THERMOCHEMICAL MODELLING OF FLUID-ROCK REACTIONS IN VERA RUBIN RIDGE, GALE CRATER, MARS.


Introduction: Vera Rubin Ridge (VRR) in Gale Crater, Mars, is a ~200 m wide ~6.5 km long northeast-southwest resistant geomorphological feature on the northern slopes of Aeolis Mons (Mt. Sharp). Analysis of Compact Reconnaissance Imaging Spectrometer for Mars (CRISM) orbital data showed that VRR has strong hematite spectral signatures [1]. Hematite was confirmed in-situ at VRR with the Curiosity rover [2],[3] and has been shown to be present throughout the Murray formation [4–7]. VRR is stratigraphically continuous with the underlying Murray formation [8,9].

Previous thermochemical modelling showed how hematite at VRR could have formed as the result of open-system weathering at high water/rock ratios [10]. Here we use thermochemical modelling to investigate possible reaction pathways for the hematite-clay-bearing assemblage observed at VRR, starting from an identified least-altered (minimum clay content) Murray composition, and a Mars basal brine [11,12].

Modelling method: We used CHIM-XPT software to conduct thermochemical modelling [13]. CHIM-XPT is a program for computing multicomponent heterogeneous chemical equilibria in aqueous-mineral-gas systems, and has been previously used to model fluid-rock reactions at Gale crater [10–12]. Every calculation step calculates equilibrium between the fluid and the dissolved rock, meaning that each step can be treated and interpreted independently from the direction from which it was reached. In CHIM-XPT, the water/rock ratio (W/R) is the ratio of incoming fluid to reacted rock. The models here focus on 1–100,000 W/R.

The Starting Fluid. Gale Portage Water (GPW) [11] was selected as the starting fluid, as this fluid was derived from equilibrium mediation between a brine and rocks of the Gale area. The solution is initially oxidizing (all S species as SO\(_4^{2-}\)), and the redox in the fluid is controlled by the SO\(_4^{2-}/HS^-\) pair. The redox of the system throughout each model is dependent on the Fe\(^{2+}/Fe^{3+}\) ratio of the host rock. pH was modelled as a free-parameter throughout.

The Starting Rock. The initial starting composition was selected from Murray drill sample compositions that are mineralogically least altered, based on clay content (assuming the clays are authigenic). The Telegraph Peak (TP) drill sample was found to be most statistically representative of a bulk lower Murray composition based on a comparison to the average APXS compositions of the Confidence Hills, Mojave 2 and TP CheMin samples [7]. Furthermore, ChemCam bulk contour analysis [14] showed TP to be nearest the overall mean Murray composition. Based on these analyses, TP compositions [7] were selected for the initial starting compositions in our modelling.

Results: Initial modelling focused on establishing a ‘parameter space’ where the bulk, crystalline and amorphous components of TP [7] were reacted with GPW [11]. Modelling progressed to investigate different dissolution scenarios. Guided by relative mineral solubility [15], we investigated dissolving TP amorphous and olivine [7]. We then progressed to mixtures of olivine, TP amorphous and TP crystalline [7] to adjust for the lack of Al in the resulting mineral phases compared to VRR drill samples [2,3].

Fig. 1: CHIM-XPT thermochemical model for the reaction of the amorphous component of TP [7] and GPW [11]. The model was run at 50 °C and 0.1 Fe\(^{1+}/Fe_{total}\).

Initial ‘Parameter Space’ Modelling. This focused on modelling the individual bulk, crystalline and amorphous components of TP [7] with GPW [11] over a range of temperatures (20 °C, 50 °C, and 100 °C). The total Fe in the starting composition was expressed in various amounts of Fe\(^2+\) and Fe\(^3+\) (100% Fe\(^2+\), 90% Fe\(^2+\) & 10% Fe\(^3+\), 50% Fe\(^2+\) & 50% Fe\(^3+\)).

The bulk composition of TP failed to produce any Fe-oxide at 20 °C and 50 °C, but did produce hematite at 100 °C for >100,000 W/R.

The crystalline composition of TP only produced Fe-oxide at 0.5 Fe\(^{1+}/Fe_{total}\), with goethite precipitating at 20 °C (<10 wt.% for ~5,000–100,000 W/R) and...
hematite precipitating at 50 °C (<10 wt.% −8,000–100,000 W/R) and 100 °C (<10 wt.% 1,000–100,000 W/R with trace amounts down to ~50 W/R).

The amorphous composition of TP produced significant amounts of Fe-oxide. At 20 °C, >10 wt.% goethite precipitated between 12,000–100,000 W/R. At 50 °C, >10 wt.% and up to ~30 wt.% hematite precipitates between 5,000–100,000 W/R (Fig. 1). At 100 °C >30 wt.% hematite precipitates between 10,000–100,000 W/R for all modelled Fe²⁺/Fe³⁺ ratios.

Our thermochemical modelling assumes that the presently observed mineralogy is what was precipitated. However, it is possible that goethite initially precipitated and underwent phase transition to hematite over time [18]. Furthermore, the presence of talc in Figs. 1–3 is not observed in drill-hole compositions to date, and presence of jarosite and akaganeite in Steer and Rock Hall [2] is not currently explained by our models. Further work will be undertaken to refine the predicted mineral assemblages and to include reaction pathways that lead to sulfate and oxide precipitation.

**Summary:** Thermochemical modelling of the reaction between TP compositions [7] and GPW [11] has shown that hematite is readily produced at >100 W/R and 50 °C, for all modelled Fe²⁺/Fe³⁺ ratios. This suggests that the hematite at VRR could have been a conduit for groundwater flow at temperatures ≥50 °C.