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Early Diagenesis at and below Vera Rubin ridge, Gale crater, Mars

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ABSTRACT

Data returned by NASA’s Mars Science Laboratory Curiosity rover showed evidence for abundant secondary materials, including Fe-oxides, phyllosilicates, and an amorphous component on and below Vera Rubin ridge in the Murray formation. We used equilibrium thermochemical modeling to test the hypothesis that altered sediments were deposited as detrital igneous grains and subsequently underwent diagenesis. Chemical compositions of the Murray formations’ altered components were calculated using data returned by the Chemistry and Mineralogy X-ray diffraction instrument and the Alpha Particle X-ray Spectrometer on board Curiosity. Reaction of these alteration compositions with a CO$_2$-poor and oxidizing dilute aqueous solution was modeled at 25 – 100 °C, with 10 – 50% Fe$^{3+}$/Fe$_{tot}$ of the host rock. The modeled alteration assemblages included abundant phyllosilicates and Fe-oxides at water to rock ratios >100. Modeled alteration abundances were directly comparable to observed abundances of hematite and clay minerals at a water to rock ratio of 10,000, for system temperatures of 50 – 100 °C with fluid pH ranging from 7.9 to 9.3. Modeling results suggest that the hematite-clay mineral assemblage is primarily the result of enhanced groundwater flow compared to the Sheepbed mudstone observed at Yellowknife Bay, and underwent further, localized alteration to produce the mineralogy observed by Curiosity.
INTRODUCTION

At Gale crater, evidence for an ancient lacustrine environment within which sediments were deposited and diagenetically altered have been found in the Murray formation by the NASA Mars Science Laboratory (MSL) Curiosity rover (e.g., Grotzinger et al., 2015; Edgar et al., 2020; see Fraeman et al., 2020 for an overview of the entire geologic context). In this study we use thermochemical modeling to test the hypothesis that the sediments were deposited as detrital igneous grains followed by diagenetic alteration conditions where groundwater interacted with the sediments. This is in continuation of previous work at the first units encountered by Curiosity at Yellowknife Bay (Bridges et al., 2015b; Schwenzer et al., 2016). In doing so, we aim to constrain associated temperature, water to rock ratios, pH, and redox conditions of the early diagenetic alteration pathways that resulted in the predominantly hematite-clay mineral assemblage in the Murray formation on and below Vera Rubin Ridge (VRR). We assume that Ca-sulfate vein formation as described in L’Haridon et al. (2020) and acidic alteration (Rampe et al., 2020b) belong to a later phase of alteration as well as Ca-sulfate formation, which has been shown to occur during a later step of fluid evolution (Schwenzer et al., 2016). The former is consistent with the sedimentological observations (Edgar et al., 2020) and observations by Curiosity’s instruments (e.g., McAdam et al., 2020).

Overview of the Murray Formation

The Murray formation was first encountered on sol 792 of the NASA MSL mission and constitutes the sedimentary units analyzed by the Curiosity rover from the foothills of Mount Sharp (Grotzinger et al., 2015). From orbit, the lower Murray formation has areas with orbital spectral signatures consistent with Fe/Mg clay, hydrated silica, iron oxides, and sulfate (Anderson and Bell, 2010; Milliken et al., 2010; Thomson et al., 2011; Fraeman et al., 2013; Fraeman et al., 2016). Distinct from orbit by its hematite spectral signature (Fraeman et al., 2013), VRR is a ~200 m wide ~6.5 km long northeast-southwest resistant geomorphological feature on the northern slopes of Mount Sharp (Fig. 2). VRR is composed of the Pettegrove Point member that is overlain by the Jura member (Fig. 1). They have a combined stratigraphic thickness of approximately 60 m and were investigated by Curiosity from sol 1809 to sol 2302, when Curiosity traversed off VRR.

At the time of writing, the Murray formation is estimated to be ~300 m thick and has been subdivided into seven members that are separated by conformable contacts, as illustrated by Edgar et al. (2020). Briefly, the seven members are the Pahrump Hills, Hartman’s Valley, Karasburg, Sutton Island, Blunts Point, Pettegrove Point and Jura, comprised of sand- silt and mudstones of lake bed and fluvial origin as detailed in the following sections (Grotzinger et al., 2015; Bristow et al., 2018; Edgar et al., 2018; Fedo et al., 2018; Gwizd et al., 2018; Rivera-Hernandez et al., 2019; Stack et al., 2019; Edgar et al., 2020).

Summary of Murray Formation stratigraphic members below Vera Rubin Ridge

Stratigraphic members below VRR of interest in this study were the Hartmann’s Valley, Karasburg, Sutton Island, and Blunts Point members (Fig. 1).

The Hartmann’s Valley member (25 m thick) has grains ranging in size between silt to medium sand and with meter-scale, trough cross-bedding indicative of either fluvial or aeolian settings (Bristow et al., 2018; Fedo et al., 2018; Gwizd et al., 2018). The Oudam sample was
drilled from this member, in fine-grained bedrock containing diagenetic Ca-sulfate mineral veins (Bristow et al., 2018).

The Karasburg member (37 m thick) in the Murray Buttes locality is characterized by a heterolithic mudstone/fine sandstone assemblage with abundant cm-scale concretions and a purple hue (Fedo et al., 2018). In this member, the samples Marimba and Quela were drilled from flat mudstone bedrock containing some small Ca-sulfate mineral veins and mm-scale Mn-enriched concretions (Bristow et al., 2018; L’Haridon et al., 2018; Meslin et al., 2018).

The Sutton Island member (98 m thick) also contains a heterolithic mudstone/sandstone assemblage that is largely red in color and has an increased abundance of Ca-sulfate features (e.g., veins and lenticular crystals) and polygonal cracks interpreted to result from desiccation at the lake margins (Stein et al., 2018). The Sebina sample was drilled from flat, fine-grained bedrock that has abundant, cm-scale nodules and Ca-sulfate mineral veins.

The Blunts Point member (~100 m thick) is exposed immediately below VRR on the north-facing side and shares a conformable boundary with the ridge (Edgar et al., 2020; Fedo et al., 2018). The Blunts Point member is characterized by well laminated mudstone with cross-cutting curviplanar Ca-sulfate veins (Edgar et al., 2020). The Duluth drill sample was acquired near the base of VRR in a well laminated bedrock with diagenetic raised ridges and Ca-sulfate veins (Fig. 1).

**Summary of Murray Formation stratigraphic members on Vera Rubin Ridge**

The lithology at VRR appears consistent with the Murray formation mudstones analyzed previously (Fedo et al., 2018; Heydari et al., 2018), with a grain size similarly dominated by mud and contains occasional intervals of fine to medium sandstone, as estimated using the Mars Hand Lens Imager on-board MSL supplemented by ChemCam LIBS data and the Gini Index Mean Score (Edgar et al., 2020). APXS and ChemCam analyses in the Murray formation members below and on VRR indicate that it is compositionally indistinguishable from the Murray formation, and that Fe concentrations are not elevated for VRR; this suggests the compositional trends from the rocks below and the rocks on VRR are largely the result of post-depositional processes, such as diagenesis (Thompson et al., 2020; Frydenvang et al., 2020).

Rocks within the Pettegrove Point member of the ridge are thin, planar laminated, fine-grained mudstones. Rocks within the Jura member at the top of the VRR stratigraphic succession are also characterized by fine-grained and planar laminated facies, but with a greater degree of variability in color (red, purple, grey and tan) and meter-scale inclined strata (Edgar et al., 2018, 2020). Generally, rocks within the Pettegrove Point member have fewer diagenetic features such as nodules, mineral veins, fractures and erosion resistant features (L’Haridon et al., 2020), and are lower in Al₂O₃, SiO₂, and higher in FeO₇ than the Jura member (Thompson et al., 2020). Both members are generally red/purple (Bennett et al., 2018) though grey bedrock patches up to ~10 m in diameter have also been detected along VRR but are more common in the Jura member (Frydenvang et al., 2020; L’Haridon et al., 2020). These grey patches are currently interpreted as resulting from diagenesis, possibly from the circulation of warm, oxidizing, fluids (e.g., Bibi et al., 2011; Peretyazhko et al., 2016, 2018; McAdam et al., 2020; Fraeman et al., 2020; Rampe et al., 2020b) or reducing groundwater (Frydenvang et al., 2020; Horgan et al., 2020; L’Haridon et al., 2020).
Three drill samples from VRR were analysed by the Chemistry and Mineralogy (CheMin) instrument, which uses X-ray diffraction to determine the mineralogy of drilled samples (Blake et al., 2012). The Stoer sample was drilled from red bedrock in the Pettegrove Point member that contained small scale (~1 mm in diameter) concretions. The drilled samples ‘Highfield’ and ‘Rock Hall’ were acquired in the Jura Grey and Red Jura facies respectively. The Grey Jura ‘Highfield’ sample was drilled at the Lake Orcadie locality, which is within a diagenetic grey patch with abundant Ca-sulfate veins, Fe-rich nodules and dark, elongate diagenetic features. MAHLI images before and after drilling showed that Highfield had potentially sampled these features. The Highfield sample is also notable for the likely presence of ‘grey hematite’, i.e., relatively coarse hematite with crystallites >5 μm in diameter (Rampe et al., 2020). On the basis of Catling and Moore’s (2003) study of grey hematite, Rampe et al. (2020b) inferred a lower temperature limit of ~100 °C for the grey hematite-bearing assemblage. The Rock Hall Jura sample was acquired just south of the grey Lake Orcadie area in a red patch of bedrock with an unusual, shiny, nodular appearance.

Mineralogy of the Murray Formation

CheMin analyzed drilled rock powders from the Murray samples Confidence Hills, Mojave2, Telegraph Peak, Buckskin, Oudam, Marimba, Quela, Sebina, and Duluth below VRR, and Stoer, Highfield, and Rock Hall on VRR itself (Rampe et al., 2020a). Of these drilled rock powders, the last eight (Oudam through Rock Hall) belong to the hematite-clay mineral group investigated in this paper. CheMin data show the presence of detrital igneous minerals and products of aqueous alteration in these eight samples. Details of the mineralogy of these samples are reported by Bristow et al. (2018), Rampe et al. (2020a), Achilles et al. (2020), and Rampe et al. (2020b). All eight samples contain abundant primary magmatic minerals in the form of plagioclase feldspar (19 – 24 wt.% of the bulk sample) with unit-cell parameters consistent with an andesine composition and lesser amounts of pyroxene (1 – 9 wt.% of the bulk sample), alkali feldspar (<3 wt.% of the bulk sample) and quartz (<1 wt.% of the bulk sample). The low angular resolution of CheMin prevents a confident identification of the type(s) of pyroxene present (Rampe et al. 2020b).

Phyllosilicates are present in every sample but vary in structure and abundance through the stratigraphy. Ferripyrophyllite (Fe₂Si₄O₁₀(OH)₂), a member of the talc-pyrophyllite group, is the most likely phyllosilicate in Oudam (3 ± 1 wt.% of the bulk), based on position and breadth of the (001) basal peak (9.6 Å in CheMin data) and the H₂O evolution profile observed from the SAM EGA analysis (Bristow et al., 2018; Achilles et al., 2020). In Marimba, Quela, and Sebina the phyllosilicate basal spacings and the positions of the (02l) band suggest a mix of dioctahedral and trioctahedral smectite (e.g., Bristow et al., 2018). Smectite comprises 28 ± 3 wt.% of the bulk of Marimba, and the (02l) peak position is consistent with a 1:2 dioctahedral:trioctahedral weight abundance ratio, with SAM EGA data consistent with Fe-montmorillonite and Mg-saponite (Bristow et al., 2018; Achilles et al., 2020). Dioctahedral:trioctahedral ratios for the 16 ± 2 wt.% smectite in Quela and the 19 ± 2 wt.% smectite in Sebina are 1:1 and 5:3, respectively, based on CheMin data (Bristow et al., 2018; Achilles et al., 2020). Data from Duluth indicate the presence of exclusively dioctahedral smectite. Analysis of CheMin data for Duluth indicates 15 ± 7 wt.% bulk Fe³⁺ clay, with both CheMin (Rampe et al., 2020b) and SAM EGA data (McAdam et al., 2020) consistent with the presence of nontronite. The basal spacings in the patterns from Stoer, Highfield, and Rock Hall are at 9.6 Å instead of the 10 Å position attributed to collapsed smectite in previous samples (Bristow et al., 2018), indicating a collapsed smectite with a small
interlayer cation or the trioctahedral phyllosilicate ferripyrophyllite (Bristow et al., 2018; Rampe et al., 2020b). Phyllosilicate abundances for Stoer, Highfield and Rock Hall were reported to be 10 ± 5, 5 ± 2 and 13 ± 6 wt.% bulk, respectively (Rampe et al., 2020b). Overall, the abundance and nature of the clay minerals in conjunction with primary magmatic minerals, allows for testing the hypothesis that all or a significant proportion of the clay minerals were formed in situ during post-depositional diagenesis.

Each drill sample contains abundant Fe-oxides and/or Fe-oxyhydroxides (Achilles et al., 2020; Rampe et al., 2020b). Hematite (α-Fe₂O₃) is present in every sample in abundances of ~3 wt.% of the bulk sample for Rock Hall up to ~16 wt.% of the bulk sample for Stoer. The colors of the drill tailings in Oudam and Highfield suggest grey or specular hematite is present in these samples. Akaganeite (α-Fe₃⁺O(OH,Cl)) is more abundant than hematite in Rock Hall, and constitutes 6 wt.% of the bulk sample. A trace amount of akaganeite is present in Stoer. Magnetite is present in Duluth, Stoer, Highfield, and Rock Hall near the detection limit of CheMin (Rampe et al., 2020b).

Ca-sulfate minerals are prevalent in every sample, but vary in type and abundance among samples. Most samples have all three varieties of Ca-sulfate (anhydrite – CaSO₄, bassanite – CaSO₄•0.5H₂O, and gypsum – CaSO₄•2H₂O), with the exception of Rock Hall. Rock Hall has the greatest abundance of Ca-sulfate of these samples, in which ~11 wt.% of the bulk sample is comprised of anhydrite. Duluth has the lowest abundance of Ca-sulfate of these samples, with ~1.5 wt.% anhydrite, ~3.5 wt.% bassanite, and gypsum is at the CheMin detection limit (~1 wt.%; Blake et al., 2012). Jarosite (KFe₃(SO₄)₂(OH)₆) is present in abundances up to a few wt.% in Marimba, Quela, Sebina, Stoer, and Rock Hall (Achilles et al., 2020; Rampe et al., 2020b).

All samples contain abundant X-ray amorphous materials, and FULLPAT analyses indicate that opal-CT comprises a portion of the X-ray amorphous materials in the Oudam and Highfield samples. Based on mass balance calculations using CheMin mineral abundances, crystal chemistry using CheMin mineral abundances, crystal chemistry, and bulk APXS measurements, the composition of the X-ray amorphous component is variably enriched in Si, Fe, and S in these samples (Achilles et al., 2020; Rampe et al., 2020b).

There are several hypotheses for the variations in mudstone mineralogy such as variations in redox within the lake during the deposition of the Murray sediments (Hurowitz et al., 2017), a combined effect of source composition, mineral transportation and alteration (Bedford et al., 2019), diagenetic alteration at cold to low-T hydrothermal temperatures (Achilles et al., 2020; Frydenvang et al., 2020; McAdam et al., 2020; Thompson et al., 2020; Fraeman et al., 2013, 2016; Rampe et al. 2017), or aqueous alteration of deposited sediments by a short-lived, acidic fluid during late diagenesis (Rampe et al., 2017). In particular, the presence and crystallinity of akaganeite on VRR indicates alteration by warm, acidic fluids (Rampe et al., 2020b).

**METHODS**

Thermochemical modeling is an important tool to define reaction pathways in environments where a fluid has interacted with a rock and changed its initial mineralogy, especially for martian environments where observations are limited compared to more accessible environments on Earth. On Mars, a wide range of reaction pathway problems have been studied using a variety of thermochemical tools. On the basis of Martian meteorite and orbital data, problems such as carbonate and acidic reactions, silicate deposition, and organic molecule
reactions have been studied and tested successfully against experimental or observational results (Griffith & Shock, 1995, 1997; Hausrath et al., 2018; McAdam et al., 2008; Tosca et al., 2004, 2008; Varnes et al., 2003; Zolotov and Mironenko, 2007, 2016; Zolotov and Shock, 1999, 2005). CHIM-XPT and its predecessor CHILLER have been used to study a wide range of Martian P-T regimes, mainly for clay forming environments (Bridges and Schwenzer, 2012; Filiberto and Schwenzer, 2013; Schwenzer and Kring, 2009, 2013; Schwenzer et al., 2012), and also successfully tested against observational ground truth in meteorites (Bridges and Schwenzer, 2012; Melwani-Daswani et al., 2016) and in situ on Mars (Bridges et al., 2015b; Schwenzer et al., 2016; Schieber et al., 2017). It is on this basis that CHIM-XPT was used to develop and test the models presented here.

Thermochemical modeling has limitations, which may, for example, cause the occurrence of phases not known to form under given pressure-temperature-composition conditions in nature. This is due to the limits in the availability of experimentally derived thermochemical data and the resulting extrapolation of the data application beyond the pressure-temperature range of available data. This is taken into account through careful cross checking of results. We refer readers to a wide range of literature (e.g., Ganguly, 2008; Holland and Powell, 1998; Künn, 2004; Reed, 1997; Rimstidt, 2014), which will provide the theoretical backing of the technique and further insights into problem solving and decision making to limit the impact of modelling limitations. This is especially the case for clay minerals due to limitations in the dataset in modeling iron-bearing environments on Mars (Catalano, 2013; Chevrier et al., 2007). Therefore, in this paper, care was taken to use all information available to constrain the modeling parameters when evaluating the results. It is important to note that in a sedimentary environment, or any dynamic geologic environment, the models describe reaction pathways of local equilibrium and not an overall equilibrium of the entire geologic unit (Kühn, 2004). Thus, such models enable the assessment of a reaction pathway for a given overall chemistry to reach the resulting alteration assemblage.

**Thermochemical code and database**

CHIM-XPT (Reed et al., 2010) is a program for computing multicomponent heterogeneous chemical equilibria in aqueous-mineral-gas systems and was chosen because of its heritage in modeling basaltic and ultramafic systems (e.g., Reed, 1983; Palandri and Reed, 2004), and modeling alteration on Mars (Schwenzer and Kring, 2009; Bridges and Schwenzer, 2012; Schwenzer et al., 2012; Filiberto and Schwenzer, 2013; Bridges et al., 2015b; Schwenzer et al., 2016; Melwani-Daswani et al., 2016; Schieber et al., 2017). The model calculation proceeds in steps of reactant rock added to the fluid, where each step calculates equilibrium between the fluid and the dissolved rock, meaning that each step can be treated and interpreted independently to the direction from which it was reached, and each step determines the properties of the minimum Gibbs free energy of the system. Step size may vary depending on task requirements. A weight ratio of the total number of fluid grams relative to the total number of grams of reactant rock is used, so model calculations are independent of the size of the system, but the calculations are usually based on 1 kg (55.5 mol) of water. Precipitates are not fractionated from the system unless stated otherwise. In CHIM-XPT, the water to rock ratio (W/R) is the ratio of incoming fluid to reacted host rock. Inputs required for the execution of CHIM-XPT in this study were the starting fluid composition, reactant rock chemical composition, system temperature and pressure. We assume that the diagenesis was disconnected from the atmosphere, as can be expected from a subsurface fluid. Therefore, we did not include a...
gas phase at the start of the calculations and did not replenish any CO$_2$ or other components typically found in the Martian atmosphere. We note that gas phases were allowed to form, should they have become stable, but none were observed at the conditions that led to clay and hematite formation.

The SOLTHERM thermodynamic database used with CHIM-XPT in this study contains the following sheet silicates: talc (Mg, Fe, Mg-Al endmembers); pyrophyllite; chlorites (clinochlore, daphnite, Mn-chlorite, Al-free chlorite); kaolinite; illite; montmorillonite (Ca, K, Mg, Na endmembers); beidellite (Ca, H, K, Mg, Na endmembers); nontronite (Ca-, H-, K-, Mg-, Na-nontronite); serpentine (antigorite, chrysotile, greenalite); sepiolite. The only member of the kaolinite group the database contains is kaolinite. There is no saponite or vermiculite in the database, which may affect the results and therefore comparisons between our modeled mineral assemblages and the observed mineral assemblages on Mars. In this study, results are plotted in mineral abundance (wt.%) vs W/R, with chloride endmembers (Fe$^{3+}$ sheet silicate and nontronite endmembers (Fe$^{3+}$ clay mineral) combined. Given the limitations of the thermochemical database with regards to the full range of possible phyllosilicates, in this study the modeled phyllosilicate assemblages are summed to calculate an average clay mineral with average composition that is then taken as a chemical analogue to the clay minerals observed in Gale crater. As discussed in discussion section ”Clay minerals and comparisons to Yellowknife Bay and other Martian terrains, including Nakhlite Martian Meterorites”, this is consistent with previous approaches by Bridges et al. (2015b) and Schwenzer et al. (2016).

Critical evaluation of the results and comparison to terrestrial mineralogy are required to further evaluate the modeling results. Extrapolation of the thermodynamic data from higher temperature might occur in the low temperature range modeled here, which can lead to the occurrence of mineral phases not known to form at the given pressure-temperature range. This problem is dealt with through careful evaluation of every phase forming and exclusion of those known to not form in the given pressure-temperature range, owing to kinetic constraints at low temperature (Melwani-Daswani et al., 2016, supporting material). The temperature transition between the Fe-oxides, goethite and hematite, was taken into consideration for the modeling. As documented by Cornell and Schwertmann (2006; and references therein) fine grained goethite is stable ≤ 25 °C, relative to coarse grained hematite. However, in laboratory environments where ferrihydrite was stored as aqueous suspensions at 24 °C and pH between 2.5 and 12, both hematite and goethite form, with the highest hematite abundance forming between pH 7 and 8 and the highest abundance of goethite forming at pH 4 and pH 12 (Schwertmann and Murad, 1983). The formation of other Fe-oxides such as ferrihydrite, goethite and akaganite may have occurred first in the sediments of Gale crater and later transformed to hematite via mechanisms documented by Cornell and Schwertmann (2006; and references therein). However, while those precursors could have been a factor in some of the rocks we are studying, such as those that had high amounts of Fe in their amorphous components, it is difficult to definitively identify poorly ordered/amorphous Fe phases such as ferrihydrite with Curiosity’s payload, and at the time of writing there is no direct evidence for ferrihydrite. For the purposes of this modeling study we assume direct precipitation of goethite or hematite from the modeled porewater-rock interactions, and not from the alteration of ferrihydrite. Magnetite was allowed to form but did not under the conditions modeled here at the W/R assumed relevant.

A list of the minerals excluded in each model can be found in the supplementary material (Table S3). The model calculations show finer details than any of the sources of information
from Mars: detection limit for the phase abundances with CheMin on *Curiosity* is ~1 wt.% (Blake et al., 2012) and the pixel footprint of CRISM is nominally 18 m, (Murchie et al., 2007) limiting the detection of trace phases in both instances. Therefore, small proportions of a mineral phase (<5 wt.% within a modeled alteration assemblage may not be easily comparable to the observed martian mineralogy. Minerals modeled to form that have not been observed in the Murray formation, either in-situ with *Curiosity* or remotely with CRISM, were allowed to form due to a lack of justification for their removal.

**Modeled water-rock ratio**

CHIM-XPT thermochemically calculated mineral assemblages are illustrated as plots of mineral abundance (weight %) vs W/R ratio (e.g. Fig. 5 – 7). W/R ratio is a progress variable which indicates limited reactant rock dissolution at high W/R with a relative increase in reactant rock dissolution at low W/R. The mass of alteration minerals precipitated ranges from 0.01 g at 100,000 W/R to 1,000 g at 1 W/R. CHIM-XPT works on the basis of complete dissolution of the defined reactant rock components in relevant W/R proportions per interaction step. While in real-world systems only partial host rock dissolution occurs (making the W/R values reported here comparably higher), modeling complete host rock dissolution enables a systematic investigation of modeled alteration minerals and associated fluids as a function of the W/R process variable. High W/R (>5,000) is representative of a system where only a small amount of the rock is dissolved in a large mass of fluid (i.e., where only a limited surface area interacts with the fluid and/or timescales are short). This can represent freshwater inflow environments such as in rock fractures or rock surfaces exposed to regular precipitation. Lower W/R ranges are more representative of minimal fresh-fluid inflow and stagnant fluids reacting with large rock surfaces. Comparison to experiments and geologic analogs show that W/R of around 1,000 is generally representative of water-rich diagenetic or small fracture settings, whereas lower W/R are comparable to low-water diagenetic and eventually metamorphic conditions. For an in-depth discussion how model and experimental water to rock ratios compare, see Olsson-Francis et al. (2017). CHIM-XPT calculates the equilibrium reaction of the fluid with the precipitating phases at an overall chemistry of the system.

**HYPOTHESIS AND MODEL SET-UP**

The purpose of this study is to constrain the formation conditions (W/R, temperature, redox, pH) under which the hematite-clay mineral assemblages observed on and below VRR precipitated. More specifically, we test the hypothesis that the formation of clays and hematite in the Murray formation on and below VRR is of groundwater type, which stands in a continuation of previous studies. For Yellowknife Bay, the lowest point in the stratigraphy encountered by *Curiosity* early in the mission, studies showed that groundwater-type, circumneutral fluids could explain the clay mineral formation with sulfate formation representing a distinct, later alteration event (Bridges et al., 2015b; Schwenzer et al., 2016). Haurath et al. (2018) came to a similar conclusion for Stimson formation alteration, where early was dominated by olivine- in circumneutral fluids and no redox-change was required to explain the observations. In this study, however, we focus on the alteration of a magmatic host rock chemistry, as could be deposited by the fluvial-lacustrine conditions described for Gale crater, in a circumneutral groundwater.

There is a wide range of fluid-related observations and phenomena observed before the *Curiosity* rover encountered VRR, such as overall lake water changes (Hurowitz et al., 2017),
nODULES (Nachon et al., 2014) and silica-rich haloes (Frydenvang et al., 2017), some of which are also observed on VRR. Other studies of VRR alteration (Fraeman et al., 2020; L'Haridon et al., 2020; McAdam et al., 2020; Rampe et al., 2020b; Yen et al., 2020) assume a multi-stage fluid evolution or a diversity of fluids to explain their observations. This leads to testing the hypothesis in this paper that the first step of alteration was one of in situ diagenesis in a dilute, circumneutral fluid. This study is in context of the late-stage phases (Ca-sulfate, jarosite, akaganeite) forming when fluid concentrations had evolved to higher sulfate concentration, or more generally higher salinity, and locally to acidic conditions (Achilles et al., 2020; L'Haridon et al., 2020; McAdam et al., 2020; Rampe et al., 2020b; Yen et al., 2020). This is further supported by SAM investigations of jarosite-bearing sediments showed a bimodal age distribution with a much younger potential formation age for the jarosite-bearing alteration (Martin et al., 2017). Therefore, modeling the reaction pathways of the late-stage Ca-sulfate, jarosite, and akaganeite is not a goal of this study.

The starting fluid

Gale Portage Water (GPW) (Bridges et al., 2015a, 2015b; Schwenzer et al., 2016; Turner et al., 2019) was used as the fluid in this study, making results presented here for the Murray formation directly comparable to the studies for Yellowknife Bay. This fluid was derived from equilibration of a fluid used previously in Mars fluid-rock interaction modeling (Schwenzer and Kring, 2009) with the basaltic Portage Soil composition sampled from the Rocknest aeolian bedform in Gale crater (Bish et al., 2013). As argued in Bridges et al. (2015b), this assumed that Rocknest can be regarded as an average crustal composition in Gale crater, and GPW is a fluid of groundwater-type concentration directly derived from local bedrock. Because of the dilute nature of this fluid, the chemistry of the rock reactants becomes the dominating factor in the water-rock reaction modeling. Thus, providing additional confidence that reaction pathways described here are directly applicable to Curiosity’s observations in the Murray Formation. The exact W/R at which the rock composition dominates the reaction is dependent on the solubility of the element considered and its concentration in the rock. Influence of the fluid is highest for elements not contained in the rock, which mostly concerns volatiles such as CO2. However, CO2 concentration is low (1.68 x 10^-4 mole/L) and does not typically lead to the formation of carbonates beyond trace level. A detailed summary of how this fluid was derived is given in Bridges et al (2015b), and the fluid composition is given in Table 3 of Bridges et al. (2015b). Chlorine concentration in GPW is about an order of magnitude higher than CO2, at 5.76 x 10^-3 mole/L, but chlorine does not participate in mineral formation. It is an important carrier of ionic strength at the very highest W/R, but at the W/R considered important here, does not cause any change of the system. It also is a potential complexing anion to increase Fe-solubility (e.g., Cornell and Schwertmann, 2006), but Cl concentrations are too low to significantly increase Fe-concentration in the fluid (Bridges et al., 2015b). Our Fe-concentrations are generally in family with other similar models (e.g., Marion et al., 2003). Cl-concentrations only influence apatite formation, which forms as Cl-apatite instead of hydroxyl-apatite, if Cl-concentrations are significant, but forms F-apatite once fluorine is present (Filiberto and Schwenzer, 2013). We note that – due to the lack of quantitative fluorine data for VRR rocks – we have not incorporated fluorine into the system. The solution is initially oxidizing with all S species as SO4^{2-}, and the redox in the fluid is controlled by the SO4^{2-}/HS^- pair. The redox of the system throughout each model is dependent on the Fe^{2+}/Fe^{3+} ratio of the total Fe in the reactant rock composition. The fluid composition is represented in each
calculated step by a set of 112 different ionic species. Unless stated otherwise, fluid pH was modeled as a free parameter.

**Host rock composition**

The *Curiosity* samples studied are mixtures of primary magmatic phases, such as pyroxene and plagioclase (Achilles et al., 2020; Rampe et al., 2020b), and alteration phases such as phyllosilicates and hematite (Rampe et al., 2020b; McAdam et al., 2020). Olivine is absent in the samples, which is in agreement with literature sources that have shown it to be the first mineral to dissolve in water-rock reactions (e.g., Gudbrandson et al., 2011; Hausrath et al., 2018). To use chemistry and mineralogy data from *Curiosity* to isochemically model the diagenetic alteration assemblage that occurred in the Murray formation, the unaltered host rock component needs to be removed from the model input data. For this, the chemistry of the observed alteration must be considered. Hereby defined in this study as “chemical alteration compositions”, the chemistry of the altered components was calculated and used as the reactant host rock composition in the thermochemical modeling as follows:

Calculating the chemical alteration compositions was undertaken by combining measurements taken by APXS and CheMin instruments on-board the *Curiosity* rover for drilled samples obtained in the Murray formation: Oudam, Marimba, Quela, Sebina, Duluth, Stoer, Highfield and Rock Hall (summarized in introduction section “Overview of the Murray formation”). Using crystal chemistries derived for the magmatic minerals (Table S1) (Morrison et al., 2018; Achilles et al., 2020; Rampe et al., 2020b) and their abundances determined using CheMin data (Bristow et al., 2018; Rampe et al., 2020a), the associated compositional oxide wt.% contributions were subtracted from the bulk APXS measurements. Ca-sulfates observed by CheMin (Achilles et al., 2020; Rampe et al., 2020b) were also removed as these have been shown to have formed during late diagenesis (Nachon et al., 2014), during a possible two-step process (Schwenger et al., 2016). This approach includes contributions from the amorphous component that, at the time of writing, is not well understood and is hypothesized to contain primary basaltic glass, nanophase Fe-oxides, amorphous sulfates, and sulfides (Bish et al., 2013; Smith et al., 2019; Rapin et al., 2019; Rampe et al., 2020a; Achilles et al., 2020; Wong et al., 2020). The approach taken here, to input the chemical composition of the host rock and not the mineral composition, raises questions as to what was the phase composition and abundances within the host rock. In discussion section “Comparison of modeled mineral phases with Murray Formation observations” we use comparisons between the modeled data in this paper and the observations made by *Curiosity* to infer compositional characteristics regarding the amorphous phase identified observed in Gale crater, including the possibility of a volcanic glassy component.

Given the young, Amazonian age inferred from K-Ar investigations of the jarosite in the Mojave drill sample (Martin et al., 2017), it is also assumed that the observed akaganeite and jarosite formed during late-stage acidic alteration after the formation of hematite and clay minerals (e.g. Achilles et al., 2020; Rampe et al., 2020b). Thus, the formation of akaganeite and jarosite is not within the scope of this study. However, their chemical composition is included in the calculated alteration composition as their formation is likely the result of localized alteration where elements were remobilized from the products of the first step of diagenesis that is being modeled here.
Pressure-Temperature space

The temperatures selected for modeling were 25 °C, 50 °C, 75 °C and 100 °C, as this broad range can be consistent with the alteration mineralogy observed in Gale crater (Bristow et al., 2018; Rampe et al., 2020a). The pressure was selected to prevent boiling for each temperature. Results were then evaluated with respect to temperature and used as a guide for thermochemical modeling where reactant rock was the calculated chemical alteration compositions for drill samples collected in the Murray formation below VRR and the overall Murray formation.

Redox considerations

No direct measurements of overall redox conditions in the rock were possible (e.g., because of the presence of the amorphous phase; Achilles et al., 2020; Rampe et al., 2020b). Thus, different Fe$^{2+}$/Fe$^{3+}$ ratios were modeled. Resultant modeled mineral assemblages were then compared to assemblages observed by Curiosity to evaluate the plausibility of the model. Models were run at host rock chemistries of 0.1 and 0.5 Fe$^{3+}$/Fe$_{tot}$ to explore the varying effect on the resultant alteration mineral assemblage. SO$_3$ was recalculated as FeS and the equivalent amount of Fe was subtracted from FeO. Cl was recalculated as NaCl and the equivalent amount of Na was subtracted from Na$_2$O.

MODEL RESULTS

Calculation of the host rock composition

The result of the magmatic mineral subtraction process (hypothesis and model set-up section “Host rock composition”) shows a similar chemical alteration composition throughout the Murray formation below VRR (Fig. 3). SiO$_2$ varies between 45 and 49 wt.% (Fig. 3), FeO$_T$ has a larger variation, between 23 and 29 wt.%, and a positive correlation with SiO$_2$. MgO varies between 5 and 7 wt.%, Al$_2$O$_3$ between 5 and 7 wt.%. Alkali oxides (Na$_2$O and K$_2$O) are below 2 wt.% in all calculated alteration compositions below VRR. The three samples acquired on VRR display a larger variability in SiO$_2$, varying from 35 to 54 wt.%. FeO$_T$ has a similar variation to below VRR, varying between 23 and 29 wt.%. MgO is also similar in range on VRR compared to below, 3 to 6 wt.%, while Al$_2$O$_3$ is lower on VRR, ranging from 3 to 6 wt.%. Alkali oxides (Na$_2$O and K$_2$O) are below 2 wt.% in all calculated chemical alteration compositions on VRR.

For the VRR samples, there are some noteworthy observations for the chemical alteration compositions (Fig. 3): The Rock Hall and Highfield drill holes sampled red and grey Jura, respectively, with the FeO$_T$ in our calculation being about 5% higher in Rock Hall than in Highfield. The red Jura target, Rock Hall, is very similar in FeO$_T$ to the red Pettiegrove Point Stoer target. When averaged over the entirety of the drilled samples, chemical alteration compositions below and on VRR are remarkably similar (Fig. 4), pointing towards local element mobility. This confirms that the fluid influence is minimal, and supports our assumption of a dilute incoming fluid, but also points towards the absence of a large-scale fluid movement in late-stage diagenesis after the formation of the hematite-clay dominant mineral assemblage.

The proportion of alteration minerals of the different samples was also assessed, with the ratio of alteration to primary mineral content detailed in Table 1. These ratios indicate that, if Ca-sulfates are excluded, samples below VRR contain a greater fraction of identified alteration products compared to VRR, with Duluth containing the least amount of alteration. This is
consistent with Mangold et al. (2019a; 2019b) who showed a decrease in the Chemical Index of Alteration above the Sutton Island member. We have thus decided to model the average alteration composition of VRR and compare those models to models which use an overall average alteration composition and an average alteration composition for below VRR (excluding Duluth as this had further diagenetic features). A comparison of the mean alteration chemical compositions for the pre-VRR Murray formation (Oudam to Sebina) and for VRR Murray (Stoer to Rock Hall) shows that they are similar (Fig. 4, see also Table S4) with the standard deviation of these mean compositions overlapping.

**Alteration mineralogy in a groundwater-dominated setting**

As described above, we hypothesize that the sediments in the Murray formation on and below VRR were altered in situ from magmatic detrital precursors, and our model with the calculated chemical alteration composition described above then allows us to compare a predicted mineral assemblage with what has been observed by Curiosity. The reasoning for this is that what currently remains as unaltered phases will not have taken part in the reactions, and what is now the alteration assemblage was once a magmatic mineral, e.g., olivine, pyroxene, plagioclase, and has been reacted into the minerals we find now. With this composition we modeled the average VRR Murray alteration composition isothermally at four different temperatures. The resulting mineral assemblages can be divided into three W/R sections. At very high W/R (above W/R of 10,000), models between 50 and 100 °C show a hematite-clay mineral assemblage whereby the dominant clay is chlorite (Fig. 5 and 6). At 25 °C, goethite forms instead of hematite (Fig. 5A and 6A). Between W/R of 10,000 and 100, nontronite dominates; below W/R of about 100 a talc-chlorite assemblage forms. This generalized pattern applies for different redox settings as well as for the VRR-only and ‘all Murray’ compositions (Fig. 5 – 7).

Assessing the influence on temperature in the system that has 10% of the FeO\(_T\) as Fe\(^{3+}\) (Fig. 5A-D) shows that the W/R range in which nontronite forms becomes smaller and the amount of nontronite forming decreases as the temperature increases from 25 °C to 100 °C. Instead of nontronite, a hematite–chlorite assemblage forms at high W/R. Note also the increased abundance of SiO\(_2\) with increasing temperature. At low W/R, changes are limited to small compositional changes, leading to the formation of a few wt.% of epidote at the highest temperature. This metamorphic assemblage is not found at Gale crater, for which reason we focus the discussion in the next section on the high and intermediate W/R.

In the more oxidizing system (50% of the FeO\(_T\) as Fe\(^{3+}\); Fig. 6A-D), changes are subtle. Thermochemical models run at 25 °C show goethite abundances vary from 7.5 wt.%, 2.0 wt.% and 0.0 wt.% for models with 10% Fe\(^{3+}\)/Fe\(_{tot}\) to 9.6 wt.% 1.6 wt.% and 5.2 wt.% for models with 50% Fe\(^{3+}\)/Fe\(_{tot}\), at 10,000 W/R, 1,000 W/R and 100 W/R, respectively. At and above 50 °C, hematite forms instead of goethite. As expected, hematite content increases with increasing Fe\(^{3+}\) availability, meanwhile Fe-sulfide decreases (Tables S5 and S6).

The nontronite field expands towards intermediate and lower W/R, and so does that of the SiO\(_2\)-phase, which forms around W/R of 1,000 – 10,000 (Fig. 5 and 6, Table S7). We note, though, that the overall pattern of a hematite-clay mineral assemblage at high W/R and nontronite at intermediate W/R remains stable (Table 2). However, at high W/R the dominant clays are chlorites, the onset of which occurs at lower W/R with increasing temperature (Fig. 5 and 6). Modeling the different Murray averages (Table S4) at 50 °C and at 10% and 50% FeO\(_T\) as Fe\(^{3+}\) returns almost indistinguishable results (Fig. 7), which demonstrates that the hematite-
clay mineral assemblage is the dominant assemblage at high water to rock ratios over a wide
range of environmental conditions.

**Summary of the modeled alteration mineral assemblages**

Thermochemical modeling undertaken for this study has focused on the hematite-clay
mineral alteration assemblage observed in the Murray formation, with emphasis on sedimentary
rocks on and below VRR. Chemical compositions of the alteration mineral assemblages
observed by *Curiosity* were calculated and used as the reactant rock chemical compositions for
thermochemical modeling. Modeled alteration mineral assemblages have been produced for a
variety of W/R, temperatures and varying Fe$^{3+}$/Fe$_{tot}$ content (Fig. 5 and 6).

For the models in this study, at and above 50 °C hematite forms, which is likely a result
of its higher thermodynamic stability compared to goethite (Cornell and Schwertmann, 2006 and
references therein) and makes these models directly comparable to observations by *Curiosity*.
Table S5 gives details of the modeled hematite abundances showing an increase in hematite
abundance with temperature at 10,000 and 1,000 W/R, with an additional increase associated
with Fe$^{3+}$ availability. Magnetite was allowed to form in the models (Fig. 5 and 6) but did not
form at the W/R assumed relevant. Table S6 demonstrates that Fe-sulfide abundance increases
with temperature at 10,000 W/R with minor wt.% variation at 1,000 W/R and 100 W/R. Fe-
sulfide abundances decrease at all W/R with increased Fe$^{3+}$ availability.

The dominant clay minerals, when formed in our thermochemical models, belong to the
smectite group (nontronite), though in some cases significant chlorites form. Clay mineral
abundance varies significantly with temperature. In summary, the overall clay abundance
decreases with temperature regardless of Fe$^{3+}$ availability (Table S8); however, there is an
increase in clay abundance at 1,000 W/R and 100 W/R from 10% Fe$^{3+}$/Fe$_{tot}$ to 50% Fe$^{3+}$/Fe$_{tot}$ at
each modeled temperature. Summed in Table S7, there is a clear decrease in nontronite
abundance with temperature and a significant increase at 100 W/R with increased Fe$^{3+}$
availability. The chlorite abundance increases with temperature and decreases with Fe$^{2+}$
availability (Table S9). The third group of clay minerals that formed in the models in this study
was talc; for each modeled temperature, talc forms with increasing abundance from 10,000 – 1
W/R (Fig. 5 and 6). Overall, talc abundance decreases with temperature and Fe$^{2+}$ availability
(Table S10).

Thermochemical modeling at 50 °C for the calculated alteration chemical compositions
for the rocks below VRR and the overall Murray formation (Fig. 7) returned near
indistinguishable results at high W/R compared to the calculated chemical alteration composition
for the rocks on VRR (Fig. 5 and 6). This implies dominance of a hematite-clay mineral
assemblage for high W/R over a range of environmental conditions.

To assess redox, all Fe- and S-bearing phases have to be taken into account. We note here
that ferrous (including magnetite) and ferric minerals form at different W/R. The main influence
on redox is expected from variations in the host rock chemical composition, especially in a
subsurface system without access to atmospheric CO$_2$. While, as stated in the results section, the
main silicate mineral composition remains largely unaffected by redox, there are differences in
the hematite/sulfide ratio (Fig. 5 and 6). Taking 75 °C as an example (Tables S5 and S6), 9.6
wt.% hematite and 10 wt.% pyrite precipitate from the system at 1,000 W/R and 10% Fe$_{tot}$ as
Fe$^{3+}$. In contrast, at 50% Fe$_{tot}$ as Fe$^{3+}$ in the host rock, the hematite/sulfide ratio increases from
approximately 1 to 1.4 (12 wt.% hematite and 8.5 wt.% pyrite). However, at lower temperatures, 
and higher and lower W/R, the situation is different.

The pH for the models shown in this paper (Fig. 5 and 6) at 10,000 W/R ranges from 7.9 
to 9.3, as shown in Fig. 8. At 10,000 W/R for the VRR Murray composition derived in Table S4, 
the pH trends from 9.4 to 8.0 for 25 to 100 °C at 10% Fe³⁺/Fe tot content and 9.2 to 7.9 for 25 to 
100 °C at 50% Fe³⁺/Fe tot content. For pre-VRR Murray and overall Murray compositions in 
Table S4, the pH is 8.8 for 10% Fe³⁺/Fe tot content and the pH is 8.7 for 50% Fe³⁺/Fe tot. We 
attribute this increase in pH with decreasing W/R to H being consumed by phyllosilicate 
formation as increasing pH is commonly observed with decreasing W/R and increasing 
phyllosilicate formation (e.g., Schwenzer and Kring 2009, Bridges and Schwenzer, 2012; 
Bridges et al., 2015b).

DISCUSSION

Our discussion first assesses the modeled mineral assemblages in context with the 
observations of the Murray formation. We then compare to other Martian-based models before 
looking at terrestrial comparisons.

Comparison of modeled mineral phases with Murray Formation observations

The modeled early diagenetic mineral assemblages (Fig. 5 – 7) were compared to mineral 
phases observed by CheMin. To compare the modeled clay minerals to observations in the 
Murray formation, their chemistry is considered in discussion section “Clay minerals and 
comparisons to Yellowknife Bay and other Martian terrains, including the Nakhlit Martian 
Meteorites”. Whilst the modeling method used in this study calculates chemical equilibrium at 
specific W/R, it is important to remember that low-temperature diagenetic processes are 
dominated by chemical kinetics (Misra, 2012). For this reason, we rely on comparisons with the 
ground truth – the observed mineralogy at VRR – for the setup of our model and the 
interpretation of our results. It is the combination of ground truth from the rover instruments with 
the models that allow us to arrive at our conclusions.

The abundances of Fe-oxides formed in the models varies with Fe³⁺/Fe tot content and, 
depending on temperature, speciation. Thermochemical models run at 25 °C do not compare well 
with Curiosity with regards to Fe-oxide abundance with clay (Tables 2 and 4), and 
as hematite, not goethite, is observed in the drilled samples (Achilles et al., 2020; Rampe et al., 
2020b). As goethite can transform to hematite over time (Cornell and Schwertmann, 2006 and references therein) there is a possibility that these lower temperature models reflect the actual 
reaction pathway, and that the hematite-clay mineral assemblage observed in the Murray 
transformed from a goethite-clay mineral assemblage with burial and diagenesis of the 
sediments. It is important to note that, like goethite, ferrihydrite or magnetite could have also 
been precursors to the hematite observed in the Murray formation. This possibility is discussed 
by Achilles et al. (2020) for the hematite observed in Oudam and by Rampe et al. (2020b) for the 
samples on VRR. The mechanisms to transform ferrihydrite to hematite include aqueous 
suspension under weakly acidic to weakly alkaline pH, and aging in a humid environment 
(Cornell and Schwertmann, 2006 and references therein). These mechanisms are discussed by 
Rampe et al. (2020b) for VRR with particular note on hematite crystallite size being an indicator 
for formation process. Magnetite was observed by Curiosity on VRR in Duluth, Stoer and 
Highfield (Rampe et al., 2020b), and is also a potential precursor that can be transformed to
hematite by oxidation. Magnetite was included in the chemical alteration compositions. Therefore, the models do not rule out the transformation of other Fe-oxides to hematite. However, in this study it is assumed that the present-day Fe-oxides observed in the Murray formation and VRR are what precipitated at the time of formation and have not since transformed, and so the thermochemical modeling results are compared to *Curiosity* observations.

Ca-sulfate veining has been observed extensively in Gale crater (introduction section) and their removal from the compositional data prior to modeling complicates the identification of Ca-rich phases in our models. In addition, trace amounts of carbonates such as calcite and rhodochrosite have not been observed with CheMin in the samples in this paper at the time of writing (January 2021). For these reasons, trace amounts of calcite, rhodochrosite, merwinite (Ca$_3$MgSi$_2$O$_8$) and spurrite (Ca$_4$Si$_2$O$_8$(CO$_3$)) were excluded from the thermochemical model plots (Fig. 5 – 7). However, siderite may be present in Rock Hall (Rampe et al., 2020b), and Fe/Mg/Fe-Mg carbonates have been identified in SAM EGA data of samples from the Stimson formation (Sutter et al., 2017). It has been suggested, using an experimental approach on martian crust simulants, that carbonates only form when olivine is abundant and would not form in olivine-free assemblages under a CO$_2$ atmosphere (Baron et al., 2019).

The hematite/sulfide ratio is mainly controlled by the variation in the onset of nontronite formation (at systematically lower W/R with increasing temperature) under high-W/R conditions, and at low-W/R by the ratio of nontronite to celadonite as Fe-phyllosilicate phase. This demonstrates that the main redox sensitivity of such systems might not be discernible via the clay minerals alone. Phyllosilicates and their role in interpretation of the Mars samples is still being refined (Hurowitz et al., 2017; Bristow et al., 2018) and models suffer from uncertainties in the thermochemical data for these minerals (Catalano, 2013). Thus, it is encouraging to see that the system changes within strictly constrained boundaries of the input redox parameter conditions of the host rock.

**Comparison of modeled mineral abundances to the observed Murray Formation mineralogy**

For comparing the modeled mineral assemblages in this paper to observations made by *Curiosity*, we also considered the relative abundance of the observed altered phases. The observed CheMin clay mineral / Fe-oxide ratios in Table 4 are most comparable with the modeled ratios at 50 °C and 10,000 W/R shown in Tables 2 and 3, as also shown in Fig. 9. Fig. 9 shows that the observed CheMin ratios are not comparable with modeled clay mineral / Fe-oxide ratios at 50 °C for 1,000 or 100 W/R. Summing all Fe-oxides reported by CheMin significantly lowers the clay mineral / Fe-oxide ratio for VRR (Table 4), largely due to akaganeite in the Rock Hall drilled sample. As previously discussed, akaganeite might belong to a later alteration phase and formed independent of the hematite-clay mineral diagenetic environment where Fe is remobilized. To quantitatively compare the modeled and observed alteration mineral assemblages, we have plotted hematite content vs clay mineral content from *Curiosity* drill samples and from our thermochemical models at 10,000 W/R (Fig. 10). In Fig. 10, CheMin abundances of clay minerals and hematite are normalized to the calculated alteration component, which includes the composition of the amorphous component. The samples analysed by CheMin show a trend, whereby samples with high clay content are generally lower in hematite. The modeled alteration assemblages also show this trend with high clay content corresponding to low
hematite abundances in the modeled alteration assemblage. Thus, the models and samples show
the same correlation between hematite and clays, although the concentration of both is generally
higher in the modeled alteration phase compared to the drilled samples. In Fig. 10, the linear
trend of clay mineral and hematite abundances at 10,000 W/R from our thermochemical models
at 50 °C is parallel to the trend for CheMin samples Marimba, Sebina, Quela, Highfield and
Oudam. Linear regression analysis revealed the comparability of these trends, with a gradient -2.35 ± 0.09 for the thermochemical models run at 50 °C and -1.88 ± 0.56 for Oudam, Marimba,
Sebina, Quela, Duluth, Highfield and Rock Hall, with respective R² values of 0.99 and 0.70. The
uncertainty associated with Duluth clay mineral and hematite abundance is within this negative
correlation, so was included in the regression analyses. Rock Hall does not clearly align with this
negative correlation. So to test whether Rock Hall can be considered an outlier, linear regression
was performed on Oudam, Marimba, Sebina, Quela, Duluth and Highfield. This linear regression
gave a gradient of -2.56 ± 0.45 with an R² value of 0.89. The improvement of the fit confirms
that Rock Hall can be considered an outlier, which is further suggestive of alteration after the
initial hematite-clay mineral main phase alteration modeled in this paper. In Fig. 10, the position
of Stoer relative to negative correlation of clay minerals and hematite in Marimba, Sebina,
Quela, Highfield and Oudam is comparable to the position of the thermochemical models run at
50 °C relative to those run at higher temperature (75 °C and 100 °C). This shows a possible
increase in temperature for the formation of the main phase alteration mineral assemblage in
Stoer.

The plotted data in Fig. 10 also show that, although the bulk alteration composition used
in the thermochemical models varies to a relatively minor extent (Table S4), such small
variations have a significant influence on the precipitated amounts of clay minerals and hematite
in the thermochemical model results. However, this variation in modeled mineral abundance
corresponds to modeled CheMin secondary mineral abundance. In Fig. 10, the shift between the
trend in the CheMin detections and the trend in the thermochemical models raises the question of
the role of the CheMin amorphous component, as chemical contributions from the amorphous
component detected by APXS were included in the chemical alteration compositions used in the
thermochemical modeling.

Here we assume the amorphous component includes sample constituents that cannot be
identified from CheMin data because crystalline materials in the samples below 1 wt.% are
included in the amorphous component. The composition of this amorphous composition is
estimated using CheMin-APXS FULLPAT analysis (Rampe et al., 2020b; Achilles et al., 2020).
The origin of the X-ray amorphous component is presently not fully understood. However,
studies have suggested the presence of nanophase Fe-oxides, amorphous sulfates, silicates and
low amounts of basaltic glass in the amorphous component (Bish et al., 2013; Smith et al., 2019;
Rapin et al., 2019; Rampe et al., 2020a). A study by Bridges et al (2015b), where Portage soil
was used in the starting composition, assumed that the amorphous component contained volcanic
or impact glass, which was then reacted together with olivine to successfully model the alteration
assemblages observed at Yellowknife Bay. As mentioned in introduction section “Overview of
the Murray Formation”, the amorphous material is rich in Fe and Si, suggesting the presence of
nanophase Fe-oxides and amorphous silica (Rampe et al., 2020a; Achilles et al., 2020).
Considering the models presented in this paper assume that all mineralogy is essentially
crystalline, the linear shift between modeled and observed trends in Fig. 10 could be attributed to
an unreacted component of the calculated alteration composition in the drill samples. The
Marimba, Quela, Sebina, Highfield and Stoer mineral abundances are clustered close to the
thermochemical models run with their alteration compositions. As previously noted, the position of Stoer is comparable to higher temperature (75 °C and 100 °C) thermochemical models.

We tested the idea that the amorphous component might only partially belong to the alteration materials by removing the amorphous component from the CheMin mineral abundances for each drill hole and re-normalizing. This improves the fit between thermochemical model results and CheMin observations with regards to the negative correlation between hematite and clay mineral abundances (Fig. 11). Although there are comparable negative correlations, albeit of different gradients, between the drilled samples analysed by CheMin and the thermochemical models in Fig. 11, the trend with temperature shown in Fig. 10 is lost. This suggests that a part of the amorphous component is reactive. We hypothesize that the CheMin alteration composition was not fully reacted during this phase of alteration. Therefore, this could suggest that the amorphous component had a reactive component and a non-reactive component or that time was insufficient to fully react the amorphous component. The latter could be indicative of the alteration of a volcanic glass (e.g. Wolff-Boenisch et al., 2004). Considering the modeling method assumes complete host rock dissolution and igneous phases are still present in Curiosity observations, an unreacted igneous glass component is favored here.

Clay minerals and comparisons to Yellowknife Bay and other Martian terrains, including the Nakhlite Martian Meteorites

The results of this work can be compared to earlier theoretical modeling involving general Martian compositions deduced from Martian meteorite compositions (e.g., Schwenzer and Kring, 2009; Filiberto and Schwenzer, 2013), the composition of rocks at Yellowknife Bay in Gale crater (Bridges et al., 2015b; Schwenzer et al., 2016), and the nakhlite Martian meteorite alteration (Bridges and Schwenzer, 2012). Those models are based on varying knowledge of the reaction path, with general assumptions on the alteration paragenesis from orbiters for the first set of models, more precise knowledge of alteration mineral occurrence from CheMin at Yellowknife Bay, and finally a detailed investigation of the alteration assemblage in the nakhlite meteorites. Our study is comparable to the Yellowknife Bay study (Bridges et al., 2015b) in the level of knowledge of the alteration assemblage, and comparisons between the Yellowknife Bay Sheepbed member of the Bradbury Group and the Murray formation are particularly important in understanding the evolution of fluids in the Gale crater sediments. The dioctahedral clay nontronite in our models is a close analog to the dioctahedral smectite identified in Murray (Bristow et al., 2018), therefore, our models have identified a possible diagenetic process to form the hematite-clay mineral assemblage in the Murray formation. The phyllosilicate-bearing assemblage analyzed in the Gale sediments contains no serpentine, chlorite and, at most, traces of illite (Bristow et al., 2018; Rampe et al., 2020b). Phyllosilicates are widespread in the Murray formation and, when present, dominated by smectites.

We start by comparing our assemblage to general Martian alteration deduced from orbiter observations and the MER rovers. The basis for these models were Martian meteorite compositions, e.g., LEW88516 for the models presented by Schwenzer and Kring (2009). Based on this poikilitic shergottite composition, the models are dominated by forsteritic olivine (57 vol.%, with contributions from pyroxene (22 vol.%) and plagioclase (16 vol.%) (Gleason et al., 1997). These models showed that there are two important observations: nontronite – which is observed from orbit (Ehlmann et al., 2009; Miliken et al., 2010; Carter et al., 2013) and used as the anchor point for this study – most frequently occurs at intermediate temperatures (~150 °C)
and is replaced by chlorite and eventually amphiboles at the higher temperatures of up to 350 °C. At lower temperatures (<90 °C) the dominance of forsteritic olivine causes talc formation to become more dominant. The system is, however, multi-dimensional, as W/R also influences the assemblage, with nontronite generally dominant around W/R of 1,000, being replaced by other phyllosilicates and eventually amphiboles towards lower W/R. At the highest W/R (e.g. 10,000) hematite dominates, with some nontronite. The results are similar to the results of this study focusing on the Murray formation of Gale crater in the comparable pattern of hematite dominance at the highest W/R. They differ in the occurrence of talc at the lower temperatures, which can be explained by the difference between LEW88516 and our calculated alteration compositions for the Murray formation in the MgO concentration (24 wt.% for LEW88516) (Schwenzer and Kring, 2009) vs. 5 wt.% (Table S4). This highlights the importance of host rock composition for the alteration mineral formation, along with the effect of temperature and different W/R ratios. The importance of host rock composition is also highlighted by the fact that models with ‘Home Plate’ MER Spirit rock ‘Fastball’ composition form nontronite at temperatures as low as 13 °C (Filiberto and Schwenzer, 2013). Filiberto and Schwenzer (2013) hypothesized that 13 °C was the temperature at 1 km depth if the surface temperature is zero and the Martian geothermal gradient is 13 °C (Babeyko and Zharkov, 2000). MgO concentration in the assumed host rock is 12 wt.%, significantly lower than for LEW88516, but higher than in this study. MgO/Al₂O₃ ratios are 7.6 for LEW88516, 1.5 for Fastball, and 0.97 for the models in this study (Table S4). This significantly influences the stability of talc compared to nontronite, but also the W/R at which the transition between nontronite and chlorite occurs. Those studies (Filiberto and Schwenzer, 2013) were based on well-known host rock compositions, but for the Gusev alteration mineral assemblage only orbiter data could be used.

Previous modeling for the Sheepbed unit (Bridges et al., 2015b) was based on Portage Soil with 13 wt% MgO, and an MgO/Al₂O₃ ratio of 0.92 (Blake et al., 2013). Clay minerals formed in the first stage of the model, because sulfate minerals were only observed in later veins (Bridges et al., 2015b; Schwenzer et al., 2016). The models showed that the secondary mineral assemblage formed by the reaction of a CO₂-poor and moderately oxidizing, dilute aqueous solution with the sedimentary rocks at 10 – 50 °C and W/R of 100 – 1,000, pH of ~7.5 – 12 (but mainly near neutral through most of the reaction range). Modeled phyllosilicates were Fe-smectite and chlorite. The bulk phyllosilicate composition was close to saponite stoichiometry, which is inferred from CheMin data (Vaniman et al., 2014) though more Fe-rich than the bulk clay mineral composition predicted here for the Murray formation at high W/R (Fig. 12).

The nakhlite meteorite alteration assemblage’s reaction pathways were assessed in a different way (Bridges and Schwenzer, 2012), because both the host rock and the alteration assemblage are known to a high level of detail, including precipitation and re-dissolution successions (e.g. Hicks et al., 2014; Bridges et al., 2019). As a consequence of that knowledge, the host rock composition could be adapted to match differences in dissolution of individual mineral phases. The model concentration has 9 wt.% MgO, and the MgO/Al₂O₃ ratio is 1.9. Furthermore, the observations of the alteration mineral succession in the meteorites indicated a two-step process, carbonate formation first, followed by the silicate-forming phase. This caused a change in the fluid during mineral precipitation, with the silicate-stage fluid being enriched in Si, Na, Al, and K. Taking this into account, Fe-smectites form over a wide range of W/R down to 100 W/R or as low as 10 W/R (Bridges and Schwenzer, 2012). This study on Martian meteorites demonstrates the importance in the accuracy of reaction pathway models presented here for
observations on the nature of the alteration phase, as well as any indication of differences in alteration conditions for different stages of alteration.

For the Sheepbed mudstone of the Bradbury Group there exists the same level of detailed mineralogical knowledge on pre and post alteration mineralogy as we have for the VRR-Murray formation rocks. CheMin X-ray diffraction patterns and SAM evolved gas analyses from mudstone samples demonstrate that phyllosilicate types and abundances vary within the stratigraphic section. Abundant (~20 wt.% of the bulk) Fe\(^{3+}\)-bearing saponite (a trioctahedral smectite) was identified in mudstone from the Yellowknife Bay formation at the base of the section (Tretiman et al., 2014; Vaniman et al., 2014). Mudstone from the Pahrump Hills member, ~60 m up section from Yellowknife Bay and at the base of the Murray formation, has little to no phyllosilicate (up to 8 wt.% of the bulk), and CheMin XRD patterns are consistent with the presence of collapsed smectite (Rampe et al., 2017). Much of the Murray formation stratigraphically above the Pahrump Hills member has abundant (up to 28 wt.% of the bulk) collapsed smectite (Bristow et al., 2018). Both CheMin and SAM data suggest that the structure of the smectite changes from being dominantly trioctahedral (i.e., saponite) lower in the section to being dominantly dioctahedral (i.e., montmorillonite and/or nontronite) higher in the section (Bristow et al., 2018). Duluth was the first sampled analyzed by Curiosity to clearly contain only dioctahedral smectite (nontronite, in this case) (Rampe et al., 2020b; McAdam et al., 2020).

Phyllosilicates on VRR are distinctly different from those detected in mudstone stratigraphically below the ridge. The abundance of phyllosilicates on VRR is relatively low compared to much of the Murray formation (e.g., Highfield contains 5 wt.% phyllosilicate) (Rampe et al., 2020b).

Based on our models, this decrease in phyllosilicates likely relates to the aqueous alteration environment being more oxidizing on the VRR relative to that below (Fig. 10). Furthermore, the phyllosilicates on VRR show a 9.6 Å basal spacing, rather than a 10 Å basal spacing observed in nearly all other phyllosilicate-bearing XRD patterns. This suggests that the phyllosilicate is either a completely collapsed smectite, possibly from alteration in acidic fluids, or that the phyllosilicate is a non-expanding clay mineral, like ferripyrophyllite. SAM data are also consistent with completely collapsed nontronite or ferripyrophyllite on VRR. A similar phyllosilicate was observed in very low abundances in the Oudam sample, drilled stratigraphically above the Pahrump Hills member (Bristow et al., 2018), and the mineralogical similarities between Oudam and Highfield from the gray Jura suggest a comparable aqueous alteration history, as also suggested by our model results (Fig. 10). However, Bristow et al. (2018) hypothesized a detrital origin for ferripyrophyllite at Oudam.

There is a notable similarity in trioctahedral saponite observed in Sheepbed mudstone (Vaniman et al., 2014) and the phyllosilicate identified in the nakhlite martian meteorites. Some of these 11 meteorites contain ferric saponite with a trioctahedral structure (Hicks et al., 2014).

Trioctahedral saponite as an alteration product of Martian mafic material can be regarded as one of the expected types. Fe/Mg smectites, whether tri or dioctahedral have been identified in numerous places from orbit by reflectance spectroscopy (e.g. Ehlimann et al., 2011; Bibring et al., 2006). In the thermochemical models presented in this paper for the Murray formation, the dioctahedral clay nontronite is dominant at W/R up to 10,000 (Fig. 5 – 7). Similar to the Gale in situ analyses, serpentine is absent and chlorite is only dominant at extreme W/R, >10,000 in our models. The slightly elevated temperatures of the models in this study (50 – 100 °C) compared to thermochemical modeling for Yellowknife Bay (10 – 50 °C; Bridges et al., 2015b). In addition, thermochemical modeling for Yellowknife Bay suggested a pH range of 7.5 – 12 (Bridges et al., 2015b), which encompasses the pH range of 7.9 – 9.3 for the alteration assemblages modeled in
this study. To further compare the results of this study to previous works, we have summed the phyllosilicate chemical composition for the models in this paper and included them on a ternary plot in Fig. 12, with comparisons to previous modelling at Yellowknife Bay (Bridges et al., 2015b) and martian meteorite studies (Hicks et al., 2014). As shown in Fig. 12, the modeled phyllosilicates at W/R 10,000 in this study are close to observed Fe-saponites from the nakhlites (Lafayette) and Sheepbed unit model saponite (Bridges et al., 2015b). Murray formation modeling in this paper at 1,000 and 100 W/R trend towards a nontronite composition, as shown in Fig. 12.

Comparison to alteration processes in terrestrial environments

Impact craters are long recognized as depressions in which lakes can form, and sediments accumulate (Cabrol and Grin, 1999; Osinski et al., 2013). Gale crater is no exception. *Curiosity* is exploring a rich stratigraphy of lakebed sediments (e.g., Edgar et al., 2020; Fedo et al., 2018; Grotzinger et al., 2014, 2015; Hurowitz et al., 2017; introduction section). Our hypothesis is that the clay-hematite assemblage is predominantly the result of the first phase of diagenetic alteration that affected the sediments brought into Gale crater. The sedimentary textures within Gale crater, such as cross-bedding, and coarser units such as conglomerates (Williams et al., 2013; introduction section) thereby support the assumption of a dynamic environment with freshwater inflow. This underpins our assumption of a dilute, circumneutral 'groundwater-type' fluid as the initial pore fluid. The main difference between most of those settings and terrestrial analogues is, unfortunately, one of host rock composition as most of Earth's examples are in more evolved rocks and/or less iron-rich than the compositionally-basaltic sequences at Gale crater.

Several terrestrial impact craters show sedimentary sequences of lake beds, starting with coarse grained sequences gradually transitioning to finer grained sequences, but with considerable variation and cyclicity, e.g., the Gardnos impact structure in Norway (Kalleson et al., 2008) and the Boltysch impact structure in the Ukraine (Gilmour et al., 2013). The 14.808 \pm 0.021 Ma (Schmieder et al., 2018), 24 km diameter Ries impact crater (Nördlingen, Germany) has been compared to Gale crater (Arp et al., 2019) as it contains a succession of lake bed sediments and was a closed basin. Interestingly, the transport distances for conglomerates at Ries are considered to be short due to impact-pre-processing (Arp et al., 2019). The sequence begins with clastic sediments with conglomerates and evidence of cyclicity including variation in geochemical properties of the lake (oxygen fugacity, salinity, acidity, and new freshwater inflow; Jankowski, 1977; Arp et al., 2013), which is similar to observations at Gale of fan deposit formation (Williams et al., 2013) and a complex lake chemistry (Hurowitz et al., 2017). This shows that sedimentation was accompanied by significant quantities of water and, therefore, a groundwater-type scenario as assumed in our study is plausible. Alteration mineralogy of the original target rock is consistent with a low-temperature alteration process (Muttik et al., 2008; Sapers et al., 2017; Caudill et al., 2021), whereby clays are montmorillonite-type smectites. This is comparable to our findings of nontronite, as the difference in clay minerals reflects the chemical differences of the host rock: gneiss fragments dominated by feldspars at Ries crater, basaltic sediments dominated by mafic phases and plagioclase at Gale crater. The higher Fe-content of the rocks at Gale crater, 23 – 30 wt.% (Table S2), compared to the Ries basement rocks, 2 – 9 wt.% (von Engelhardt, 1997), explains the different clay mineral chemistry but maintains the low-temperature, groundwater-driven clay formation environmental conditions.
We next compare to a setting more comparable in chemistry while noting that Fe-concentrations at VRR are still higher than those of most terrestrial basalts; Lonar crater (Hagerty and Newsom, 2003) and the basalt alteration widely observed in the Deccan trap basalts. In the Northeastern part of the Deccan Traps, near Jabalpur, unweathered basalts generally contain between 11 and 15 wt.% FeO$_{tot}$ (Peng et al., 1998). Investigation of sedimentary and weathered basalt samples by Salil et al. (1997) in the same region near Jabalpur show that clay minerals have a similar FeO$_{tot}$ content compared to the basalts: between 10 and 17 wt.% with a dominance of Fe$_2$O$_3$. Iron concentrations between rock and alteration assemblage appear constant, if not slightly higher in the sediments and basaltic alteration assemblages, which is comparable to our observations where Fe in the fluid is generally very low (~$10^8$) at 10,000 W/R (Fig. S1). For the Deccan Traps, Fe-concentrations in runoff water of the region appear comparably very low as they are not reported for the region (Gupta et al., 2011), or more widely in cold and warm spring waters as well as runoff across the Deccan (Minissale et al., 2000). Clay-mineralogy in the sediments and Deccan alteration is Fe, Mg-rich smectites, which the authors describe as similar to but not quite nontronitic as the Fe-content is below 50% (Salil et al., 1997). This shows that water-dominated, groundwater-rock reactions such as proposed here occur in weathering or diagenetic environments and are similar to the observed clay mineralogy at the Gale site and in our models.

**CONCLUSIONS**

In this paper we showed that local element mobility rather than large scale, basin-wide fluid movement is the most likely alteration mechanism for the main phase of alteration in the Murray formation. For this, we derived chemical alteration compositions using data from CheMin (Achilles et al., 2020; Rampe et al., 2020b) and APXS (Thompson et al., 2020) and found them to be very similar for the Murray formation below as well as on Vera Rubin ridge. After this main phase, based on the age dating and the local nature of its occurrence, a younger alteration event caused the observed akaganeite and jarosite, whereby we assume that Fe was remobilized from the mineral assemblages in our models (i.e. Fe-sulfides).

Our thermochemical modeling results indicate that the alteration that produced the early diagenetic hematite-clay mineral assemblage in the Murray formation, below and on VRR, can be explained by reaction of dilute brine with the calculated alteration compositions at temperatures between 50 and an upper limit of 100 °C, and at a notably high W/R of approximately 10,000 with pH ranging from 7.9 to 9.3. Modeling at 25 °C produces goethite, which could transition to hematite at a later stage however, this phase together with the relative abundances with the clay content are not directly comparable to *Curiosity* observations. We interpret the high W/R as enhanced groundwater flow through this part of the Gale sedimentary sequence compared to that in the Bradbury Group and its Yellowknife Bay mudstone, where lower temperature alteration (20 – 50 °C) with a broader pH range of 7.5 – 12 (Bridges et al., 2015b). This further constraint on the fluid temperature and pH in the models presented in this paper is likely to have important implications for microbial habitability.

The trends between the modeled and observed phyllosilicate and hematite abundances are comparable, but not exact due to our model not precipitating metastable phases such as the CheMin amorphous component. We interpret the amorphous component to have partially contributed to the phase of alteration modeled in this study. Future work is necessary to determine the composition and origin of the CheMin amorphous component, and thus how its
chemistry can be most appropriately used in thermochemical modeling studies. We will continue modeling the Gale crater sedimentary mineral assemblage which in future work we aim to extend from the Bradbury and Murray-VRR, into the clay-bearing Glen Torridon and overlying Sulfate Units.
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Fig. 1. Stratigraphic column detailing the morphological characteristics of the stratigraphic groups and units encountered by the Curiosity rover up to and including the Jura member. Drill holes of interest in this study are annotated: Oudam (OU), Marimba (MB), Quela (QL), Sebina (SB), Duluth (DU), Stoer (ST), Highfield (HF), and Rock Hall (RH). Image credit: The MSL sedimentology and stratigraphy working group.
Fig. 2. HiRISE orbital overview of the NASA Mars Science Laboratory *Curiosity* rover’s traverse from Bradbury Landing up to sol 2359. Select locations are annotated as well as sol 1809 when *Curiosity* ascended on to Vera Rubin ridge (VRR), and sol 2302 when *Curiosity* traversed off VRR and into the Clay Bearing Unit. Adapted image from NASA-JPL/Fraeman.
Table 1. Ratio of alteration to primary mineral content per drill hole, including and excluding amorphous component as an alteration component. Ca-sulfates were excluded. Errors propagated from uncertainty in CheMin mineral abundances. Pyroxenes and feldspars were considered to be primary minerals. Fe-oxides, jarosite, quartz, halite, fluorapatite and phyllosilicates were considered to be alteration. Source data take from Achilles et al. (2020) and Rampe et al. (2020b).

<table>
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<tr>
<th>Drill hole</th>
<th>Including Amorphous</th>
<th>Excluding Amorphous</th>
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<tbody>
<tr>
<td>Oudam</td>
<td>2.1 ± 0.4</td>
<td>0.6 ± 0.1</td>
</tr>
<tr>
<td>Marimba</td>
<td>4.9 ± 0.8</td>
<td>2.3 ± 0.3</td>
</tr>
<tr>
<td>Quela</td>
<td>4.1 ± 0.8</td>
<td>1.3 ± 0.1</td>
</tr>
<tr>
<td>Sebina</td>
<td>5.3 ± 1.0</td>
<td>1.8 ± 0.2</td>
</tr>
<tr>
<td>Duluth</td>
<td>2.9 ± 0.3</td>
<td>0.8 ± 0.3</td>
</tr>
<tr>
<td>Stoer</td>
<td>2.3 ± 0.2</td>
<td>1.0 ± 0.2</td>
</tr>
<tr>
<td>Highfield</td>
<td>2.5 ± 0.2</td>
<td>0.6 ± 0.1</td>
</tr>
<tr>
<td>Rock Hall</td>
<td>2.0 ± 0.6</td>
<td>0.9 ± 0.2</td>
</tr>
</tbody>
</table>
Fig. 3. Main-phase alteration assemblage compositions for Oudam, Marimba, Quela, Sebina, Duluth, Stoer, Highfield and Rock Hall. Calculated using APXS and CheMin results (Achilles et al., 2020; Rampe et al., 2020b), as explained in hypothesis and model set-up section “Host Rock Composition”. Samples Oudam, Marimba, Quela, Sebina, and Duluth are from pre-VRR Murray formation localities. Stoer, Highfield and Rock Hall represent VRR Murray. *indicates drill samples acquired form VRR.

Fig. 4. Mean alteration compositions for the Pre-VRR Murray formation (Oudam, Marimba, Quela, Sebina), VRR Murray Formation (Stoer, Highfield and Rock Hall), the overall Murray Formation. Calculated using APXS and CheMin results. Error bars show standard deviation for each calculated mean, illustrating the statistical similarity between the calculated alteration compositions.
Fig. 5. CHIM-XPT results for the calculated VRR alteration composition (Table S4) reacted with GPW (Bridges et al., 2015b). Reactions at 25, 50, 75 and 100 °C, and 10% of the FeO$_T$ is Fe$^{3+}$, as indicated. Goethite only precipitated at 25 °C (A), and hematite did not precipitate at that temperature. K-nontronite precipitated at 25 °C (A), whereas Na-nontronite did not precipitate at 100 °C (D) and Ca-nontronite precipitated at all temperatures. Chlor-apatite formed at 25 °C (A), 50 °C (B) and 75 °C (C), whereas apatite formed at 75 °C (C) and 100 °C (D). Trace abundances of calcite, spurrite and merwinite are not included on the plots.
Fig. 6. CHIM-XPT results for the calculated VRR alteration composition (Table S4) reacted with GPW (Bridges et al., 2015b). Reactions at 25, 50, 75 and 100 °C, and 50% of the FeO_T is Fe^{3+}, as indicated. Goethite only precipitated at 25 °C (A), and hematite did not precipitate at that temperature. K-nontronite precipitated at 25 °C (A) and 50°C (B), whereas Na-nontronite did not precipitate at 100 °C (D) and Ca-nontronite precipitated at all temperatures. Rhodochrosite precipitated at 25 °C (A) and 50 °C (B), and trace wt.% of calcite precipitated at all temperatures for 50% Fe^{3+}/Fe_{tot} (A,B,C,D). Trace abundances of calcite, rhodochrosite, spurrite and merwinite are not included on the plots.
Table 2. Summed VRR clay mineral wt.% / Fe-oxide wt.% ratio at high (10,000), intermediate (1,000) and low (100) W/R ratios in thermochemical models (Fig. 6 and 7) using the calculated VRR mean alteration composition (Table S4) as the host rock (at both 10 and 50% Fe$^{3+}$/Fe$_{\text{tot}}$) reacted with GPW (Bridges et al., 2015b) at 25, 50, 75 and 100 °C using CHIM-XPT (Reed et al., 2010).

<table>
<thead>
<tr>
<th>Temperature °C</th>
<th>10% Fe$^{3+}$/Fe$_{\text{tot}}$ Models</th>
<th>50% Fe$^{3+}$/Fe$_{\text{tot}}$ Models</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>10,000 W/R</td>
<td>1,000 W/R</td>
</tr>
<tr>
<td>25 °C</td>
<td>10.4</td>
<td>43.2</td>
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<tr>
<td>50 °C</td>
<td>1.2</td>
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<tr>
<td>75 °C</td>
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<td>6.9</td>
</tr>
<tr>
<td>100 °C</td>
<td>1.2</td>
<td>3.8</td>
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Fig. 7. CHIM-XPT results for the calculated alteration composition for the Murray formation and the Murray and VRR (overall) (Table S4) reacted with GPW (Bridges et al., 2015b). Reactions are at 50 °C and the Fe$^{3+}$ content of FeO$_r$ is at 10% and 50%, as indicated. Trace abundances of rhodochrosite, calcite, spurrite and merwinite are not included on the plot.
Table 3. Summed clay wt.% / Fe-oxide wt.% ratio at high (10,000), intermediate (1,000) and low
(100) W/R ratios in models for the pre-VRR Murray mean and overall mean calculated alteration
compositions (Table S4) as the host rock, at 10 and 50% Fe$^{3+}$/Fe$_{tot}$, reacted with GPW (Bridges
et al., 2015b) at 50 °C using CHIM-XPT (Reed et al., 2010).

<table>
<thead>
<tr>
<th></th>
<th>10% Fe$^{3+}$/Fe$_{tot}$ Models</th>
<th>50% Fe$^{3+}$/Fe$_{tot}$ Models</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>10,000 W/R 1,000 W/R 100 W/R</td>
<td>10,000 W/R 1,000 W/R 100 W/R</td>
</tr>
<tr>
<td>Pre-VRR</td>
<td>3.0 -- --</td>
<td>2.6 -- --</td>
</tr>
<tr>
<td>Overall</td>
<td>1.9 -- --</td>
<td>1.7 -- --</td>
</tr>
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</table>

Fig. 8. Plot showing pH variation with W/R for the CHIM-XPT models shown in Fig. 5 – 7.
Starting compositions for Pre-VRR Murray, VRR Murray and Overall Murray used in the
modeling are shown in Table S4.
Table 4. Calculated clay mineral / hematite and clay mineral / Fe-oxide sum for drilled samples in the pre-VRR Murray formation and VRR, determined using CheMin wt.% mineral abundances (Achilles et al., 2020; Rampe et al., 2020b). Means are included for pre-VRR and VRR samples, as well as an overall mean. Errors shown are propagated errors calculated from errors in mineral abundances.

<table>
<thead>
<tr>
<th>Drill Hole</th>
<th>Clay mineral / Hematite</th>
<th>Clay mineral / Fe-oxide sum</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oudam</td>
<td>0.3 ± 0.1</td>
<td>0.3 ± 0.1</td>
</tr>
<tr>
<td>Marimba</td>
<td>4.4 ± 0.9</td>
<td>4.4 ± 0.9</td>
</tr>
<tr>
<td>Quela</td>
<td>2.3 ± 0.4</td>
<td>2.3 ± 0.4</td>
</tr>
<tr>
<td>Sebina</td>
<td>2.8 ± 0.6</td>
<td>2.8 ± 0.6</td>
</tr>
<tr>
<td>Duluth</td>
<td>2.4 ± 1.1</td>
<td>2.1 ± 1.0</td>
</tr>
<tr>
<td><strong>Pre-VRR mean</strong></td>
<td><strong>2.4 ± 0.3</strong></td>
<td><strong>2.4 ± 0.3</strong></td>
</tr>
<tr>
<td>Stoer</td>
<td>0.6 ± 0.3</td>
<td>0.6 ± 0.3</td>
</tr>
<tr>
<td>Highfield</td>
<td>0.5 ± 0.2</td>
<td>0.5 ± 0.2</td>
</tr>
<tr>
<td>Rock Hall</td>
<td>4.5 ± 2.1</td>
<td>1.5 ± 0.7</td>
</tr>
<tr>
<td><strong>VRR mean</strong></td>
<td><strong>1.9 ± 0.7</strong></td>
<td><strong>0.8 ± 0.3</strong></td>
</tr>
<tr>
<td><strong>Overall mean</strong></td>
<td><strong>2.2 ± 0.3</strong></td>
<td><strong>1.8 ± 0.2</strong></td>
</tr>
</tbody>
</table>

Fig. 9. Plot comparing calculated clay mineral / hematite and clay mineral / Fe-oxide sum for drill samples in the pre-VRR Murray formation and VRR, determined using CheMin wt.% mineral abundances (Achilles et al., 2020; Rampe et al., 2020b). Means are included for pre-VRR and VRR samples, as well as an overall mean. Errors shown are propagated errors calculated from errors in mineral abundances. Values for models run at 50 °C at 10,000 W/R, 1,000 W/R and 100 W/R from Tables 2 and 3 are included for comparison.
Fig. 10. Clay mineral abundance vs hematite abundance from CheMin (Achilles et al., 2020; Rampe et al., 2020b) normalized to the alteration component (including amorphous phase) compared to CHIM-XPT results in this study (Figs 5–7; Tables S5 and S8) at high (10,000) W/R. Drill holes Oudam, Marimba, Quela, Sebina, Duluth, Stoer, Highfield and Rock Hall are annotated with OU, MB, QL, SB, DU, ST, HF and RH, respectively. Linear regression carried out on thermochemical models at 50 °C in the Fig. revealed the trend $y = (-2.35 \pm 0.09)x + (102.15 \pm 2.16)$ with an $R^2$ value of 0.99. Comparative linear regression carried out on CheMin drill samples Oudam, Marimba, Sebina, Quela, Duluth, Highfield and Rock Hall in the Fig. gave a trend of $y = (-1.88 \pm 0.56)x + (39.23 \pm 6.24)$ with an $R^2$ value of 0.70. This trend changed to $y = (-2.56 \pm 0.45)x + (48.38 \pm 5.33)$ with an $R^2$ value of 0.89 with Rock Hall removed from the analysis.
Fig. 11. Clay mineral abundance vs hematite abundance from CheMin (Achilles et al., 2020; Rampe et al., 2020b) normalized to the alteration component (excluding the amorphous component) compared to CHIM-XPT results in this study (Fig. 5 – 7; Tables S5 and S8) at high (10,000) W/R. The comparable negative correlations of different gradients for the CheMin drill holes and the thermochemical models and the trend with temperature shown in Fig. 10 indicates that only a part of the derived alteration composition is reactive. Drill hole names and abbreviations detailed in Fig. 10 caption.
Fig. 12. Mg-Si+Al-Fe\textsubscript{tot} at. ternary plot. Results at 10,000 (red), 1,000 (green) and 100 (blue) W/R from the thermochemical models in this paper are plotted with some additional data from other sources for comparison. Lafayette saponite data taken from Hicks et al. (2014). CHIM-XPT results for the Sheepbed unit from Bridges et al. (2015b) at 100 W/R are shown for comparison.
Table S1. Table outlining the crystal chemistries for the magmatic phases in Oudam, Marimba, Quela, Sebina, Duluth, Stoer, Highfield and Rock Hall.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Plagioclase</th>
<th>Alkali feldspar</th>
<th>Orthopyroxene</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oudam</td>
<td>An40(7)(^a)</td>
<td>-</td>
<td>Mg(<em>{0.79(7)})Fe(</em>{1.20(8)})Ca(_{0.01(4)})Si(_2)O(_6)(^c)</td>
</tr>
<tr>
<td>Marimba</td>
<td>An39(5)(^a)</td>
<td>K(<em>{0.76(13)})Na(</em>{0.24})Al(_3)Si(_3)O(_8)(^b)</td>
<td>Mg(<em>{0.79(7)})Fe(</em>{1.20(8)})Ca(_{0.01(4)})Si(_2)O(_6)(^c)</td>
</tr>
<tr>
<td>Quela</td>
<td>An39(6)(^a)</td>
<td>K(<em>{0.65(18)})Na(</em>{0.35})Al(_3)Si(_3)O(_8)(^b)</td>
<td>Mg(<em>{0.79(7)})Fe(</em>{1.20(8)})Ca(_{0.01(4)})Si(_2)O(_6)(^c)</td>
</tr>
<tr>
<td>Sebina</td>
<td>An42(6)(^a)</td>
<td>K(<em>{0.65(18)})Na(</em>{0.35})Al(_3)Si(_3)O(_8)(^b)</td>
<td>Mg(<em>{0.79(7)})Fe(</em>{1.20(8)})Ca(_{0.01(4)})Si(_2)O(_6)(^c)</td>
</tr>
<tr>
<td>Duluth</td>
<td>An34(4)(^d)</td>
<td>K(<em>{0.65(18)})Na(</em>{0.35})Al(_3)Si(_3)O(_8)(^b)</td>
<td>Mg(<em>{0.79(7)})Fe(</em>{1.20(8)})Ca(_{0.01(4)})Si(_2)O(_6)(^c)</td>
</tr>
<tr>
<td>Stoer</td>
<td>An41(4)(^d)</td>
<td>K(<em>{0.65(18)})Na(</em>{0.35})Al(_3)Si(_3)O(_8)(^b)</td>
<td>Mg(<em>{0.79(7)})Fe(</em>{1.20(8)})Ca(_{0.01(4)})Si(_2)O(_6)(^c)</td>
</tr>
<tr>
<td>Highfield</td>
<td>An28(3)(^d)</td>
<td>K(<em>{0.65(18)})Na(</em>{0.35})Al(_3)Si(_3)O(_8)(^b)</td>
<td>Mg(<em>{0.79(7)})Fe(</em>{1.20(8)})Ca(_{0.01(4)})Si(_2)O(_6)(^c)</td>
</tr>
<tr>
<td>Rock Hall</td>
<td>An44(7)(^d)</td>
<td>K(<em>{0.65(18)})Na(</em>{0.35})Al(_3)Si(_3)O(_8)(^b)</td>
<td>Mg(<em>{0.79(7)})Fe(</em>{1.20(8)})Ca(_{0.01(4)})Si(_2)O(_6)(^c)</td>
</tr>
</tbody>
</table>

\(^a\) derived by Achilles et al. (2020). \(^b\) values for average Murray formation for alkali feldspar derived by Morrison et al. (2018b). \(^c\) values for average Yellowknife Bay for orthopyroxene derived by Morrison et al. (2018b). \(^d\) derived by Rampe et al. (2020).
Table S2. Table detailing calculated alteration compositions for Oudam, Marimba, Quela, Sebina, Duluth, Stoer, Highfield and Rock Hall. Errors propagated from APXS accuracy, uncertainty in CheMin mineral abundances and uncertainties associated with magmatic phase crystal chemistry. Source data take from Achilles et al. (2020) and Rampe et al. (2020).

<table>
<thead>
<tr>
<th></th>
<th>Oudam</th>
<th>Marimba</th>
<th>Quela</th>
<th>Sebina</th>
</tr>
</thead>
<tbody>
<tr>
<td>SiO₂</td>
<td>48.83 ± 2.55</td>
<td>47.33 ± 1.32</td>
<td>44.85 ± 1.47</td>
<td>45.38 ± 1.44</td>
</tr>
<tr>
<td>TiO₂</td>
<td>1.77 ± 0.07</td>
<td>1.37 ± 0.04</td>
<td>1.38 ± 0.05</td>
<td>1.30 ± 0.04</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>5.08 ± 0.49</td>
<td>6.18 ± 0.40</td>
<td>5.79 ± 0.49</td>
<td>6.45 ± 0.39</td>
</tr>
<tr>
<td>Cr₂O₃</td>
<td>0.54 ± 0.02</td>
<td>0.42 ± 0.01</td>
<td>0.38 ± 0.01</td>
<td>0.45 ± 0.02</td>
</tr>
<tr>
<td>FeO₇</td>
<td>29.30 ± 0.98</td>
<td>28.47 ± 0.61</td>
<td>23.79 ± 0.56</td>
<td>22.91 ± 0.70</td>
</tr>
<tr>
<td>MnO</td>
<td>0.37 ± 0.02</td>
<td>0.11 ± 0.01</td>
<td>0.29 ± 0.01</td>
<td>0.25 ± 0.01</td>
</tr>
<tr>
<td>MgO</td>
<td>7.42 ± 0.40</td>
<td>5.74 ± 0.25</td>
<td>4.99 ± 0.18</td>
<td>5.75 ± 0.31</td>
</tr>
<tr>
<td>CaO</td>
<td>0.40 ± 0.65</td>
<td>2.19 ± 0.38</td>
<td>5.47 ± 0.32</td>
<td>4.57 ± 0.30</td>
</tr>
<tr>
<td>Na₂O</td>
<td>1.56 ± 0.26</td>
<td>1.51 ± 0.12</td>
<td>1.51 ± 0.21</td>
<td>1.89 ± 0.20</td>
</tr>
<tr>
<td>K₂O</td>
<td>1.46 ± 0.08</td>
<td>0.71 ± 0.12</td>
<td>0.65 ± 0.10</td>
<td>0.71 ± 0.08</td>
</tr>
<tr>
<td>P₂O₅</td>
<td>0.88 ± 0.09</td>
<td>1.34 ± 0.09</td>
<td>1.45 ± 0.10</td>
<td>0.86 ± 0.07</td>
</tr>
<tr>
<td>SO₃</td>
<td>1.81 ± 0.31</td>
<td>4.02 ± 0.47</td>
<td>8.14 ± 0.27</td>
<td>8.12 ± 0.32</td>
</tr>
<tr>
<td>Cl</td>
<td>0.59 ± 0.02</td>
<td>0.61 ± 0.03</td>
<td>1.33 ± 0.03</td>
<td>1.35 ± 0.26</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th></th>
<th>Duluth</th>
<th>Stoer</th>
<th>Highfield</th>
<th>Rock Hall</th>
</tr>
</thead>
<tbody>
<tr>
<td>SiO₂</td>
<td>38.34 ± 1.33</td>
<td>42.91 ± 2.24</td>
<td>53.84 ± 2.18</td>
<td>35.91 ± 3.25</td>
</tr>
<tr>
<td>TiO₂</td>
<td>1.49 ± 0.05</td>
<td>1.64 ± 0.06</td>
<td>1.31 ± 0.05</td>
<td>1.69 ± 0.09</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>3.49 ± 0.34</td>
<td>3.36 ± 0.64</td>
<td>6.66 ± 0.59</td>
<td>3.10 ± 1.09</td>
</tr>
<tr>
<td>Cr₂O₃</td>
<td>0.48 ± 0.04</td>
<td>0.46 ± 0.05</td>
<td>0.42 ± 0.05</td>
<td>0.52 ± 0.03</td>
</tr>
<tr>
<td>FeO₇</td>
<td>29.71 ± 0.86</td>
<td>29.46 ± 0.99</td>
<td>23.46 ± 1.07</td>
<td>28.19 ± 2.28</td>
</tr>
<tr>
<td>MnO</td>
<td>0.36 ± 0.04</td>
<td>0.27 ± 0.02</td>
<td>0.22 ± 0.02</td>
<td>0.13 ± 0.02</td>
</tr>
<tr>
<td>MgO</td>
<td>3.28 ± 0.31</td>
<td>6.17 ± 0.36</td>
<td>5.54 ± 0.44</td>
<td>4.65 ± 1.11</td>
</tr>
<tr>
<td>CaO</td>
<td>6.72 ± 0.37</td>
<td>3.32 ± 0.67</td>
<td>1.29 ± 0.54</td>
<td>3.67 ± 1.17</td>
</tr>
<tr>
<td>Na₂O</td>
<td>1.09 ± 0.20</td>
<td>1.25 ± 0.25</td>
<td>1.28 ± 0.24</td>
<td>1.40 ± 0.28</td>
</tr>
<tr>
<td>K₂O</td>
<td>0.93 ± 0.11</td>
<td>1.10 ± 0.27</td>
<td>1.01 ± 0.13</td>
<td>1.05 ± 0.06</td>
</tr>
<tr>
<td>P₂O₅</td>
<td>1.56 ± 0.10</td>
<td>1.31 ± 0.08</td>
<td>1.17 ± 0.08</td>
<td>1.47 ± 0.10</td>
</tr>
<tr>
<td>SO₃</td>
<td>11.25 ± 0.45</td>
<td>7.30 ± 0.85</td>
<td>2.44 ± 0.72</td>
<td>14.69 ± 1.55</td>
</tr>
<tr>
<td>Cl</td>
<td>1.30 ± 0.05</td>
<td>1.45 ± 0.06</td>
<td>1.37 ± 0.07</td>
<td>3.54 ± 0.16</td>
</tr>
</tbody>
</table>
**Table S3.** Table detailing rationale for minerals prevented from forming in CHIM-XPT (Reed et al., 2010) thermochemical modelling. For acmite, akermanite, antigorite, Fe-anthophyllite, ferroactinolite, bredigite, grunerite, jennite, pyroxmangite, riebeckite, tobermorite, tremolite, wollastonite see Melwani Daswani et al. (2016).

<table>
<thead>
<tr>
<th>Mineral</th>
<th>Rationale</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al-free_chlorite</td>
<td>Caused convergence issues during modeling</td>
<td>Tantawy et al., 2014</td>
</tr>
<tr>
<td>Belite</td>
<td>Industrial mineral in cement</td>
<td>Tantawy et al., 2014</td>
</tr>
<tr>
<td>Clinozoisite</td>
<td>Clinozoisite-epidote series, members occur from ~130 bar and 320°C.</td>
<td>Deer et al., 2013 and references therein</td>
</tr>
<tr>
<td>Deerite</td>
<td>Forms from 200 °C and 8 kbar.</td>
<td>Vernié et al., 1986 and references therein</td>
</tr>
<tr>
<td>Diopside</td>
<td>Ca clinopyroxene. In series with hedenbergite</td>
<td>Deer et al., 2013 and references therein</td>
</tr>
<tr>
<td>Epidote, Fe-epidote</td>
<td>Clinozoisite-epidote series, members occur from ~130 bar and 320°C.</td>
<td>Deer et al., 2013 and references therein</td>
</tr>
<tr>
<td>Hedenbergite</td>
<td>Ca clinopyroxene. In series with diopside</td>
<td>Deer et al., 2013 and references therein</td>
</tr>
<tr>
<td>Prehnite</td>
<td>Forms at 200-300 °C.</td>
<td>Robinson &amp; Bevans, 1999</td>
</tr>
<tr>
<td>Rankinite</td>
<td>High temperature Ca-silicate phase.</td>
<td>Treiman &amp; Essene, 1983</td>
</tr>
<tr>
<td>Rhodonite</td>
<td>Pyroxenoid</td>
<td>Pinckney &amp; Burnham, 1988</td>
</tr>
<tr>
<td>Tephroite</td>
<td>Mn-olivine</td>
<td>Deer et al., 2013 and references therein</td>
</tr>
<tr>
<td>Tilleyite</td>
<td>Occurs in contact metamorphic zone between igneous rocks and limestones. Low pressure, high-T.</td>
<td>Harker, 1959</td>
</tr>
<tr>
<td>tr.625ac.25prg.125</td>
<td>Solid solution of tremolite, actinolite and paragonite</td>
<td>Melwani Daswani et al., 2016 and references therein</td>
</tr>
<tr>
<td>Vesuvianite</td>
<td>Requires T &gt; 360 °C and P&gt; 50 bar.</td>
<td>Deer et al., 2013 and references therein</td>
</tr>
</tbody>
</table>
Table S4. Details of mean alteration compositions for the Murray formation, VRR, and an overall composition. Errors propagated from APXS accuracy errors, uncertainty in CheMin mineral abundances and uncertainties associated with magmatic phase crystal chemistry. Standard deviations for each mean is also detailed. Source data take from Achilles et al. (2020) and Rampe et al. (2020).

<table>
<thead>
<tr>
<th></th>
<th>Pre-VRR Murray*</th>
<th>VRR Murray</th>
<th>Overall*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean</td>
<td>Standard Deviation</td>
<td>Mean</td>
</tr>
<tr>
<td>SiO₂</td>
<td>46.60 ± 3.48</td>
<td>1.83</td>
<td>44.22 ± 4.79</td>
</tr>
<tr>
<td>TiO₂</td>
<td>1.45 ± 0.10</td>
<td>0.21</td>
<td>1.55 ± 0.12</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>5.87 ± 0.92</td>
<td>0.60</td>
<td>4.37 ± 1.70</td>
</tr>
<tr>
<td>Cr₂O₃</td>
<td>0.45 ± 0.03</td>
<td>0.07</td>
<td>0.47 ± 0.07</td>
</tr>
<tr>
<td>FeO</td>
<td>26.12 ± 1.45</td>
<td>3.23</td>
<td>27.04 ± 2.73</td>
</tr>
<tr>
<td>MnO</td>
<td>0.25 ± 0.04</td>
<td>0.11</td>
<td>0.21 ± 0.04</td>
</tr>
<tr>
<td>MgO</td>
<td>5.97 ± 0.56</td>
<td>1.03</td>
<td>5.45 ± 1.27</td>
</tr>
<tr>
<td>CaO</td>
<td>3.16 ± 5.19</td>
<td>2.30</td>
<td>2.76 ± 2.12</td>
</tr>
<tr>
<td>Na₂O</td>
<td>1.62 ± 0.41</td>
<td>0.18</td>
<td>1.31 ± 0.42</td>
</tr>
<tr>
<td>K₂O</td>
<td>0.88 ± 0.23</td>
<td>0.39</td>
<td>1.05 ± 0.29</td>
</tr>
<tr>
<td>P₂O₅</td>
<td>1.13 ± 0.18</td>
<td>0.30</td>
<td>1.32 ± 0.16</td>
</tr>
<tr>
<td>SO₃</td>
<td>5.52 ± 1.19</td>
<td>3.14</td>
<td>8.14 ± 3.00</td>
</tr>
<tr>
<td>Cl</td>
<td>0.97 ± 0.20</td>
<td>0.43</td>
<td>2.12 ± 0.15</td>
</tr>
</tbody>
</table>

*Not including Duluth.

Table S5. Hematite wt.% at high (10,000), intermediate (1,000) and low (100) W/R ratios in thermochemical models (Figures 5 and 6) and using the calculated VRR mean alteration composition (Table S4) as the host rock (at both 10 and 50% Fe³⁺/Fe tot) reacted with GPW (Bridges et al., 2015b) at 50, 75 and 100 °C using CHIM-XPT (Reed et al., 2010).

<table>
<thead>
<tr>
<th></th>
<th>10% Fe³⁺/Fe tot Models</th>
<th>50% Fe³⁺/Fe tot Models</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>10,000 W/R</td>
<td>1,000 W/R</td>
</tr>
<tr>
<td>50 °C</td>
<td>28.6</td>
<td>2.1</td>
</tr>
<tr>
<td>75 °C</td>
<td>35.8</td>
<td>9.6</td>
</tr>
<tr>
<td>100 °C</td>
<td>36.4</td>
<td>14.5</td>
</tr>
</tbody>
</table>
Table S6. Pyrite wt.% at high (10,000), intermediate (1,000) and low (100) W/R ratios in thermochemical models (Figures 5 and 6) using the calculated VRR mean alteration composition (Table S4) as the host rock (at both 10 and 50% Fe\(^{3+}/Fe_{tot}\)) reacted with GPW (Bridges et al., 2015b) at 25, 50, 75 and 100 °C using CHIM-XPT (Reed et al., 2010).

<table>
<thead>
<tr>
<th></th>
<th>10% Fe(^{3+}/Fe_{tot}) Models</th>
<th>50% Fe(^{3+}/Fe_{tot}) Models</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>10,000 W/R</td>
<td>1,000 W/R</td>
</tr>
<tr>
<td>25 °C</td>
<td>10.8</td>
<td>9.9</td>
</tr>
<tr>
<td>50 °C</td>
<td>12.4</td>
<td>10.0</td>
</tr>
<tr>
<td>75 °C</td>
<td>15.4</td>
<td>10.0</td>
</tr>
<tr>
<td>100 °C</td>
<td>15.1</td>
<td>10.0</td>
</tr>
</tbody>
</table>

Table S7. Summed nontronite wt.% at high (10,000), intermediate (1,000) and low (100) W/R ratios in thermochemical models (Figures 5 and 6) using the calculated VRR mean alteration composition (Table S4) as the host rock (at both 10 and 50% Fe\(^{3+}/Fe_{tot}\)) reacted with GPW (Bridges et al., 2015b) at 25, 50, 75 and 100 °C using CHIM-XPT (Reed et al., 2010).

<table>
<thead>
<tr>
<th></th>
<th>10% Fe(^{3+}/Fe_{tot}) Models</th>
<th>50% Fe(^{3+}/Fe_{tot}) Models</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>10,000 W/R</td>
<td>1,000 W/R</td>
</tr>
<tr>
<td>25 °C</td>
<td>55.1</td>
<td>62.0</td>
</tr>
<tr>
<td>50 °C</td>
<td>0.0</td>
<td>60.1</td>
</tr>
<tr>
<td>75 °C</td>
<td>0.0</td>
<td>36.8</td>
</tr>
<tr>
<td>100 °C</td>
<td>0.0</td>
<td>23.4</td>
</tr>
</tbody>
</table>

Table S8. Summed clay wt.% at high (10,000), intermediate (1,000) and low (100) W/R ratios in models for the VRR mean composition (Table S4), at both 10 and 50% Fe\(^{3+}/Fe_{tot}\), reacted with GPW (Bridges et al., 2015b) at 25, 50, 75 and 100 °C using CHIM-XPT (Reed et al., 2010).

<table>
<thead>
<tr>
<th></th>
<th>10% Fe(^{3+}/Fe_{tot})</th>
<th>50% Fe(^{3+}/Fe_{tot})</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>10,000 W/R</td>
<td>1,000 W/R</td>
</tr>
<tr>
<td>25 °C</td>
<td>77.8</td>
<td>84.5</td>
</tr>
<tr>
<td>50 °C</td>
<td>34.4</td>
<td>84.3</td>
</tr>
<tr>
<td>75 °C</td>
<td>42.9</td>
<td>65.8</td>
</tr>
<tr>
<td>100 °C</td>
<td>42.9</td>
<td>54.9</td>
</tr>
</tbody>
</table>

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**Table S9.** Summed chlorite wt.% at high (10,000), intermediate (1,000) and low (100) W/R ratios in models for the VRR mean composition (Table S4), at both 10 and 50% Fe\textsuperscript{3+}/Fe\textsubscript{tot}, reacted with GPW (Bridges et al., 2015b) at 25, 50, 75 and 100 °C using CHIM-XPT (Reed et al., 2010).

<table>
<thead>
<tr>
<th></th>
<th>10% Fe\textsuperscript{3+}/Fe\textsubscript{tot}</th>
<th>50% Fe\textsuperscript{3+}/Fe\textsubscript{tot}</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>10,000 W/R</td>
<td>1,000 W/R</td>
</tr>
<tr>
<td>25 °C</td>
<td>15.1</td>
<td>11.9</td>
</tr>
<tr>
<td>50 °C</td>
<td>34.4</td>
<td>12.9</td>
</tr>
<tr>
<td>75 °C</td>
<td>42.9</td>
<td>19.7</td>
</tr>
<tr>
<td>100 °C</td>
<td>42.9</td>
<td>23.8</td>
</tr>
</tbody>
</table>

**Table S10.** Summed talc wt.% at high (10,000), intermediate (1,000) and low (100) W/R ratios in models for the VRR mean composition (Table S4), at both 10 and 50% Fe\textsuperscript{3+}/Fe\textsubscript{tot}, reacted with GPW (Bridges et al., 2015b) at 25, 50, 75 and 100 °C using CHIM-XPT (Reed et al., 2010).

<table>
<thead>
<tr>
<th></th>
<th>10% Fe\textsuperscript{3+}/Fe\textsubscript{tot}</th>
<th>50% Fe\textsuperscript{3+}/Fe\textsubscript{tot}</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>10,000 W/R</td>
<td>1,000 W/R</td>
</tr>
<tr>
<td>25 °C</td>
<td>7.4</td>
<td>10.5</td>
</tr>
<tr>
<td>50 °C</td>
<td>0.0</td>
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<tr>
<td>75 °C</td>
<td>0.0</td>
<td>9.3</td>
</tr>
<tr>
<td>100 °C</td>
<td>0.0</td>
<td>7.7</td>
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</tbody>
</table>
Figure S1. Modeled fluid composition at high (10000) W/R for all three alteration derived alteration compositions (Table S4) for all modeled temperatures and Fe\textsuperscript{3+}/Fe\textsubscript{tot}.

Supplementary Material References


