an inductively
234	coupled plasma-optical emission spectrometer (ICP-OES, Thermo iCAP [™] 6300). Analytical
235	precision for analysis of Si and Mg was better than 3% and 1% RSD at 1σ , respectively. The Mg

236	concentrations of a subset of samples from 2012 and 2014 were determined using both the IC
237	and ICP-OES. Both instruments yielded consistent results (see Figure S1b). Reported Mg values
238	for the 2014 samples came from the IC while the rest came from the ICP-OES. For both the ICP-
239	MS and ICP-OES, samples were stored HDPE bottles that were acid-washed and rinsed in
240	deionized water. Before going into the field, bottles were spiked with trace metal grade HNO_3^- to
241	acidify the samples to pH <2. For calibration, a multi-element standard (High-Purity Standards
242	(HPS), North Charleston, SC) was used for the ICP-MS while single element standards for Mg
243	and Si (PerkinElmer [®] , Waltham, MA) were used for the ICP-OES.
244	Measurements of dissolved inorganic carbon (DIC) concentrations were performed using
245	a wet oxidation total carbon analyzer (OI Analytical 1010 TOC Analyzer) coupled with a
246	Thermo Delta Plus Advantage mass spectrometer. Samples were stored in VWR CS24 amber
247	glass vials with butyl rubber septa, which has low permeability to gases. Each sample was heated
248	followed by acidification with H_3PO_4 to drive off the DIC as CO_2 and analyzed using the mass
249	spectrometer. Further details of the method can be found in Gilles St-Jean (2003) and Robinson
250	(2018). Stable isotope (δ^2 H and δ^{18} O) values were determined using an Off Axis - Integrated
251	Cavity Output Spectrometer (OA-ICOS, LGR DLT-100) using methods similar to those reported
252	in Meyer-Dombard et al. (2015). Accounting for instrument drift and normalization to the
253	Vienna Standard Mean Ocean Water (VSMOW) was done following van Geldern and Barth
254	(2012). Precision was usually within $\pm 0.5\%$ and $\pm 0.1\%$ for $\delta^2 H$ and $\delta^{18}O$, respectively.
255	Water isotope analysis was conducted at the Sabo Laboratory at ASU School of Life
256	Sciences. IC and ICP-MS analyses were conducted at the ASU GEOPIG Laboratory while ICP-
257	OES and total carbon analyses where performed at Metals, Environmental and Terrestrial
258	Analytical Laboratory (METAL) at ASU (formerly, ASU Goldwater Environmental Laboratory

259	and ASU W. M. Keck Foundation Laboratory for Environmental Biogeochemistry). Results of
260	select field and laboratory measurements (pH, temperature, conductivity, water isotope, and
261	dissolved concentrations of major rock-forming elements) are shown in Table 1, while the full
262	dataset is compiled in Table S1.

Results from instrumental analysis were further processed to calculate charge balance, 263 264 ionic strength, mineral saturation indices, and aqueous speciation with the EQ3/6 program (Wolery and Jarek, 2003) using the thermodynamic dataset described in the supplementary 265 266 materials. Charge balance and ionic strength of fluids are documented in Table S1, while 267 saturation indices for some minerals are reported in Table S2 in the supplementary files accompanying this work. Results of calculations of aqueous speciation are depicted in Figure S2. 268 Analytical results for circumneutral to slightly alkaline fluids are mostly within 5% of 269 electroneutrality. Results for a few pH <11 well samples have high positive charge imbalance 270 271 due to unmeasured DIC or, if measured, have unusually low values for this pH range. 272 Unexpectedly low DIC values could be attributed to carbonate precipitation during well sampling or sample storage. Analytical results for hyperalkaline fluids are mostly within 5% of 273 electroneutrality, with some near 20%. At extremes in pH, poor charge balance can be attributed 274 275 to small uncertainties in pH measurements as OH⁻ and H⁺ are major components of basic and acidic solutions, respectively (e.g., see Nordstrom et al., 2009 for charge imbalance in acidic 276 277 examples). Shifting field-measured pH values by about 0.1 to 0.2 units, which is within the range 278 of observed fluctuations in the field, results in close to complete charge balance.

Analytical data of most samples from 2010, except for isotopic and dissolved Mg and Si values, were previously reported in the work of Canovas et al. (2017) that focuses on quantifying energy supplies available to chemotrophic microorganisms in serpentinizing fluids. Some fluids

282	from the 2009, 2010, and 2012 fieldwork were also concomitantly sampled by collaborators
283	(Paukert et al., 2012; 2019). DIC and dissolved Al values determined as described above are
284	reported in Paukert et al. (2012) and repeated here, while concentrations of all other solutes were
285	determined independently. Details on which samples correspond to these previous studies can be
286	found in Table S1. Overall, this work expands analytical data previously reported by the above
287	studies by supplying additional data from 79 fluid samples.

Table 1

Temperature, pH, Conductivity, Water Isotope, and Major Element Compositions of Samples														
Sample Number	Site	pН	Т	Cond	δ ¹⁸ Ο	δ²H	DIC	Cl	Na ⁺	 K+	$\underset{(a)}{Mg^{+2}}$	Ca ⁺² (b)	Si ⁺⁴ (c)	
			°C	µS/cm	(‰ VSMOW)	(‰ VSMOW)	mmolal	mmolal	mmolal	mmolal	µmolal	mmolal	µmolal	
Fluids with pH between 7 to 9														
100112AI	AB	8.5	29	467	3.97	15.35	2.62	1.16	0.81	0.03	1602	0.37	200	
140110B	D	8.4	23.5	568	1.49	6.99	3.63	1.58	1.31	0.04	1840	0.49	183	
090118P	D	8.8	20.2	973	2.58	13.43	3.68	5.53	4.35	0.10	2148	0.42	169	
140114S	F	7.7	21	787	-0.28	3.63	4.39	3.19	3.20	0.11	2010	0.60	200	
100107C	М	9.3	25.4	778	-0.86	-0.59	1.93	3.71	3.81	0.09	1445	0.33	289	
100107D	М	8.6	25.6	809	-0.81	-1.16	3.08	2.98	3.24	0.07	1522	0.84	347	
090120W	М	8.7	24.2	811	-0.70	-3.00	2.83	3.40	3.36	0.09	1691	0.72	397	
140111G	Q	8.9	22.6	586	0.25	4.94	4.11	1.53	1.53	0.04	2140	0.36	276	
1201140	Q	8.7	23	454	0.62	3.91	2.82	1.30	1.19	0.04	1451	0.30	138	
100109X	Q	8.9	23.3	664	-0.34	3.39	4.49	1.71	1.76	0.03	2290	0.40	283	
090119S	Q	9.0	21.6	493	1.43	7.80	2.05	2.08	2.03	0.06	1227	0.17	116	
090119T	Q	8.6	24.8	527	1.37	6.31	2.78	1.94	1.84	0.05	1541	0.26	239	
140116B	SH	7.9	26.5	760	0.33	0.39	5.13	1.28	1.06	0.10	2810	0.56	303	
140116C	SH	8.7	27.3	777	0.07	0.35	4.75	1.54	1.37	0.10	2680	0.63	279	
120116Q	SH	7.7	25	744	1.07	7.47	5.22	1.28	1.00	0.08	2908	0.51	309	
100113AP	SH	7.9	27	723	0.94	4.91	5.17	1.43	1.16	0.08	2680	0.58	288	
090123E	SH	8.2	23.9	700	0.93	5.38	4.96	1.37	1.08	0.09	2664	0.51	305	
100114AR	SU	8.8	21.5	923	0.14	3.00	3.04	5.19	5.43	0.10	1485	0.45	78.5	
090122B	SU	8.8	17.1	1440	0.05	1.54	3.25	9.36	8.90	0.17	1777	0.12	97.2	
100115AW	U	8.5	17.5	911	0.26	-1.26	n.d.	7.36	4.51	0.14	314	1.65	33.9	
					Fluids wi	ith pH bet	ween 9 d	and 11						

						-							
140110D	D	10.4	21.8	778	0.69	3.73	0.81	4.79	4.92	0.14	782	0.48	91.4
140112L	D	9.8	21.3	659	1.06	6.17	1.88	3.75	3.72	0.11	1150	0.31	111
090118N	D	9.7	17.5	864	1.15	8.10	2.32	6.61	6.22	0.17	1115	0.26	74.6

	Contra	nucu)											
100107E	М	10.1	26.9	910	-1.75	-4.63	0.68	4.86	4.89	0.10	885	0.14	214
090120X	М	10.6	27.7	962	-0.98	-4.45	0.15	5.40	5.05	0.12	810	0.95	186
140111H	Q	10.2	20.2	597	0.20	4.46	1.80	3.08	3.59	0.08	1170	0.21	145
140111I	Q	10.9	18.8	683	0.61	6.36	0.85	3.69	4.36	0.10	807	0.37	100
120114N	Q	9.8	21.8	667	-0.32	0.65	2.24	3.17	3.78	0.08	1082	0.28	133
100109T	Q	9.6	23.7	685	-0.19	3.04	3.56	2.38	2.83	0.05	1924	0.04	226
100109W	Q	10.3	20.8	696	0.24	5.83	1.93	3.35	4.15	0.07	1192	0.27	119
090117J	Q	10.6	18.8	823	-0.75	2.41	1.78	4.44	5.16	0.11	994	0.04	104
140116D	SH	9.1	27.1	777	-0.02	0.40	3.98	1.92	1.86	0.10	2430	0.72	260
120116R	SH	10.6	25.8	767	-0.72	1.89	0.76	3.51	4.26	0.12	807	0.19	83.4
100113AN	SH	10.2	28.6	720	-0.27	1.53	1.66	3.32	3.83	0.11	1433	2.17	157
100113AO	SH	10.4	27.4	734	-0.01	2.31	1.40	3.56	4.18	0.11	1298	0.52	138
090123G	SH	10.6	27.6	712	-0.45	1.17	0.59	3.54	3.94	0.12	1535	0.60	176
090122A	SU	10.9	17.4	1215	-0.72	-2.38	0.46	7.83	8.22	0.17	860	0.49	36.3
090122C	SU	10.8	18.5	1154	-1.10	-3.04	0.35	7.95	8.79	0.19	305	1.35	17.8
				Hy	peralkali	ne Spring	g Fluids	(pH >11	!)				
140115X	AB	11.4	29.5	2949	-0.95	-2.31	0.03	11.4	13.9	0.32	0.45	1.89	2.22
140115Y	AB	11.6	24.5	2778	-0.83	-1.55	0.11	11.8	14.3	0.33	1.26	1.79	2.16
140115Z	AB	11.3	32.2	4050	-0.10	-2.08	0.03	12.6	17.1	0.33	0.41	2.37	1.38
120118Y	AB	11.5	32.5	3343	0.04	-2.47	0.12	6.80	9.24	0.19	1.02	1.88	2.36
120118Z	AB	11.4	30.6	2140	-0.75	-0.70	0.12	9.85	12.1	0.27	0.70	1.50	2.96
120129G	AB	11.0	37.3	1455	-1.50	-4.77	0.05	5.11	5.19	0.15	0.63	1.48	2.82
120129H	AB	11.4	38.2	1616	-1.60	-5.61	0.04	3.41	3.79	0.10	0.51	1.41	2.89
100111AA	AB	11.7	28.5	3400	-0.94	-2.33	0.04	12.0	15.1	0.26	0.28	1.66	1.79
100111AB	AB	11.7	25.4	2770	-1.02	-1.94	0.13	12.3	15.7	0.28	0.79	1.61	2.45
100111AC	AB	11.6	16.5	1873	0.20	3.18	0.69	11.6	14.6	0.29	181	0.33	38.1
100111AD	AB	11.6	32.4	3235	-0.29	-2.64	0.05	13.0	18.1	0.27	0.28	2.18	0.55
100111AE	AB	11.9	18.9	2468	0.03	-0.68	0.07	13.4	18.3	0.29	1.46	0.73	7.84
100111AF	AB	11.9	24.2	2850	-0.74	-1.98	0.10	11.0	15.4	0.22	1.74	1.64	2.49
100112AG	AB	11.1	37.7	1421	-1.24	-3.38	0.04	5.73	6.14	0.15	0.65	1.73	1.37
100112AH	AB	11.2	38.4	1537	-1.72	-4.16	0.05	5.77	6.75	0.14	0.34	1.58	1.43
090116F	AH	11.8	26.3	2250	0.14	4.93	0.11	9.34	11.4	0.31	0.81	1.22	4.28
090116G	AH	11.8	27.1	2230	0.17	3.77	2.80	9.09	11.1	0.27	0.79	1.12	3.35
090116H	AH	11.5	24.5	1890	1.26	9.70	0.57	10.6	13.0	0.28	0.87	0.30	27.8
090116I	AH	11.9	20.7	2200	0.29	6.34	0.18	9.35	11.4	0.27	0.42	0.85	14.7
140110C	D	11.4	27	2011	-0.10	-0.07	0.05	7.45	7.91	0.22	1.30	1.96	4.26
140112K	D	11.4	26.9	1870	-0.77	-2.10	0.04	7.16	7.59	0.21	2.24	1.89	3.88
140112M	D	11.4	28.2	2058	-0.64	-0.45	0.05	7.36	7.89	0.22	0.67	1.98	5.60
090118O	D	11.5	30.3	1905	-0.60	-0.42	0.13	7.56	8.09	0.22	1.82	1.07	11.3
1401130	F	11.4	28.4	2329	-0.52	0.83	0.06	7.92	9.40	0.25	1.07	2.09	3.68
140113P	F	11.5	25.9	2224	-0.54	0.98	0.05	7.90	9.53	0.25	1.28	2.03	1.94
140114R	F	11.6	21.1	1844	-0.72	0.62	0.04	7.10	8.10	0.23	1.46	1.85	2.44

	\ \	/											
140114T	F	11.4	27.2	2061	-0.71	1.58	0.04	6.89	7.97	0.22	1.83	1.91	1.84
140114U	F	11.4	21.7	1479	0.24	4.27	0.17	4.72	8.09	0.22	16.1	1.17	7.43
140114V	F	11.4	24.4	1803	-0.21	2.90	0.23	6.98	8.43	0.22	5.60	1.71	2.75
120111A	F	11.6	22.3	1737	-0.07	2.02	0.09	4.62	5.33	0.14	2.04	1.31	7.88
120111B	F	11.4	29.4	1912	-0.41	1.46	0.09	6.39	7.38	0.20	1.32	1.71	4.46
120111C	F	11.3	31.9	1889	-0.45	0.81	0.07	4.94	5.41	0.15	1.07	1.72	6.40
120113I	F	11.4	30	2276	-0.30	0.70	0.14	7.24	8.85	0.22	2.42	1.71	3.29
120113J	F	11.6	25.8	2343	-0.21	1.87	0.09	3.56	4.20	0.10	0.59	1.72	3.82
120113K	F	11.4	30.9	2364	0.72	4.97	0.12	6.10	7.53	0.19	0.45	1.60	20.4
120113L	F	11.3	28.4	2297	1.66	9.71	0.16	10.3	12.79	0.33	0.96	0.73	26.4
100108J	F	11.7	23.9	2214	-0.33	1.91	0.05	8.58	10.4	0.22	1.97	1.89	1.73
100108K	F	11.7	24.7	2266	-0.47	2.00	0.07	8.35	10.4	0.22	1.78	1.87	1.60
100108L	F	11.6	29.2	2442	0.49	4.01	0.07	8.44	10.6	0.22	0.70	1.94	0.89
100108M	F	11.7	26	n.d.	0.27	3.55	0.08	9.99	12.7	0.30	0.46	1.95	7.02
100108N	F	11.6	27.6	2380	-0.05	1.83	0.04	8.28	10.4	0.21	0.89	1.75	0.43
100108O	F	11.8	22	2179	0.16	2.33	0.07	8.70	10.8	0.23	1.12	1.92	0.98
100108P	F	11.5	23.3	1507	0.67	5.48	0.06	7.92	9.61	0.21	15.3	0.56	7.48
100108Q	F	11.6	22.9	1752	0.18	4.44	0.08	7.59	9.19	0.20	6.36	1.19	2.47
100108R	F	11.6	27.9	1967	0.99	3.57	0.06	7.28	8.89	0.18	1.27	1.70	0.78
090115A	F	11.5	29.1	2251	-0.42	0.16	0.10	8.26	9.66	0.25	34.7	2.07	21.9
090115B	F	11.7	24.1	2154	-0.40	0.97	0.12	8.44	9.81	0.27	1.23	1.69	3.68
090115C	F	11.7	25.1	2280	-0.73	0.89	0.06	8.20	9.61	0.25	41.0	2.21	10.6
090115D	F	11.8	20.7	2270	-0.09	2.14	0.16	8.45	9.90	0.26	4.44	1.55	5.13
090115E	F	11.8	18	2240	-0.36	1.00	0.12	8.53	10.0	0.27	1.86	1.70	3.49
100107A	М	11.3	32.4	1900	-2.19	-8.41	0.11	7.25	7.18	0.12	0.21	2.06	37.2
100107B	М	11.3	30.3	1848	-0.92	-5.13	0.11	7.31	7.16	0.14	6.31	1.91	39.3
090120U	М	11.4	31.4	1676	-1.74	-7.86	0.11	7.10	6.58	0.14	102	1.95	50.5
090120V	М	11.4	30.6	1616	-1.68	-8.84	0.05	7.13	6.62	0.16	22.3	1.67	45.0
140111F	Q	11.6	23.8	1470	-0.61	1.07	0.04	4.66	5.86	0.14	3.82	1.69	10.5
120112E	Q	11.3	30.1	1762	-0.56	1.06	0.07	3.38	4.20	0.10	0.68	1.74	5.56
120112F	Q	11.4	27.4	1781	-0.70	0.82	0.11	4.32	5.45	0.13	1.93	1.72	4.62
120112H	Q	11.3	31.9	1893	-0.56	1.41	0.07	4.79	6.33	0.14	0.74	1.80	31.5
100109S	Q	11.8	22.4	1685	-0.37	1.30	0.10	5.32	7.03	0.13	0.54	1.59	3.77
100109U	Q	11.7	24.9	1778	-0.80	0.90	0.12	5.06	6.90	0.11	1.23	1.51	0.47
100109V	Q	11.2	23.7	904	0.41	6.03	0.66	4.25	5.60	0.10	347	0.47	51.7
090117K	Q	11.7	18.5	1074	-0.32	4.18	0.24	5.30	6.60	0.14	74.5	0.25	25.3
090117M	Q	11.7	22.6	1649	-0.62	2.41	n.d.	5.22	6.41	0.15	6.65	1.52	3.82
090119Q	Q	11.7	23.2	1797	-0.52	-0.82	0.13	5.29	6.76	0.15	56.5	1.82	24.0
090119R	Q	11.6	16.6	911	n.d.	n.d.	0.24	5.27	6.89	0.15	10.5	0.20	20.2
140117F	SH	11.5	26.2	1868	-2.04	-6.01	0.03	5.35	6.41	0.15	1.94	1.85	2.65
140117G	SH	11.4	30.5	2015	-2.20	-6.28	0.02	5.33	6.27	0.15	0.79	1.90	1.50
140117H	SH	11.5	29.6	2067	-2.06	-4.91	0.03	5.77	7.14	0.16	1.04	2.08	1.66
140117I	SH	11.3	32.3	2000	-2.22	-5.23	0.05	5.32	6.28	0.15	8.09	1.86	1.96

Table 1 (continued)

		,											
140117J	SH	11.3	31.6	1387	-1.79	-4.14	0.07	4.71	5.83	0.14	19.5	0.83	48.9
140117K	SH	11.5	27.4	1980	-2.13	-4.51	0.03	5.76	7.16	0.16	1.48	2.08	2.26
140117L	SH	11.6	27.3	2087	-2.15	-4.33	0.02	5.81	7.33	0.16	1.47	2.14	1.87
120116P	SH	11.2	34.3	1745	-2.10	-4.28	0.06	4.49	5.45	0.13	2.49	1.67	1.70
120116S	SH	11.4	29.5	1729	-1.87	-3.95	0.08	4.58	5.61	0.13	4.54	1.70	2.23
120117T	SH	11.3	32.6	1721	-2.12	-4.06	0.04	3.28	4.01	0.09	1.48	1.65	2.08
120117U	SH	11.3	32	1720	-2.05	-4.19	0.04	5.15	6.25	0.15	0.87	1.65	1.77
120117V	SH	11.7	23.6	1951	-1.67	-2.21	0.08	5.40	6.81	0.14	0.68	1.75	3.63
120117X	SH	11.3	32.8	1681	-2.18	-5.07	0.01	5.10	6.04	0.15	0.56	1.67	2.37
100112AJ	SH	11.5	33.4	1782	-2.22	-5.33	0.05	5.58	7.13	0.12	1.33	1.90	0.59
100112AK	SH	11.5	33.3	1793	-2.63	-5.79	0.04	5.57	7.06	0.12	0.45	1.86	0.73
100112AL	SH	11.5	32.6	1772	-2.45	-5.20	0.04	5.54	7.12	0.12	0.44	1.82	0.57
100112AM	SH	11.5	31.5	1772	-1.97	-4.66	0.04	5.51	7.13	0.12	0.62	1.92	0.53
090123D	SH	11.5	34	1778	-1.93	-4.70	0.09	5.35	6.48	0.15	1.80	1.46	1.91
090123F	SH	11.6	28.1	1770	-1.57	-3.96	0.14	5.61	6.80	0.16	1.86	1.50	1.55
100114AS	SU	11.3	18.2	1013	0.91	0.48	0.47	6.12	7.42	0.14	291	0.30	22.7
100114AT	SU	11.6	30.4	1893	-1.88	-7.27	0.04	6.33	8.40	0.14	0.70	1.72	3.53
100114AU	SU	11.6	20.08	1258	-0.91	-3.64	0.07	6.24	7.80	0.14	157	0.29	16.1
090122Z	SU	11.6	29.5	1897	-2.20	-7.87	0.11	6.01	7.68	0.22	0.54	0.21	4.66
100115AV	U	11.6	21.4	1442	-1.19	-8.79	n.d.	6.99	4.75	0.12	4.15	2.48	0.39
						Well Flı	ıids						
120119B	N14	10.1	24	855	-0.21	-1.71	0.07	4.27	2.31	0.08	32.6	1.14	5.95
120119A	N14	11.1	29.1	1920	0.64	0.89	0.09	7.68	5.40	0.12	5.68	3.42	2.78
120119C	N14	11.0	33.7	2296	0.31	-0.05	0.08	8.12	5.29	0.12	4.19	3.47	2.72
120123E	N04	10.7	28.3	3327	-1.38	-3.65	0.39	7.58	4.05	0.11	10.2	7.73	8.29
120126F	W16	8.0	31.8	492	-0.48	0.11	0.28	0.80	1.17	0.05	1420	0.38	456
120127G	W17	9.2	31	532	-0.56	0.76	2.35	1.16	0.85	0.04	2218	0.05	1.48
120130I	W5	9.3	32.7	595	-0.76	-0.72	1.90	2.03	1.32	0.07	2458	0.05	4.53
120130J	W5	9.4	32.1	610	-0.49	0.21	1.80	2.00	1.31	0.06	2419	0.05	4.42
100119AX	S4B	7.2	33.8	598	-2.54	-10.09	n.d.	1.15	1.26	0.04	1273	1.06	375
100119AY	S4A	7.6	34	491	-3.11	-9.94	n.d.	0.76	0.78	0.04	1123	0.96	366
100120AZ	S 3	9.4	34.1	586	-1.00	-4.91	n.d.	2.24	1.58	0.05	2100	0.03	4.38

Table 1 (continued)

Notes: b.d.l. - below detection limit; n.d. - not determined; Sites: AB - Al Bana, AL - Al Hilayw, D - Dima, F - Falaij, M - Masibt, Q - Qafifah, SH - Shumayt, SU - Sudari, U - Uqaybah, N14 - NSHQ-14, N04 - NSHQ-04, W16 - WDA-16, W17 - WDA17, W05 - WDA-05, S4A - SJA-04A, S4B - SJA-4B, SJ3 - SJA-03.

^aAll samples were analyzed using the ICP-OES except for those with sample number starting with "14" which were analyzed using the IC.

^bAll samples were analyzed using the ICP-MS, except for these with sample number starting with "14" which were analyzed using the IC.

^cAll samples were analyzed using the ICP-OES except for those with sample number starting with "14" which were analyzed using the ICP-MS.

289 **3. Results**

290 3.1. Model Results

Model results are described in Leong and Shock (2020) and a worked example is provided in the supplementary files of this paper and illustrated in Figure S4. Specifically, reaction paths depicting various stages of the serpentinization process – primary mineral dissolution (Stage 1), incipient serpentinization (Stage 2), mineral carbonation (Stage 3), brucite formation (Stage 4), H₂ formation (Stage 5), and equilibrium with diopside (Stage 6) – are summarized in the supplementary materials.

297 3.2. Analytical Results

Results of field and laboratory analyses are summarized in Figure 3 where samples are 298 classified based on their pH: circumneutral (pH 7-9), intermediate (pH 9-11), and 299 hyperalkaline (pH > 11). Fluids sampled from wells are indicated by the solid black squares 300 regardless of their pH. Samples from Masibt, WDA-16, and fluids from previous studies 301 associated with gabbroic rocks were not included in Figure 3 as simulations on gabbro-water 302 interactions were not conducted in this study. Data from gabbro-hosted fluids are plotted in 303 Figures S7 and S8 and preliminary discussions on their similarities and differences with 304 305 ultramafic-hosted fluids can be found in the supplementary files accompanying this work.

306 Ultramafic-hosted fluids sampled during this study have pH values ranging from 7.2 to 307 11.9 and conductivities from 450 to 4050 μ S/cm. Circumneutral to slightly alkaline fluids are 308 enriched in dissolved O₂ and DIC (Figures 3a and 3b). The dissolved oxygen concentration 309 measured from surface and mixed fluids is constrained by the solubility of atmospheric O_{2(g)}, 310 which at ambient conditions (25°C) is close to 275 μ molal. On the other hand, dissolved O₂ 311 concentrations of hyperalkaline fluids vary from low concentrations (~10 μ molal) to values close



312

Figure 3. Measured dissolved O₂ (a) and conductivity (b), as well as concentrations of total
dissolved inorganic carbon or DIC (c), Na (d), K (e), Cl (f), Ca (g), Mg (h), and Si (i) plotted
against pH. Green and blue squares indicate stream and hyperalkaline spring fluids, respectively.
Orange squares indicate where fluids resulting from significant mixing between these endmember fluids was physically observed. Black squares show fluids sampled from wells
regardless of their pH. Various symbols in grey are values reported in the literature. Dashed grey
line in (a) depicts the solubility of O₂ at current atmospheric compositions.

321	to those measured from circumneutral fluids. This suggests that some of the deep-seated
322	hyperalkaline fluids interact with the atmosphere either during ascent or upon sampling.
323	Hyperalkaline fluids are generally more conductive (Figure 3c) due to elevated total
324	concentrations of Na, Cl, K, and Ca (Figures 3d –3g). Aside from these species, OH ⁻ also
325	contributes to the high conductivities measured in basic solutions. On the other hand, these high-
326	pH fluids are strongly depleted in total dissolved Mg (Figure 3h) and Si (Figure 3i) by
327	approximately 2–3 orders of magnitude relative to circumneutral fluids. Speciation of dissolved
328	solutes are depicted in Figure S2. As shown in Figure S2a, the most common solutes in
329	circumneutral fluids are Mg^{+2} and HCO_3^- . Hyperalkaline fluids are mostly composed of Na^+ and
330	Cl ⁻ , followed by OH ⁻ and Ca ⁺² . As depicted in Figure S2b, most of the dissolved Si in
331	circumneutral fluids exists as $SiO_{2(aq)}$, while the anion $HSiO_3^-$ tends to dominate in hyperalkaline
332	solutions followed by the neutral complex, $NaHSiO_{3(aq)}$. The dominant Mg and Ca species at all
333	pH values of the investigated fluids are Mg^{+2} (Figure S2c) and Ca^{+2} (Figure S2d), respectively.
334	Following Mg^{+2} , the MgOH ⁺ complex can comprise one-third of the total dissolved Mg in
335	hyperalkaline solutions. The dominant DIC component in circumneutral solutions is HCO ₃ ⁻ ,
336	while both CO_3^{-2} and the neutral complex $CaCO_{3(aq)}$ are predominant in hyperalkaline fluids
337	(Figure S2e).

Hyperalkaline fluids sampled from sites close to the basal thrust of the ophiolite (Falaij, Al Bana, Al Hilayw) have higher concentrations, by at most a factor of two, of total dissolved Na, K, and Cl than fluids sampled from other sites (see Figure S3). Results of Mann-Whitney Utest for equal medians show that hyperalkaline fluids sampled from both geological settings are significantly different from each other at p < 0.01 (see Table S3 and the supplementary materials for more details on the statistical analysis). We offer possible origins for the elevated

344	concentrations of Na, Cl, and K of samples collected near the basal thrust in the discussion
345	below. On the other hand, pH and total dissolved Mg, Si, Ca, and inorganic carbon are
346	statistically similar (see Table S3) among high pH samples retrieved from all ultramafic-hosted
347	sites despite their variable geological settings (basal thrust, near or far from the ultramafic-
348	gabbro contact). This suggests similar underlying reactions that constrain these concentrations.
349	We expound further on these reactions in the following section.
350	Stable isotopic data, δ^{18} O vs δ^{2} H, of H ₂ O for fluids sampled in this study are shown in
351	Figure 4, together with relevant local meteoric water lines determined by Weyhenmeyer et al.
352	(2002) for moisture sources coming from the north (Mediterranean, LMWL-N) and the south
353	(Indian Ocean, LMWL-S). Note that fluids hosted in the Samail ophiolite plot between the two
354	meteoric water lines. The Mediterranean Sea is the dominant source of precipitation in modern
355	Oman while the southern moisture source is much rarer and comprised of cyclones and
356	depressions generated in the Indian Ocean that only occasionally reach the northern parts of
357	Oman where the ophiolite is located (Weyhenmeyer et al., 2002). However, Paukert et al. (2019)
358	showed that fluids sampled from the Samail ophiolite are too enriched in ¹⁸ O and/or depleted in
359	² H relative to calculated Raleigh distillation curves that model evaporation from the northern
360	Mediterranean source. Our dataset extends that shown by Paukert et al. (2019) and further
361	supports conclusions by previous investigators that fluids hosted in the Samail ophiolite are
362	likely recharged by a combination of northern and southern sources (Weyhenmeyer et al., 2002;
363	Matter et al., 2005; Paukert et al., 2019). Through noble gas geothermometry, Paukert et al.
364	(2019) also showed that some hyperalkaline fluids in Oman were derived from fluids recharged
365	at low temperatures during the last glacial age where the dominant moisture source was likely
366	coming from the south.



367

Figure 4. δ^{18} O vs δ^{2} H crossplot. Filled circles and squares represent well fluids and hyperalkaline spring fluids, respectively. Colors indicate sample sites of hyperalkaline and well sites (sites in legend are arranged from southeast to northwest in location). Samples in open green squares represents stream samples regardless of location. Local meteoric water lines are plotted based on the southern (LMWL-S) and northern (LMWL-N) moisture sources for Oman (Weyhenmeyer et al., 2002). The grey dashed line depicts the Global Meteoric Water Line (GWML).

Isotopic data from circumneutral to slightly alkaline fluids and hyperalkaline fluids 376 overlap, though some of the latter samples are more depleted in ¹⁸O and ²H. In addition, the ¹⁸O 377 and ²H compositions of hyperalkaline fluids seem to exhibit geographical trends, that is, fluids 378 from the Miskin Massif (Shumayt and Al Bana) located in the central part of the Samail ophiolite 379 are more depleted in both ²H and ¹⁸O relative to those sampled at the Bahla (Wells WDA-05, 16 380 and 17) and Wadi Tayin (Falaij, Dima, and Qafifah) Massifs, which is further south. Fluids from 381 the northern massifs Wuqbah (Uqaybah) and Hilti (Sudari) and fluids sampled from the northern 382 portions of the Samail Massif (Masibt, and wells SJA-03 and 04) are depleted in ²H relative to 383 the other samples. Fluids sampled from Al Hilayw, located in the southern portion of the Samail 384

Massif, have isotopic values that plot closely with those from a nearby site from the Wadi Tayin 385 386 Massif (Falaij). Results of analysis of similarities (ANOSIM) calculations show that many sites have isotopic compositions that are not significantly distinct, though the water isotopic 387 compositions of a given site can be significantly different from another site that is further away 388 (at p < 0.05, see Table S4). Results of statistical analyses indicate that samples from Al Bana and 389 390 Shumayt are the most isotopically unique. However, variabilities in the isotopic compositions from a given study site, especially those trending toward heavier ¹⁸O and ²H, can be accounted 391 392 for by variable degrees of evaporation and can result in compositions that can overlap with those 393 from another study site. Evaporation can thus overprint a much more distinct geographical disparity that can be informed by future isotopic and hydrologic investigations. 394

395

4. Reconciling Predictions with Reality

In the following section, compositions measured from environmental samples are 396 397 compared with those predicted from equilibrium calculations. Deviations from predictions may 398 occur and are likely to be consequences of slower approaches to equilibrium due to kinetic inhibition at low temperatures. Furthermore, several transport processes such as leaching from 399 minerals, the input of infiltrating atmospheric $CO_{2(g)}$, and shallow groundwater mixing with 400 401 surfacing deep-seated fluids can shift compositions away from those predicted by equilibrium models. We simulate these transport processes, investigate their effect on fluid compositions, and 402 403 test if these processes can account for natural observations that deviate from equilibrium 404 predictions. We first discuss trends involving pH and dissolved Si, Mg, Ca, and inorganic carbon 405 and then proceed to discuss those for dissolved Na and Cl.

406 4.1. pH and dissolved Si, Mg, Ca, and inorganic carbon

407	Thermodynamic simulations, depicted in Figure S4, predict trends in the dissolved
408	concentrations of Si, Mg, Ca, and inorganic carbon as an outcome in the progress of the
409	subsurface serpentinization reactions. Comparison of predictions with analytical measurements
410	for Si, Mg, Ca, and DIC are shown in pH vs. concentration plots in Figures 5a-5d and on activity
411	diagrams representing the MgO-SiO ₂ -H ₂ O (Figure 5e) and MgO-CaO-SiO ₂ -H ₂ O (Figure 5f)
412	systems. Overall, compositions of fluids measured from environmental samples are generally
413	inconsistent with model results, as most fluid samples do not coincide with predicted reaction
414	paths (solid dark blue curves in Figure 5). Trends from environmental samples show that fluids
415	evolve from relatively dilute rainwater into fluids more concentrated in Si, Mg, and DIC (mainly
416	speciated as HCO_3^{-}) at pH 7 to 9. Compositions of fluids then evolve into hyperalkaline fluids
417	(pH > 11) with low concentrations of Si (Figure 5a) and Mg (Figure 5b). At these highly basic pH
418	values, sampled fluids have high Ca (Figure 5c) and low DIC (Figure 5d) concentrations. The
419	reaction paths generally follow similar trajectories but at different solute concentrations than
420	those measured from the environmental samples. A few environmental samples are consistent
421	with equilibrium predictions. As an example, measured compositions of hyperalkaline fluids
422	seem to be bounded by equilibrium constraints as shown by the end of reaction paths depicted in
423	Figure 5. Specifically, the most depleted Si (Figure 5a) and Mg (Figure 5b) values measured
424	from end-member hyperalkaline fluids in Oman seem to be constrained by equilibrium fluid-
425	mineral reactions (end of the dark blue curves in Figures 5a and 5b), as discussed below. Except
426	for these few cases, most environmental samples cannot be accounted for by equilibrium
427	calculations alone. Hence, we discuss fluid mixing (dashed yellow lines) as a process to account
428	for variations in samples. Compositions of end-member fluids used in our mixing calculations



Figure 5. Comparison of results from reaction path and mineral solubility calculations with total 430 431 dissolved concentrations of Si (a), Mg (b), Ca (c), and dissolved inorganic carbon (DIC) (d) of samples as well as calculated activities from environmental measurements plotted in activity 432 433 diagrams for the MgO-SiO₂-H₂O (e) and CaO-MgO-SiO₂-H₂O (f) systems at ambient conditions $(25^{\circ}C, 1 \text{ bar})$. Symbols for environmental samples are similar to those in Figure 3, with the 434 addition of rainwater data from Oman and surrounding areas (yellow cross symbols, Ahmed et 435 al., 2000; Matter et al., 2006). Solid dark blue curves represent reaction paths. Dashed grey lines 436 in (a) to (f) show calculated saturation conditions for indicated minerals or mineral assemblages 437 while the solid black lines in (f) depict phase boundaries between minerals. Dashed red lines in 438 (d) depict calculated DIC at indicated $CO_{2(g)}$ fugacity values. Dashed grey line lines in (f) 439 represent calcite saturation at indicated DIC concentrations. Circles with text represent fluids 440 used in the calculations: rainwater (r), representative surface circumneutral fluid (s), 441 representative deep-seated fluids in equilibrium with Ctl-Brc-Di (d₁). Ctl-Brc-Cal at 10 µmolal 442 DIC (d₂), and 20 µmolal DIC (d₃). The latter four compositions (s, d₁, d₂, and d₃) were used as 443 end-member fluids in mixing calculations. Dashed yellow lines represent three mixing paths 444 between these end-member fluids (s and three deep endmembers: d_1 , d_2 , and d_3) while filled 445 vellow circles along these mixing lines in (a), (e), and (f) represent various mixing extents in %. 446 Upper, middle, and lower dashed yellow curves in (e) indicate mixing between fluid s and fluids 447 d_1 , d_2 , and d_3 , respectively. All three calculated deep-seated fluids (d_1 , d_2 , and d_3) are coincident 448 449 in the MgO-SiO₂-H₂O activity diagram and are hence just labelled as d in (e). Acronyms for minerals are based on the recommendations of Whitney and Evans (2010): Fo- forsterite, En-450 enstatite, Ctl- chrysotile, Brc- brucite, Tlc- talc, Oz- quartz, Di- diopside, Cal- calcite. 451

452

can be found in Table S6 and details of the calculations are expounded upon below. In additional terms of the calculations are expounded upon below.	ition,
--	--------

454 we also discuss dissolution of other minerals (*e.g.*, brucite, carbonates) aside from those modeled

455 above (olivine and pyroxene) that can account for some compositional trends observed in natural

samples. In the following discussions we first examine pH 7 to 9 fluids enriched in Mg and DIC

457 (green squares) and then proceed to discuss fluids with hyperalkaline (pH >11, blue squares) and

- 458 intermediate (pH 9 to 11, orange squares) pH. Ultimately, these discussions lead to a
- 459 classification scheme for these fluid types guided by a combination of constraints from fluid-
- 460 mineral equilibria and relevant transport processes.
- 461 4.1.1. pH 7–9, Mg⁺²-HCO₃⁻ Type 1 fluids

462 Fluids of this type, referred to as Type 1, mainly occur in streams (green squares in

463 Figure 5) and shallow groundwater (some black squares in Figure 5). These fluids are

464	characterized by neutral to slightly alkaline pH (pH 7–9) and elevated concentrations of DIC,
465	Mg, and Si relative to rainwater and hyperalkaline fluids. As shown in Figure S2a, the most
466	common solutes in these fluids are Mg^{+2} and HCO_3^- , and hence are also referred as Mg^{+2} - HCO_3^-
467	type fluid. In the model, Type 1 compositions are attained at early stages of the overall reaction
468	progress where solutes accumulate in the fluid through initial dissolution of primary minerals but
469	before Si, Mg, and DIC are consumed to form serpentine, carbonates, and brucite.
470	Concentrations of Si in these fluids are consistent with the maximum Si concentrations (Figure
471	5a) and $aSiO_2$ (Figure 5e) predicted by the reaction path calculations (black solid curves) when
472	fluids attain equilibrium with serpentine during the start of Stage 2 (see Figure S4d) of the
473	overall reaction progress. As depicted in Figure 5e, environmental samples rarely plot below the
474	chrysotile saturation line, implying constraints on the Si concentrations of ultramafic-hosted
475	fluids by serpentine precipitation. Most of the samples, however, are oversaturated with respect
476	to chrysotile (see Table S2), which implies a kinetic inhibition to serpentine precipitation.
477	Furthermore, most circumneutral to slightly alkaline fluids plot above quartz and talc saturation
478	(see Figure 5e), which suggests kinetic inhibition may apply to most silicates.
479	While there is close agreement between the equilibrium predictions and the measured Si
480	of Type 1 fluids, the agreement does not hold true for Mg, Ca, and DIC. The modeled
481	concentrations, while showing an increase in these constituents (solid dark blue curves in Figures
482	5b to 5d), do not increase enough to meet measured concentrations of Mg, Ca, and DIC. Cipolli

483 et al. (2004) showed that these elevated concentrations can be attained during reactions at higher

484 fCO_2 , up to two orders of magnitude higher (fCO_2 up to $10^{-1.5}$) than that imposed by the present

485 atmosphere. Results constrained by elevated $fCO_{2(g)}$ levels are indicated by the reaction paths

486 depicted in Figure 5d as dashed red curves that depict more dissolved CO_2 at higher $fCO_{2(g)}$

values at a given pH. Alternatively, instead of being the result of elevated $fCO_{2(g)}$, high 487 concentrations of Ca, Mg, and DIC observed in environmental samples in this pH range can also 488 489 be attained through interactions with minerals that are abundant in serpentinities or highly altered ultramafic rocks that comprise the shallow aquifer. As shown by the dashed grey curves in 490 Figures 5b and 5d, respectively, the lower bounds in the Mg and DIC concentrations of natural 491 492 fluids with pH values between 7 and 9 are close to those dictated by the solubility of magnesite. In addition, Mg concentrations of Type 1 fluids are between values predicted for fluids in 493 494 equilibrium with magnesite or brucite (Figure 5b), while the trend in Ca concentrations of the 495 natural samples follows the V-shaped trend of fluids in equilibrium with calcite (Figure 5c). Overall, it can be seen that the compositions of these circumneutral to slightly alkaline fluids are 496 more consistent with the dissolution and precipitation of minerals (serpentine, brucite, magnesite, 497 and calcite) common in the highly altered rocks that compose the shallow aquifer than they are 498 499 with initial stages of the dissolution of primary minerals in aquifers composed of fresh or 500 relatively less altered ultramafic rocks. This observation is consistent with Neal and Stanger (1986) and Marques et al. (2008), who argued that Mg- and HCO₃⁻-rich (Type 1) fluids could 501 have been generated in shallow aquifers consisting of highly altered ultramafic rocks. Fluids that 502 503 plot between the brucite and magnesite solubility curves in Figure 5b can be described by a process involving the dissolution of brucite and subsequent precipitation of magnesite given by, 504

$$Mg(OH)_{2(brucite)} + Mg^{+2} + 2HCO_3^{-} \rightarrow 2MgCO_{3(magnesite)} + 2H_2O.$$
 (1)

It is possible for reaction (1) to proceed readily at shallow aquifer settings, as brucite is highly
soluble at circumneutral pH (see dashed grey curve for brucite in Figure 5b) and is
experimentally known to dissolve rapidly at these pH conditions (Pokrovsky and Schott, 2004).
Dissolution of brucite likely accounts for the depleted Mg/Si ratios observed from some highly

509	altered ultramafic rocks relative to those that are less altered (Malvoism, 2015; de Obeso and
510	Kelemen, 2018; 2020), though these observations can likewise be accounted for by enrichment
511	in silica (de Obeso and Kelemen, 2018). It is of note that natural brucite commonly exists in
512	solid solutions of Mg^{+2} and Fe^{+2} , and the dissolution, mobilization, and eventual oxidation of
513	Fe ⁺² -bearing brucite could supply H ₂ independent of the H ₂ generated during the serpentinization
514	of fresher ultramafic rocks (Frost et al., 2013; Miller et al., 2016; Mayhew et al., 2018;
515	Templeton and Ellison, 2020). However, the production of H ₂ from the alteration of brucite-
516	bearing serpentinites can only proceed at O ₂ -free regions in the aquifer, as Fe ⁺² released from
517	brucite dissolution will be otherwise oxidized by O ₂ to form ferric oxyhydroxides.
518	4.1.2. Hyperalkaline (pH >11) Ca-OH Type 2 fluids
519	Hyperalkaline fluids were sampled from discharging springs (blue squares in Figure 5)
520	and one well (NSHQ-14). These are classified as Type 2 fluids and are characterized by their
521	hyperalkaline pH, elevated Ca concentrations, and extremely low Si and Mg concentrations. As
522	shown in Figure S2a, the most common solutes in these fluids, after Na ⁺ and Cl ⁻ , are Ca ⁺² and
523	OH ⁻ , and hence they are also referred to as Ca ⁺² -OH ⁻ fluids. Despite Na ⁺ and Cl ⁻ comprising
524	most of the total dissolved solutes, these fluids are far less saline than seawater and hence are not
525	often classified as Na ⁺ -Cl ⁻ fluids. Compositions of Type 2 fluids are characteristic of fluids
526	constrained by the chrysotile-brucite-diopside assemblage (Ctl-Brc-Di, see end of reaction path
527	in Figure S4e or reaction S7) predicted by equilibrium calculations. However, upon closer
528	examination, Clt-Brc-Di equilibrium (point d_1 in Figure 5) yields lower Si (Figure 5a) and Mg
529	(Figure 5b) and higher Ca (Figure 5c) concentrations than those measured from environmental
530	samples. Analogous comparisons can be made with the activity diagrams shown in Figures 5e
531	and 5f, where the $aSiO_{2(aq)}$ values of hyperalkaline samples approach that dictated by Ctl-Brc

532	equilibrium but plot below the Ctl-Brc-Di equilibrium value. We have yet to sample fluids with
533	Si concentrations or $aSiO_{2(aq)}$ lower than values dictated by Ctl-Brc equilibrium, implying
534	constraints by this mineral assemblage on the silica content of serpentinization-generated fluids.
535	Despite being close to or approaching predicted compositions, the variabilities in pH and
536	concentrations of dissolved Si and Mg observed for hyperalkaline fluids cannot be fully
537	accounted for by equilibrium predictions. Specifically, pH values of hyperalkaline fluids range
538	from 11 to 12, and at these pH values, Si and Mg concentrations can vary by up to three orders of
539	magnitude. Trends resulting from variabilities in host rock compositions shown by Leong and
540	Shock (2020) cannot account for such wide variations within this small range of pH. Accounting
541	for mineral solid solutions predicts slightly higher Si concentrations (Leong and Shock, 2020)
542	but does not fully reconcile models with actual measurements.
543	Reaction paths and solubility curves depicted in Figure S4 simulate rainwater-ultramafic
544	rock interactions at 25°C. However, temperatures of fluids during sampling varied from 15° to
545	40°C. We calculated the pH constrained by the Ctl-Brc-Di equilibrium at temperatures ranging
546	from 0–60°C and results are plotted in Figure 6 (solid black curve). Most environmental samples
547	from Oman, however, have pH values below those predicted by the Ctl-Brc-Di equilibrium. Only
548	a few samples from Oman (grey triangles and x-marks from Chavagnac et al., 2013b and
549	Giampouras et al., 2020, respectively) and a few from other low-temperature serpentinizing
550	environments on Earth (filled black circles in Figure 6) plot near this predicted trend. However,
551	it is of note that pH values higher than those predicted by Ctl-Brc-Di equilibrium are rare,
552	implying that upper limits on the pH of serpentinization-generated fluids are constrained by
553	diopside dissolution. Extremely high pH values (12.5 to >13) were observed in a few continental
554	sites such as in Maqarin, Jordan (Khourey et al., 1985, 1992; Alexander et al., 1992; Pederson et

al., 2004) and Lake Calumet, Chicago (Roadcap et al., 2005; Ohlsson et al., 2019). These
localities, however, are not hosted by ultramafic lithologies and the extremely alkaline pH values
are likely results of reactions akin to that of groundwater-concrete interactions (Gaucher and
Blanc, 2006).



559

560 Figure 6. Calculated pH in equilibrium with the Ctl-Brc-Di (solid black curve, reaction 7) and Ctl-Brc-Cal (solid grey curves, reactions 4 and 9) mineral assemblages at temperatures ranging 561 from 0 to 60°C. For the Ctl-Brc-Cal equilibrium assemblages, the DIC concentrations in µmolal 562 are indicated above the solid grey curves. Black circles represent the highest pH values measured 563 in low-temperature serpentinizing systems: SCS- South Chamorro Seamount in the Izu-Bonin-564 Mariana arc (Mottl, 2009); BA- Dinaride ophiolite in Bosnia and Herzegovina (Etiope et al., 565 566 2017); US - Coast Range ophiolites microbial observatory or CROMO, California, USA 567 (Crespo-Medina et al., 2014); CA – Tableland ophiolite, Canada (Szponar et al., 2013); IT – Voltri Massif, Italy (Schwarzenbach et al., 2013); TU – Amik Basin, Turkey (Yuce et al., 2014); 568 and JP – Hakuba Happo spring, Japan (Homma and Tsukahara, 2008; Suda et al., 2014). The pH 569 of these fluids trend closely with and rarely exceed the pH predicted by the Ctl-Brc-Di 570 equilibrium, suggesting that upper limits on the pH of serpentinizing fluids are constrained by 571 diopside dissolution. In contrast, most hyperalkaline (pH >11) fluids from Oman, represented by 572 573 the various grey symbols, plot below these theoretical maxima but above those predicted for by Ctl-Brc-Cal equilibrium with DIC concentrations of 20 µmolal. Grey square symbols depict data 574 resulting from this work, while those in triangles, diamonds, and crosses represent data reported 575 in Chavagnac et al. (2013b), Rempfert et al. (2017), and Giampouras et al. (2020), respectively. 576

Leong and Shock (2020) discuss various mineral-fluid reactions (e.g., andradite, calcite) 577 that can shift fluid compositions in equilibrium with Ctl-Brc-Di to lower $aCa^{+2}/(aH^{+})^{2}$ and hence 578 to lower pH and Ca concentrations. One such process is the introduction of DIC along the fluid 579 pathway, such as when ascending deep-seated hyperalkaline fluids are modified by infiltrating 580 atmospheric CO_{2(g)}, mix with DIC-rich shallow groundwaters, and/or come in contact with 581 582 carbonate veins in the shallow subsurface. Magnesite and dolomite are highly and slightly undersaturated, respectively, in these Mg-poor fluids (see Table S2) and will dissolve upon 583 584 interaction with surfacing deep-seated fluids. With these additional DIC inputs, Ca-rich 585 hyperalkaline fluids can precipitate calcite, driving fluid compositions away from Ctl-Brc-Di equilibrium to conditions set by chrysotile-brucite-calcite (Ctl-Brc-Cal) equilibrium. The pH 586 values and Ca concentrations constrained by Ctl-Brc-Cal equilibrium vary depending on the 587 amount of DIC supplied by the transport processes mentioned above. Equilibrium with calcite, as 588 shown through the reaction 589

$$Ca^{+2} + HCO_3^{-} \Leftrightarrow CaCO_{3(calcite)} + H^+,$$
(2)

dictates that a higher amount of DIC will result in lower $aCa^{+2}/(aH^{+})^{2}$. Calculated $aCa^{+2}/(aH^{+})^{2}$ 590 values at variable DIC concentrations up to 50 µmolal are plotted in the activity diagram shown 591 in Figure 5f. At 25°C and 1 bar, DIC concentrations exceeding 8 μ molal result in $aCa^{+2}/(aH^{+})^{2}$ 592 values lower than those constrained by Ctl-Brc-Di equilibrium. DIC concentrations of <20 593 µmolal can account for the $aCa^{+2}/(aH^{+})^{2}$ of most hyperalkaline samples. Results of calculations 594 of pH constrained by the Ctl-Brc-Cal equilibrium accounting for both variable temperatures (0-595 60°C) and DIC concentrations (10–50 µmolal) are plotted in Figure 6 (solid grey curves). Most 596 597 hyperalkaline fluids in Oman have pH values close to those predicted by equilibrium with calcite 598 at DIC concentrations close to 10 µmolal and not more than 20 µmolal, suggesting a DIC-limited

environment in the deep subsurface. Minor DIC input from the atmosphere or through mixing 599 600 with DIC-rich shallow groundwater can easily saturate calcite and consequently shift the pH of surfacing deep-seated hyperalkaline fluids. Calcite and dolomite were observed in active spring 601 sites in Oman (Kelemen and Matter, 2008) and are evidence that this shift could be actively 602 occurring in the shallow subsurface beneath hyperalkaline discharge zones. Moreover, the 603 604 extremely low DIC (<20 µmolal) predicted for the subsurface can inform biogeochemical studies on the amount of inorganic carbon available for subsurface microbes. In these carbon-limited 605 606 environments, dissolution of carbonates can be a robust inorganic carbon source for deep-seated 607 microbes (Miller et al., 2018), and investigations into the subsurface occurrence of these minerals provide insights into how deep the biosphere extends into ultramafic aquifers. 608 4.1.3. Mixed Fluids 609

The Si and Mg concentrations of hyperalkaline fluids vary by nearly three orders of 610 magnitude and cannot be fully accounted for by a shift toward Ctl-Brc-Cal equilibrium or by 611 612 variabilities in temperature. At temperatures between 0-100 °C, predicted total dissolved Si only varies by an order of magnitude and Mg varies even less (Leong and Shock, 2020). Leong and 613 Shock (2020) showed that reaction with more orthopyroxene-rich or more serpentinized 614 615 ultramafic rocks can lead to higher Si concentrations. However, the same models also predict lower trends in Mg concentrations. Instead, variabilities observed in almost all sampled fluids 616 617 can be accounted for by mixing (dashed yellow lines in Figure 5) between deep-seated fluids 618 close to equilibrium with the rock as represented by Ctl-Brc-Di (d₁) or Ctl-Brc-Cal (d₂ and d₃) 619 equilibria and a representative surface fluid (s, Sample 140116B). Further details on the 620 composition of end-member fluids used in our mixing calculations can be found in Table S6.

621	Deep-seated hyperalkaline springs often discharge where neutral to slightly alkaline
622	shallow aquifer fluids discharge as well. Thus, mixing between these two fluid types at discharge
623	zones is likely to occur. As surficial and shallow subsurficial fluids have Si and Mg
624	concentrations that are several orders of magnitude elevated relative to deep-seated hyperalkaline
625	fluids, even minor amounts of mixing can lead to large variabilities in the Si and Mg
626	concentrations. Mixing calculations reveal that a majority of the deep-seated hyperalkaline fluids
627	samples were modified slightly by an input of <10% of surficial and/or shallow surficial fluids
628	(see Figure 5a). Some samples have less influence (<1%) and can be interpreted as the most
629	pristine representatives of the deep subsurface. Using tritium (³ H) analysis, Paukert et al. (2019)
630	suggested that hyperalkaline spring fluids in Oman are influenced by up to 6% modern
631	groundwater. This is consistent with our calculated mixing extents for most hyperalkaline fluids
632	(<10%, see Figure 5a).

We use Si concentrations to quantify mixing extents (Figure 5a) because we have 633 constraints on the Si concentration of at least one of the end-member fluids, and it allows us to 634 take advantage of the highly contrasting Si concentrations of end-member fluid types and the 635 generally conservative nature of Si during mixing. In studies involving submarine hydrothermal 636 637 fluids, the extent of seawater input to venting fluids can be quantified using Na, Cl, and Mg owing to known constraints on end-member concentrations. However, unlike seawater where Na 638 639 and Cl concentrations are known, end-member Na and Cl concentrations are more variable in 640 continental serpentinizing environments. In the Samail ophiolite, end-member hyperalkaline fluids have Na and Cl concentrations that vary among sample sites, with elevated concentrations 641 642 observed at sites closest to the basal thrust of the ophiolite. Si behaves more conservatively than 643 Mg, as Mg-bearing minerals such as brucite and layered doubled hydroxides are commonly

644	observed in hyperalkaline pools (Neal and Stanger, 1984; Paukert et al., 2012; Chavagnac et al.,
645	2013a; Giampouras et al., 2020) and in shallow aquifers (Taylor et al., 1991). Most Type 2
646	hyperalkaline fluids have $aMg^{+2}/(aH^{+})^{2}$ values that lie above the brucite solubility line, and few
647	cross below it toward the chrysotile saturation line, as shown in Figure 5e.
648	The Mg vs Si concentrations of samples are plotted in Figure S5, along with the
649	calculated mixing pathway depicted in Figure 5. Trends shown in Figure S5 suggest that brucite
650	precipitation at variable pH values above 10, where brucite can be stable or metastable, can
651	cause Mg concentrations to decrease from concentrations along the mixing pathway toward those
652	dictated by the saturation of brucite. In addition, as shown in Figure S5, the Si concentrations of
653	fluids does not decrease to values lower than those depicted along the mixing curves. Despite the
654	thermodynamic drive to precipitate, most fluids remain oversaturated with respect to chrysotile.
655	Tutolo et al. (2018) postulated kinetic inhibition for serpentine precipitation (see reaction S2) or
656	the brucite silicification reaction (see reaction S4), resulting in Si concentrations that are
657	persistently above those dictated by chrysotile solubility (dashed grey lines in Figures 5a and 5e).
658	Mixing can transport Si into a Si-poor deep-seated fluid, and the sluggish rate of serpentine
659	formation makes Si a reliable mixing indicator in comparison to other aqueous species. Though
660	rare, secondary silicate minerals such as suolunite (Ca ₂ Si ₂ O ₅ (OH) ₂ :H ₂ O) are known to precipitate
661	in outcrops (Stanger and Neal, 1984) and hyperalkaline pools (Chavagnac et al., 2013a) in
662	Oman, and determining the rate of Si incorporation into solids at low-temperature conditions will
663	help improve its use as a tracer for mixing in serpentinizing systems. Another advantage of using
664	Si is the ease of measurement of even dilute concentrations (<1 ppm $SiO_{2(aq)}$) in the field using a
665	colorimetric spectrometer. Field measurements can quantify mixing extents during sampling and
666	exploration. Mixing calculations and field measurements of Si concentrations provide a rapid
and robust method to quantify how much the deep-seated signature of serpentinization-generatedfluids is affected by shallow subsurface processes.

As a result of mixing, most fluids with pH values greater than 9 have DIC concentrations 669 that imply supersaturation with respect to magnesite and calcite (Figure 5d) and precipitation of 670 these minerals can be favorable. However, at pH > 11 calculated DIC levels in equilibrium with 671 672 magnesite (dashed grey line in Figure 5d) would require Mg values (dashed grey line in Figure 5b) that exceed those measured in our samples, resulting in conditions that are not favorable for 673 674 precipitation of magnesite (see Table S2). Thus, only calcite (or aragonite, see Table S2) is 675 favored to precipitate in hyperalkaline environments, consistent with studies on travertine and other precipitates associated with hyperalkaline seeps (e.g., Kelemen and Matter, 2008; 676 Chavagnac et al., 2013a; Mervine et al., 2014; Giampouras et al., 2020). At pH values between 9 677 and 11, Mg (Figure 5b) and DIC (Figure 5d) concentrations of sampled fluids plot above those 678 679 dictated by magnesite solubility, indicating that magnesite is favored to precipitate in these 680 fluids. Their lack of occurrence in spring sites, however, suggests that there are kinetic inhibitions to precipitate magnesite in surficial mixing environments. These kinetic inhibitions 681 could lead to the precipitation of hydrated magnesium carbonates (e.g., hydromagnesite, 682 683 dypingite, nesquehonite) that are known to occur in hyperalkaline and mixed fluids in Oman (Giampouras et al., 2020) instead of magnesite. Our mixing calculations also highlight the strong 684 685 buffering capacity of hyperalkaline fluids resulting from mineral-fluid reactions that generate 686 Ca-OH solutions (Ctl-Brc-Di equilibrium, or Ctl-Brc-Cal equilibrium at low DIC concentrations 687 of $<20 \mu$ molal). A significant amount of mixing is required to lower the pH below 11, amounting 688 to 40% surface/shallow water input if pH is constrained by Ctl-Brc-Cal equilibrium at 10 µmolal 689 DIC or up to 80% if constrained to higher pH by Ctl-Brc-Di equilibrium.

690	In addition to fluid mixing, the elevated Si and Mg concentrations relative to those
691	predicted by equilibrium calculations could also result from olivine and pyroxene dissolving
692	faster than serpentine precipitating, as argued by Nesbitt and Bricker (1978). As shown in Figure
693	5e, most hyperalkaline fluids are undersaturated with respect to forsterite and plot between the
694	forsterite and brucite saturation lines. Observations from environmental samples likely reflect
695	complementary primary mineral dissolution and secondary mineral precipitation processes.
696	Quantifying how much the elevated Si and Mg concentrations of hyperalkaline fluids can be
697	attributed to either or both dissolution and mixing processes remains to be solved.
698	4.1.4. pH 9–11, low Si fluids
699	Most samples with pH ranging from 9 to 11 can be attributed to mixing of end-member
700	fluids with the exception of a few samples with dilute Si concentrations (<10 μ molal) that are
701	comparable to those measured in hyperalkaline fluids. These unusually low concentrations of Si
702	at these intermediate pH ranges can be attributed to various processes discussed below.
703	As depicted in Figure 5a, fluids with pH values from 9.0 to 9.5 plot close to the predicted
704	reaction path controlled by equilibria with both chrysotile and magnesite, which signifies fluid
705	compositions indicative of incipient serpentinization (Stage 2) and mineral carbonation (Stage 3)
706	stages of the overall alteration progress. Most fluids of this type were sampled in wells, depicted
707	in Figure 5a as solid black squares (this study) and grey diamonds (Rempfert et al., 2017),
708	suggesting that these fluids could be representative of intermediate stages of the overall
709	serpentinization progress where shallow aquifer fluids are infiltrating into the deep subsurface in
710	the recharge zone. Actual Si concentrations of sampled well fluids could be lower and plot closer
711	to values predicted by equilibrium calculations if upper aquifer fluids, which are Si-rich,
712	contaminate deep-seated samples during sampling of wells.

713	Alternatively, these fluid compositions could indicate an alteration process involving
714	gabbroic rocks, as many of these fluids were sampled in wells located close to the ultramafic-
715	gabbro contact in the Samail ophiolite. Models simulating water interacting with gabbro or a
716	mixture of gabbroic and ultramafic rocks at low temperatures could assess if these fluids
717	correspond closely with predictions. However, many of the hyperalkaline spring fluids depicted
718	in Figure 5a are also located close to the ultramafic-gabbro transition and do not exhibit any
719	compositional differences from fluids located far from this transition. Like the springs, the
720	composition of these well fluids may be more strongly influenced by fluid-mineral processes
721	associated with ultramafic rocks than with gabbroic rocks.
722	4.1.5. Classification of ultramafic-hosted fluids
723	The discussions above provide a framework for classifying low-temperature, ultramafic-
724	hosted fluids as constrained by predictions of the serpentinization reaction path, the solubility of
725	serpentinization-relevant minerals, and the subsequent mixing process that occurs as fluids are
726	discharged back to the surface. Specifically, by drawing on trends in pH and total dissolved Si,
727	as shown in Figure 7, we can classify these fluids into four types (dotted red fields in Figure 7):
728	<i>Type 1 Mg</i> ⁺² - <i>HCO</i> ₃ ⁻ – As first proposed by Barnes and O'Neill (1969), these fluids have
729	compositions that are enriched in Mg and bicarbonate, and could be products of the first stage of
730	the alteration progress that leads to enrichment of solutes in fluids through the dissolution of
731	primary minerals in systems open to atmospheric influences (Leong and Shock, 2020). However,
732	as discussed above, these fluids are likely to result from interactions of rain with serpentinites or
733	highly altered ultramafic rocks that are common in the shallow aquifer. Equilibria with
734	serpentine, brucite, and magnesite in the presence of the atmosphere results in fluids with pH
735	values between 7 and 9 that are enriched in Si (~200–500 μ molal, mostly speciated as SiO _{2(aq)}),

736	as depicted in Figure 7. Furthermore, as shown in Figures 5b and 5d, these fluids are enriched
737	relative to other fluid types in Mg (~1000–10,000 μ molal, mostly speciated as Mg ⁺² , see Figure
738	S2c) and DIC (2,000–5,000 μ molal, almost entirely speciated as HCO ₃ ⁻ , Figure S2e),

respectively.

Type 2 Ca^{+2} -OH⁻ – This fluid type, also first proposed by Barnes and O'Neill (1969), is 740 hyperalkaline and otherwise known as Ca⁺²-OH⁻-type fluids. Type 2 fluids are characterized by 741 their high pH (>11), elevated concentrations of Ca (>1,000 μ molal, mostly speciated as Ca⁺², 742 Figure S2d), and very low concentrations of Mg and Si (<3 µmolal), which are consequences of 743 744 equilibria with Ctl-Brc-Di or with Ctl-Brc-Cal at limited DIC concentrations (<20 µmolal) during advanced stages of serpentinization (Stages 4 to 6). As shown in Figure 7, a few 745 hyperalkaline samples have Si concentrations that are $<3 \mu$ molal ($<180 \text{ ppb SiO}_2$) and can be 746 interpreted to be close to end-member Type 2 compositions (<1% mixing). Most other 747 hyperalkaline fluids, however, have higher Si concentrations and can be accounted for by greater 748 749 degrees of fluid mixing.

Mixed Fluids – In this classification scheme, this fluid type encompasses fluids with pH 750 >9 that are too alkaline to be in equilibrium with serpentine, brucite, and carbonates in the 751 752 presence of the atmosphere. Note that fluids with pH <9 can also be influenced by mixing, as most fluids sampled in this study plot along the mixing trend shown in Figure 7. At higher pH, 753 754 this fluid type also includes hyperalkaline fluids that are Si-enriched compared with serpentine + 755 brucite equilibrium. Mixing between circumneutral stream fluids and hyperalkaline springs is 756 commonly observed at the surface, as they often occur together. On the other hand, it is likely 757 that subsurface mixing between end-member Type 1 and 2 fluids in discharge zones accounts for 758 the large variabilities in Si concentration observed in hyperalkaline fluids. The mixing trend

759	shown in Figure 7 depicts that between end-member Type 1 and 2 fluids. However, mixing can
760	also occur between any fluid type (e.g., between one end-member or mixed fluids with
761	intermediate fluids) that can account for fluids outside the dotted red fields in Figure 7 that
762	represent the four different fluid types discussed in this work.
763	Intermediate Fluids – These fluids, also with pH values ranging from 9 to 11, share
764	similar characteristics with mixed fluids. However, unlike mixed fluids, these fluids have low Si
765	concentrations (<10 µmolal) that are comparable to most hyperalkaline fluids. As discussed
766	above, fluids of this composition may indicate incipient serpentinization (Stage 2) and mineral
767	carbonation (Stage 3) during intermediate stages of the alteration progress.
768	Multivariate statistical analyses (Non-metric Multidimensional Scaling/NMDS and
769	analysis of similarities/ANOSIM) were conducted to evaluate if the overall major element
770	compositions of these four fluid types are significantly different from each other (see
771	supplementary materials for details on statistical methods). Results of NMDS ordination are
772	shown in Figure S6 and depict distinct clustering of circumneutral (Type 1), intermediate, and
773	hyperalkaline (Type 2) fluids. Mixed fluids, especially those with hyperalkaline pH, mostly
774	overlap with end-member Type 2 fluids and indicate that overall compositions of mixed fluids
775	(not just their pH and Si concentrations) are indistinguishable from end-member hyperalkaline
776	solutions. Results of ANOSIM calculations, shown in Table S5, reveal that compositions of
777	intermediate and mixed fluids, despite the overlapping pH, are statistically significant at $p < 0.05$.
778	Mixed and Type 2 fluids, however, are not significantly different at $p < 0.05$. At hyperalkaline
779	conditions, precipitation of brucite and calcite can be fast and drive the Mg and Ca
780	concentrations, respectively, of mixed and Type 2 fluids to values dictated by the saturation
781	curves of these minerals (see Figures 5b and 5c). In addition, ANOSIM results show that the

782	overall compositions of circumneutral Type 1 and intermediate fluids are not statistically
783	significant at $p < 0.05$ (see Table S5). While the pH and Si concentrations of intermediate fluids
784	are distinct from those of Type 1 fluids, their Mg values are not. As shown in Figure 5b, the Mg
785	concentrations of intermediate fluids (black symbols with pH between 9 and 10) plot close to
786	those dictated by the solubility of brucite. At this pH range, brucite is unstable and dissolution is
787	rapid (Pokrovsky and Schott, 2004), more than 2 and 4 orders of magnitude faster, respectively,
788	than dissolution of olivine (Pokrovsky and Schott, 2000) and chrysotile (Thom et al., 2013). The
789	rapid dissolution of brucite is likely to bring the Mg concentrations of both Type 1 and
790	intermediate fluids to similar values. Both fluid types have the potential to precipitate magnesite,
791	and brucite could have provided additional Mg for mineral carbonation aside from that derived
792	from the primary minerals.

793 Overall, fluid interaction with both the unaltered and altered aquifer host rocks would result in a diverse composition of fluids that can deviate from equilibrium expectations. 794 However, note that some fluids have Si concentrations that track along the equilibrium reaction 795 path depicting progress of the overall serpentinization reactions, and that the kinetic barriers for 796 precipitation of Si-bearing minerals during mixing (as opposed to brucite and calcite) can 797 798 provide information on the overall mixing progress. The trends shown in Figure 7 can allow immediate and robust classification of fluids while in the field using a pH meter and a portable 799 spectrometer for colorimetric measurements of dissolved SiO_{2(aq)}. These data provide real-time 800 801 information on the extent of the serpentinization and mixing progress and can concurrently feed into decisions involving exploration, sampling, and experiments while in the field. 802



803

Figure 7. Key serpentinization reactions and resulting fluid types summarized in a plot of Si 804 concentration vs pH. Fluid samples from the Oman ophiolite analyzed in this work (square 805 symbols) as well as well fluids (black diamonds) from Rempfert et al. (2017) are plotted relative 806 to the reaction and mixing paths. Analogous to Figures 3 and 5, colored square symbols depict 807 surface samples where green, orange, and blue symbols represent samples with pH values of <9, 808 between 9 and 11, and >11, respectively. Black squares and diamonds indicate subsurface 809 samples (wells) regardless of their pH. Overall, fluids can be accounted for by four processes: (1) 810 mineral dissolution into rain which leads to Type 1 fluids (see Stage 1), (2) incipient 811 serpentinization and mineral carbonation (see Stages 2-3) which leads to intermediate (Int) fluid 812 types, (3) advanced stages of serpentinization (see Stages 4–6) which lead to Type 2 fluids, and 813 (4) mixing (Mix) between deep-seated Type 2 and surficial or near-surficial Type 1 fluids. 814 Dotted red lines depict suggested pH and dissolved Si values that characterized these four 815 different fluid types. Note that many hyperalkaline fluids (pH >11) are classified as mixed fluids 816

817 in this scheme.

818 4.2. Dissolved Na and Cl

Petrological studies of serpentinized ultramafic rocks reveal that fluid-mobile elements can be sequestered during serpentinization in submarine and subduction environments (Scambelluri et al., 2001; Sharp and Barnes, 2004; Deschamps et al., 2013). The behavior of these elements during serpentinization in continents, where water-rock interactions are facilitated by meteoric fluids instead of seawater, is less well known. Trends in the Na and Cl concentrations of fluids from Oman were investigated to provide insights into their behavior during continental serpentinization.

826 Hyperalkaline fluids are at least twice as enriched in Na (Figure 8a) and Cl (Figure 8b) relative to circumneutral to slightly alkaline fluids and are orders of magnitude enriched relative 827 to rainwater. Similar trends can be observed for K (Figure 3e). The relative enrichment in the Na, 828 K, and Cl content of hyperalkaline fluids can result from two processes: (1) enrichment due to 829 loss of H₂O during the hydration of ultramafic rocks, and/or (2) leaching from host rocks during 830 water-rock interactions. Both can occur concomitantly, as hyperalkaline fluids are products of 831 large extents of water-rock interactions that lead to both water loss and possibly substantial 832 inputs from the reacting rocks. Moreover, the fluids most enriched in Na, K, and Cl are 833 834 hyperalkaline fluids sampled close to the basal thrust of the Samail ophiolite. These sample locations are typically located at topographic lows which implies longer fluid pathways that lead 835 836 to more hydration and leaching. Simulations of H₂O loss and leaching were performed to 837 evaluate the influence of hydration and leaching processes in accounting for natural 838 concentrations. The amount of H₂O lost during the overall reaction progress is shown in Figure 839 8c. At log rock-to-water ratio ~0 (*i.e.*, reaction of 1 kg of rock in 1 kg of water), usually inferred 840 as the maximum extent of reactions from deep-sea hydrothermal vent fluids (Taylor, 1974;



842 843 844 845 846 847 848 849 850 851 852 853 854	Figure 8. Reaction paths (dark blue curves) depicting predicted trends in the concentration of dissolved Na (a) and Cl (b) due to mineral leaching and amount of H ₂ O loss (c) as serpentinization progresses. Measured dissolved Na (a) and Cl (b) from environmental samples are plotted for comparison. Symbols in (a) and (b) are the same as those in Figures 3, 5 and 7. Reaction paths start from rainwater (r) and ends at fluids in equilibrium with either the Ctl-Brc-Di (d ₁) or the Ctl-Brc-Cal at 10 µmolal DIC (d ₂) assemblages. Solid white squares (with bold black outline) indicate extents of water-rock interactions with values of rock-to-water ratios given at the right. The dashed red curve in (b) depicts results of simulations with a serpentinite (100% chrysotile or serpentine). Solid red squares in (b) indicate extents of reactions with values of rock-to-water ratios given at the left. Dark blue curve in (c) represent reaction path for model ultramafic rock with composition 85% olivine-14% orthopyroxene-1% clinopyroxene. Grey curves in (c) represent trends for olivine-only or orthopyroxene-only scenarios. Dashed red lines in (c) show extent of reactions required to remove 15% and 50% of the starting reacting fluid.
855	
856	Edmond et al., 1979; Foustoukos et al., 2008), the loss of H_2O due to hydration is ~15%. It
857	follows that mineral hydration cannot solely account for the enrichment of Na and Cl observed in
858	sampled fluids. Accounting for an enrichment of at least a factor of two observed from
859	hyperalkaline fluids relative to lower-pH samples requires H_2O loss of >50%, which can only be
860	attained at extremely high rock-to-water ratios (log rock-to-water ratio of >0.5 , or reaction of >3
861	kg of rock in 1 kg of water). Similarly high extents of water-rock interactions are needed
862	regardless of compositional variabilities of the reacting rock (grey curves in Figure 8c).
863	Serpentinization of olivine-rich rocks tend to consume more H ₂ O as it precipitates more brucite,
864	which contains ~ 30 wt.% H ₂ O as hydroxide in its structure. Increased contributions from
865	orthopyroxene favors the formation of serpentine (~13 wt.% H_2O) and talc (~5 wt.% H_2O) which
866	are less hydrous than brucite and would require higher degrees of interaction to result in similar
867	amounts of H ₂ O loss.
868	In addition to increases in concentrations due to H ₂ O loss, Na, K, and Cl can be leached
869	from host rocks during water-rock interactions. Using chemical data from Hanghoj et al. (2010)
870	on the Na content of partially serpentinized Oman harzburgites (~0.03 mole Na ⁺ /kg rock or 0.1

 $wt.\% Na_2O; Cl^-$ was calculated to balance Na⁺ by charge), we can simulate leaching of Na and Cl

872	during serpentinization. Results of this simulation, depicted as solid dark blue curves in Figures
873	8a and 8b, show that if Na and Cl were leached from the host rocks during water-rock
874	interactions, it would entail rock-to-water ratios of between 0.2 and 0.5 (<i>i.e.</i> , reaction of 0.2 to
875	0.5 kg of rock in 1 kg of water) to account for environmental measurements. The above range in
876	rock-to-water ratios depicts lower and more plausible extents of water-rock reactions than those
877	required if Na and Cl enrichment is accounted for by mineral hydration only. Neal and Stanger
878	(1985) also postulated in favor of a leaching origin of chloride in hyperalkaline fluids in Oman,
879	as large amounts of H ₂ O incorporated into serpentine and brucite would result in fluids that are
880	extremely ² H-rich and ¹⁸ O-depleted. In contrast, as shown in Figure 4, about half of the
881	hyperalkaline fluid analyses (all filled colored symbols in Figure 4), which are indicative of large
882	extents of water-rock interactions, are ² H-depleted relative to circumneutral fluids (open green
883	squares in Figure 4). As for the source of these solutes, Neal and Stanger (1985) indicated that
884	chloride could be leached from mineral surfaces and secondary precipitates such as salts and
885	metal hydroxychlorides. Boschetti and Toscani (2008) also discussed leaching from saline fluid
886	inclusions to account for the elevated concentrations of Na and Cl in hyperalkaline fluids seeping
887	from serpentinite outcrops in Northern Italy. All of the above suggests that the host rocks in
888	these continental aquifers interacted previously with saline fluids, likely during seawater-driven
889	serpentinization before the ophiolite was emplaced onto the continent.
890	Sodium and chloride concentrations of circumneutral (Type 1) fluids, on the other hand,

cannot be accounted for by leaching from rocks with concentrations typical of those from
unaltered or partially serpentinized Oman harzburgite, as shown by the solid dark blue curves in
Figures 8a and 8b. Since many circumneutral fluids were sampled in surface stream runoffs,
evaporation could contribute to enrichment of Na and Cl. However, extreme evaporation of

>50% water loss to account for enrichment from typical rainwater concentrations would result in 895 fluids highly enriched in ²H and ¹⁸O, which was not observed. Additional input of Na and Cl 896 could have come from interactions with rocks that are more enriched in these elements than 897 typical partially altered ultramafic rocks. The shallow aquifers where these fluids occur are 898 composed of highly altered rocks, which may be more enriched in Na and Cl. We simulated 899 900 leaching of Cl from serpentinites using the average Cl content (~3000 ppm Cl or 80 millimole Cl/kg rock) of serpentinities or completely serpentinized ultramafic rocks compiled by Deschamp 901 902 et al. (2013). Results of these calculations, depicted in Figure 8b as the reaction path in dashed 903 red lines, show that only ~7 to 20 grams of serpentinite would need to react with a kilogram of water (log water-to-rock ratio ~-2 to -1.5) to account for the Cl content (0.8-2 millimolal) of 904 most Type 1 fluids. Note that the reaction path shown in Figure 8b depicts reaction of rainwater 905 with a serpentinite and hence the maximum pH at ~ 8.2 is attained when the rainwater approaches 906 907 equilibrium with serpentine.

Further evidence that supports leaching during the evolution of rainwater into Type 1 circumneutral fluids and eventually into Type 2 hyperalkaline fluids is shown in Figure 9a. There is a general increase in the Na/K ratio of fluids as pH increases. The Na/K ratios of all fluid types would remain constant if hydration and evaporation were the only processes accounting for the increasing Na, K, and Cl content. The increasing Na/K ratio with pH could be attributed to leaching from precipitates that formed during previous seawater-rock interactions, as seawater is enriched in Na relative to K.

As shown in Figure 9b, the trend in the Na/Cl ratio, like the trend in the Na/K ratio, is not constant with pH. Type 1 fluids (pH <9) are depleted in Na relative to Cl, while Type 2 fluids (pH >11) have Na/Cl ratios above unity or those typical of seawater (0.8). The relative

918	enrichment of Na with reaction progress could be due to the release of sodium from Na-bearing
919	minerals such as plagioclase and alkali-bearing phyllosilicates (Boschetti and Toscani, 2008;
920	Chavagnac et al., 2013b). Another source could be Na-bearing clinopyroxene, though Chavagnac
921	et al. (2013b) argued that this contribution is likely minor, as these minerals are typically not
922	abundant in harzburgites from Oman. Conversely, another process that could account for the
923	trends shown in Figure 9b is the removal of Cl through incorporation into minerals where
924	structural substitutions for hydroxide are possible, such as in apatite and some phyllosilicates.
925	Layered doubled hydroxides (LDHs) can also attract anions into their positively charged
926	interlayers and are known to occur in hyperalkaline pools and in mixing zones between Type 1
927	and Type 2 fluids (Paukert et al., 2012; Chavagnac et al., 2013a), as well as in ultramafic-hosted
928	aquifers in Oman (Taylor et al., 1991). Taylor et al. (1991) characterized pyroaurite
929	$[Mg_6Fe^{+3}_2(CO_3)(OH)_{16}:4H_2O]$, the ferric iron end-member of hydrotalcite, from aquifer rocks in
930	Oman that mainly hosts CO ₃ ⁻² but also contains minor amounts of Cl ⁻ and SO ₄ ⁻² . These authors
931	proposed that the pyroaurite was generated when Fe ⁺² mobilized during silicate dissolution was
932	partially oxidized during mixing with oxic, shallow fluid sources. Alternatively, Fe-bearing
933	brucite in altered ultramafic rocks could be partially oxidized to form a LDH, as exemplified by
934	the reaction

$$8Mg_{0.75}Fe(II)_{0.25}(OH)_{2(brucite)} + 0.5O_{2(aq)} + 2Cl^{-} + 3H_2O + 2H^{+}$$
(10)
$$\rightarrow Mg_6Fe(III)_2Cl_2(OH)_{16}: 4H_2O_{(iowaite)}$$

for iowaite, the Cl⁻ end-member of pyroaurite. Analogous reactions can also be written for other
LDHs such as green rust or fougerite, the former of which has numerous properties that are
invoked as ideal for the emergence of life on Earth (Russell, 2018). Note that most water samples

938 retrieved from wells remain chloride-rich (black squares in Figure 9b), implying that alteration

939 processes involving LDHs may be limited to deep-seated fluids surfacing as hyperalkaline seeps.



940

Figure 9. Trends in the total (a) Na/K and (b) Na/Cl ratios of fluids sampled from Oman.
Symbols are the same as those in Figures 3, 5, 7, and 8. The increasing Na/K ratio in (a) suggests
leaching from host rocks that were altered by seawater-rock interaction. Modern seawater has an
average Na/K ratio of ~46 (dashed grey line, Millero et al., 2008). The increasing Na/Cl ratio in
(b) suggests additional sources of Na or sinks for Cl. Type 2 fluids have Na/Cl ratios that are
above unity (dashed black line) and that typical of seawater (0.86: dashed grey line, Millero et al., 2008).

948

949 **5. Timescales of Reactions**

In the preceding discussions, the extents of water-rock reactions are depicted in terms of 950 rock-to-water ratios that are independent of time. To gain insights into the timescales required to 951 reach the various stages of serpentinization, we conducted reaction path calculations that 952 incorporate dissolution rates of primary minerals at ambient conditions (~25 °C). Rates for 953 forsterite, enstatite, and diopside dissolution are based on experiments of Pokrovsky and Schott 954 (2000), Oelkers and Schott (2001), and Golubev et al. (2005), respectively, which include the 955 pH-dependence of the dissolution rates of the reacting minerals. Calculations were done 956 assuming constant surface area (1 m^2) of fresh ultramafic rock interacting with 1 kg of fluid. 957

958 Results are shown in Figure 10a, which depicts both the timescales and the degree of water-rock959 interactions required to attain the various stages of the overall serpentinization process.





Figure 10. (a) Timescales (in log years) required to attain the various pH and stages of
serpentinization. (b) Timescales (in log years) required to attain saturation of three secondary
phases common in the altered ultramafic rocks in the shallow subsurface. Numbers inside circles
in (a) indicate the starting points of the six stages of serpentinization. Black filled circles in (b)
depict end of reaction path depicting dissolution of indicated mineral. Vertical dashed lines
indicate various timepoints (1 day, month, and year) as a reference.

```
Fluids with circumneutral pH (7 to 9) are attained with minimal reaction progress at
968
       timescales from a few months to at most a decade. Attaining serpentine saturation (pH ~8, Stage
969
       2) is quick (a few months) while attaining carbonate saturation (Stage 3) entails a few years.
970
       However, as discussed above, these circumneutral Type 1 fluids are likely to result from fluid
971
972
       interacting with the highly altered host rocks of shallow aquifers composed mostly of serpentine,
973
       brucite, and carbonates. Calculations simulating the dissolution of these secondary phases at
974
       ambient conditions were conducted using data from Thom et al. (2013), Pokrovsky and Schott
       (2004), and Pokrovsky and Schott (1999) for chrysotile, brucite, and magnesite, respectively.
975
976
       The pH-dependences of the dissolution rates were also taken into consideration. Calculations for
```

each of these minerals were done separately and were terminated when fluids reached saturation 977 with the mineral involved. As with the reaction path involving unaltered ultramafic rocks, it is 978 assumed that each kg of water reacts with 1 m² surface area of the involved mineral. Although 979 shallow aquifer rocks could be made up of variable proportions of these secondary phases, this 980 idealized assumption allows us to evaluate how rapidly each of these minerals can attain 981 982 saturation. Results are depicted in Figure 10b, which shows that brucite saturation can be attained relatively quickly owing to its rapid dissolution rates (2 orders of magnitude faster than 983 forsterite dissolution). Fluids would require a few months to attain equilibrium with magnesite. 984 985 Serpentine saturation can be attained either through the dissolution of primary minerals in ultramafic rocks (Figure 10a, Stage 2) or serpentinites (Figure 10b). Both reach similar fluid 986 chemistry (pH \sim 8) but the latter would require almost a decade, two orders of magnitude slower 987 than the former due to the sluggish dissolution rate of chrysotile relative to those of the primary 988 minerals. 989

990 Compositions characteristic of Type 2 hyperalkaline fluids (Figure 10a) require much longer durations to attain. As shown in Figure S4, a rock-to-water ratio of at least 0.1 is needed 991 to attain the maximum pH via Ctl-Brc-Di equilibrium (Stage 6), and this requires timescales 992 993 close to 10,000 years. About 100,000 years would be needed to attain higher extents of reaction at a rock-to-water ratio of 1. Calculated timescales between 10,000–100,000 years correspond to 994 995 the last glacial age, which is consistent with the hypothesis of Paukert et al. (2019) that some 996 hyperalkaline fluids in Oman were recharged during this period. Our kinetic results also 997 complement the experimental results of McCollom and Donaldson (2016) that show very slow 998 rates of H₂ and CH₄ generation during low-temperature (<100 °C) serpentinization.

999 Inferred timescales for various stages of serpentinization, however, should be taken 1000 cautiously as discrepancies between laboratory-determined dissolution rates and those 1001 determined through field studies are well known. Field-determined rates are far slower than those measured in the laboratory, often entailing differences of several orders of magnitude (White and 1002 1003 Brantley, 2003). For example, field-determined weathering rates of ultramafic rocks in the 1004 Amazon are at least two orders of magnitude more sluggish than laboratory-determined rates 1005 (Freyssinet and Farah, 2000). Studies of other lithological settings also report discrepancies 1006 between field and laboratory-determined rates (e.g., Velbel, 1993; White and Brantley, 2003; 1007 Maher, 2010), with the latter yielding rates at most four orders of magnitude faster than the former. These studies attribute the observed differences to numerous factors such as estimations 1008 1009 of reacting surface areas and their evolution during the weathering process, fluid residence times, 1010 and the age of the rock. Taking the discrepancies into account, extrapolated timescales from our 1011 calculations to natural systems lead to much longer estimates, perhaps even requiring millions of 1012 years to attain hyperalkaline conditions if the deviation is at least 2 orders of magnitude. Better constraints on the natural rate at which low-temperature serpentinization occurs would provide 1013 powerful insights into the flux of reduced volatiles as well as the rate of energy transfer between 1014 1015 the lithosphere and the biosphere in deep ultramafic aquifers here on Earth and other rock bodies 1016 in our solar system.

1017 **6.** Conclusions

Overall, calculations identify the reactions that lead to the contrasting pH, fluid
compositions, and redox potentials of end-member Type 1 circumneutral (pH 7 to 9) and Type 2
hyperalkaline (pH >11) fluids. The former are outcomes of mineral dissolution reactions and
fluid interactions with highly altered rocks that are common in shallow ultramafic aquifers, while

the latter approach equilibrium with serpentine and brucite, which actively form during advanced
stages of serpentinization. The most alkaline fluids documented in low-temperature
serpentinizing systems follow constraints dictated by diopside dissolution and some fluids in
Oman approach this upper limit. The pH values of most hyperalkaline fluids in Oman, however,
are lower than this constraint and can be attributed to a shift from diopside dissolution to calcite
precipitation driven by the addition of dissolved inorganic carbon (DIC) as surfacing deep-seated
hyperalkaline fluids encounter the atmosphere.

1029 Most fluids with pH values between those of Type 1 and Type 2 can be attributed to 1030 mixing of these end-member fluids. Our calculations highlight potential imprints of *shallow* subsurficial processes on the deep-seated signatures of discharging hyperalkaline fluids through 1031 1032 mixing. Shallow portions of the *discharge* zone could be hotspots for subsurface life that may 1033 bloom where reduced deep-seated fluids encounter oxidized shallow aquifer fluids. As systems 1034 with contrasting oxidation-reduction potentials meet, habitability is defined by the extent to 1035 which such systems fail to equilibrate abiotically, leaving opportunities for microbial 1036 communities to exploit.

1037 In contrast, some fluids with pH values between 9 and 11 do not correspond to mixing 1038 trends between end-member fluids. Rather than forming through mixing, these fluids may be indicative of intermediate stages of serpentinization. As this stage of reaction progress 1039 1040 corresponds to incipient serpentinization and the carbonation of ultramafic rocks, further focus 1041 on this fluid type could help inform potential studies targeting ultramafic aquifers as hosts for 1042 deep storage of sequestered atmospheric CO₂ (Kelemen and Matter, 2008; Kelemen et al., 2011; 1043 National Academies, 2019). In addition, targeting fluids with these compositions will also reveal 1044 information on reactions that facilitate the transformation of oxic surface fluids into reduced,

deep-seated, hyperalkaline fluids as they infiltrate deeper into the aquifer. Conditions for 1045 1046 habitable environments may exist through this fluid transformation and future exploration of this fluid type will provide insights into the habitability of recharge zones of the fluid pathway and 1047 how they differ from discharge zones and their surface expressions. The Oman Drilling Project 1048 (omandrilling.ac.uk, Kelemen et al., 2013) recently drilled several wells that tap into fluids 1049 1050 indicative of various stages of serpentinization and is poised to reveal a comprehensive assessment of microbial communities powered by rock and fluid interactions in the ultramafic 1051 1052 subsurface. Models from this work can be used to assess these samples. We also propose the 1053 practicality of measuring both pH and Si concentrations with field-portable equipment, allowing rapid assessment of fluid type and extents of mixing while in the field, which can augment 1054 1055 exploration with real-time information.

We demonstrated that the compositions of serpentinization-generated fluids can be partly 1056 accounted for by predictions of fluid-mineral equilibria and showed how processes such as 1057 1058 leaching and fluid mixing can account for compositions that deviate from equilibrium expectations. Both water and ultramafic rocks are believed to abound in ice-covered ocean 1059 worlds in our Solar System such as Europa and Enceladus, and serpentinization is thought to 1060 1061 enable habitable environments in these terrestrial bodies as it does on Earth (Vance et al., 2007; 1062 Holms et al., 2015; Glein et al., 2015; Waite et al., 2017; Glein and Zolotov, 2020). By 1063 demonstrating that simulations of low-temperature serpentinization can be successfully applied 1064 to analogous processes here on Earth, this work substantiates these calculations as useful tools in 1065 exploring serpentinization reactions beyond our own planet.

1066 Finally, further work on the kinetics of low-temperature serpentinization, whether1067 determined through laboratory experiments or in the field, will ultimately provide stronger

1068	constraints on the rate of energy transfer between the lithosphere and the subsurface biosphere.
1069	Rates will also feed into reactive transport simulations that could improve upon the calculations
1070	described here and reveal how various transport processes contribute to the overall compositions
1071	of serpentinization-generated fluids.
1072	7. Acknowledgments
1073	Samples used in this study were gathered with the help of Peter Kelemen, Juerg Matter,
1074	Amelia Paukert Vankeuren, Lisa Streit, Peter Canovas, Jeff Havig, and Peter Marsala. We also
1075	would like to thank Tracy Lund, Natasha Zolotova and Roy Erikson for help in analysis of fluid
1076	samples at the ASU GEOPIG laboratory, W.M. Keck Foundation Laboratory for Environmental
1077	Biogeochemistry, and Goldwater Environmental Laboratory (the latter two now merged as the
1078	Metals, Environmental and Terrestrial Analytical Laboratory (METAL)). Thanks also to Tucker
1079	Ely, Vincent Milesi, and Grayson Boyer for helpful discussions on computational tools used in
1080	this study. This work was supported by NASA Exobiology grant NNX12AB38G, the NASA
1081	Astrobiology Institute Rock-Powered Life (RPL) and Exploring Ocean Worlds (ExOW) projects,
1082	and NSF grant EAR-1515513. By acceptance, all analytical and theoretical results (Tables S1
1083	and S2) will be stored and uploaded at the EarthChem Library portal.
1084	8. References
1085	Ahmed, A. F. M., Singh, R. P., & Elmubarak, A. H. (1990). Chemistry of atmospheric
1086	precipitation at the Western Arabian Gulf Coast. Atmospheric Environment. Part A.
1087	General Topics, 24(12), 2927–2934. https://doi.org/10.1016/0960-1686(90)90473-Z
1088	Alexander, W. R., Dayal, R., Eagleson, K., Eikenberg, J., Hamilton, E., Linklater, C. M.,
1089	McKinley, I. G., & Tweed, C. J. (1992). A natural analogue of high pH cement pore
1090	waters from the Maqarin area of northern Jordan. II: results of predictive geochemical

- 1091 calculations. *Journal of Geochemical Exploration*, 46(1), 133–146.
- 1092 https://doi.org/10.1016/0375-6742(92)90104-G
- 1093 Amend, J. P., McCollom, T. M., Hentscher, M., & Bach, W. (2011). Catabolic and anabolic
- 1094 energy for chemolithoautotrophs in deep-sea hydrothermal systems hosted in different
- 1095 rock types. *Geochimica et Cosmochimica Acta*, 75(19), 5736–5748.
- 1096 https://doi.org/10.1016/j.gca.2011.07.041
- 1097 Barnes, I., O'Neil, J. R., & Trescases, J. J. (1978). Present day serpentinization in New
- 1098 Caledonia, Oman and Yugoslavia. *Geochimica et Cosmochimica Acta*, 42(1), 144–145.
- 1099 https://doi.org/10.1016/0016-7037(78)90225-9
- 1100 Bath, A., Christofi, N., Neal, C., Philp, J., Cave, M., McKinley, I., & Berner, U. (1987). Trace
- element and microbiological studies of alkaline groundwaters in Oman, Arabian Gulf: a
 natural analogue for cement pore-waters. British Geological Survey, Fluid Processes
- 1103 Research Group.
- 1104 Boschetti, T., & Toscani, L. (2008). Springs and streams of the Taro–Ceno Valleys (Northern
- 1105 Apennine, Italy): Reaction path modeling of waters interacting with serpentinized
- 1106 ultramafic rocks. *Chemical Geology*, 257(1), 76–91.
- 1107 https://doi.org/10.1016/j.chemgeo.2008.08.017
- 1108 Boulart, C., Chavagnac, V., Monnin, C., Delacour, A., Ceuleneer, G., & Hoareau, G. (2013).
- 1109 Differences in gas venting from ultramafic-hosted warm springs: The example of Iman 1110 and Voltri Ophiolites. *Ofioliti*, *38*(2), 142–156. https://doi.org/10.4454/ofioliti.v38i2.423
- 1111 Boyd, E. S., Amenabar, M. J., Poudel, S., & Templeton, A. S. (2020). Bioenergetic constraints
- 1112 on the origin of autotrophic metabolism. *Philosophical Transactions of the Royal Society*
- 1113 *A: Mathematical, Physical and Engineering Sciences, 378*(2165), 20190151.

- 1114 https://doi.org/10.1098/rsta.2019.0151
- 1115 Bruni, J., Canepa, M., Chiodini, G., Cioni, R., Cipolli, F., Longinelli, A., Marini, L., Ottonello,
- 1116 G., & Vetuschi Zuccolini, M. (2002). Irreversible water–rock mass transfer
- 1117 accompanying the generation of the neutral, Mg–HCO₃ and high-pH, Ca–OH spring
- 1118 waters of the Genova province, Italy. *Applied Geochemistry*, 17(4), 455–474.
- 1119 https://doi.org/10.1016/S0883-2927(01)00113-5
- 1120 Canovas, P. A., Hoehler, T., & Shock, E. L. (2017). Geochemical bioenergetics during low-
- 1121 temperature serpentinization: An example from the Samail ophiolite, Sultanate of Oman.
- 1122 *Journal of Geophysical Research: Biogeosciences*, *122*(7), 1821–1847.
- 1123 https://doi.org/10.1002/2017JG003825
- 1124 Charlou, J. L., Donval, J. P., Fouquet, Y., Jean-Baptiste, P., & Holm, N. (2002). Geochemistry of
- high H2 and CH4 vent fluids issuing from ultramafic rocks at the Rainbow hydrothermal
- 1126 field (36°14'N, MAR). *Chemical Geology*, 191(4), 345–359.
- 1127 https://doi.org/10.1016/S0009-2541(02)00134-1
- 1128 Chavagnac, V., Ceuleneer, G., Monnin, C., Lansac, B., Hoareau, G., & Boulart, C. (2013a).
- 1129 Mineralogical assemblages forming at hyperalkaline warm springs hosted on ultramafic
- 1130 rocks: A case study of Oman and Ligurian ophiolites: Mineral Precipitate at Alkaline
- 1131 Springs. Geochemistry, Geophysics, Geosystems, 14(7), 2474–2495.
- 1132 https://doi.org/10.1002/ggge.20146
- 1133 Chavagnac, V., Monnin, C., Ceuleneer, G., Boulart, C., & Hoareau, G. (2013b). Characterization
- 1134 of hyperalkaline fluids produced by low-temperature serpentinization of mantle
- 1135 peridotites in the Oman and Ligurian ophiolites: Hyperalkaline Waters in Oman and
- 1136 Liguria. Geochemistry, Geophysics, Geosystems, 14(7), 2496–2522.

1137	https://do	oi.org/10.1	002/ggge.2014'	7
------	------------	-------------	----------------	---

- 1138 Cipolli, F., Gambardella, B., Marini, L., Ottonello, G., & Vetuschi Zuccolini, M. (2004).
- 1139 Geochemistry of high-pH waters from serpentinites of the Gruppo di Voltri (Genova,
- 1140 Italy) and reaction path modeling of CO₂ sequestration in serpentinite aquifers. *Applied*
- 1141 *Geochemistry*, 19(5), 787–802. https://doi.org/10.1016/j.apgeochem.2003.10.007
- 1142 Crespo-Medina, M., Twing, K. I., Kubo, M. D. Y., Hoehler, T. M., Cardace, D., McCollom, T.,
- 1143 & Schrenk, M. O. (2014). Insights into environmental controls on microbial communities
- in a continental serpentinite aquifer using a microcosm-based approach. *Frontiers in*

1145 *Microbiology*, *5*, no. 604. https://doi.org/10.3389/fmicb.2014.00604

- Deschamps, F., Godard, M., Guillot, S., & Hattori, K. (2013). Geochemistry of subduction zone
 serpentinites: A review. *Lithos*, *178*, 96–127. https://doi.org/10.1016/j.lithos.2013.05.019
- 1148 Dewandel, B., Lachassagne, P., Boudier, F., Al-Hattali, S., Ladouche, B., Pinault, J.-L., & Al-
- 1149 Suleimani, Z. (2005). A conceptual hydrogeological model of ophiolite hard-rock
- aquifers in Oman based on a multiscale and a multidisciplinary approach. *Hydrogeology*
- 1151 *Journal*, 13(5–6), 708–726. https://doi.org/10.1007/s10040-005-0449-2
- 1152 Edmond, J. M., Measures, C., McDuff, R. E., Chan, L. H., Collier, R., Grant, B., Gordon, L. I.,
- 1153 & Corliss, J. B. (1979). Ridge crest hydrothermal activity and the balances of the major
- and minor elements in the ocean: The Galapagos data. *Earth and Planetary Science*

1155 *Letters*, 46(1), 1–18. https://doi.org/10.1016/0012-821X(79)90061-X

- 1156 Etiope, G., & Whiticar, M. J. (2019). Abiotic methane in continental ultramafic rock systems:
- 1157 Towards a genetic model. *Applied Geochemistry*, *102*, 139–152.
- 1158 https://doi.org/10.1016/j.apgeochem.2019.01.012
- 1159 Etiope, G., Samardžić, N., Grassa, F., Hrvatović, H., Miošić, N., & Skopljak, F. (2017). Methane

- and hydrogen in hyperalkaline groundwaters of the serpentinized Dinaride ophiolite belt,
- 1161 Bosnia and Herzegovina. *Applied Geochemistry*, 84, 286–296.
- 1162 https://doi.org/10.1016/j.apgeochem.2017.07.006
- 1163 Fecteau, K. M. (2016). Organic Carbon in Hydrothermal Systems: from Phototrophy to
- 1164 *Aldehyde Transformations*. Arizona State University.
- 1165 Fones, E. M., Colman, D. R., Kraus, E. A., Nothaft, D. B., Poudel, S., Rempfert, K. R., Spear, J.
- 1166 R., Templeton, A. S., & Boyd, E. S. (2019). Physiological adaptations to serpentinization
- in the Samail Ophiolite, Oman. *The ISME Journal*, *13*(7), 1750–1762.
- 1168 https://doi.org/10.1038/s41396-019-0391-2
- 1169 Foustoukos, D. I., Savov, I. P., & Janecky, D. R. (2008). Chemical and isotopic constraints on
- 1170 water/rock interactions at the Lost City hydrothermal field, 30°N Mid-Atlantic Ridge.
- 1171 *Geochimica et Cosmochimica Acta*, 72(22), 5457–5474.
- 1172 https://doi.org/10.1016/j.gca.2008.07.035
- 1173 Freyssinet, P., & Farah, A. S. (2000). Geochemical mass balance and weathering rates of
- 1174 ultramafic schists in Amazonia. *Chemical Geology*, *170*(1), 133–151.
- 1175 https://doi.org/10.1016/S0009-2541(99)00245-4
- 1176 Frost, B. R., Evans, K. A., Swapp, S. M., Beard, J. S., & Mothersole, F. E. (2013). The process

1177 of serpentinization in dunite from New Caledonia. *Lithos*, *178*, 24–39.

- 1178 https://doi.org/10.1016/j.lithos.2013.02.002
- 1179 Früh-Green, G. L., Orcutt, B. N., Rouméjon, S., Lilley, M. D., Morono, Y., Cotterill, C., Green,
- 1180 S., Escartin, J., John, B. E., McCaig, A. M., Cannat, M., Ménez, B., Schwarzenbach, E.
- 1181 M., Williams, M. J., Morgan, S., Lang, S. Q., Schrenk, M. O., Brazelton, W. J., Akizawa,
- 1182 N., Boschi, C., Dunkel, K. G., Quéméneur, M., Whattam, S. A., Mayhew, L., Harris, M.,

- 1183 Bayrakci, G., Behrmann, J.-H., Herrero-Bervera, E. Hesse, K., Liu, H.-Q., Ratnayake, A.
- 1184 S., Twing, K., Weis, D., Zhao, R., & Bilenker, L. (2018). Magmatism, serpentinization
- and life: Insights through drilling the Atlantis Massif (IODP Expedition 357). *Lithos*,
- 1186 *323*, 137–155. https://doi.org/10.1016/j.lithos.2018.09.012
- 1187 Gaucher, E. C., & Blanc, P. (2006). Cement/clay interactions A review: Experiments, natural
- analogues, and modeling. *Waste Management*, 26(7), 776–788.
- 1189 https://doi.org/10.1016/j.wasman.2006.01.027
- 1190 Geldern, R. van, & Barth, J. A. C. (2012). Optimization of instrument setup and post-run
- 1191 corrections for oxygen and hydrogen stable isotope measurements of water by isotope
- 1192 ratio infrared spectroscopy (IRIS). *Limnology and Oceanography: Methods*, 10(12),
- 1193 1024–1036. https://doi.org/10.4319/lom.2012.10.1024
- 1194 Giampouras, M., Garrido, C. J., Bach, W., Los, C., Fussmann, D., Monien, P., & García-Ruiz, J.
- 1195 M. (2020). On the controls of mineral assemblages and textures in alkaline springs,
- 1196 Samail Ophiolite, Oman. *Chemical Geology*, 533, no. 119435.
- 1197 https://doi.org/10.1016/j.chemgeo.2019.119435
- 1198 Glein, C. R., Baross, J. A., & Waite, J. H. (2015). The pH of Enceladus' ocean. Geochimica et
- 1199 *Cosmochimica Acta*, *162*, 202–219. https://doi.org/10.1016/j.gca.2015.04.017
- Glein, C. R., & Zolotov, M. Y. (2020). Hydrogen, hydrocarbons, and habitability across the solar
 system. *Elements*, *16*(1), 47–52. https://doi.org/10.2138/gselements.16.1.47
- 1202 Golubev, S. V., Pokrovsky, O. S., & Schott, J. (2005). Experimental determination of the effect
- 1203 of dissolved CO_2 on the dissolution kinetics of Mg and Ca silicates at 25 °C. *Chemical*
- 1204 *Geology*, 217(3), 227–238. https://doi.org/10.1016/j.chemgeo.2004.12.011
- 1205 Hammer, O., Harper, D. A. T., & Ryan, P. D. (2001). PAST: Paleontological statistics software

1206	package for education and data analysis. <i>Palaeontologia Electronica</i> , 4(1), 9pp.
1207	Hanghøj, K., Kelemen, P. B., Hassler, D., & Godard, M. (2010). Composition and genesis of
1208	depleted mantle peridotites from the Wadi Tayin Massif, Oman Ophiolite; Major and
1209	trace element geochemistry, and Os isotope and PGE systematics. Journal of Petrology,
1210	51(1-2), 201-227. https://doi.org/10.1093/petrology/egp077
1211	Helgeson, H. C., Delaney, J. M., Nesbitt, H. W., & Bird, D. K. (1978). Summary and critique of
1212	the thermodynamic properties of rock-forming minerals. American Journal of Science,
1213	278, 1–229.
1214	Holm, N. G., Dumont, M., Ivarsson, M., & Konn, C. (2006). Alkaline fluid circulation in
1215	ultramafic rocks and formation of nucleotide constituents: A hypothesis. Geochemical
1216	Transactions, 7(1), no. 7. https://doi.org/10.1186/1467-4866-7-7
1217	Holm, N. G., Oze, C., Mousis, O., Waite, J. H., & Guilbert-Lepoutre, A. (2015). Serpentinization
1218	and the formation of H_2 and CH_4 on celestial bodies (planets, moons, comets).
1219	Astrobiology, 15(7), 587-600. https://doi.org/10.1089/ast.2014.1188
1220	Johnson, J. W., Oelkers, E. H., & Helgeson, H. C. (1992). SUPCRT92: A software package for
1221	calculating the standard molal thermodynamic properties of minerals, gases, aqueous
1222	species, and reactions from 1 to 5000 bar and 0 to 1000°C. Computers & Geosciences,
1223	18(7), 899–947. https://doi.org/10.1016/0098-3004(92)90029-Q
1224	Kelemen, P., Al Rajhi, A., Godard, M., Ildefonse, B., Köpke, J., MacLeod, C., Manning, C.,
1225	Michibayashi, K., Nasir, S., Shock, E., Takazawa, E., & Teagle, D. (2013). Scientific
1226	drilling and related research in the Samail Ophiolite, Sultanate of Oman. Scientific
1227	Drilling, 15, 64–71. https://doi.org/10.5194/sd-15-64-2013

1228 Kelemen, P. B., & Matter, J. (2008). In situ carbonation of peridotite for CO₂ storage.

1229 *Proceedings of the National Academy of Sciences*, *105*(45), 17295–17300.

- 1230 https://doi.org/10.1073/pnas.0805794105
- 1231 Kelemen, P. B., Matter, J., Streit, E. E., Rudge, J. F., Curry, W. B., & Blusztajn, J. (2011). Rates
- and mechanisms of mineral carbonation in peridotite: Natural processes and recipes for
- 1233 enhanced, in situ CO₂ capture and storage. *Annual Review of Earth and Planetary*
- 1234 *Sciences*, *39*(1), 545–576. https://doi.org/10.1146/annurev-earth-092010-152509
- 1235 Kelley, D. S., Karson, J. A., Blackman, D. K., Früh-Green, G. L., Butterfield, D. A., Lilley, M.
- 1236 D., Olson, E. J., Schrenk, M. O., Roe, K. K., Lebon, G. T., Rivizzigno, P., & the AT3-60
- 1237 Shipboard Party. (2001). An off-axis hydrothermal vent field near the Mid-Atlantic Ridge

1238 at 30° N. *Nature*, 412(6843), 145–149. https://doi.org/10.1038/35084000

- 1239 Khoury, H. N., Salameh, E., Clark, I. D., Fritz, P., Bajjali, W., Milodowski, A. E., Cave, M. R.,
- 1240 & Alexander, W. R. (1992). A natural analogue of high pH cement pore waters from the
- 1241 Maqarin area of northern Jordan. I: Introduction to the site. *Journal of Geochemical*

1242 *Exploration*, 46(1), 117–132. https://doi.org/10.1016/0375-6742(92)90103-F

- 1243 Khoury, H. N., Salameh, E., & Abdul-Jaber, Q. (1985). Characteristics of an unusual highly
- 1244 alkaline water from the Maqarin area, northern Jordan. *Journal of Hydrology*, 81(1), 79–
- 1245 91. https://doi.org/10.1016/0022-1694(85)90168-4
- 1246 Klein, F., Bach, W., & McCollom, T. M. (2013). Compositional controls on hydrogen generation
 1247 during serpentinization of ultramafic rocks. *Lithos*, *178*, 55–69.
- 1248 https://doi.org/10.1016/j.lithos.2013.03.008
- 1249 Klein, F., Humphris, S. E., Guo, W., Schubotz, F., Schwarzenbach, E. M., & Orsi, W. D. (2015).
- 1250 Fluid mixing and the deep biosphere of a fossil Lost City-type hydrothermal system at the
- 1251 Iberia Margin. *Proceedings of the National Academy of Sciences*, *112*(39), 12036–12041.

- 1252 https://doi.org/10.1073/pnas.1504674112
- 1253 Lang, S. Q., Butterfield, D. A., Schulte, M., Kelley, D. S., & Lilley, M. D. (2010). Elevated
- 1254 concentrations of formate, acetate and dissolved organic carbon found at the Lost City
- 1255 hydrothermal field. *Geochimica et Cosmochimica Acta*, 74(3), 941–952.
- 1256 https://doi.org/10.1016/j.gca.2009.10.045
- 1257 Leong, J. A. M., & Shock, E. L. (2020). Thermodynamic constraints on the geochemistry of low-
- 1258 temperature, continental, serpentinization-generated fluids. *American Journal of Science*,
- 1259 320(3), 185–235. https://doi.org/10.2475/03.2020.01
- 1260 Lindsay, M. R., Amenabar, M. J., Fecteau, K. M., Debes, R. V., Martins, M. C. F., Fristad, K. E.,
- 1261 Xu, H., Hoehler, T. M., Shock, E. L., & Boyd, E. S. (2018). Subsurface processes
- influence oxidant availability and chemoautotrophic hydrogen metabolism in
- 1263 Yellowstone hot springs. *Geobiology*, *16*(6), 674–692. https://doi.org/10.1111/gbi.12308
- 1264 Maher, K. (2010). The dependence of chemical weathering rates on fluid residence time. *Earth*
- 1265 *and Planetary Science Letters*, 294(1), 101–110.
- 1266 https://doi.org/10.1016/j.epsl.2010.03.010
- 1267 Malvoisin, B. (2015). Mass transfer in the oceanic lithosphere: Serpentinization is not
- isochemical. *Earth and Planetary Science Letters*, 430, 75–85.
- 1269 https://doi.org/10.1016/j.epsl.2015.07.043
- 1270 Marques, J. M., Carreira, P. M., Carvalho, M. R., Matias, M. J., Goff, F. E., Basto, M. J., Graça,
- 1271 R. C., Aires-Barros, L., & Rocha, L. (2008). Origins of high pH mineral waters from
- 1272 ultramafic rocks, Central Portugal. *Applied Geochemistry*, 23(12), 3278–3289.
- 1273 https://doi.org/10.1016/j.apgeochem.2008.06.029
- 1274 Martin, W., & Russell, M. J. (2007). On the origin of biochemistry at an alkaline hydrothermal

- 1275 vent. *Philosophical Transactions of the Royal Society B: Biological Sciences*, 362(1486),
- 1276 1887–1926. https://doi.org/10.1098/rstb.2006.1881
- 1277 Martin, W., Baross, J., Kelley, D., & Russell, M. J. (2008). Hydrothermal vents and the origin of
- 1278 life. *Nature Reviews Microbiology*, 6(11), 805–814. https://doi.org/10.1038/nrmicro1991
- 1279 Matter, J. M., Waber, H. N., Loew, S., & Matter, A. (2006). Recharge areas and geochemical
- 1280 evolution of groundwater in an alluvial aquifer system in the Sultanate of Oman.

1281 *Hydrogeology Journal*, *14*(1), 203–224. https://doi.org/10.1007/s10040-004-0425-2

- 1282 Matter, J. M., & Kelemen, P. B. (2009). Permanent storage of carbon dioxide in geological
- reservoirs by mineral carbonation. *Nature Geoscience*, 2(12), 837–841.
- 1284 https://doi.org/10.1038/ngeo683
- 1285 Mayhew, L. E., Ellison, E. T., Miller, H. M., Kelemen, P. B., & Templeton, A. S. (2018). Iron
- 1286 transformations during low temperature alteration of variably serpentinized rocks from
- 1287 the Samail ophiolite, Oman. *Geochimica et Cosmochimica Acta*, 222, 704–728.
- 1288 https://doi.org/10.1016/j.gca.2017.11.023
- 1289 McCollom, T. M. (2007). Geochemical constraints on sources of metabolic energy for
- 1290 chemolithoautotrophy in ultramafic-hosted deep-sea hydrothermal systems. *Astrobiology*,
- 1291 7(6), 933–950. https://doi.org/10.1089/ast.2006.0119
- McCollom, T. M. (2016). Abiotic methane formation during experimental serpentinization of
 olivine. *Proceedings of the National Academy of Sciences*, *113*(49), 13965–13970.
- 1294 https://doi.org/10.1073/pnas.1611843113
- 1295 McCollom, T. M., & Donaldson, C. (2016). Generation of hydrogen and methane during
- 1296 experimental low-temperature reaction of ultramafic rocks with water. *Astrobiology*,
- 1297 *16*(6), 389–406. https://doi.org/10.1089/ast.2015.1382

- 1298 McCollom, T. M., & Seewald, J. S. (2007). Abiotic synthesis of organic compounds in deep-sea
- 1299 hydrothermal environments. *Chemical Reviews*, *107*(2), 382–401.
- 1300 https://doi.org/10.1021/cr0503660
- 1301 Mervine, E. M., Humphris, S. E., Sims, K. W. W., Kelemen, P. B., & Jenkins, W. J. (2014).
- 1302 Carbonation rates of peridotite in the Samail Ophiolite, Sultanate of Oman, constrained
- through 14C dating and stable isotopes. *Geochimica et Cosmochimica Acta*, 126, 371–

1304 397. https://doi.org/10.1016/j.gca.2013.11.007

- 1305 Meyer-Dombard, D. R., Woycheese, K. M., Yargıçoğlu, E. N., Cardace, D., Shock, E. L.,
- 1306 Güleçal-Pektas, Y., & Temel, M. (2015). High pH microbial ecosystems in a newly
- 1307 discovered, ephemeral, serpentinizing fluid seep at Yanartaş (Chimera), Turkey.

1308 *Frontiers in Microbiology*, *5*, no. 723. https://doi.org/10.3389/fmicb.2014.00723

- 1309 Miller, H. M., Matter, J. M., Kelemen, P., Ellison, E. T., Conrad, M. E., Fierer, N., Ruchala, T.,
- 1310 Tominaga, M., & Templeton, A. S. (2016). Modern water/rock reactions in Oman
- 1311 hyperalkaline peridotite aquifers and implications for microbial habitability. *Geochimica*
- 1312 *et Cosmochimica Acta*, 179, 217–241. https://doi.org/10.1016/j.gca.2016.01.033
- 1313 Miller, H. M., Chaudhry, N., Conrad, M. E., Bill, M., Kopf, S. H., & Templeton, A. S. (2018).
- 1314 Large carbon isotope variability during methanogenesis under alkaline conditions.
- 1315 *Geochimica et Cosmochimica Acta*, 237, 18–31.
- 1316 https://doi.org/10.1016/j.gca.2018.06.007
- 1317 Mottl, M. J. (2009). Highest pH. *Geochemical News*.
- 1318 Mottl, M. J., Komor, S. C., Fryer, P., & Moyer, C. L. (2003). Deep-slab fluids fuel extremophilic
- 1319 Archaea on a Mariana forearc serpentinite mud volcano: Ocean Drilling Program Leg
- 1320 195. *Geochemistry, Geophysics, Geosystems, 4*(11), no. 9009.

- 1321 https://doi.org/10.1029/2003GC000588
- 1322 Mottl, M. J., Wheat, C. G., Fryer, P., Gharib, J., & Martin, J. B. (2004). Chemistry of springs
- across the Mariana forearc shows progressive devolatilization of the subducting plate.
- 1324 *Geochimica et Cosmochimica Acta*, 68(23), 4915–4933.
- 1325 https://doi.org/10.1016/j.gca.2004.05.037
- 1326 National Academies of Sciences, Engineering, and Medicine. (2019). Chapter 6: Carbon
- 1327 Mineralization of CO₂. In *Negative Emissions Technologies and Reliable Sequestration:*
- 1328 *A Research Agenda* (pp. 247–318). Washington, D.C.: National Academies Press.
- 1329 https://doi.org/10.17226/25259
- 1330 Neal, C., & Stanger, G. (1983). Hydrogen generation from mantle source rocks in Oman. *Earth*

1331 and Planetary Science Letters, 66, 315–320. https://doi.org/10.1016/0012-

- 1332 821X(83)90144-9
- 1333 Neal, C., & Stanger, G. (1984). Calcium and magnesium hydroxide precipitation from alkaline
- 1334 groundwaters in Oman, and their significance to the process of serpentinization.
- 1335 *Mineralogical Magazine*, 48(347), 237–241.
- 1336 https://doi.org/10.1180/minmag.1984.048.347.07
- 1337 Neal, C., & Stanger, G. (1985). Past and present serpentinisation of ultramafic rocks; an example
- 1338from the Semail Ophiolite Nappe of Northern Oman. In The Chemistry of Weathering
- 1339 (pp. 249–275). Springer.
- 1340 Nesbitt, H. W., & Bricker, O. P. (1978). Low temperature alteration processes affecting
- 1341 ultramafic bodies. *Geochimica et Cosmochimica Acta*, 42(4), 403–409.
- 1342 https://doi.org/10.1016/0016-7037(78)90271-5
- 1343 Nicolas, A., Boudier, F., Ildefonse, B., & Ball, E. (2000). Accretion of Oman and United Arab

1344	Emirates ophiolite – Discussion of a new structural map. Marine Geophysical
1345	Researches, 21(3), 147-180. https://doi.org/10.1023/A:1026769727917
1346	Nordstrom, K. D., McCleskey, B. R., & Ball, J. W. (2009). Sulfur geochemistry of hydrothermal
1347	waters in Yellowstone National Park: IV Acid-sulfate waters. Applied Geochemistry,
1348	24(2), 191–207. https://doi.org/10.1016/j.apgeochem.2008.11.019
1349	de Obeso, J. C., & Kelemen, P. B. (2018). Fluid rock interactions on residual mantle peridotites
1350	overlain by shallow oceanic limestones: Insights from Wadi Fins, Sultanate of Oman.
1351	Chemical Geology, 498, 139-149. https://doi.org/10.1016/j.chemgeo.2018.09.022
1352	de Obeso, J. C., & Kelemen, P. B. (2020). Major element mobility during serpentinization,
1353	oxidation and weathering of mantle peridotite at low temperatures. Philosophical
1354	Transactions of the Royal Society A: Mathematical, Physical and Engineering Sciences,
1355	378(2165), 20180433. https://doi.org/10.1098/rsta.2018.0433
1356	Oelkers, E. H., & Schott, J. (2001). An experimental study of enstatite dissolution rates as a
1357	function of pH, temperature, and aqueous Mg and Si concentration, and the mechanism
1358	of pyroxene/pyroxenoid dissolution. Geochimica et Cosmochimica Acta, 65(8), 1219-
1359	1231. https://doi.org/10.1016/S0016-7037(00)00564-0
1360	Ohlsson, J. I., Osvatic, J. T., Becraft, E. D., & Swingley, W. D. (2019). Microbial community in
1361	hyperalkaline steel slag-fill emulates serpentinizing springs. Diversity, 11(7), 103.
1362	https://doi.org/10.3390/d11070103
1363	Okland, I., Huang, S., Dahle, H., Thorseth, I. H., & Pedersen, R. B. (2012). Low temperature
1364	alteration of serpentinized ultramafic rock and implications for microbial life. Chemical

1366 Olsson, J., Stipp, S. L. S., & Gislason, S. R. (2015). Element scavenging by recently formed

Geology, 318–319, 75–87. https://doi.org/10.1016/j.chemgeo.2012.05.015

- travertine deposits in the alkaline springs from the Oman Semail Ophiolite.
- 1368 *Mineralogical Magazine*, 78(6), 1479–1490.
- 1369 https://doi.org/10.1180/minmag.2014.078.6.15
- 1370 Palandri, J. L., & Reed, M. H. (2004). Geochemical models of metasomatism in ultramafic
- 1371 systems: serpentinization, rodingitization, and sea floor carbonate chimney precipitation.
- 1372 *Geochimica et Cosmochimica Acta*, 68(5), 1115–1133.
- 1373 https://doi.org/10.1016/j.gca.2003.08.006
- 1374 Paukert, A. N., Matter, J. M., Kelemen, P. B., Shock, E. L., & Havig, J. R. (2012). Reaction path
- 1375 modeling of enhanced in situ CO_2 mineralization for carbon sequestration in the
- 1376 peridotite of the Samail Ophiolite, Sultanate of Oman. *Chemical Geology*, 330–331, 86–

1377 100. https://doi.org/10.1016/j.chemgeo.2012.08.013

- 1378 Paukert Vankeuren, A. N., Matter, J. M., Stute, M., & Kelemen, P. B. (2019). Multitracer
- determination of apparent groundwater ages in peridotite aquifers within the Samail
- 1380 ophiolite, Sultanate of Oman. *Earth and Planetary Science Letters*, *516*, 37–48.
- 1381 https://doi.org/10.1016/j.epsl.2019.03.007
- 1382 Pedersen, K., Nilsson, E., Arlinger, J., Hallbeck, L., & O'Neill, A. (2004). Distribution, diversity
- and activity of microorganisms in the hyper-alkaline spring waters of Maqarin in Jordan.

1384 *Extremophiles*, 8(2), 151–164. https://doi.org/10.1007/s00792-004-0374-7

- 1385 Pfeifer, H. R. (1977). A model for fluids in metamorphosed ultramafic rocks : observations at
- 1386 surface and subsurface conditions (high pH spring waters). *Schweizerische*
- 1387 *Mineralogische Und Petrographische Mitteilungen*, 57(3), 361–396.
- 1388 https://doi.org/10.5169/seals-44441
- 1389 Pokrovsky, O. S., & Schott, J. (1999). Processes at the magnesium-bearing carbonates/solution

- 1390 interface. II. Kinetics and mechanism of magnesite dissolution. *Geochimica et*
- 1391 *Cosmochimica Acta*, 63(6), 881–897. https://doi.org/10.1016/S0016-7037(99)00013-7
- 1392 Pokrovsky, O. S., & Schott, J. (2000). Kinetics and mechanism of forsterite dissolution at 25°C
- and pH from 1 to 12. *Geochimica et Cosmochimica Acta*, 64(19), 3313–3325.
- 1394 https://doi.org/10.1016/S0016-7037(00)00434-8
- Pokrovsky, O. S., & Schott, J. (2004). Experimental study of brucite dissolution and precipitation
 in aqueous solutions: surface speciation and chemical affinity control. *Geochimica et*
- 1397 *Cosmochimica Acta*, 68(1), 31–45. https://doi.org/10.1016/S0016-7037(03)00238-2
- 1398 Rempfert, K. R., Miller, H. M., Bompard, N., Nothaft, D., Matter, J. M., Kelemen, P., Fierer, N.,
- 1399 & Templeton, A. S. (2017). Geological and geochemical controls on subsurface
- 1400 microbial life in the Samail Ophiolite, Oman. *Frontiers in Microbiology*, 8, no. 56.
- 1401 https://doi.org/10.3389/fmicb.2017.00056
- 1402 Roadcap, G. S., Kelly, W. R., & Bethke, C. M. (2005). Geochemistry of extremely alkaline (pH
- 1403 >12) ground water in slag-fill aquifers. *Groundwater*, 43(6), 806–816.
- 1404 https://doi.org/10.1111/j.1745-6584.2005.00060.x
- 1405 Robinson, K. J. (2017). *Modeling Aqueous Organic Chemistry in Experimental and Natural*1406 *Systems*. Arizona State University, Tempe, AZ.
- 1407 Russell, M. J., Daniel, R. M., Hall, A. J., & Sherringham, J. A. (1994). A hydrothermally
- precipitated catalytic iron sulphide membrane as a first step toward life. *Journal of Molecular Evolution*, *39*(3), 231–243. https://doi.org/10.1007/BF00160147
- 1410 Russell, M. J., Hall, A. J., & Martin, W. (2010). Serpentinization as a source of energy at the
- 1411 origin of life. *Geobiology*, 8(5), 355–371. https://doi.org/10.1111/j.1472-
- 1412 4669.2010.00249.x

- 1413 Russell, M. J. (2018). Green Rust: The simple organizing 'seed' of all life? *Life*, 8(3), no. 35.
- 1414 https://doi.org/10.3390/life8030035
- 1415 Sader, J. A., Leybourne, M. I., McClenaghan, M. B., & Hamilton, S. M. (2007). Low-
- 1416 temperature serpentinization processes and kimberlite groundwater signatures in the
- 1417 Kirkland Lake and Lake Timiskiming kimberlite fields, Ontario, Canada: implications for
- 1418 diamond exploration. *Geochemistry: Exploration, Environment, Analysis*, 7(1), 3–21.
- 1419 https://doi.org/10.1144/1467-7873/06-900
- 1420 Sano, Y., Urabe, A., Wakita, H., & Wushiki, H. (1993). Origin of hydrogen-nitrogen gas seeps,
- 1421 Oman. Applied Geochemistry, 8(1), 1–8. https://doi.org/10.1016/0883-2927(93)90053-J
- Schrenk, M. O., Brazelton, W. J., & Lang, S. Q. (2013). Serpentinization, carbon, and deep life. *Reviews in Mineralogy and Geochemistry*, 75(1), 575–606.
- 1424 https://doi.org/10.2138/rmg.2013.75.18
- 1425 Schulte, M., Blake, D., Hoehler, T., & McCollom, T. (2006). Serpentinization and its
- 1426 Implications for life on the Early Earth and Mars. *Astrobiology*, *6*(2), 364–376.
- 1427 https://doi.org/10.1089/ast.2006.6.364
- 1428 Schwarzenbach, E. M., Lang, S. Q., Früh-Green, G. L., Lilley, M. D., Bernasconi, S. M., &
- 1429 Méhay, S. (2013). Sources and cycling of carbon in continental, serpentinite-hosted
- 1430 alkaline springs in the Voltri Massif, Italy. *Lithos*, *177*, 226–244.
- 1431 https://doi.org/10.1016/j.lithos.2013.07.009
- 1432 Shock, E., & Canovas, P. (2010). The potential for abiotic organic synthesis and biosynthesis at
- seafloor hydrothermal systems. *Geofluids*, *10*(1–2), 161–192.
- 1434 https://doi.org/10.1111/j.1468-8123.2010.00277.x
- 1435 Shock, E. L. (1992). Chemical environments of submarine hydrothermal systems. In N. G. Holm

- (Ed.), *Marine Hydrothermal Systems and the Origin of Life: Report of SCOR Working Group 91* (pp. 67–107). Dordrecht: Springer Netherlands. https://doi.org/10.1007/978-94011-2741-7_5
- 1439 Shock, E. L., & Helgeson, H. C. (1988). Calculation of the thermodynamic and transport
- 1440 properties of aqueous species at high pressures and temperatures: Correlation algorithms
- 1441 for ionic species and equation of state predictions to 5 kb and 1000°C. *Geochimica et*
- 1442 *Cosmochimica Acta*, 52(8), 2009–2036. https://doi.org/10.1016/0016-7037(88)90181-0
- 1443 Shock, E. L., Helgeson, H. C., & Sverjensky, D. A. (1989). Calculation of the thermodynamic
- and transport properties of aqueous species at high pressures and temperatures: Standard
- 1445 partial molal properties of inorganic neutral species. *Geochimica et Cosmochimica Acta*,
- 1446 53(9), 2157–2183. https://doi.org/10.1016/0016-7037(89)90341-4
- 1447 Shock, E. L., Oelkers, E. H., Johnson, J. W., Sverjensky, D. A., & Helgeson, H. C. (1992).
- 1448 Calculation of the thermodynamic properties of aqueous species at high pressures and
- 1449 temperatures. Effective electrostatic radii, dissociation constants and standard partial
- 1450 molal properties to 1000 C and 5 kbar. *Journal of the Chemical Society, Faraday*
- 1451 *Transactions*, 88(6), 803–826. https://doi.org/10.1039/FT9928800803
- 1452 Shock, E. L., Sassani, D. C., Willis, M., & Sverjensky, D. A. (1997). Inorganic species in
- 1453 geologic fluids: Correlations among standard molal thermodynamic properties of aqueous
- ions and hydroxide complexes. *Geochimica et Cosmochimica Acta*, 61(5), 907–950.
- 1455 https://doi.org/10.1016/S0016-7037(96)00339-0
- 1456 Sleep, N. H., Meibom, A., Fridriksson, T., Coleman, R. G., & Bird, D. K. (2004). H₂-rich fluids
- from serpentinization: Geochemical and biotic implications. *Proceedings of the National Academy of Sciences*, 101(35), 12818–12823. https://doi.org/10.1073/pnas.0405289101
 - 71
manuscript submitted to Journal of Geophysical Research-Solid Earth

1459	Stanger, G. (1986). The hydrogeology of the Oman Mountains. The Open University, UK.
1460	St-Jean, G. (2003). Automated quantitative and isotopic (¹³ C) analysis of dissolved inorganic
1461	carbon and dissolved organic carbon in continuous-flow using a total organic carbon
1462	analyser. Rapid Communications in Mass Spectrometry, 17(5), 419–428.
1463	https://doi.org/10.1002/rcm.926
1464	Suda, K., Ueno, Y., Yoshizaki, M., Nakamura, H., Kurokawa, K., Nishiyama, E., Yoshino, K.,
1465	Hongoh, Y., Kawachi, K., Omori, S., Yamada, K., Yoshida, N., & Maruyama, S. (2014).
1466	Origin of methane in serpentinite-hosted hydrothermal systems: The CH ₄ -H ₂ -H ₂ O
1467	hydrogen isotope systematics of the Hakuba Happo hot spring. Earth and Planetary
1468	Science Letters, 386, 112-125. https://doi.org/10.1016/j.epsl.2013.11.001
1469	Sverjensky, D. A., Shock, E. L., & Helgeson, H. C. (1997). Prediction of the thermodynamic
1470	properties of aqueous metal complexes to 1000°C and 5 kb. Geochimica et
1471	Cosmochimica Acta, 61(7), 1359–1412. https://doi.org/10.1016/S0016-7037(97)00009-4
1472	Szponar, N., Brazelton, W. J., Schrenk, M. O., Bower, D. M., Steele, A., & Morrill, P. L. (2013).
1473	Geochemistry of a continental site of serpentinization, the Tablelands Ophiolite, Gros
1474	Morne National Park: A Mars analogue. Icarus, 224(2), 286–296.
1475	https://doi.org/10.1016/j.icarus.2012.07.004
1476	Taylor, H. P. (1974). The application of oxygen and hydrogen isotope studies to problems of
1477	hydrothermal alteration and ore deposition. Economic Geology, 69(6), 843-883.
1478	https://doi.org/10.2113/gsecongeo.69.6.843
1479	Taylor, R. M., Hansen, H. C. B., Stanger, G., & Koch, C. B. (1991). On the genesis and

- 1480 composition of natural pyroaurite. *Clay Minerals*, *26*(3), 297–309.
- 1481 https://doi.org/10.1180/claymin.1991.026.3.01

- 1482 Templeton, A. S., & Ellison, E. T. (2020). Formation and loss of metastable brucite: does Fe(II)-
- 1483 bearing brucite support microbial activity in serpentinizing ecosystems? *Philosophical*
- 1484 Transactions of the Royal Society A: Mathematical, Physical and Engineering Sciences,

1485 *378*(2165), 20180423. https://doi.org/10.1098/rsta.2018.0423

- 1486 Thom, J. G. M., Dipple, G. M., Power, I. M., & Harrison, A. L. (2013). Chrysotile dissolution
- rates: Implications for carbon sequestration. *Applied Geochemistry*, *35*, 244–254.
 https://doi.org/10.1016/j.apgeochem.2013.04.016
- 1489 Tutolo, B. M., Luhmann, A. J., Tosca, N. J., & Seyfried, W. E. (2018). Serpentinization as a
- reactive transport process: The brucite silicification reaction. *Earth and Planetary Science Letters*, 484, 385–395. https://doi.org/10.1016/j.epsl.2017.12.029
- 1492 Vacquand, C., Deville, E., Beaumont, V., Guyot, F., Sissmann, O., Pillot, D., Arcilla, C., &
- 1493 Prinzhofer, A. (2018). Reduced gas seepages in ophiolitic complexes: Evidences for
- 1494 multiple origins of the H₂-CH₄-N₂ gas mixtures. *Geochimica et Cosmochimica Acta*, 223,
- 1495 437–461. https://doi.org/10.1016/j.gca.2017.12.018
- 1496 Vance, S., Harnmeijer, J., Kimura, J., Hussmann, H., de Martin, B., & Brown, J. M. (2007).
- 1497 Hydrothermal systems in small ocean planets. *Astrobiology*, 7(6), 987–1005.
- 1498 https://doi.org/10.1089/ast.2007.0075
- Velbel, M. A. (1993). Constancy of silicate-mineral weathering-rate ratios between natural and
 experimental weathering: implications for hydrologic control of differences in absolute
 rates. *Chemical Geology*, *105*(1), 89–99. https://doi.org/10.1016/0009-2541(93)90120-8
- 1502 Waite, J. H., Glein, C. R., Perryman, R. S., Teolis, B. D., Magee, B. A., Miller, G., Grimes, J.,
- 1503 Perry, M. E., Miller, K. E., Bouquet, A., Lunine, J. I., Brockwell, T., & Bolton, S. J.
- 1504 (2017). Cassini finds molecular hydrogen in the Enceladus plume: Evidence for

- 1505 hydrothermal processes. *Science*, *356*(6334), 155–159.
- 1506 https://doi.org/10.1126/science.aai8703
- 1507 Wang, D. T., Reeves, E. P., McDermott, J. M., Seewald, J. S., & Ono, S. (2018). Clumped
- 1508 isotopologue constraints on the origin of methane at seafloor hot springs. *Geochimica et*
- 1509 *Cosmochimica Acta*, 223, 141–158. https://doi.org/10.1016/j.gca.2017.11.030
- 1510 Weyhenmeyer, C. E., Burns, S. J., Waber, H. N., Macumber, P. G., & Matter, A. (2002). Isotope
- 1511 study of moisture sources, recharge areas, and groundwater flow paths within the eastern
- 1512 Batinah coastal plain, Sultanate of Oman. *Water Resources Research*, *38*(10), no. 1184.
- 1513 https://doi.org/10.1029/2000WR000149
- 1514 Wheat, C. G., Fryer, P., Fisher, A. T., Hulme, S., Jannasch, H., Mottl, M. J., & Becker, K.
- 1515 (2008). Borehole observations of fluid flow from South Chamorro Seamount, an active
 1516 serpentinite mud volcano in the Mariana forearc. *Earth and Planetary Science Letters*,
- 1517 267(3), 401–409. https://doi.org/10.1016/j.epsl.2007.11.057
- 1518 Wheat, C. G., Seewald, J. S., & Takai, K. (2020). Fluid transport and reaction processes within a
- 1519 serpentinite mud volcano: South Chamorro Seamount. *Geochimica et Cosmochimica*
- 1520 Acta, 269, 413–428. https://doi.org/10.1016/j.gca.2019.10.037
- 1521 White, A. F., & Brantley, S. L. (2003). The effect of time on the weathering of silicate minerals:
- 1522 why do weathering rates differ in the laboratory and field? *Chemical Geology*, 202(3),
- 1523 479–506. https://doi.org/10.1016/j.chemgeo.2003.03.001
- 1524 Wolery, T., & Jarek, R. (2003). EQ3/6, Version 8.0, Software User's Manual (No. 10813- UM-
- 1525 8.0–00) (p. 376). Las Vegas, Nevada: US Department of Energy, Office of Civilian
- 1526 Radioactive Waste Management, Office of Repository Development.
- 1527 Wolery, T. J., & Jove-Colon, C. F. (2004). Qualification of thermodynamic data for geochemical

1528	modeling of mineral-water interactions in dilute systems (No. ANL-WIS-GS-000003) (p.
1529	212). Las Vegas, Nevada: US Department of Energy, Office of Civilian Radioactive
1530	Waste Management, Office of Repository Development.
1531	Yuce, G., Italiano, F., D'Alessandro, W., Yalcin, T. H., Yasin, D. U., Gulbay, A. H., Ozyurt, N.
1532	N., Rojay, B., Karabacak, V., Bellomo, S., Brusca, L., Yang, T., Fu, C. C., Lai, C. W.,
1533	Ozacar, A., & Walia, V. (2014). Origin and interactions of fluids circulating over the
1534	Amik Basin (Hatay, Turkey) and relationships with the hydrologic, geologic and tectonic
1535	settings. Chemical Geology, 388, 23-39. https://doi.org/10.1016/j.chemgeo.2014.09.006
1536	Zgonnik, V., Beaumont, V., Larin, N., Pillot, D., & Deville, E. (2019). Diffused flow of
1537	molecular hydrogen through the Western Hajar mountains, Northern Oman. Arabian

Journal of Geosciences, *12*(3), no. 71. https://doi.org/10.1007/s12517-019-4242-2



Journal of Geophysical Research: Solid Earth Supporting Information for Theoretical predictions vs environmental observations on serpentinization fluids:

Lessons from the Samail ophiolite in Oman

J. A. M. Leong^{1,2}, A. E. Howells^{1,3}, K. J. Robinson^{1,4}, A. Cox⁵, R. V. Debes II^{1,2}, K. Fecteau^{1,4}, P. Prapaipong^{1,2}, and E. L. Shock^{1,2,4}

¹Group Exploring Organic Processes In Geochemistry (GEOPIG) ²School of Earth & Space Exploration ³School of Life Sciences ⁴School of Molecular Sciences Arizona State University Tempe, AZ 85287, USA

⁵Laboratory Exploring Geobiochemical Engineering and Natural Dynamics (LEGEND) Montana Technological University Butte, MT 59701, USA

Contents of this file

Text S1 to S3 Figures S1 to S8 Tables S3 to S7

Additional Supporting Information (Files uploaded separately)

Captions for Tables S1 and S2

Introduction

Computational models used in this work and previously described in Leong and Shock (2020) are summarized in this supplementary document (Section S1). Furthermore, additional details on statistical methods applied to analytical results are also presented (Section S2). Fluids sampled from gabbroic bodies in the Samail ophiolite, which were not discussed in the main text, are described in this supporting document (Section S3). Supplementary figures and tables that depict results of analytical (Figures S1–S3, S5, and S7–S8; Tables S1 and S2), theoretical (Figures S2, S5, and S7–S8; Tables S1 and S2), theoretical (Figures S2, S5, and S7–S8; Tables S2), and statistical methods (Figure S6; Tables S3–S5) are also presented. Compositions of end-member fluids used in mixing calculations conducted in this work are compiled in Table S6.

S1. Computational Model

S1.1. Model Setup

Reactions that drive the transformation of rainwater to fluids that are reduced and hyperalkaline occur through progressive reactions with ultramafic rocks, which can be tracked using reaction-path calculations. These calculations simulate reactions of minerals with a fluid and determine the compositions of coexisting solid phases and fluid constituents attained at thermodynamic equilibrium at various extents of overall reaction progress. Progress of the alteration process is related to an increase in the rock-to-water ratio, simulating a given mass of fluid reacting with more rock as it infiltrates deeper into the subsurface. Following Cipolli et al. (2003) and Paukert et al. (2012), the model starts as a system open to atmospheric exchange and then transitions into a system closed to input from the atmosphere, simulating fluid pathways infiltrating deeper into the aquifer. Simulations were run with a fluid-centered, flow-through physical system that tracks compositions as fluid reacts with rock. For the composition of the starting fluid, a global average rainwater composition compiled by Hao et al. (2017) from Berner and Berner (2012) was used. Reacting rocks have starting compositions typical of harzburgites common in the Oman ophiolite (Hanhoj et al., 2010) and are composed mostly of olivine (>60 mol%) with some orthopyroxene (<40 mol%) and minor amounts of clinopyroxene (<5 mol%). In the initial models, inputs from primary minerals are controlled by relative dissolution rates. Calculations were then performed where experimentally-derived dissolution rates of the primary minerals are used to assess timescales needed to reach various stages of the overall serpentinization progress.

Aside from reaction paths, concentrations of solutes at a given pH in equilibrium with various serpentinization-relevant minerals (brucite, magnesite, calcite, serpentine, etc.) were calculated. Dissolved Mg concentrations in equilibrium with brucite at a given pH was determined, and predicted Mg concentrations constrained by the solubility of magnesite were also calculated. DIC levels in equilibrium with magnesite at given pH and Mg values were determined using charge balance. Similar calculations were performed to determine the dissolved Ca and DIC concentrations of fluid in equilibrium with calcite. Dissolved Si concentrations in equilibrium with serpentine (chrysotile) were determined using calculated Mg values in equilibrium with magnesite at pH <10 and brucite at pH >10.

The solubility-reaction path program EQ3/6 (Wolery and Jarek, 2003) with a customized thermodynamic database was used for these calculations. Thermodynamic data used in the calculations were calculated with the SUPCRT code (Johnson et al., 1992) using standard state thermodynamic data for aqueous species taken from Shock and Helgeson (1988), Shock et al. (1989, 1992, 1997), and Sverjensky et al. (1997), together with the revised Helgeson-Kirkham-Flowers equations of state (Shock et al., 1992). Data for minerals were mostly taken from Helgeson et al. (1978) and Wolery and Jove-Colon (2004) with the addition of estimated standard Gibbs energies of formation of Fe end-members of several serpentinization-relevant Mg- and Fe-bearing minerals. Data for these minerals were added, while maintaining internal thermodynamic consistency, to explore solid solution behavior during serpentinization. These data, including discussion of data sources, can be found in Leong and Shock (2020), which includes a wider range

of model results that encompass the effects of variable temperatures (0–100°C), compositions of reacting ultramafic rocks, fluid salinity, and buffering capacities of serpentinization-relevant minerals.

S1.2. Model Results: Stages of the serpentinization reaction progress

Results of calculations simulating the serpentinization of an ultramafic rock of the composition 85% olivine (Forsterite₉₀, Mg_{1.8}Fe_{0.2}SiO₄), 14% orthopyroxene (Enstatite₉₀, Mg_{0.9}Fe_{0.1}SiO₃), and 1% clinopyroxene (Diopside₉₅, CaMg_{0.95}Fe_{0.05}Si₂O₆) at ambient conditions (25°C, 1 bar) are shown in Figure S4. Results of simulations with other olivine-rich compositions yield similar trends as demonstrated by Leong and Shock (2020). Consequences of variable temperatures (0–100°C) and increasing pyroxene abundance on the compositions of the coexisting fluid and solid precipitates are also described in detail in the same study. The calculated increase in pH with reaction progress is shown in Figure S4a, and changes with increasing pH in the total concentrations of Ca, Mg, Si, Fe, and dissolved inorganic carbon (DIC), as well as the number of moles of secondary minerals produced or consumed, are depicted in Figures S4b and S4c, respectively. Calculated reaction paths are depicted in activity diagrams that show the stabilities of minerals relative to the activities of aqueous solutes in the MgO-SiO₂-H₂O (Figure S4d) and CaO-MgO-SiO₂-H₂O (Figure S4e) systems.

The predicted reaction path shown in Figure S4 can be divided into six general stages as indicated by the circled numbers that depict the starting points of these various stages. These stages are discussed in detail by Leong and Shock (2020) and summarized below where the phrases in italics encapsulate each stage.

(1) The dissolution of primary minerals into rainwater begins when rainwater reacts with ultramafic rocks that are formed in the Earth's mantle. When transported to the surface through tectonic events, ultramafic rocks are unstable and are readily altered in the presence of aqueous fluids. Rain is slightly acidic and contains protons derived from atmospheric $CO_{2(q)}$ via

$$CO_{2(g)} + H_2 0 \Leftrightarrow HCO_3^- + H^+$$
(S1)

which drive the dissolution of primary minerals present in ultramafic rocks through hydrolysis reactions. Consumption of protons during mineral hydrolysis increases pH as illustrated in Figure S4a. Dissolution of minerals yields solutes to the fluid, driving an increase in the total Mg, Si, and Ca concentrations, as depicted in Figure S4b. The fluid pathway is open to atmospheric exchange and consequently, reaction S1 dictates that the DIC must increase with pH to maintain equilibrium with a constant atmospheric $fCO_{2(g)}$ of ~10^{-3.5}. The increase in DIC during this stage is depicted in Figure S4b. As shown in Figure S4c, no minerals except for an oxyhydroxide (goethite) precipitate at this stage. Also shown in the activity diagrams depicted in Figures S4d and S4e, the increase in pH and total Mg, Si, and Ca will drive the reaction path toward higher $aMg^{+2}/(aH^+)^2$ and $aCa^{+2}/(aH^+)^2$ (activity ratios) as well as higher $aSiO_{2(aq)}$ (silica activity). This stage marks the transition from a slightly acidic and dilute rainwater to a circumneutral (pH 7-8) fluid enriched in Mg^{+2} and HCO₃⁻.

(2) The *onset of serpentinization* occurs as the solution is concentrated enough to precipitate serpentine, as represented by the mineral chrysotile. Quartz and talc can also be saturated at this stage, but the formation of these minerals was suppressed in the

model to evaluate the maximum silica activity and total Si concentration attained when serpentine saturation is achieved, which occurs when the reaction path reaches the chrysotile saturation line as shown in Figure S4d. The precipitation of chrysotile from solutes, as depicted by the reaction

 $3Mg^{+2} + 2SiO_{2(aq)} + 5H_2O \rightarrow Mg_3Si_2O_5(OH)_{4(chrysotile)} + 6H^+$, (S2) consumes Si and causes a decrease in its concentration as depicted in Figure S4b. Precipitation of chrysotile also causes the reaction path depicted in Figure S4d to shift to lower silica activity along the chrysotile saturation line. Despite the consumption of Mg through reaction (S2), continuing dissolution of olivine-dominated rocks results in a surplus of Mg with respect to Si consumed during serpentinization as depicted by

 $2Mg_2SiO_{4(\text{forsterite})} + 2H^+ + H_2O \rightarrow Mg_3Si_2O_5(OH)_{4(\text{chrysotile})} + Mg^{+2}$, (S2) leading to further increases in aqueous Mg at this stage. The pH continues to increase despite the release of protons during serpentine formation (reaction S2), as it is not enough to counter their consumption during the dissolution of primary minerals as exemplified by reaction (S3). Nevertheless, in addition to reaction (S1), the release of protons through precipitation reactions (e.g., reaction S2) can drive further dissolution and thus overall reaction progress.

(3) As fluids infiltrate deeper into the subsurface, a *transition from an open to a closed system* occurs. As shown in Figure S4a, a sharp increase in pH occurs once the atmosphere no longer supplies protons via reaction (S1). The increase in pH, as well as the increasing concentrations of Mg and Ca, will result in the *precipitation of carbonates*. As shown in Figure S4c, the first carbonate to precipitate is dolomite followed by magnesite, the latter forming more abundantly than the former. As shown in Figure S4b, the precipitation of these Mg- and Ca-bearing carbonates drives a decrease in the DIC and the total Ca and Mg concentrations.

(4) *Brucite saturation* is attained once the pH is high enough (pH ~ 10). As shown in Figure S4d, brucite precipitation occurs at extremely low $aSiO_{2(aq)}$ where the trend of decreasing $aSiO_{2(aq)}$, observed since the onset of serpentinization at Stage 2, stops at the intersection of the brucite and chrysotile saturation lines. In Figure S4e, this corresponds to where the reaction path encounters the boundary between the chrysotile and brucite stability fields. From this point onward in overall reaction progress, the $aSiO_{2(aq)}$ of the solution will be fixed by an equilibrium assemblage of chrysotile and brucite (CtI-Brc) depicted by the reaction

$$BMg(OH)_{2(brucite)} + 2SiO_{2(aq)} \Leftrightarrow Mg_3Si_2O_5(OH)_{4(chrysotile)} + H_2O.$$
(S4)

Despite the $aSiO_{2(aq)}$ being fixed by reaction (S4), the speciation of Si allows for an increase in total dissolved Si, as depicted in Figure S4b, because the abundance of $HSiO_3^-$ and consequently the total Si concentration increases with increasing pH to maintain equilibrium with the reaction

$$\text{SiO}_{2(aq)} + \text{H}_20 \Leftrightarrow \text{HSiO}_3^- + \text{H}^+.$$
 (S5)

The precipitation of brucite causes an abrupt decrease in the total Mg concentration, as depicted in Figure S4b. Together with deceasing DIC, decrease in Mg concentration causes magnesite and, eventually, dolomite to stop precipitating as shown in Figure S4c. On the other hand, the total Ca concentration increases as more Ca is

released into the fluid from the dissolution of clinopyroxene relative to its uptake through the precipitation of dolomite. This stage marks the transition from a fluid enriched in Mg and DIC to an increasingly alkaline and Ca-rich solution.

(5) In addition to becoming more alkaline, *fluids can become more reduced* as H_2 is generated from the oxidation of ferrous iron from the primary minerals to ferric iron in secondary minerals coupled with the reduction of water to H_2 as depicted by the generalized reaction

 $2\text{Fe}(\text{II})O_{(\text{primary minerals})} + H_2O \rightarrow \text{Fe}(\text{III})_2O_{3(\text{secondary minerals})} + H_{2(aq)}.$ (S6) At these reduced conditions, the main iron-hosting secondary mineral transitions from goethite to magnetite (Figure S4c). The results shown in Figure S4 are from a calculation without considering the incorporation of Fe into solid solutions of serpentine and brucite. If allowed, formation of magnetite is less favored, and most of the iron is incorporated into ferrous and ferric iron-bearing serpentine and ferrous iron-bearing brucite (see McCollom and Bach, 2009; Klein et al., 2009; 2013; 2014; Leong and Shock, 2020). Formation of methane is also favored at these reduced conditions. When not kinetically inhibited, the formation of methane will drive DIC to extremely low concentrations (<10⁻¹² molal). If abiogenic methane formation is kinetically inhibited, as it is likely to be at low-temperature conditions (Shock, 1992; McCollom, 2016; Wang et al., 2018) or in gas-poor systems (McCollom, 2016; Etiope and Whiticar, 2019), DIC can persist in solution at low but detectable concentrations (~10 µmolal) as depicted in Figure S4b. Increasing pH and Ca concentration will saturate calcite (Figure S4c). As the DIC concentration is minimal at this stage of the reaction progress, the amount of precipitating calcite will not be enough to counter the increasing Ca concentration brought about by the increasing amount of clinopyroxene dissolved. With increasing pH and total Ca concentration, the reaction path depicted in Figure S4e follows an increasing trajectory in $aCa^{+2}/(aH^{+})^{2}$ along the boundary between the chrysotile and brucite stability fields until it reaches equilibrium with diopside.

(6) Chrysotile-Brucite-Diopside (Ctl-Brc-Di) equilibrium is attained when the reaction path depicted in Figure S4e reaches the point where the stability fields of chrysotile, brucite and diopside meet. So far, since the beginning of Stage 1, release of protons through precipitation reactions is not enough to counter its consumption through dissolution reactions, hence pH continues to increase. Throughout overall reaction progress, releases of Mg and Si through the dissolution of primary minerals are mostly incorporated into precipitating serpentine, brucite, magnesite, and dolomite. On the other hand, especially during the latter stages of overall reaction progress, dissolved Ca will continue to accumulate in the fluid despite its incorporation into dolomite and calcite owing to the very low amount of DIC remaining in solution. The release of Ca⁺² into the solution from the dissolution of diopside is accommodated by the loss of H⁺ (or depending on how the precipitation and dissolution reactions are written, release of OH⁻) and causes the continuous increase in pH. The increasing pH trend stops when the fluid reaches equilibrium with diopside, as shown by point 6 in the activity diagram depicted in Figure S4e. At this point, diopside dissolution is no longer favorable. Moreover, the equilibrium assemblage of chrysotile, brucite and diopside (Ctl-Brc-Di), depicted by the reaction

$CaMgSi_2O_{6(diopside)} + 2Mg(OH)_{2(brucite)} + 2H^+ \Leftrightarrow Ca^{+2} +$

$$Ig_3Si_2O_5(OH)_{4(chrysotile)} + H_2O$$
,

corresponds to an invariant point in the CaO-MgO-SiO₂-H₂O system. Thus, not only the pH (Figure S4a) but also the Ca, Mg, and Si concentrations (Figure S4b) will remain unchanged despite continuing reaction progress, until diopside is completely reacted. As Ctl-Brc equilibrium (Stage 4, Figure S4d) ensures that the fluids stay undersaturated or out of equilibrium with forsterite (olivine) and enstatite (orthopyroxene) at low temperatures, fluids attaining equilibrium with diopside (clinopyroxene) are as close as possible to being in equilibrium with ultramafic rocks, and thus define the final stage, Stage 6, of overall serpentinization progress. Reaction S7 also constrains the maximum pH allowed in serpentinizing environments (~ 12.2 at 25°C and 1 bar), explaining why extremely alkaline environments (pH >13) are rare during ultramafic weathering (Leong & Shock, 2020).

Trends depicted in Figure S4 apply to reactions with olivine-rich lithologies. Greater contributions from orthopyroxene will generate fluids that first equilibrate with chrysotile and diopside before attaining brucite saturation. In the activity diagram depicted in Figure S4e, these reaction paths will track along the chrysotile-diopside boundary before reaching the Ctl-Brc-Di invariant point. Eventually, significant contributions from orthopyroxene (50-60 mole %) will yield fluids that end up at the chrysotile-tremolite-diopside (Ctl-Tr-Di) invariant point. Even greater (>60% mole %) orthopyroxene contribution will yield fluids in equilibrium with chrysotile-talc-tremolite (Ctl-Tlc-Tr). The latter two equilibrium assemblages imply higher aSiO_{2(aq)} and lower $aCa^{+2}/(aH^{+})^{2}$ activity ratios compared to Ctl-Brc-Di equilibrium, as depicted in Figure S4e, generating more Si-rich and less alkaline fluids. Dissolution of secondary minerals such as serpentine and brucite and relict clinopyroxene in partially serpentinized ultramafic rocks can also lead to Ctl-Brc-Di equilibrium, especially if brucite is still abundant. Lower amounts of reacting brucite can lead to reaction paths ending at the Ctl-Tr-Di or the Ctl-TIc-Tr equilibrium points as in orthopyroxene-rich settings. Further details on the consequences of variable compositions of the reacting rocks on the compositions of fluids generated during low temperature and high temperature alteration of ultramafic rocks can be found in Leong and Shock (2020) and Klein et al. (2013), respectively.

S2. Statistical Analyses

Statistical analyses were conducted to evaluate if compositional differences between different fluid types are significant. All calculations were performed using the Past *v4.03* statistical software (Hammer et al., 2001). Non-parametric Mann-Whitney Utests were conducted to evaluate differences in the median values of dissolved species measured from hyperalkaline fluids (pH > 11) collected at sites near or far from the basal thrust of the Samail ophiolite. The U-values, z-score, and *p*-values calculated using the Mann-Whitney U-test are compiled in Figure S2. Results of calculations show that the Cl, Na, and K concentrations of fluids collected from sites near and far from the basal thrust are significantly different at *p* < 0.01. Results from the Mann-Whitney U-test calculations, however, show that the DIC concentrations of fluids hosted near the basal thrust, which

(S7)

are typically higher, are significantly different (at p < 0.05) from those hosted far from the basal thrust.

Multivariate statistical analyses (analysis of similarities or ANOSIM, at 9,999 permutations) were conducted to evaluate if the O and H stable isotopic compositions of fluids sampled from different study sites are distinct from each other. Results of calculations are shown in Table S4, where study sites are arranged according to their locations. Most sites have isotopic compositions that are not significantly distinct, though the stable isotopic compositions of water at a given site can be significantly different from another site that is geographically further away. Samples collected from Shumayt and Al Bana have stable isotopic compositions that are the most distinct amongst the study sites.

Both Non-metric Multidimensional Scaling (NMDS) and ANOSIM were conducted to evaluate distinctions among various fluid types informed by simulations of subsurface reaction pathways and subsequent mixing processes. The pH and the square root of the molal concentrations of Mg, Si, Ca, DIC, Na, K, and Cl of investigated fluids identified as Type 1, intermediate, Type 2 and mixed in Figure 7 were used in the statistical analyses. Results of NMDS calculations are shown in a scatter plot in Figure S6 and *p*-values derived from ANOSIM calculations (at 9999 permutations) are summarized in Table S5. Both analyses use the Euclidian similarity matrix. Stress value from the NMDS is low at 0.0043.

S3. Differences in the compositions of ultramafic- and gabbro-hosted fluids

Most simulations of gabbro alteration are focused at high temperature and pressure conditions such as those occurring in submarine hydrothermal systems (McCollom and Shock, 1998; Palandri and Reed, 2004; Bach and Klein, 2009; Bach and others, 2013, Seyfried and others, 2015). Predictions of fluid compositions arising during interactions between meteoric water and gabbroic rocks in the continental subsurface are far scarcer. Gabbroic rocks comprise approximately a third of the surface exposures of the Samail ophiolite (Nicolas et al., 2000) and are also common in many other ophiolites. Gabbro-hosted hyperalkaline fluids are also documented in several studies (Neal and Stanger, 1985; Dewandel et al., 2005; Paukert et al., 2012; Chavagnac et al., 2013b; Cardace et al., 2013; Rempfert et al., 2017), and many of these springs occur near the gabbro-ultramafic contact. The influence of ultramafic rocks underlying the gabbroic rocks on the compositions of hyperalkaline fluids seeping out from gabbroic outcrops is largely unknown.

Reported compositions of gabbro-hosted fluids in the Samail ophiolite, together with those from this work, are plotted as orange symbols in Figure S7, where they can be compared with fluids documented in ultramafic environments (grey symbols). Compositions of most gabbro-hosted fluids overlap those hosted in ultramafic rocks. However, there are some notable differences that could be attributed to fluid-mineral reactions enabled when fluids encounter gabbroic rocks. Mineral solubility calculations were conducted to provide preliminary insights into how various minerals or mineral assemblages can account for these aqueous trends. Solubility calculations for 25°C and 1 bar were conducted using the same software package and thermodynamic data described in the method section of this paper. Results of calculations are plotted as red curves in Figure S7.

Gabbro- and ultramafic-hosted Type 1 fluids (circumneutral, pH 7-9) have overlapping total dissolved Si values as shown in Fig S7a. In contrast, some of the highest dissolved Si values measured in Type 2 fluids (hyperalkaline, pH > 11) are from gabbro-hosted fluids. This could be attributed to the higher silica content of gabbros, which can lead to the precipitation of secondary minerals with higher silica contents than serpentine and brucite such as talc, quartz, chlorite, zeolites and clay minerals. As shown in Figure S7a, all fluids hosted in either gabbroic or ultramafic rocks are below amorphous silica saturation. Type 1 fluids hosted in both lithologies are typically above guartz saturation while Type 2 fluids are usually well below saturation with respect to quartz and the chrysotile-talc equilibrium assemblage. Type 2 fluids hosted in ultramafic rocks approach equilibrium with serpentine and brucite. In contrast, despite overlapping with fluids hosted in ultramafic rocks as shown in Figure S7a, the Si concentrations of gabbro-hosted fluids do not approach the low values dictated by chrysotile-brucite equilibrium. This trend is likely to be attributed to the geological setting of gabbrohosted hyperalkaline springs, many of which occur near the gabbro-ultramafic lithological contact. As gabbros are stratigraphically above peridotites in the Samail ophiolite, it is likely that deep-seated fluids would have first reacted with ultramafic rocks before encountering gabbroic rocks on their way to discharge at the surface. Such a scenario is analogous to ultramafic to gabbro mass transfer processes at high temperatures that have been proposed to facilitate the formation of rodingites (Bach and Klein, 2009; Bach et al., 2013).

As shown in Figure S7b, there are no noticeable differences in the Mg concentrations of Type 1 fluids hosted in either ultramafic or gabbroic rocks. As in the case of ultramafic-hosted fluids, Mg concentrations of Type 1 gabbro-hosted fluids rarely fall below values dictated by magnesite solubility. On the other hand, there are some noticeable differences between the Mg concentrations of Type 2 hyperalkaline fluids in these two lithological environments. Some of the most Mg-depleted Type 2 fluids were sampled from gabbroic-hosted springs. Unlike most ultramafic-hosted samples, some gabbro-hosted samples have Mg values below those set by brucite solubility. These lower Mg concentrations could be attributed to the precipitation of relatively Si-rich Mg-silicate minerals (e.g., talc, saponite, chlorite) that commonly occur in altered mafic rocks. As an example, as depicted in Figure S7b, lower Mg concentrations can be attained if fluids are in equilibrium with chrysotile and talc. As with Si, more constraints can be provided on the Mg concentrations of fluids dictated by the solubility of aluminumbearing silicates such as chlorite, zeolite and clay minerals that are commonly observed in gabbroic rocks altered at low temperatures (Bach et al., 2001).

Dissolved Ca concentrations of Type 2 fluids from both gabbro and ultramafic bodies have similar trends that follow the solubility of calcite as shown in Figures 5c and S4c. However, total calcium concentrations in Type 1 fluids are somewhat more enriched in gabbro-hosted fluids than ultramafic-hosted fluids. Type 1 fluids, as discussed in the main text of this work, are products of early stages of reaction progress dictated by the dissolution of protolith minerals. Gabbroic rocks are more enriched in Ca compared to

ultramafic rocks due to the more abundant presence of Ca-plagioclase (anorthite) and Ca-clinopyroxene (diopside).

Like Ca, dissolved Na concentrations of Type 1 fluids hosted in gabbros are elevated compared to those hosted in ultramafic rocks as shown in Figure S7d. This could be attributed to the relatively elevated Na content of gabbroic rocks. However, similar observations can also be made for the dissolved Cl values of gabbro-hosted fluids plotted in Figure S7e, suggesting that leaching from salt minerals could be an alternative source for both Na and Cl. Figure S7f, however, shows no apparent differences between K trends of both gabbro- and ultramafic-hosted fluids, suggesting that leaching could be attributed only to halite (NaCl). In addition, leaching from other Na-bearing phases (*e.g.*, plagioclase and pyroxenes) can also occur concomitantly, as gabbro-hosted Type 1 fluids are typically more enriched in Na relative to Cl compared to Type 1 fluids hosted in ultramafic rocks as shown in Figure S7g.

As shown in Figure S7h, the total dissolved aluminum concentrations of Type 1 gabbro-hosted fluids overlap with those hosted in ultramafic rocks, but gabbro-hosted Type 2 fluids have elevated total dissolved Al values, closely following those dictated by the solubility of gibbsite, as shown in Figure S7h. In contrast, all ultramafic-hosted Type 2 fluids remain highly undersaturated with respect to gibbsite, which is not surprising as ultramafic rocks are typically depleted in Al. Chavagnac and others (2013b) documented Al-bearing layered double hydroxides such as hydrotalcite and indigirite in gabbro-hosted hyperalkaline pools marked by elevated dissolved Al content.

Further insight into the underlying mineral controls on the compositions of gabbro-hosted fluids, in comparison to those hosted in ultramafic rocks, can be revealed through mineral activity diagrams. Activities of several aqueous species (SiO_{2(aq)}, Ca⁺², H⁺) were evaluated from the measured values using the same thermodynamic database as all other calculations in this study and are plotted in an activity diagram for the CaO-Al₂O₃-SiO₂-H₂O (phase boundaries as orange lines, balanced on Al₂O₃) and CaO-MgO-SiO₂-H₂O system (black lines, balanced on MgO) systems shown in Figure S8. Calculated activities $(aCa^{+2}/(aH^{+})^2, aSiO_2)$ of gabbro- (orange symbols) and ultramafic-hosted (grey symbols, values similar to Figure 5f) fluids are overlain in the activity diagram. Gabbrohosted hyperalkaline fluids typically have similar calcium to hydrogen activity ratios $(aCa^{+2}/(aH^{+})^{2})$ as those hosted in ultramafic bodies but are characterized by having somewhat higher silica activities (aSiO₂). As shown in Figure S8, these gabbro-hosted fluids seem to be drawn to equilibrium with gibbsite, grossular (garnet), and prehnite, unlike ultramafic-hosted fluids that approach equilibrium with chrysotile and brucite resulting in lower aSiO₂. Similarly, Type 1 fluids hosted in gabbroic rocks mostly have overlapping $aCa^{+2}/(aH^{+})^{2}$ with those hosted in ultramafic rocks but tend to have higher aSiO₂. These gabbro-hosted fluids are within the stability field of scolecite and beidellite, which are minerals that belong to the zeolite and smectite groups, respectively. In addition to ultramafic rocks, the Oman Drilling Project (Kelemen et al., 2013) has recovered gabbroic rocks. Examination of low-temperature secondary phases from recovered cores would provide constraints on which minerals dictate the evolution of fluids during fluid-gabbro interactions. Thermodynamic simulations of low-temperature fluid-gabbro interactions and eventually testing these models with more comprehensive

and in-depth investigations of gabbro-hosted fluids will bridge our current knowledge gap in this poorly explored fluid-rock system. Gabbros are the most common rocks that comprise the oceanic crust and are also common in ophiolites. Results of this future work can potentially inform future biogeochemical studies on the likelihood of these vast lithological systems to host habitable environments.



Figure S1. Comparison of analytical results of dissolved Ca (a) and Mg (b) determined through the ICP-MS and ICP-OES, respectively, versus that determined through the IC.



Figure S2. (a) Total dissolved solutes (TDS, solid black line) and ionic strength (IS, dashed black line) of sampled fluids (values at right of plot), arranged with increasing pH from left to right. Component species of the total dissolved solutes (in mole %) are represented by the colored field. Also shown are component species that comprise the total dissolved Si (b), Mg (c), Ca (d), and inorganic carbon (e).



Figure S3. Dissolved Na and Cl concentrations of hyperalkaline fluids (pH >11) sampled at study sites close to the basalt thrust (blue circles) and those sampled far within the ophiolite (brown circles, mostly near the peridotite-gabbro contact).



Figure S4. Reaction paths simulating serpentinization of an ultramafic rock with the composition 85% olivine-14% orthopyroxene-1% clinopyroxene. (a) depicts the increase in pH as serpentinization progresses, highlighting six stages discussed in the text. Progress of the serpentinization process is depicted in rock-to-water ratio (*e.g.*, log value of -3 represents reaction of 1 g of reacted ultramafic rock in 1000 g water). (b) and (c) depict changes in the concentrations of aqueous species (in log molality) and amount of minerals precipitated (in log moles) with pH, respectively. Reaction paths are plotted on activity diagrams for the (d) MgO-SiO₂-H₂O and (e) CaO-MgO-SiO₂-H₂O systems. Dashed lines in (d) and (e) indicate saturation of indicated minerals, while solid lines in (e) represent boundaries for the stability fields of minerals. Dark blue curves show results of reaction path calculations. Numbers indicate the starting points of the six stages of serpentinization.



Figure S5. Plot showing the Mg versus Si concentrations of sampled fluids, as well as the mixing path described in Figure 5. Symbols shown are similar to that depicted in Figures 3, 5, 8, and 9. In addition, the horizontal gray dashed lines depict saturation line for brucite, given the pH indicated above these lines. Note that in cases where the Si concentrations of fluids exceed that depicted by the brucite-chrysotile equilibrium line (lower left dashed black curve), brucite is metastable with respect to chrysotile, talc and quartz (not shown). At pH = 9, quartz can be saturated when Si concentrations. Note that while Mg concentrations of samples tend to plot from the mixing line towards the brucite saturation line at variable pH, the Si concentrations that are far lower than that depicted in the mixing curve are rare. A few samples with low Si (<10⁻⁵ molal) and high Mg (>10⁻³ molal) concentrations can be accounted for by intermediate stages of the subsurface serpentinization pathway (see dark blue curve in Figure 5a).



Figure S6. Results of Non-metric Multidimensional Scaling (NMDS) ordination depicted in a scatter biplot. A circle depicts an individual sample, and its color represents its classification as shown in Figure 7. Samples that are more similar to one another are ordinated closer together. The axes as well as the orientation of the plot are arbitrary. Stress value is low at 0.0043.



Figure S7. Comparison between ultramafic-hosted (grey symbols) and gabbroic-hosted (orange symbols) fluids in Oman. Symbol shapes and data sources are the same as in Figure 3. Trends in the total dissolved Si (a), Mg (b), Ca (c), Na (d), Cl (e), K (f), ratio between total dissolved Na and Cl (g), and total dissolved Al (h) with pH are shown. Dashed red lines and curves show solubility conditions for various minerals: Am – amorphous silica, Qz – quartz, Ctl – chrysotile, Tlc – talc, Brc – brucite, Mgs – magnesite, Cal – calcite, Gbs – gibbsite.



Figure S8. Log $aCa^{+2}/(aH^{+})^2 vs aSiO_2$ of gabbro-hosted fluids (various symbols in orange) plotted on an activity diagram for the CaO-Al₂O₃-SiO₂-H₂O system (phase boundaries as orange lines) at 25°C, 1 bar. Underlying it is an activity diagram for the CaO-MgO-SiO₂-H₂O system (black lines) and plotted activities of ultramafic-hosted fluids (various symbols in grey), similar to those shown in Figure 5f. Symbols are the same as in Figure S1. Note that the lines represent stability boundaries between the indicated minerals: Brc – brucite; Ctl – chrysotile; Tlc – talc; Tr – tremolite; Di – diopside; Qz – quartz; Am – amorphous SiO₂, Gbs – gibbsite; Kln – kaolinite; Prl – pyrophyllite; Bei – beidellite; Sco – scolecite: Lmt – laumontite; Prh – prehnite; Grs – grossular; Ka – katoite. Thermodynamic properties of minerals are taken from Helgeson et al. (1978) and Wolery and Jove-Colon (2004).

Table S1. Field and laboratory measurements for samples collected in this study.

Table S2. Saturation states of various relevant minerals.

	Mann-Whitney U-test				
Species	U value	z-score	p-value		
Cl	294	5.5143	< 0.0001		
Na	292	5.5313	< 0.0001		
Κ	248	5.9055	< 0.0001		
Si	824	1.0076	0.31363		
Mg	713	1.9516	0.05099		
Ca	836	0.90564	0.36513		
DIC	641	2.2533	0.02424		
pН	727	1.8325	0.06687		

Table S3. Results of Mann-Whitney U-test of hyperalkaline fluids (pH > 11) sampled closed to or far from the ophiolite's basalt thrust.

Sites	NSHQ14	Dima	Qafifah	Falaij	Al Hilayw	Misbit	SJA	WDA	Al Bana	Shumayt	Sudari
NSHQ14		1	1	1	1	1	1	1	1	0.0495	1
Dima	1		1	1	1	1	1	1	1	0.011	1
Qafifah	1	1		1	0.4675	0.033	0.1925	1	0.0055	0.0055	0.275
Falaij	1	1	1		0.4345	0.0055	0.022	1	0.0055	0.0055	0.0165
Al Hilayw	1	1	0.4675	0.4345		1	1	1	0.0165	0.011	1
Misbit	1	1	0.033	0.0055	1		1	1	0.055	0.022	1
SJA	1	1	0.1925	0.022	1	1		1	0.198	0.11	1
WDA	1	1	1	1	1	1	1		1	0.011	1
Al Bana	1	1	0.0055	0.0055	0.0165	0.055	0.198	1		0.0055	0.7975
Shumayt	0.0495	0.011	0.0055	0.0055	0.011	0.022	0.11	0.011	0.0055		0.033
Sudari	1	1	0.275	0.0165	1	1	1	1	0.7975	0.033	

Note 1: Sites are arranged in this table according to their location, with those sampled from the southeastern part of the ophiolite (starting with NSHQ14) at the top/left to those sampled in the northern part of the ophiolite at the bottom/left.

Note 2: p-values in blue indicate sites where isotopic compositions are significantly different at p < 0.05.

Table S4. *p*-values derived from ANOSIM calculations of the water isotopic compositions of fluids collected from various sites.

Fluid Type	circumneutral	intermediate	mixed	hyperalkaline
circumneutral		0.0618	0.0006	0.0006
intermediate	0.0618		0.0006	0.0024
mixed	0.0006	0.0006		1
hyperalkaline	0.0006	0.0024	1	

Note: *p*-values indicating statistical significance at p < 0.05 are marked as blue.

Table S5. Bonferroni-corrected p-values derived from ANOSIM between different fluid types.

	Surface Fluid (s) ¹	Deep Subsurface Fluid 1 (d ₁) ²	Deep Subsurface Fluid 2 $(d_2)^3$	Deep Subsurface Fluid 3 (d ₃) ⁴
pН	7.9	12.3	11.7	11.1
Na (mmolal) ⁵	1.14	10	10	10
Ca (mmolal)	0.56	14.4	3.2	0.7
Mg (µmolal)	2810	0.03	0.2	2.6
Cl (mmolal) ⁵	1.28	10	10	10
Si (µmolal)	303	1.3	0.3	0.1
DIC (µmolal)	5120	8	10	20

¹composition taken from sample 140116B. Sample is closest to predicted compositions in equilibrium with secondary phases common in the shallow subsurface.

²fluid in equilibrium with the Ctl-Brc-Di and calcite assemblage at 25°C and 1 bar.

³fluid in equilibrium with the Ctl-Brc-Cal assemblage, given 10 μ molal DIC, at 25°C and 1 bar.

⁴fluid in equilibrium with the Ctl-Brc-Cal assemblage, given 20 μ molal DIC, at 25°C and 1 bar.

⁵Deep fluids were given Na and Cl concentrations of 10 mmolal as an average of the range of concentrations (3-20 mmolal) observed in environmental samples

Table S6. Composition of fluids used in mixing calculations.