Dataset for H₂, CH₄ and organic compounds formation during experimental serpentinization

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Abstract
Serpentinization refers to the alteration of ultramafic rocks that produces serpentines and secondary (hydr)oxides under hydrothermal conditions. Serpentinization can generate H₂, which in turn can potentially reduce CO/CO₂ and produce organic molecules via Fischer–Tropsch type (FTT) and Sabatier type reactions. Over the last two decades, serpentinization has been extensively studied in laboratories, mainly due to its potential applications in prebiotic chemistry, origin of life in extreme environments, development of carbon-free energies and CO₂ sequestration. However, the production of H₂ and organics during experimental serpentinization is hugely variable from one publication to another. The experiments span over a large range of pressure and temperature conditions, and starting compositions of fluid and solid phases are also highly variable, which collectively adds up to more than a hundred variables and leads to controversial results. Therefore, it is extremely difficult to compare results between studies, explain their variability and identify key parameters controlling the reactions. To overcome these limitations, we collected and analysed 30 peer-reviewed articles including over 100 experimental parameters and ca. 30 mineral and organic products, hence building up a database can be completed and

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Dataset
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1 | INTRODUCTION

Earth’s mantle is predominantly composed of peridotites, a type of rock made of Mg- and Fe-rich silicate minerals such as olivine and pyroxene. Though modern Earth's crust is not ultramafic, plate tectonics bring ultramafic rocks to the surface of Earth at various geologic settings, such as slow mid-ocean ridges (e.g. Mid Atlantic Ridge, Southwest Indian Ridge), oceanic transform faults (e.g. Vema OTF in the Atlantic, São Pedro and São Paulo Archipel OTFs in the Equatorial Atlantic, Shaka and Prince Edward OTFs in the South-West Indian Ocean) and at convergent margins (e.g. Oman, New Caledonia). The alteration and hydration of peridotite result in the formation of serpentine group minerals (e.g. lizardite, chrysotile and antigorite) and secondary (hydr)oxides (e.g. brucite, magnetite). These serpentine-forming reactions are called serpentinization. During serpentinization, the ferrous iron in olivine and pyroxene is often oxidized to ferric iron, which produces H₂ through the reduction of water. As a consequence, CO or CO₂ could be reduced by H₂ through Sabatier type (R1) or Fischer–Tropsch type (R2) reaction to form CH₄ and/or other organic compounds.

\[
4H_2 + CO_2 = CH_4 + 2H_2O \quad (1)
\]

\[
(3n + 1)H_2 + nCO_2 = C_nH_{2n+2} + 2nH_2O \quad (2)
\]

Analyses of many natural samples have shown abundant release of H₂, CH₄ and other organic compounds in fluids from natural serpentinization areas and questioned the exact reactions mechanisms involved (Barnes et al., 1967; Wenner and Taylor, 1973; Charlou et al., 2002; Proskurowski et al., 2006, 2008; Konn et al., 2009). For instance, ultramafic rocks were abundant on the primitive Earth and possibly other rocky planetary bodies (Ehmann et al., 2010; Zahnle et al., 2011; Holm et al., 2015; Etiope et al., 2018), so their observation raises several major scientific questions related to serpentinization (Sleep et al., 2004; Schulte et al., 2006; Russell et al., 2010; Hellevang et al., 2011; Guillot and Hattori, 2013; Mayhew et al., 2013; McCollom and Seewald, 2013; Brazil, 2017; Ménez et al., 2018): What is the role of serpentinization in the origin of life—on Earth, and elsewhere? Could the serpentinization reaction sustain microbial communities in the primitive and modern ocean? Could our modern societies use the H₂ produced by serpentinization reactions to help reduce anthropogenic CO₂ emission?

To address these questions, there is an urgent need to understand the serpentinization process and more specifically its capacity to generate reducing conditions and produce abiotic organics. Therefore, tens of experiments have attempted to provide answers to this question. Although they all agree on the production of H₂ by serpentinization (Sleep et al., 2004; Seyfried et al., 2007), the production of CH₄ and more complex hydrocarbons (e.g. C₂H₆, C₃H₈) via FTT or Sabatier reactions has always been and is still highly debated (e.g. Evans et al., 2013; McCollom and Seewald, 2013; McCollom et al., 2015). Despite the tremendous collaborative efforts of the community all over the world, we still do not fully understand why similar experimental incentives lead to so contrasted, if not contradictory results. A strong limitation is that those numerous serpentinization experiments have been run under very different conditions using various creative protocols. In order to understand the similarities and discrepancies between results, it requires us to compare more than a hundred variables not even fully identified before the present study. Therefore, we carefully read 30 peer-reviewed publications that described experimental serpentinization and other publications for comparison, analysed 195 experiments and compiled parameters in the dataset described in Section 2. This dataset will be continuously updated as new results become available and used for various purposes related to serpentinization and associated reactions.

2 | DATASET DESCRIPTION

2.1 | Overview of the dataset

We have collected the data in 30 relevant experimental articles that report measured H₂ and organic compounds (OC) production related to the serpentinization reaction (Berndt et al.,
1996; Horita and Berndt, 1999; McCollem and Seewald, 2001; McCollem and Seewald, 2003; Allen and Seyfried, 2003; Foustoukos and Seyfried, 2004; Seewald et al., 2006; Seyfried et al., 2007; Fu et al., 2007, 2008; Ji et al., 2008; Dufaud et al., 2009; Jones et al., 2010; McCollem et al., 2010, 2016; Marcaillou et al., 2011; Neubeck et al., 2011, 2014; Lafay et al., 2012; Lazar et al., 2012, 2015; Klein and McCollem, 2013; Okland et al., 2014; Klein et al., 2015; Huang et al., 2015, 2016; McCollom, 2016; McCollom and Donaldson, 2016; Miller et al., 2017; Grozeva et al., 2017). The vast majority of these studies report on experimental serpentinization starting with olivine or peridotites, but a couple of studies are included in the dataset on purpose, to investigate FTT reactions without olivine as a starting mineral (Foustoukos and Seyfried, 2004; Fu et al., 2007; Ji et al., 2008). The latter bypasses the serpentinization reaction and uses formic acid, which decomposes into H₂ and CO/CO₂ at temperatures and pressures similar to those typical of hydrothermal serpentinization and could react to form abiotic organic compounds.

The reported experiments covered a large range of experimental conditions, including the temperature (T), pressure (P), experiment duration, chemical compositions of both reactants and products, as well as types of reactors, origins of mineral samples. We summarized the information into a single large Excel spreadsheet of 133 columns and 733 rows. The spreadsheet columns are divided into 3 main sections (Figure 1): article information (green header), experimental conditions (blue header); and results (yellow header). The section on article information includes details of all published articles of this dataset: data ID, article title, year of publication, authors and DOI numbers, which help the readers of the present contribution to trace back the original studies. The 733 rows describe 195 experiments that include sometimes multiple samplings on the course of the experiments to evaluate the reaction kinetics.

Before moving into the details of each section of the dataset, there is some important general information. In the section dedicated to experimental conditions, it is important to keep in mind that most parameters are independent of each other, but a few of them are dependent. For example, the magnesium content of olivine ‘Mg#(Ol)’ displays a value only for experiments that include olivine as a starting mineral (Foustoukos and Seyfried, 2004; Fu et al., 2007; Ji et al., 2008). The latter bypasses the serpentinization reaction and uses formic acid, which decomposes into H₂ and CO/CO₂ at temperatures and pressures similar to those typical of hydrothermal serpentinization and could react to form abiotic organic compounds.

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![FIGURE 1](Screenshot of a representative version of the Excel spreadsheet for experimental parameters and results. The current dataset has 134 columns (parameters/variables) and 733 rows (measurements). The dataset is divided into three parts—article information, experimental conditions and results).
Some programs cannot handle dataset with empty cells, so we assigned ‘nan’ (not a number) to values that were either not measured or not reported, and ‘0’ to measurements that were below detection limits or reported as ‘not observed’ in the original paper. We also paid attention to assign to each parameter a dedicated format that is given in Tables 1-9 of the present contribution:

1. Numeric: data that are integers and float numbers. For example, temperature data are mostly integers and NaCl concentration data are float numbers.
2. Categorical: data that are not quantitative but categories, such as the rock type and degree of alteration.
3. Binary: Data that are either 0 (no) or 1 (yes), such as the reactor types.
4. Ternary: Data that are 0 (no), 1 (yes) or 2 (yes with $^{13}$C). For example, in the Carbon_in_solid column, 2 means carbon is $^{13}$C labelled.

### 2.2 Experimental parameters

#### 2.2.1 Reaction conditions and reactor information

The reaction conditions are listed first in Table 1: temperature (T) ranges between 25 and 500°C, pressure (P) between 0.1 and 350 MPa, and experimental durations are between 0 and 20,499 hrs. In most experiments, intermediate sampling was performed to study reaction kinetics and thus is reported in the dataset as a parameter—0 means no intermediate sampling performed whilst 1 means that the experiments were sampled through time. With 733 measurements and 195 experiments, each experiment on average takes 3.75 measurements.

Different types of reactors are used depending on the experimental P-T conditions, experimental protocols and sample volumes (Table 2). The reactor materials are also indicated since they are made of metals, whose catalytic role has often been suggested. For each experiment, one of the columns dedicated to the nature of the reactor has a value of 1, others are 0. For instance, a reactor made of both Au and Ti corresponds to ‘1’ in the column Reactor_Au/Ti, other columns display a ‘0’ for that experiment.

#### 2.2.2 Starting mineral, fluid and gas compositions

The information on the composition of rocks, minerals, solutes in the aqueous phase and gas is summarized in Tables 3-5. Table 3 contains the provenance of samples, the type of rock, the degree of alteration (when indicated by the authors), the composition of the rocks (expressed in weight per cent of each mineral normalized to reach 100%), the Mg# (olivine only), the grain size and other information on the starting mineral phases that we considered useful. The provenance of a sample is either its original geological location or the name of the company where it was synthesized. The estimated degree of alteration is based on
mineral descriptions in the original articles—fresh (nearly 0% altered), altered (nearly 100% altered) and medium altered. The missing values for the degree of alteration are mainly from experiments started without a mineral phase, which are dedicated to carbon speciation at high P and T. Information of grain size (max and min) and surface area (SSA, cm²/g of rock) that are very important for the kinetics of the reaction is also reported in this section when available, otherwise they are labelled as ‘nan’. Simple statistics analyses of these parameters are displayed in Tables 3-5, which helps readers to decide if this dataset contains useful data for them.

The chemical composition of the starting aqueous solution is described in Table 4. The solutes include many organic and inorganic salts whose concentrations are given in mol/kg (molal) as volumes change significantly under hydrothermal conditions. When articles reported in their experimental section that milli-Q water (18.2 MΩ/cm) was used, the column ‘precised_water_clean’ displays a value of ‘1’, otherwise the value is ‘0’. Values of the pH of the starting solution are also given as ‘initial pH’ when available. The initial amount of carbon among starting chemicals is one important aspect in this study, which allows comparing between studies the produced reduced carbon compounds through FTT or Sabatier reactions or any other reaction. We created a column ‘CO₂_initial’ that tells whether authors flushed their system or not. If not, we assumed that fluids were at equilibrium with present day atmospheric CO₂, which leads to a CO₂ concentration of 0.01 millimole per kg H₂O.

When an experiment contained a headspace filled with gas, we reported as much as we could the gas composition. It is given in relative volume percentage of N₂, CO₂, H₂, CO, CH₄.

### Table 3: Summary of starting mineral compositions and relevant parameters

| Parameters                | Min   | Max   | Average | Number of ‘nan’ | Data type   |
|---------------------------|-------|-------|---------|-----------------|-------------|
| Provenance_of_sample      | –     | –     | –       | 0               | Categorical |
| Rock_type                 | –     | –     | –       | 0               | Categorical |
| Degree_of_alteration      | –     | –     | –       | 73              | Categorical |
| Olivine_wt%               | 0     | 100.00| 49.50   | 33              | Numeric     |
| Mg/(Ol)                   | 0     | 92    | 70.28   | 231             | Numeric     |
| Clinopyroxene_wt%         | 0     | 40    | 1.52    | 5               | Numeric     |
| Orthopyroxene_wt%         | 0     | 100   | 4.13    | 5               | Numeric     |
| Pyroxene_wt%              | 0     | 100   | 5.38    | 0               | Numeric     |
| Spinel_wt%                | 0     | 5     | 0.25    | 0               | Numeric     |
| Magnetite_wt%             | 0     | 100   | 4.30    | 0               | Numeric     |
| Haematite_wt%             | 0     | 92.6  | 0.76    | 0               | Numeric     |
| Serpentine_wt%            | 0     | 100   | 3.37    | 0               | Numeric     |
| Brucite_wt%               | 0     | 6     | 0.30    | 0               | Numeric     |
| Talc_wt%                  | 0     | 1     | 0.05    | 0               | Numeric     |
| Carbonate_wt%             | 0     | 0.25  | 0.01    | 0               | Numeric     |
| Amphibole_wt%             | 0     | 5     | 0.09    | 0               | Numeric     |
| Chlorite_wt%              | 0     | 1     | 0.02    | 0               | Numeric     |
| Phlogopite_wt%            | 0     | 1     | 0.05    | 0               | Numeric     |
| Chromite_wt%              | 0     | 100   | 2.00    | 0               | Numeric     |
| Clinohlore_wt%            | 0     | 1     | 0.04    | 0               | Numeric     |
| GLASS                     | 0     | 75    | 1.39    | 0               | Numeric     |
| SiO₂_wt%                  | 0     | 100   | 0.82    | 0               | Numeric     |
| Fe_wt%                    | 0     | 100   | 5.59    | 0               | Numeric     |
| NiFe_wt%                  | 0     | 100   | 6.92    | 0               | Numeric     |
| FeO_wt%                   | 0     | 100   | 1.34    | 0               | Numeric     |
| FeS_wt%                   | 0     | 50    | 0.35    | 0               | Numeric     |
| NiO_wt%                   | 0     | 50    | 0.34    | 0               | Numeric     |
| Grain_size_min            | 0     | 300   | 51.88   | 195             | Numeric     |
| Grain_size_max            | 0     | 3,000 | 213.04  | 207             | Numeric     |
| SSA                       | 0     | 71,000| 7832.77 | 535             | Numeric     |
and Ar in the headspace, as the description in the experiment methods section in the articles seldom provided detailed information on the gas composition (Table 5). For each experiment, the initial concentrations of these gases are therefore reported as vol%, and the sum of these species in the headspace equals arbitrarily 100% or 0 when information is missing.

### 2.2.3 Potential catalysts and carbon sources

Previous studies (Fu et al., 2008; Andreani et al., 2013; Mayhew et al., 2013) have shown that the serpentinization reaction and the production of H₂ and organic species (CH₄, formate, acetate, etc.) can be largely influenced by the presence of cations in solution or solid catalysts (accessory mineral surfaces). In industrial H₂ and CH₄ production, metal-bearing catalysts are critical. Platinum, rhodium, ruthenium, cobalt and other metallic materials are well-known catalysts for methanization of dry gas by FTT or Sabatier reaction (McKee, 1967; Melael et al., 2014; Stangeland et al., 2017). In natural environments, a large number of metallic phases are present in ultramafic rocks and minerals, and also in experimental materials as impurities. Variabilities of the kinetics of serpentinization and H₂ and CH₄ formation, both in nature and experiments, could be due to the effect of catalysts either in their mineral form or as solute (e.g. Andreani et al., 2013; Mayhew et al., 2013; Etiope and Ionescu, 2015). Therefore, we created a list of minerals with potential or expected catalytic effects and recorded their presence or absence in each experiment (Table 6).

The carbon source(s) are also recorded in the dataset in a ‘ternary’ data format (definition in section 2) for two main goals here: (a) identify if the amount of reduced carbon products is favoured by certain carbon-bearing reactants; (b) help readers to easily locate experiments labelled with ¹³C and further analyse the influence of background contaminations.
### Table 6  Potential catalysts and carbon source

| Parameters          | Number of 0 | Number of 1 | Number of 2 | Average | Number of 'nan' | Data type |
|---------------------|-------------|-------------|-------------|---------|-----------------|-----------|
| Catalyst_Spinel     | 571         | 162         | –           | –       | 0               | Binary    |
| Catalyst_Magnetite  | 603         | 130         | –           | –       | 0               | Binary    |
| Catalyst_Chromite   | 584         | 149         | –           | –       | 0               | Binary    |
| Catalyst_Ni_bearing | 628         | 105         | –           | –       | 0               | Binary    |
| Catalyst_Fe_bearing | 624         | 109         | –           | –       | 0               | Binary    |
| Carbon_in_Solid     | 688         | 45          | 0           | –       | 0               | Ternary   |
| Carbon_in_Liquid    | 172         | 429         | 132         | –       | 0               | Ternary   |
| Carbon_in_Gas       | 584         | 121         | 28          | –       | 0               | Ternary   |

### Table 7  Other information related to experimental conditions

| Parameters         | Number of 0 | Number of 1 | Average | Number of 'nan' | Data type |
|--------------------|-------------|-------------|---------|-----------------|-----------|
| isBlank            | 609         | 124         | –       | 0               | Binary    |
| Add_Solids         | 140         | 593         | –       | 0               | Binary    |
| Add_Liquids        | 0           | 733         | –       | 0               | Binary    |
| Add_Gases          | 478         | 255         | –       | 0               | Binary    |
| Intermediate_sampling | 167       | 566         | –       | 0               | Binary    |

| Parameters             | Min  | Max     | Average | Number of 'nan' | Data type |
|-----------------------|------|---------|---------|-----------------|-----------|
| Total_volume_ml        | 0.31 | 250.00  | 81.82   | 412             | Numeric   |
| Mass_solids           | 0    | 120     | 10.70   | 42              | Numeric   |
| Mass_liquids          | 0    | 180     | 37.58   | 92              | Numeric   |
| Water_Rock            | 0    | 400     | 13.26   | 170             | Numeric   |

### Table 8  Mineral products of experimental results

| Parameters             | Number of 0 | Number of 1 | Average | Number of 'nan' | Data type |
|-----------------------|-------------|-------------|---------|-----------------|-----------|
| Chrysotile_product    | 330         | 74          | –       | 328             | Binary    |
| Lizardite_product     | 444         | 28          | –       | 261             | Binary    |
| Antigorite_product    | 471         | 0           | –       | 262             | Binary    |
| Phyllosilicate_product| 325         | 134         | –       | 274             | Binary    |
| Talc_product          | 293         | 10          | –       | 430             | Binary    |
| Brucite_product       | 354         | 44          | –       | 335             | Binary    |
| Magnetite_product     | 219         | 96          | –       | 418             | Binary    |
| Haematite_product     | 180         | 0           | –       | 553             | Binary    |
| Chromite_product      | 249         | 15          | –       | 469             | Binary    |
| Pentlandite_product   | 263         | 2           | –       | 468             | Binary    |
| Heazlewoodite_product | 225         | 4           | –       | 504             | Binary    |
| Carbonate_product     | 145         | 67          | –       | 521             | Binary    |
| Siderite_product      | 129         | 7           | –       | 597             | Binary    |
| Iowaite_product       | 261         | 10          | –       | 462             | Binary    |
| Ni_Fe_phase_product   | 170         | 62          | –       | 501             | Binary    |
| Reduced_carbon_product| 52          | 17          | –       | 664             | Binary    |
| Unidentified_phase_product | 132    | 17          | –       | 584             | Binary    |
2.2.4 Other information

All other information regarding the starting materials is listed in Table 7. The ‘isBlank’ parameter indicates whether an experiment is blank or control (1 for no mineral within a series of experiments), in order to distinguish them from experiments and avoid any inaccurate interpretation of the data. The ‘Add_solids’, ‘Add_liquids’ and ‘Add_gases’ are used to indicate the presence of a solid, liquid or gas phase, respectively (0 means absent). The ‘mass_solids’, ‘mass_liquids’ and ‘Water_rock’ ratio are also reported when available, as well as the total volume of the reaction cells.

3 RESULTS

The results section of the dataset is divided into two subsections—mineral and gas/fluid products. The columns that describe mineral products are less populated, since most contributions focused on the production of gas species (H₂, CH₄, etc.) and only a few of them also described the kinetics of serpentinization and analysed the final mineral composition.

3.1 Mineral products

Analyses of mineral products are reported in Table 8. It includes all the minerals identified by the authors of the 30 peer-reviewed articles. Each mineral produced during experiments is defined as a parameter, and reading through the columns, one can see that the data available in the literature are sparse. As a consequence, statistics on the minerals produced during the reaction is poorly constrained. The presence or absence of secondary minerals is not always clearly established in the articles and strongly depends on the details provided by the authors and characterization technique used in those contributions. In some cases, descriptions of the solid phase did not mention the experiment number they were referring to. In addition, in most experimental settings dedicated to the understanding of H₂ and CH₄ production during serpentinization, solids are accessible only at the end of the experiments. The mineral feature is assigned ‘nan’ if the information is not clearly stated. Hence, this part of the dataset should be used with great care and we encourage the reader to refer to the original article as necessary.

3.2 H₂ and hydrocarbon products in final fluids

This subsection focuses on the experimental measurements of H₂, CH₄ and OC during experimental serpentinization. We assigned an individual parameter to each compound. The details on the analysis of the composition of final fluids are described in Table 9. It is important to take into account that the measurement precision and detection limit of those compounds are very different across studies and potentially evolved through time as analytical tools and methods improve. Obviously, the
Most commonly measured species are H2 and CH4, given the topic. Other organic species are measured only in a few studies and have therefore many missing values (nan) in the dataset. Though each organic compound is listed in its own column, all species other than H2 and CH4 shall be combined in order to perform some meaningful data analysis.

4 | DISCUSSIONS AND PERSPECTIVES

This dataset provides an up-to-date collection of experimental results until early 2019 that can be used to address the implications of serpentinization and related processes for the production of H2, CH4 and higher hydrocarbons. It is well known that P-T conditions largely control experimental results, but they alone could not explain the large variability of the measured concentrations of H2, CH4 and higher hydrocarbons. The other parameters investigated by the experimental community at large are so numerous that their investigation cannot be done easily and requires a computing approach that can deal with many parameters at the same time. We hope that the database is progressively enriched with upcoming experimental results. As an example, we have used data science techniques to extract embedded information and identify key experimental parameters, which cannot be accessed by checking a limited number of parameters.

In our recent paper (Barbier et al., 2020), we used network analysis and machine-learning algorithms to analyse a processed version of this dataset. We found that, as previously known, pressure and temperature are the two most important parameters that govern the production of H2 and OC. However, we did not find evidences to support the occurrence of R1 or R2 reactions. Moreover, by comparing the concentrations of final OC with initial carbon input, we found that the OC products in several studies are from unidentified sources and likely result from contamination, in agreement with the scarce 13C-labelled studies (e.g. McCollom et al., 2010; Grozeva et al., 2017). Also, the measurements of initial and final pH values are often not reported, despite the important role of pH on the serpentinization kinetics (e.g. Huang et al., 2019; McCollom et al., 2020). More information and the detailed analytical methods can be found in Barbier et al. (2020). The dataset can also be extended to other reactions under similar conditions using different solid reactants, such as mine-tailing products, to produce H2, CH4 and other carbon species (e.g. Kularatne et al., 2018; Michiels et al., 2018; Brunet, 2019). We hope that this dataset and the analysis by Barbier et al. (2020) stimulate complementary experiments that could fill the identified gaps; this would allow editing future versions of this dataset in a few years and potentially help people better understand the fate of carbon under highly reducing conditions such as the one produced during serpentinization.

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CONFLICT OF INTEREST

The authors declare no conflict of interest.

OPEN PRACTICES

This article has earned an Open Data badge for making publicly available the digitally-shareable data necessary to reproduce the reported results. The data is available at http://info.deepcarbon.net/individual/n1819 Learn more about the Open Practices badges from the Center for OpenScience: https://osf.io/tvyxz/wiki

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