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sediment source regions with time. We conclude that Gale sediment provenances were much more varied than suggested by the largely homogeneous, globally-distributed Martian basalt inferred from orbit, showing that complex magmatic assemblages exist within the ancient highland crust surrounding Gale.

1. INTRODUCTION

Mars’ global sedimentary record has been generated by fluvial, lacustrine, glacial, and aeolian processes (e.g., Malin and Edgett, 2000; Hynek and Phillips, 2001; 2003; Stack et al., 2014a). Throughout geological time, these processes have eroded sediments from their source rocks in the Martian crust, and redeposited them elsewhere as sedimentary units. Understanding the origin of these sediments can potentially provide information on Mars’ crustal compositions, and the igneous processes that generated them (e.g., McSween et al., 2009; Mangold et al., 2017b). Orbital mappimg with Mars Global Surveyor and Mars Odyssey data has shown that the mineralogy of Mars’ surface is dominated by primary mafic igneous minerals such as olivine and pyroxene (Bandfield et al., 2000; Hamilton and Christensen, 2005; Chevrier and Mathé, 2007; Hanna et al., 2016). This predominance of primary mafic minerals on the planet’s surface suggests that a relatively low rate of chemical weathering has been present throughout Mars’ geological history, and indicates that Mars’ crust is chiefly basaltic in composition (McSween et al., 2009). However, some occurrences of silica-rich and alkali-rich material have been observed in Martian samples such as in Martian meteorites NWA 6963 (Filiberto et al., 2014), NWA 7034 (Agee et al., 2013) and NWA 7533 (Humayun et al., 2013). Orbital data have also detected silica-rich material on the NW flank of Syrtis Major (Bandfield et al., 2004), though it is debated whether this relates to evolved crustal material (Bandfield et al., 2004; Bandfield, 2006) or late-stage aqueous alteration (Smith and Bandfield, 2012). Silica-rich and alkali-rich igneous float and sediments have been identified at the surface in Gale crater (Sautter et al., 2014; 2015; Morris et al., 2016; Edwards et al., 2017) and show that evolved magmatic compositions could be more widespread on Mars than that which is indicated by Near Infrared spectroscopy from orbit. Several factors have impeded the remote detection of these evolved compositions on the surface, for instance; hydrothermal alteration of the Martian crust (Bibring et al., 2006; Ehlimann et al., 2013), global dust which obscures much of the planet (McSween et al., 2009), and widespread, mixed Martian basaltic soil identified from ground-based observations (Chevrier and Mathé, 2007). Hence, identifying evolved crustal compositions requires landed missions such as the NASA Pathfinder, Mars Exploration Rovers (MER), and Mars Science Laboratory (MSL).

Since landing in August 2012, MSL has been geochemically and mineralogically analysing the sedimentary succession within Gale crater along its >19.6 km drive to date (Fig. 1). The Early Hesperian-age Gale crater was selected as the MSL landing site due to its low elevation as a closed basin, and the ~5 km high, stratified sedimentary mound Aeolis Mons, informally named “Mt Sharp”, which was seen from orbit to have hydrated clay- and sulfate-bearing units (Milliken et al., 2010; Fraeman et al., 2016) on its lower slopes. These hydrated units made Gale a good candidate site to better understand the role water played in Mars’ early history, and to determine whether Gale crater was habitable at some point in its past (Grotzinger et al., 2014). A key MSL mission objective is to analyse the geology and geochemistry of this preserved geological record in order to assess paleo-environmental change in relation to the depositional environments of the sedimentary units (Grotzinger et al., 2012). The paleo-environment of deposition is not the only contributor to sediment geochemistry; source region composition and the weathering conditions of the sediment host rock are also key components (Weltje, 2012). In addition to its primary mission aim, the MSL’s campaign therefore presents a unique opportunity to investigate Martian source region characteristics recorded in the sediments using the extensive geochemical dataset acquired by its instruments with, in particular for our study, ChemCam (Laser-Induced Breakdown Spectroscopy) and CheMin (X-ray Diffraction). The ChemCam instrument aboard MSL analysed ~1200 sedimentary targets within the period of the study (up to sol 1482), generating the most in-depth Martian chemostratigraphic record to date on another planet.

Terrestrial provenance studies have shown that the mineralogy and geochemistry of fine-grained clastic sedimentary units are largely characteristic of their source region, provided the extent of aqueous alteration, transport fractionation, and the effects of grain size variation have been taken into account (Nesbitt and Young, 1982; Roser and Korsch, 1986; Herron, 1988; Nesbitt and Wilson, 1992; Kroonenberg, 1994; Jin et al., 2006; Tolosana-Delgado and von Eynatten, 2009; Bloemsma et al., 2012; Weltje, 2012). Many terrestrial studies utilised ratios of major element oxides such as SiO2/Al2O3 and Na2O/K2O (e.g., Pettijohn et al., 1972; Roser and Korsch, 1986; Jin et al., 2006) to classify sediments according to magmatic source regions (Roser and Korsch, 1986; Herron, 1988; Kroonenberg, 1994), or determine the extent of chemical weathering they have undergone (Nesbitt and Young, 1982; Nesbitt and Wilson, 1992; Babechuk et al., 2014). If the mineralogical differences are acknowledged between the Earth and Mars (i.e., quartz is abundant in Earth’s sedimentary record and not in that of Mars), these terrestrial methods for determining sediment source region variations have the potential to be applied to ChemCam instrument major element compositions of sedimentary targets. Therefore, in this study we have grouped ChemCam major element analyses according to stratigraphic position, grain
size, and target textural characteristics, to distinguish the extent of post-depositional aqueous alteration on our samples, and delineate the effects of grain size on geochemistry. Then, we have utilised the mineralogical data provided by the Chemistry and Mineralogy (CheMin) instrument alongside the ChemCam dataset to gain some understanding of the compositional diversity of the Martian crust in the source regions around Gale.

1.1. Gale crater’s sedimentary record

The Gale crater succession encountered by MSL consists of mudstone-, fine to coarse sandstone-, and conglomerate-rich units. These units were interpreted to have been deposited in a dominantly fluvial-lacustrine paleoenvironment (Grotzinger et al., 2014; Grotzinger et al., 2015) following the formation of the crater ~3.8 ± 0.3 Ga (Thomson et al., 2011). Stratigraphic analysis showed that sediments infilled the crater and were subsequently exhumed and eroded by aeolian processes, leaving the present day ~5 km high, stratified Mt. Sharp (Grotzinger et al., 2015). Stratigraphic subdivisions of Gale crater along the rover traverse (Figs. 2, 3) are:

- The Bradbury Group, consisting of mostly fluvially-deposited conglomerate and sandstone, associated with deposition during delta progradation and sampled by the rover during sols 1–750 (Grotzinger et al., 2015).
The Mt. Sharp Group (analysed from sol 755; Fig. 1), largely composed of finely-laminated mudstone (Murray Formation), deposited within the standing water body located at the end of Gale’s ancient fluviolacustrine system (Grotzinger et al., 2014, 2015; Stack et al., 2016).

The Siccar Point Group (which constitutes the Stimson Formation, and was analysed by ChemCam during sols 990–1352), a lithified aeolian capping unit that unconformably overlies the Mt. Sharp Group (Fraeman et al., 2016; Banham et al., 2018).

In this study, we have solely analysed the data collected from the Bradbury and Mt Sharp stratigraphic groups in order to focus our geochemical interpretation on sediments deposited in the fluviolacustrine setting, as well as to examine the composition of the most ancient sedimentary units so far analysed by MSL within Gale crater.

This study aims to expand upon the key areas of geochemical interest within the Bradbury (Yellowknife Bay formation, Darwin waypoint, Kimberley formation) and Mt Sharp Groups (Pahrump Hills, Marias Pass, and Murray Buttes localities) that have provided a basis for previous studies into sediment source regions (McLennan et al., 2014; Vaniman et al., 2014; Anderson et al., 2015; Sautter et al., 2015; Mangold et al., 2015; Le Deit et al., 2016; Mangold et al., 2016; Morris et al., 2016; Thompson et al., 2016; Treiman et al., 2016; Edwards et al., 2017; Hurowitz et al., 2017; Mangold et al., 2017b; Rampe et al., 2017a; Siebach et al., 2017).

These studies show the sediments to be largely basaltic in nature, though alkali- and silica-enrichments have also been identified in Gale’s sediments, possibly relating to distinct, evolved magmatic source regions (Le Deit et al., 2016; Morris et al., 2016; Thompson et al., 2016; Treiman et al., 2016; Siebach et al., 2017). MSL has also identified evidence for fractionated crustal compositions in over 60 ChemCam targets of float and conglomerate clasts that show clear textural evidence – porphyritic, cumulate or vesicular – for an igneous origin (Sautter et al., 2014; Mangold et al., 2016; Cousin et al., 2017; Edwards et al., 2017). Their compositions ranged from basalt (most abundant) to trachybasalts (Edwards et al., 2017).

1.1.1. The Bradbury Group

Three key waypoints were analysed by MSL within the Bradbury Group: Yellowknife Bay, Darwin and Kimberley. Yellowknife Bay (sols 57–324) consists of a 4–6 m thick lacustrine mudstone (Sheepbed Member) overlain by cross-bedded, fluvial sandstone and conglomerate (Gillespie Lake and Glenelg Members; Figs. 2 and 3B). Widespread diagenetic features – which will be discussed in more detail in Section 1.2 – were identified in the sedimentary succession at this waypoint, such as raised ridges (Leveille et al., 2014; Fig. 4A), filled and hollow nodules (Stack et al., 2014b; Fig. 4B), and sulfate veins (Rapin et al., 2016; Fig. 4C). Two samples (John Klein and Cumberland) were drilled from the Sheepbed mudstone and delivered to ChemCam for mineralogical analysis (Vaniman et al., 2014).

After Yellowknife Bay, MSL arrived at the Darwin outcrop (sols 392–401) which is comprised of multiple exposures of conglomerate and sandstone bedrock, overlain by sand drifts and dark float accumulations (Vasavada et al., 2014; Mangold et al., 2016). Exposures of these conglomerates were also identified on the traverse leading up to and beyond the Kimberley formation described below (Mangold et al., 2016). Several clasts of igneous rock were identified at this locality within conglomerate units, or as igneous float (Mangold et al., 2016). These have been used as a comparison dataset in our study.

The final key waypoint investigated by MSL in the Bradbury Group is the Kimberley formation (sols 574–632). The Kimberley formation is the stratigraphically youngest key waypoint of the Bradbury Group (Grotzinger et al., 2015), and constitutes a fining upwards
sequence of polymict conglomerate to fine to medium-grained sandstone, capped by a massive coarse sandstone member (Thompson et al., 2016; Treiman et al., 2016). This formation was subdivided into the Point Coulomb and Liga conglomerate, Square Top fluvial sandstone, Dillinger trough cross-bedded sandstone, and Mt. Remarkable massive sandstone Members. These Members were analysed by ChemCam and the Alpha Particle X-Ray Spectrometer (APXS) instrument (Grotzinger et al., 2015; Le Deit et al., 2016; Thompson et al., 2016), and the Windjana sample was drilled from the Dillinger Member and delivered to CheMin (Treiman et al., 2016).

1.1.2. The Mt Sharp Group (Murray formation)

The lower Murray formation (lowermost unit of the Mt. Sharp Group from the Pahrump Hills to Marias Pass locality; Figs. 1–3A) is a laminated, fine-grained mudstone that appears stratigraphically ~20 m above the Kimberley formation (Fig. 2; Grotzinger et al., 2015). Drilled samples from the lower Murray formation Pahrump Hills include Confidence Hills, Mojave 2, and Telegraph Peak (Rampe et al., 2017a), along with Buckskin from Marias Pass (Morris et al., 2016). The change from the Bradbury Group to the Mt. Sharp Group was marked by a transition from fluvial sandstone and conglomerate, detected at the base (e.g., Bald Mountain, sols 775–778), to thinly laminated mudstone with extensive diagenetic features such as erosion resistant nodules, raised ridges, and dark and light mineral veins (Nachon et al., 2017). Cross-laminated mudstone/siltstone and fine-grained sandstone facies are also present in this unit, but overall grain size varies much less than in the Bradbury Group (Grotzinger et al., 2015; Hurowitz et al., 2017; Rampe et al., 2017a; Siebach et al., 2017).

From orbit, the Murray Formation appears homogeneous, at least up to the Vera Rubin (Hematite) Ridge, and has intermittent areas with notable infrared signatures from orbit of phyllosilicate and sulfate (Fraeman et al., 2013, 2016). From rover observations, a distinction can be made within the Murray formation between Murray units containing abundant hematite, and Murray units with high abundances of magnetite and crystalline silica (Morris et al., 2016; Hurowitz et al., 2017; Rampe et al., 2017a). The apparent interfingering boundary between Bradbury Group sediments and the Murray formation implies coeval deposition as the rivers transporting material from the crater rim reached a perennial lake (Grotzinger et al., 2015). Moving up the Mt. Sharp stratigraphic succession towards the Murray Buttes, the Murray formation was seen to be punctuated by a section of trough cross-bedded, fine-grained sandstone and siltstone, followed by a return to well-
laminated mudstone, with some interstratified sandstone. CheMin samples were drilled from the Murray formation encountered after the Naukluft plateau (sols 1373–1482) are Oudam and Marimba 2 (Bristow et al., 2017; Rampe et al., 2017).  

1.2. Aqueous alteration within Gale crater sediments

The variety of alteration features seen within the Gale crater stratigraphy – especially in the mudstone members – suggests that aqueous alteration occurred after deposition and lithification of the sediments (Leveille et al., 2014; Nachon et al., 2014; Bridges et al., 2015; Rapin et al., 2016; Schwenzer et al., 2016; Frydenvang et al., 2017; Nachon et al., 2017). Some features were identified throughout the stratigraphic record, such as calcium-sulfate mineral veins (Nachon et al., 2014; Rapin et al., 2016; L’Haridon et al., 2018). Meanwhile others were seen in specific areas (Fig. 4) i.e., raised ridges in Yellowknife Bay (Leveille et al., 2014), Mg-rich dendritic concretions at Pahrump Hills (Kah et al., 2015), dark mineral veins at Garden City (Nachon et al., 2017; L’Haridon et al., 2018), and alteration halos between Marias Pass and Bridger Basin (Frydenvang et al., 2017; Yen et al., 2017).

ChemCam analyses taken across diagenetic features show that chemical remobilisation appears localised to weaknesses in the lithological units such as fractures (Frydenvang et al., 2017; Nachon et al., 2017; L’Haridon et al., 2018). These weaknesses were exploited by the groundwater causing the dissolution and precipitation of soluble minerals and mobile elements along them. Hence, the low permeability of the units appears to have limited the open system alteration in bulk rock compositions away from the structural weaknesses of the bedrock (McLennan et al., 2014; Anderson et al., 2015).

In addition to the obvious diagenetic features, so far all Gale crater CheMin analyses of host rock and soil targets have contained a substantial proportion (16–60 wt%) of an amorphous and phyllosilicate (AP) component (Bish et al., 2013; Blake et al., 2013; Dehouck et al., 2014; Vaniman et al., 2014; Treiman et al., 2016; Rampe et al., 2017a). A best estimate of the AP abundance and composition can be modelled using a full-pattern FULLPAT fitting method which compares known terrestrial natural and synthetic amorphous and clay samples to Mars CheMin analyses (Chipera and Bish, 2002, 2013). Then AP compositions are calculated through a mass balance calculation using the bulk sample geochemistry analysed by the APXS instrument combined with the modelled FULLPAT sample phase abundances (Morrison et al., 2018). The amorphous component in the lithified sedimentary units – whether basaltic or silicic – has largely been hypothesised to relate to diagenetic cement derived from the aqueous alteration of the volcanic minerals or glass that constitute Gale’s sedimentary units (Vaniman et al., 2014; Treiman et al., 2016; Rampe et al., 2017a). In general, the calculated chemical composition of the AP component has been associated with material initially derived from a basaltic source (Bish et al., 2013; Blake et al., 2013; Dehouck et al., 2014; Vaniman et al., 2014), except for the high-silica amorphous
component in the Buckskin drill hole that was instead associated with an evolved volcanic source and/or enhanced leaching (Morris et al., 2016; Frydenvang et al., 2017).

In this study we will first characterise the extent of the geochemical remobilisation associated with the observed aqueous alteration features and calculated AP components up to sol 1482. If bulk rock geochemical compositions are shown to be uncorrelated with Gale crater alteration trends, sediment source characteristics can be identified. As the AP component may relate to diagenesis, and as it constitutes a substantial proportion of Gale’s sedimentary targets, it is crucial to understand the geochemical effect of the amorphous component on ChemCam analyses. Thus, we have also compared the CheMin and APXS derived compositions of the AP component to the bulk host rock geochemistry. If the compositions are similar between the AP component and local host rock, the AP component can be identified as a product of closed system chemical weathering, and sediment source characteristics can be constrained.

2. METHODS

2.1. ChemCam instrument

The remote sensing Chemistry and Camera (ChemCam) instrument suite uses a Laser-Induced Breakdown Spectroscopy (LIBS) instrument to ablate a target host rock or soil in order to acquire its major, minor and trace element compositions (Maurice et al., 2012; Wiens et al., 2012; Clegg et al., 2017). A series of 14 mJ laser pulses is directed at a target between 2.2–7.0 m from the rover mast to induce a short-lived plasma on the incident sample (Maurice et al., 2016; Wiens et al., 2012; Maurice et al., 2016). The excited electrons of the atoms and ions within the sample then emit at diagnostic wavelengths as they decay to lower energy levels, which are subsequently detected by the telescopic imager and passed to three spectrometers in the 240–340 nm ultra-violet light (UV), 380–470 nm violet (VIO), and 470–900 nm visible and near infra-red (VNIR) spectral ranges (Wiens et al., 2012).

Each target is analysed by a raster pattern of observation points which in turn consist of 30–50 spectral analyses. The repeated firing of the laser at a specific point on the incident target removes dust in the first five laser pulses with further pulses generating a sample depth profile (Wiens et al., 2012). Observation point analyses are averages of the spectra conducted at a given raster point. Raster geometries occur in either a matrix (2 × 2, 3 × 3, 4 × 4, or 5 × 5), or a linear (5 × 1, 10 × 1, or 20 × 1) configuration. Point footprints range from 350–750 μm in size depending on distance from the rover (Wiens et al., 2012; Maurice et al., 2016). This small footprint enables the identification and analysis of small-scale features such as diagenetic mineral veins and clasts, which make it imperative that the context of each analysis is documented. As such, the ChemCam baseline analysis sequence collects Remote Micro Imager (RMI) images with a submillimetre resolution of ~90 μrad before and after the LIBS analysis (Maurice et al., 2012; Le Mouélic et al., 2015), which have been used in this study for target classification alongside the rover’s other on-board cameras Mast Camera (Mastcam) (Malin et al., 2017) and Mars Hand Lens Imager (MAHLI) (Edgett et al., 2012).

Quantitative analysis is achieved for each observation point by processing the LIBS spectra through several pre-processing steps to yield clean atomic emission spectra (Wiens et al., 2013). The spectra averaged from all the laser shots minus the first five (to remove the effects of surface dust) is then calibrated against a set of more than 400 standards, including on-board calibration targets (Vaniman et al., 2012; Blaney et al., 2014) in a process that combines sub-model Partial Least Squares regression (PLS) (Anderson et al., 2017) and an Independent Component Analysis (ICA) algorithm (Clegg et al., 2017). This yields the abundances of eight major elements (SiO₂, TiO₂, Al₂O₃, FeO_total, MgO, CaO, Na₂O, and K₂O) in the form of oxide wt%. Accuracy and precision for each element oxide are noted in Table A.1 of Appendix A. In our study, the totals were not normalized to 100 % to account for any non-determined contributions of S, P, F, Cl, and H which ChemCam does not routinely analyse.

Mean accuracies for the observation points presented here are given in Table A.1 of Appendix A, and are used for comparison with absolute abundance values, such as mineral compositions, or for comparisons with measurements made by other instruments. ChemCam instrument precision corresponds to the error presented with APXS results (Blaney et al., 2014). It is used to compare different ChemCam observations. The ChemCam LIBS precision is significantly less than the standard deviations of the mean compositions of the various units discussed here (Blaney et al., 2014). In addition to the geochemical data from the ChemCam instrument, we also use Gale crater mineralogical data derived by the Chemistry and Mineralogy instrument suite (CheMin). CheMin data was acquired from the NASA Planetary Data System (PDS) and peer-reviewed articles (Vaniman et al., 2014; Morris et al., 2016; Treiman et al., 2016; Rampe et al., 2017a; Morrison et al., 2018) to provide complementary mineralogical data to the ChemCam major element data.

2.2. Statistical approach to ChemCam data

This study encompasses sols 1–1482 of the MSL mission. Within this period the ChemCam instrument analysed almost 1200 targets, and close to 9500 observation points. Due to the small sampling footprint, 350–750 μm (Maurice et al., 2016), on targets of varying grain size and heterogeneity, target analyses for coarse-grained lithologies (>1 mm) were often not representative of whole-rock geochemistry. To counter this effect, we analysed the dataset in its entirety through compositional trends displayed on a series of scatter and density contour plots, in addition to simple and multiple linear regression models. The scatter plots highlight any compositional trends between two major element oxides. The density contour plots are essentially histograms in two dimensions which show the major element distribution as isolines (contours) of x and y. The two-dimensional histograms are smoothed according to the methods of Eilers and Goeman (2004). The smoothing factor used in this study is 20. After smoothing of the
two-dimensional histograms, the contours are generated, and are representative of the proportion of data point density within each smoothed ‘pixel’ of a 100 × 100 grid. Density contour plots show the compositional spread of the data population and can be used with histograms to highlight compositional foci and geochemical trends across stratigraphic units (Edwards et al., 2017). For our ChemCam dataset of the sedimentary rock record, the density contour plots have the added benefit of indicating bulk rock compositions as they are not as easily affected by outlying compositions from individual minerals or alteration features that could skew the mean (see Appendix A.1 for an example).

Major element oxide Simple Linear Regression (SLR) models (Neter et al., 1996) were determined for the Bradbury and Mt Sharp stratigraphic groups (see Appendix B) using Minitab v17. Statistically significant major element correlations for the dataset were calculated using Pearson’s correlation coefficients (r) for each plotted major element oxide, which were then compared to the p-value of their associated student’s t-test statistic. The t-test statistics and p-values were calculated for a significance level \( \alpha = 0.05 \) based on the null hypothesis that there is no correlation between the two major element variables \((H_0: \rho = 0)\). Appendix B shows tables of each major element oxide’s relative \( r \) and \( p \)-values for each of the stratigraphic groups.

As an increase in the proportion of \( \text{Al}_2\text{O}_3 \) to \( \text{CaO}, \text{Na}_2\text{O}, \text{K}_2\text{O}, \text{MgO}, \) and \( \text{FeO}_2 \) has been associated with greater sediment chemical weathering and aqueous alteration (Nesbitt and Young, 1982; Nesbitt and Wilson, 1992; McLennan et al., 2014; McLennan et al., 2015; Hurowitz et al., 2017); a Multiple Linear Regression (MLR) model (Neter et al., 1996) was constructed to show the variation in \( \text{Al}_2\text{O}_3 \) as a response to predictors \( \text{SiO}_2, \text{TiO}_2, \text{FeO}_2, \text{MgO}, \text{CaO}, \text{Na}_2\text{O}, \) and \( \text{K}_2\text{O} \). This can statistically distinguish which major elements contribute the most to aluminium’s variation, and help ascertain whether or not this is indeed related to chemical weathering regimes, or source region characteristics. The statistical significance of how the predictor variables contribute to the variation in \( \text{Al}_2\text{O}_3 \) is shown via the p-value calculated using the student’s t-test statistic. Similar to the SLR models outlined above, the model is built around the null hypothesis that there is no fit between the response and predictors \((H_0: \rho = 0)\). The results and residual plots are shown in Appendix B. Residual plots (probability and order) are also given in the appendix to illustrate the data statistics. When discussing the initial major element correlations within the dataset, Pearson’s correlation coefficients are used to show the strength of the major element correlation (assuming it is linear) and whether the correlation in question is positive or negative. \( R^2 \) values are provided in the SLR and MLR model outputs (Appendix B) and are also given to show the fit of the models to the dataset.

2.3. Removing the effects of alteration from the bulk rock dataset

ChemCam has observed a variety of geological targets (i.e., soils, mineral veins, sandstone, igneous float). As our study focuses on source region inferences from lithified sedimentary units, we grouped ChemCam observation points based on Mastcam, Navcam, MAHLI, and RMI images into the “constrained bulk rock”, “alteration”, “unconsolidated sediment”, and “float” datasets (Fig. A.2). The constrained bulk rock dataset consists of ChemCam observations that targeted sedimentary outcrop away from obvious aqueous alteration features. We further refined the constrained bulk rock dataset by removing observation points with total sum of oxides outside the range considered acceptable for the purpose of this study, 95–105%. This minimised and effectively removed any sediments associated with high quantities of S and H, volatiles which are not included in standard ChemCam analyses. Restricting the constrained bulk rock dataset to 95–105% total sum of oxides also helps isolate and remove geochemical trends relating to sulfate salts. In order to discuss how source region compositions varied relative to geological time, the constrained bulk rock dataset has been subdivided into the main stratigraphic groups (Bradbury and Mt Sharp), as well as the key waypoints discussed in Section 1.1 where MSL conducted detailed investigations into the surrounding bedrock (see Section 2.5 for more details).

In order to minimise the influence of post-depositional alteration on the constrained bulk rock geochemistry, any LIBS pit observed to overlap a diagenetic feature in RMI images was removed from the main dataset and grouped separately. The “alteration feature dataset” consists of LIBS points that have sampled the many aqueous alteration features identified in the Bradbury and Mt Sharp stratigraphic Groups that formed through the interaction of groundwater and rock (see Section 1.2 and Fig. 4). We carefully documented all those targets, which on the basis of their morphology were of secondary (diagenetic) origin, e.g., raised ridges, nodules, veins. As the alteration feature dataset investigates the geochemical trends of alteration (including concentration of S and H, among other elements not tracked quantitatively by ChemCam) on Gale crater host rock, we do not restrict alteration feature LIBS analyses to the 95–105% total range that we placed on the constrained bulk rock dataset. This alteration feature dataset was then compared to that of the constrained bulk rock dataset in order to assess the extent that these aqueous alteration processes may have affected our estimates of the pre-alteration bulk rock geochemistry.

Observation points taken at the Marias Pass locality may demonstrate silica enrichment associated with diagenesis, as well as detrital tridymite and cristobalite (Morris et al., 2016; Frydenvang et al., 2017). To minimise the influence of alteration in Marias Pass host rock targets we only included ChemCam observation points taken at the Lion Outcrop where the Buckskin drill hole was conducted and where tridymite was detected (Morris et al., 2016). Tridymite only forms in high-temperature, low-pressure conditions, and so was assumed to relate to an igneous protolith and not alteration, although subsequent alteration could have led to the formation of opaline silica detected in the Buckskin target (Morris et al., 2016). As diagenetic halos and associated silica-enrichment were usually accompanied by a strong
light-toned colour change in the bedrock (Frydenvang et al., 2017), light-toned targets in this area were assumed to mark altered bedrock and have been excluded.

Observation points identified in RMI images to show soil contamination were removed along with targets disturbed by the rover (e.g., drill cuttings and dump piles formed as a product of drilling and CheMin analysis). Non-igneous float and pebbles were identified based on the presence of sedimentary features (such as laminations and cross-bedding) or a lack of convincing igneous textures (i.e., porphyritic, vitreous, vesicular) and were excluded from the results presented here. Finally, we also removed all igneous clasts and float from the refined dataset and placed them into a comparison dataset.

2.4. Categorising the constrained bulk rock dataset according to grain size

Terrestrial sediment compositions have been shown to strongly correlate with grain size due to the processes that generate the sediments from crystalline rocks (i.e., chemical and physical weathering) (Pettijohn, 1954; Bloemsma et al., 2012; Weltje, 2012; Siebach et al., 2017). Therefore, in order to separate sedimentary processes from source rock endmember compositions, it is necessary to group and discuss samples according to grain size dependent subpopulations (Weltje, 2012). Average grain sizes of the analysed geological units and key waypoints were obtained from literature studies which used MAHLI, RMI and Mastcam images (Meslin et al., 2013; Grotzinger et al., 2014; Anderson et al., 2015; Grotzinger et al., 2015; Mangold et al., 2016; Treiman et al., 2016; Siebach et al., 2017; Mangold et al., 2017b). Average grain sizes are used as grain size determination is dependent on a well-focused image taken of a relatively dust free target which is not always possible (Mangold et al., 2017b).

Coarse-grained units were defined as those with resolvable grain sizes greater than 1 mm (Mangold et al., 2016; Mangold et al., 2017b) and were usually seen in the Bradbury Group to typically consist of clast-supported conglomerates. These were poorly sorted, with sub-rounded to sub-angular pebbles. Sand grains (defined as 0.062–1 mm) can be resolved with RMI and MAHLI images but are only identifiable using Mastcam if larger than 0.15 mm (Mangold et al., 2017b). The minimum grain size that MSL cameras can identify requires a resolution of at least 3 pixels (Mangold et al., 2017b). Therefore, at the optimum MAHLI working distance (~2 cm), grains finer than very fine sand (<0.062 mm) are largely unresolvable and constitute the mudstone grain size category (Mangold et al., 2017b). Average grain sizes for each stratigraphic group and unit are provided in Fig. A.2. of Appendix A.

2.5. Categorising the constrained bulk rock dataset according to stratigraphic position

The constrained ChemCam bulk rock database was divided into stratigraphic subgroups to help ascertain any change in endmember source compositions. The stratigraphic group divisions (Bradbury and Mt. Sharp up to sol 1482) were used to study the geochemical variation between distinct paleoenvironmental settings. A more detailed study into the compositional variation within stratigraphic groups was then provided with the analysis of stratigraphic units and waypoints, such as those outlined in Section 1.1. Fig. A.2. of Appendix illustrates each of the subgroup divisions used in our study and Table A.2 provides the total number of observation points included for each stratigraphic unit and area.

3. RESULTS

In this section, we discuss the geochemistry of the constrained bulk rock dataset (Section 3.1) followed by the geochemical trends identified in the alteration dataset (Section 3.2). Overall, the Bradbury and Mt. Sharp (Murray formation to sol 1482) stratigraphic groups exhibit distinct geochemical foci and trends across the major element oxide compositions of the ChemCam observation points (Figs. 5, 6). The constrained bulk rock dataset – subdivided according to stratigraphic group – are plotted in Fig. 5 as frequency histograms and as a series of Harker density contour plots in Figs. 6–8. Basic statistics for the constrained Bradbury and Mt Sharp bulk rock datasets are provided in Table 1. Alteration targets (ridges, veins, etc.), calculated amorphous and phyllosilicate (AP) components, and CheMin modelled primary igneous mineral phases are plotted alongside the constrained stratigraphic group density contours in Figs. 7 and 8. The CheMin AP compositions are shown in Figs. 7 and 8 to help discern the influence these components have on the modelled constrained bulk rock geochemistry. Compositions of the AP components for various drilled samples were calculated by Vaniman et al. (2014), Treiman et al. (2016), Morris et al. (2016), Rampe et al. (2017a), and Morrison et al. (2018) based on CheMin abundances and APXS-derived compositions (see Section 1.2 for details on how the AP compositions are derived).

3.1. Major element variation and correlation within the stratigraphic groups

The Bradbury Group demonstrates a clear bimodality in its histograms for MgO, Na2O, and Al2O3 (Fig. 5). Peak MgO compositions occur at 0.8–1.3 and 7.1–7.6 wt%; Na2O peaks occur at 2.9–3.1 and 5.2–5.4 wt%; and Al2O3 peaks occur at 10.4–11.0 and 17.8–18.3 wt%. The Bradbury Group shows a unimodal distribution in the histograms for SiO2, TiO2, FeOtot, CaO, and K2O (Fig. 5). These unimodal distributions are generally asymmetric, with the exception of CaO that is symmetrical (Fig. 5). This asymmetry is particularly apparent for K2O which is strongly left skewed, SiO2 and TiO2 that are weakly left skewed, and FeO which is weakly right skewed, suggesting an influence by extreme outliers on the mean compositions (Figs. 5, 6). Density contours of the Bradbury Group are generally present as concentric circles about the mean (Fig. 6), but with a slight deviation for the outermost edges towards higher Al2O3, FeOtot, Na2O, and K2O, and lower FeOtot, MgO, and Na2O concentrations (Fig. 6). These deviations roughly
correspond to the pronounced tails in the histograms or, in the cases of Al₂O₃ and MgO, to the secondary histogram peaks (Fig. 5).

The Mt. Sharp Group (Murray formation, sols 755–1482) has contours centred on a single focus for FeO_T, K₂O, Na₂O, and TiO₂ when plotted against SiO₂ (Fig. 6), though slight bimodality is shown in the FeO_T histogram with a small peak at 5.6–6.6 wt%, and a large peak at 18.9–19.9 wt%. SiO₂ and Al₂O₃ density contours and histograms show slight bimodality (peaks at 53.1–54.1 and 67.7–68.7 wt% SiO₂, 11.5–12.0 and 15.7–16.2 wt% Al₂O₃), as does MgO versus SiO₂ to a lesser extent. The low MgO subpopulation at high SiO₂ in the Harker plot contours deviate from the respective modal compositions (Fig. 6D) and correlate with the minor SiO₂ peak of ~70 wt% shown in Fig. 5A.

Calcium oxide contents are notably lower in the Mt Sharp Group with mean, peak and focal compositions all <3.4 wt% (Figs. 5 & 6 and Table 1). Mt Sharp Group total FeO and Na₂O focal compositions are similar to those of the Bradbury Group (Figs. 5, 6), but with less variability between these oxides and SiO₂, indicated by the contours centred more uniformly on the foci. The Mt. Sharp Group has a higher mean SiO₂ content (54.4 wt% vs. 49.0 wt% for the Bradbury Group, Table 1), which is reflected in its density contours with average focal compositions centred around 53.1 ± 1.9 wt% SiO₂ (Fig. 6 and Table 3). Though not identified in mean compositions, the density contour plots also show that Mt Sharp Group focal compositions are more elevated in Al₂O₃ (12.1 ± 0.8 wt%) and K₂O (1.2 ± 0.2 wt%) than Bradbury (foci at 10.6 ± 1.2 wt% and 0.5 ± 0.3 wt% respectively). Overall, the Mt Sharp
Fig. 6. Harker density contour plots of the constrained ChemCam bulk rock dataset plotted according to related stratigraphic groups; Bradbury (light green) and Mt Sharp (dark green), with SiO$_2$ on the x-axis and major element oxide on the y-axis. The light blue cross represents ChemCam accuracy (RMSEP), the dark blue cross shows precision (1$\sigma$). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
Fig. 7. Harker density contour plots of Bradbury Group constrained bulk composition (light green density contours) and alteration trends, with SiO$_2$ on the x-axis and major element oxide on the y-axis. Each scatter point for the alteration features represents a ChemCam observation point composition for mineral vein, fracture-associated halo, nodule and raised ridge targets. The light blue cross shows ChemCam accuracy (RMSEP), the dark blue cross shows precision (standard deviation). Calculated CheMin amorphous and phyllosilicate compositions (Vaniman et al., 2014; Treiman et al., 2016) are plotted as squares with Gale primary igneous minerals refined from the CheMin unit-cell parameters (olivine, augite, plagioclase and sanidine) shown as stars. RR = Raised ridges, Am + Phyll = amorphous + phyllosilicate component, CaSO$_4$ = calcium-sulfate which may be in various hydration states, Silica = silica-enrichment. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
Fig. 8. Harker density contour plots of Mt Sharp Group constrained bulk composition (dark green density contours) and alteration trends, with SiO$_2$ on the x-axis and major element oxide on the y-axis. Each scatter point for the alteration features represents a ChemCam observation point composition for mineral vein, fracture-associated halo, nodule and raised ridge targets. The light blue cross shows ChemCam accuracy (RMSEP), the dark blue cross shows precision (standard deviation). Calculated CheMin amorphous and phyllosilicate compositions (Rampe et al., 2017a) are plotted as squares, Gale primary igneous minerals refined from the CheMin unit-cell parameters (olivine, augite, plagioclase and sanidine) are shown as stars. RR = Raised ridges, Am + Phyll = amorphous + phyllosilicate component, CaSO$_4$ = calcium-sulfate which may be in various hydration states, Silica = silica-enrichment. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
Group has a smaller, more uniform, range in major element compositions than the Bradbury Group (Table 1 and Table 3).

### 3.1.1. Constrained bulk rock Simple Linear Regression (SLR) models and correlation coefficients

We use SLR models and Pearson correlation coefficients to calculate element correlations for the constrained stratigraphic group and unit datasets in order to show which minerals dominate sediment geochemistry and whether or not alteration has affected these sediments on a scale not identifiable by the MSL imaging instruments. SLR models of the constrained bulk rock dataset for the Bradbury and Mt Sharp stratigraphic groups both show strong negative correlations between FeO and SiO₂ (r values of 0.85 and 0.49 respectively; see Figs. 6C & 7C). In these models, the null hypothesis tests for no correlation between the variables in question (H₀: $\rho = 0$). Therefore, the strong negative correlations between FeO and SiO₂ are statistically significant as their p-values are much lower than the significance level at $\alpha = 0.05$ (Fig. B.1, and Tables B.1 and B.2. of Appendix B). Strong positive correlations ($r > 0.5$) also exist in both groups for Al₂O₃ and Na₂O, with the Bradbury Group demonstrating the best fit ($r = 0.86$), and the Mt Sharp Group substantially lower ($r = 0.60$) due to scatter towards elevated Na₂O concentrations (see Fig. 7 and Appendix Fig. B.2).

Both the Bradbury and Mt Sharp Groups have positive correlations between MgO versus FeO, and negative correlations between MgO against Al₂O₃ and MgO versus Na₂O. When plotted, MgO displays a curvilinear trend for MgO against Al₂O₃ (Fig. B.3 in the Appendix). This curvilinear trend is fitted best to Bradbury Group data ($r^2 = 75.5\%$), and worst to the Mt Sharp Group ($r^2 = 0.6\%$), indicative of no fit in the latter case. The Mt Sharp Group SLR models with the weakest fit to the curvilinear regression lie towards the origin of Al₂O₃ and MgO. These data points with residuals that do not fit the regression line are targets in the Marias Pass, silica-rich area.

### 3.1.2. Constrained bulk rock Al₂O₃ Multiple Linear Regression (MLR) model results

Results from the stratigraphic group Al₂O₃ MLR models (Section 2.2) show that all predictors (SiO₂, TiO₂, FeO, MgO, CaO, Na₂O, and K₂O) significantly contribute to the Bradbury Group MLR model (p-value < $\alpha$), and all predictors except K₂O (p-value = 0.60) contribute significantly to the Mt Sharp Group MLR model. This suggests a weak association of K₂O with the other predictors in the Mt Sharp Group model. Removal of K₂O from the model does not significantly improve the fit, so we do not exclude K₂O as a predictor. Both models have high adjusted r² values with Bradbury r² = 87.1% and Mt Sharp r² = 67.3% which shows the goodness-of-fit of the MLR models to the dataset.

For the Bradbury Group, the modelled percentage contributions of Na₂O (74.5%), K₂O (4.7%), and MgO (4.6%) demonstrate that they contribute the most to Al₂O₃ variation while FeO (0.3%) and TiO₂ (0.04%) contribute the least in this regression model. All predictors have negative coefficients in relation to the response with the exception of Na₂O and K₂O that are positive. In the Mt Sharp Group, SiO₂ (22.2%), FeO (20.9%), and Na₂O (9.9%) contribute the most to the model, with K₂O (0.01%) and CaO (3.7%) providing the smallest contributions. For the Mt Sharp Group, predictor coefficients show that as Al₂O₃ increases, all major elements except Na₂O decrease.

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### Table 1

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<th>Variable (wt%)</th>
<th>Unit</th>
<th>Mean</th>
<th>StDev</th>
<th>Minimum</th>
<th>Median</th>
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3.2. Alteration feature major element variation and correlations

Figs. 7 and 8 highlight the major element trends associated with alteration features identified within each stratigraphic group, and isolated from the constrained bulk rock dataset (see Section 2.3). The stratigraphic group density contours, CheMin derived amorphous and phyllosilicate (AP) components, and common primary, detrital, silicate minerals (olivine, plagioclase, alkali feldspar, and augite) are also plotted to help distinguish alteration trends from detrital igneous mineral mixing trends. Alteration features were identified within each stratigraphic group, and are presented as scatter points over the bulk rock density contours.

3.2.1. Sulfate mineral veins and fracture-associated alteration halos

Sulfate mineral veins (Fig. 4C) exhibit high calcium enrichment (Figs. 7 and 8), low abundances of other oxides, and low totals. Meanwhile, diagenetic halos (Fig. 4D), ~50 cm in width (Yen et al., 2017), are associated with an increase in silica content of the host rock (Frydenvang et al., 2017; Yen et al., 2017) and share similar elemental compositions to the high silica targets analysed at the Buckskin drill hole on sol 1057 (Fig. 8; Morris et al., 2016). The halos mostly occurred in the Mount Sharp Group, particularly between Marias Pass and Bridger Basin (Frydenvang et al., 2017; Yen et al., 2017). However, one possible halo was observed in the Bradbury Group, with the target Kukri at Darwin, on Sol 392 (Williams et al., 2014).

Within fracture-associated alteration halos FeO T and Na2O possess the best-defined negative correlations against SiO2, giving strong Pearson’s correlation coefficients of −0.90 and −0.82 respectively. The only element to show a (weak) positive correlation with halo SiO2 enrichment is TiO2 (r = 0.29, t-test p-value < α). Al2O3 and Na2O show the strongest positive correlation (r = 0.92), and display the best-defined negative trends when plotted against SiO2 (Figs. 8, 9). MgO concentrations appear to be relatively uniform at ~3.0 wt%, except at SiO2 > 80 wt% where concentrations remain at ~1.0 wt%.

Stratigraphic group density contours in our constrained bulk dataset do not appear to have been influenced by either the calcium-sulfate mineral veins or the silica-rich halos (Figs. 7, 8). However, despite measures taken to remove the effects of diagenetic silica enrichment in the Marias Pass area, scatter plots of the Marias Pass observation point compositions follow a silica-rich alteration trend (e.g., Fig. C.1. in Appendix C). The Marias Pass observation points were not removed from the bulk dataset as they constitute a small proportion of data points and hence, have not affected the overall stratigraphic group density contour focal compositions.

3.2.2. Raised ridges

Erosion resistant raised ridge features (Fig. 4A) occur across both stratigraphic groups (Figs. 7 and 8). Bradbury raised ridges (BRR), are present in both the Sheepbed Member and Kimberley formation and show
3.2.3. Nodules

Bradbury Group nodules (Fig. 4B) were predominately identified in the Sheepbed Member mudstone within “nodule-rich” bedrock (Stack et al., 2014b). Sheepbed nodules are spheroidal, at the mm-scale, and can be divided into three groups (solid, hollow, or filled) depending on their internal structure (Stack et al., 2014b). Solid nodules do not show obvious internal structures, are similar in texture and colour to the host rock and contribute the majority of CheMin nodule analyses. Hollow nodules possess raised rims and internal depressions. Finally, filled nodules classify those that contain light-toned, sulfate minerals (Stack et al., 2014b). ChemCam targets from the Sheepbed nodule-rich bedrock show minimal deviation from bulk rock composition (Fig. 7). Similar to Bradbury Group nodules, the nodule features identified in the Mt Sharp Group are also mm-scale and of a similar colour to the surrounding bedrock, though these are less frequent and have more irregular shapes (Nachon et al., 2017). Unlike the Sheepbed nodules, Mt Sharp Group nodules do not show any obvious internal structure. Nodules of the Mt Sharp Group have similar geochemical trends to the calcium-sulfate mineral veins (major element depletion in SiO₂ and the alkalis, plus most points showing Al₂O₃ and FeO_T depletion with CaO enrichment).

4. DISCUSSION

4.1. Chemical alteration of Gale crater sediments

Chemical weathering on Mars has been relatively low compared to Earth for most of the planet’s history as evidenced by the high abundance of ancient unweathered basaltic material on the Martian surface (Hoeven et al., 2003; Hauslath et al., 2008; McSween et al., 2009), and the paucity of chemical weathering products in Amazonian-aged terrains (e.g., Ehlmann et al., 2013). Instead, mechanical and physical weathering processes have dominated sediment generation (Bandfield et al., 2000; Christensen et al., 2001; McLennan et al., 2014). The majority of our constrained bulk rock major element correlations point towards the importance of detrital primary minerals in Gale’s sedimentary record. The dominant negative correlation trend for both stratigraphic groups in our constrained bulk rock dataset is FeO_T against SiO₂. The Bradbury Group also has notably strong negative correlations between MgO against Al₂O₃ and MgO against Na₂O. The only significant positive correlation trend is Na₂O versus Al₂O₃. These correlations extend throughout the stratigraphic succession. They are expressed through the elongation of density contours in the direction of ChemMin-derived olivine, augite, and feldspar compositions (Figs. 7 and 8) and are therefore likely related to mafic versus felsic mineral abundances. The strong positive association of Al₂O₃ in the MLR models with Na₂O (and K₂O for Bradbury Group) and negative association with FeO_T and MgO are also indicative that felsic/mafic mineral proportions are the controlling factors on bulk stratigraphic unit geochemistry.

The minerals that appear to dominate compositional variation within Gale’s sedimentary units constitute on average ~40–70% in all CheMin analyses and are of primary igneous origin (for CheMin mineral compositions, see Appendix A). The other ~30–60% in CheMin analyses relate to the amorphous and phyllosilicate component (Fig. 9 and discussed in Section 4.1.2; for compositions see Appendix A). Secondary minerals suggest that chemical weathering is recorded to some extent in the ChemCam target analyses included in our constrained bulk rock dataset. The abundance of alteration features in Gale crater’s stratigraphy is also evidence of aqueous alteration, likely the result of diagenetic episodes. Distinguishing alteration trends relating to these diagenetic episodes and chemical weathering from source rock characteristics is essential before any conclusions on source region chemistry can be made.

4.1.1. Element mobility and feature-related weathering

We have used major element trends of observed diagenetic features to determine the extent of alteration that has occurred in our constrained bulk rock dataset (Figs. 7, 8). As described in Section 3.2.2, the late-stage diagenetic features (mineral veins and silica-rich halos) which cross-cut stratigraphic units show clear compositional trends away from all stratigraphic group compositions (Figs. 7, 8).

For the silica-rich halos, ChemCam analyses plot away from constrained contour compositions suggesting that element mobilisation was restricted to the associated fracture zones (Frydenvang et al., 2017; Yen et al., 2017). The one exception is the Marias Pass area of the Murray formation presented in Section 3.2.2 whose compositions follow a similar trend to the silica-rich halos. This could be related to the SiO₂ polymorph grains that are abundant in the
Buckskin drill hole (Morris et al., 2016), and which would plot at a similar position on the Harker diagrams to the fracture-associated halo alteration trends (Fig. 8). Alternatively, the Marias Pass Lion outcrop (sols 1053–1066) where the Buckskin drill hole was conducted and ChemCam observation points acquired, does possess the near white, light-toned, visual characteristic of the alteration halos. These outcrops are also located near light-toned, fracture-associated halos in the overlying Stimson formation (Frydenvang et al., 2017). Therefore, it is difficult to distinguish whether the Marias Pass target igneous mineralogy and source composition have been overprinted by secondary alteration processes of the fracture-associated halo alteration. As Marias Pass analyses represent a small set of outlying Mt Sharp Group compositions that plot far from the main Mt Sharp Group contours (Fig. 8), this silica-rich endmember and halo diagenesis has not influenced the density contour compositions for the constrained bulk Murray dataset. Hence, we have not recalculated the Marias Pass data in the Chemical Index of Alteration analysis of Section 4.1.3.

Many nodule compositions reflect bulk rock composition, particularly those of the Bradbury Group (Stack et al., 2014b), which could either indicate that these observation point analyses have proportionally sampled more host rock, or that these features have formed by closed system alteration preserving geochemical source characteristics. The nodules that show geochemical variation away from bulk composition, such as those at the base of the Mt Sharp Group, appear to relate to sulfate precipitation (Fig. 8 and Nachon et al., 2017). They follow similar major element trends as the mineral veins, but this time with magnesium- as well as calcium-sulfates dominating nately calcium-sulfate mineral veins (Nachon et al., 2014; Rapin et al., 2016; L’Haridon et al., 2018). They are thought to be dominated by bassanite in ChemCam surface analyses (Rapin et al., 2016) but some appear more hydrated below the surface as exposed by the drill (L’Haridon et al., 2018; Vaniman et al., 2018). The total sum of oxides also becomes lower as a result of not directly analysing sulfur in the Ca-sulfate veins (Rapin et al., 2016). Overall, the Bradbury Group and Mt Sharp Group density contours do not appear to have been influenced by this set of alteration features (Figs. 7, 8) indicating that the compositional influence relating to these diagenetic features was successfully removed from the constrained bulk rock dataset.

Scatter trends associated with the light-toned mineral vein features demonstrate a clear trend away from host rock compositions towards CaO enrichment (Figs. 7, 8) which supports the interpretation that these are predominantly calcium-sulfate mineral veins (Nachon et al., 2014; Rapin et al., 2016; L’Haridon et al., 2018). They are thought to be dominated by bassanite in ChemCam surface analyses (Rapin et al., 2016) but some appear more hydrated below the surface as exposed by the drill (L’Haridon et al., 2018; Vaniman et al., 2018). The total sum of oxides also becomes lower as a result of not directly analysing sulfur in the Ca-sulfate veins (Rapin et al., 2016). Overall, the Bradbury Group and Mt Sharp Group density contours do not appear to have been influenced by this set of alteration features (Figs. 7, 8) indicating that the compositional influence relating to these diagenetic features was successfully removed from the constrained bulk rock dataset.

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Many nodule compositions reflect bulk rock composition, particularly those of the Bradbury Group (Fig. 7, Stack et al., 2014b), which could either indicate that these observation point analyses have proportionally sampled more host rock, or that these features have formed by closed system alteration preserving geochemical source characteristics. The nodules that show geochemical variation away from bulk composition, such as those at the base of the Mt Sharp Group, appear to relate to sulfate precipitation (Fig. 8 and Nachon et al., 2017). They follow similar major element trends as the mineral veins, but this time with magnesium- as well as calcium-sulfates dominating.
the geochemical composition (Figs. 7 and 8; Nachon et al., 2017). In the Mt Sharp Group, a few nodule analyses show high FeO\textsubscript{T} compositions $>25.0$ wt% (Fig. 8C). We interpret these FeO\textsubscript{T}-rich nodule analyses as potentially showing iron-oxides, as they are low in concentration for the other major elements with respect to bulk Mt Sharp. This is in contrast to the high-FeO\textsubscript{T} composition of the ChemCam target Hayden Peak (sol 785) that Nachon et al. (2017) interprets to be Na-jarosite due to its high Na\textsubscript{2}O compared to other Pahrump diagenetic features. As the nodules of the Bradbury Group do not deviate significantly from the Bradbury density contours, it is likely that they resulted from closed system alteration at the scale of the stratigraphic units. However, the association of sulfates with nodules in the Mt Sharp Group indicates more open system conditions, similar to the sulfate veins, to contribute the excess CaO and MgO. With these nodules removed from the constrained bulk rock dataset, the geochemical effect of open system alteration on the Mt Sharp Group constrained dataset is minimised.

The geochemical compositions of raised ridges in Gale crater vary according to stratigraphic unit (Figs. 7 and 8). Bradbury Group raised ridges show enrichments in either MgO or K\textsubscript{2}O, Na\textsubscript{2}O, and Al\textsubscript{2}O\textsubscript{3}, while raised ridges identified in the Mt Sharp Group appear to be geochemically related to calcium-sulfates comparable to the Mt Sharp Group nodules (Fig. 8). Aqueous alteration models of raised ridges in the Bradbury Group Sheepbed mudstone show that their high MgO concentrations likely occurred as a result of olivine dissolution in a closed system with a low water-rock ratio (Leveille et al., 2014; McLennan et al., 2014; Bridges et al., 2015). The enrichment in K\textsubscript{2}O, Na\textsubscript{2}O, and Al\textsubscript{2}O\textsubscript{3} for the Kimberley formation raised ridges could relate to a similar process but with feldspar dissolution, particularly due to the high total mass abundance (26.8%) of felsic minerals in the Windjana drilled sample (Treiman et al., 2016; Morrison et al., 2018). As the raised ridge features have been removed from the constrained database, the local, post-depositional remobilisation of these elements is minimised.

4.1.2. Gale crater amorphous and phyllosilicate components

Due to the lithified nature of the ChemCam host rock targets, some diagenesis will have occurred on a scale smaller than that of MAHLI image resolution (13.9 μm at a standoff distance of 2 cm; Edgett et al., 2012) during the cementation of the mud and sandstone units. For this reason, we compare the geochemical compositions modelled from CheMin and APXS analyses for the amorphous and phyllosilicate (AP) components (Vaniman et al., 2014; Treiman et al., 2016; Rampé et al., 2017a) to our constrained bulk rock dataset. In Figs. 7 and 8, we represent the composition of the X-ray amorphous and phyllosilicate components as a single point for each sample, based on the bulk composition and the calculated composition of the crystalline phases identified by CheMin (e.g., Morrison et al., 2018). In actuality, there is a range of potential AP compositions because of the uncertainties in mineral and amorphous abundances and refined crystal chemistry of major minerals calculated from CheMin data (e.g., Dehouck et al., 2014). We plot the best estimate of the AP compositions, rather than the range of potential compositions, to simplify the diagrams.

All of the Bradbury and Mt Sharp Group’s 9 drilled samples in the region included here possess varying abundances of an amorphous component, up to 60 $\%$ in the Buckskin drill hole (Morris et al., 2016). The origin of this amorphous component in Gale crater has been hypothesised to either relate to detrital, and then subsequently altered soil or aeolian material (Dehouck et al., 2014; Vaniman et al., 2014), diagenetic cement (Treiman et al., 2016), or chemically altered and weathered volcanic glass (Bish et al., 2013; Rampe et al., 2017a). Excluding the Marias Pass area, the Mt Sharp Group has $\sim$10–15$\%$ greater amorphous component than the Bradbury Group Sheepbed mudstones, which additionally possess a $\sim$10$\%$ greater abundance of phyllosilicates than the Mt Sharp mudstones reported here (Murray formation; Fig. 9).

Most AP components in mudstone units (John Klein, Mojave 2, Telegraph Peak, Confidence Hills, Marimba, with the exception of Cumberland and Buckskin) are within ChemCam instrument error of their associated stratigraphic group constrained bulk rock geochemistry (Figs. 7, 8), with some slight variations relative to their Group focus. For the purposes of our study we note that AP in the mudstone units does not demonstrate consistent enrichments in Al\textsubscript{2}O\textsubscript{3} or SiO\textsubscript{2}, or a consistent loss of divalent and monovalent cations relative to either the constrained ChemCam calculated bulk compositions presented here or in the APXS bulk compositions (Rampe et al., 2017a). Hence, Gale crater mudstone AP component may largely reflect local variations in mineralogy of the stratigraphic units that have undergone relatively closed system diagenesis as calculated AP compositions do not show the expected trends for open system aqueous alteration.

MgO-enrichment in the Cumberland AP component of the Sheepbed mudstone relative to the Bradbury bulk rock composition has been attributed to olivine saponitisation (Dehouck et al., 2014; Vaniman et al., 2014). The mean phyllosilicate abundance in the Sheepbed Member is $\sim$20$\%$ (Vaniman et al., 2014). Different 001 spacing within CheMin for the John Klein and Cumberland smectites suggests that the elevated MgO component in Cumberland most likely relates to a Mg(OH)\textsubscript{2}-rich interlayer site in the smectite component that is absent in the collapsed smectite analysed at John Klein (Vaniman et al., 2014; Bristow et al., 2015). In addition, MgO enrichment in the Cumberland AP component follows the trends of enrichment similar to the Sheepbed raised ridges and nodules (Fig. 7; Leveille et al., 2014), supporting the conclusions of Vaniman et al. (2014) and Bristow et al. (2015) that the Cumberland drill hole has sampled one of these diagenetic features. Therefore, the greater abundance of Mg\textsuperscript{2+} in the Cumberland phyllosilicates is likely a consequence of the aqueous alteration that formed these diagenetic features. This MgO-rich AP component will have consequently been removed from the constrained bulk rock dataset with the exclusion of raised ridge features.

The other mudstone sample whose AP component demonstrated a distinct composition relative to the bulk...
Mt Sharp Group focal composition is the Marias Pass Buckskin drilled sample. The Buckskin sample contains no phyllosilicate (Morris et al., 2016; Rampe et al., 2017a) and possesses strong enrichment in SiO₂ and TiO₂, as well as a depletion in FeO₇, relative to the Mt Sharp Group density contours. These major element trends are similar to those of its host rock whose composition may either relate to the silica-rich volcanic source region that formed the tridymite (Morris et al., 2016), or the fracture-associated alteration halos. Hence, this mudstone amorphous component may have also been derived from a similar process (Section 4.1.1).

The Windjana drill hole AP composition of the Kimberley formation is distinct from the mudstone and bulk Bradbury geochemistry, likely as a result of its sandstone grain size (Treiman et al., 2016) resulting in greater porosity and enabling a higher permeability and pore water content. Windjana AP possesses strong enrichments in FeO₇ and MgO with depletions in SiO₂, CaO, and K₂O relative to the bulk Bradbury Group compositions (Fig. 7). The higher pore water content of the Windjana sandstone relative to the mudstone units may have inhibited pressure solution upon compaction, more so than in the finer grained units. This would have restricted extended element remobilisation within the unit similar to cementation of terrestrial sandstones (McKinley et al., 2011; Henares et al., 2014), making the compositions of the AP component more distinct compared to that of the bulk rock. High concentrations of FeO₇ have been observed in all of Gale crater’s sedimentary units (McLennan et al., 2014; McLennan et al., 2015; Le Deit et al., 2016; Schieber et al., 2017; Siebach et al., 2017; Hurowitz et al., 2017; Mangold et al., 2017a; Mangold et al., 2018). On Earth, where carbonates represent a significant fraction of the sedimentary budget, CIA values are corrected to only account for silicate-bound Ca (Nesbitt and Wilson, 1992). Carbonates are more localised on Mars in comparison to Earth (Ehlmann et al., 2008; Bridges et al., 2019), and only traces of carbonates have been found in Gale crater (Archer et al., 2014). Ca-sulfates may, however, have been introduced to the sediment from external sources during burial and diagenesis (Schieber et al., 2017). To correct for this, targets containing sulfates have been removed from the bedrock dataset, as described in Sections 2.3 and 3.2.1., although indigenous sulfate minerals in the bedrock appear to be rare for the part of the rover traverse considered in this study (sols 1–1482). Some minor phosphates abundances have been described (Forni et al., 2015; Rampe et al., 2017a; Meslin et al., 2018). Because of the possibility of some non-Si-bound Ca in these calculations, the CIA values determined here should be considered as a minimum (Mangold et al., 2017a). CIA values for the stratigraphic groups and units (along with equations) are given in Table 2 based on both APXS and ChemCam data. Despite Marias Pass targets being included in the constrained bulk rock dataset for the Mt Sharp Group, they are not included in the CIA calculations due to their apparent very different mineralogy (e.g., tridymite; Morris et al., 2016) and/or post-depositional aqueous alteration related to the silica-rich fracture-associated halos (see above, Section 4.1.1). CIA values have also been calculated for ChemCam data by Mangold et al. (2018) for various portions of the Murray formation of the Mt. Sharp group.

Constrained ChemCam bulk rock analyses in the Bradbury Group demonstrate a low average CIA (43 ± 9, Table 2) indicative of minimal chemical alteration at the source, a similar conclusion to the CIA values obtained for Bradbury APXS analyses (McLennan et al., 2014; Hurowitz et al., 2017; Siebach et al., 2017). In contrast, constrained bulk rock analyses for the Mt Sharp Group have a relatively elevated average CIA value (52 ± 8) compared to Bradbury, though it is within error of Bradbury’s average (Table 2). These similar averages suggest that the difference...
in chemical weathering for constrained bulk rock analyses has not been strong for either stratigraphic group. However, average CIA values do vary between the different stratigraphic units and waypoints within the Bradbury and Mt Sharp Groups (Table 2). Notably, the coarser grained units (Glenelg, Gillespie Lake, Darwin and Kimberley) generally demonstrate higher CIA values, particularly Darwin (CIA 44 ± 8) compared to the fine-grained, mafic Sheepbed mudstone (CIA 40 ± 4). This difference may either reflect contributions from a relatively felsic sedimentary source region, or the mineral sorting regime preferentially preserving felsic minerals over mafic minerals in coarser grained units (Fedo et al., 2015; Siebach et al., 2017), as felsic material has a naturally higher CIA value compared to sediments rich in mafic minerals (Nesbitt and Wilson, 1992).

Previous CIA studies (Hurowitz et al., 2017; Siebach et al., 2017; Mangold et al., 2018) on Gale crater sediments have highlighted a variation between the Bradbury Group’s Sheepbed Member mudstone and Mt Sharp Group’s Murray formation mudstone. APXS analyses (Hurowitz et al., 2017; Siebach et al., 2017) and ChemCam analysis (Mangold et al., 2018) show the Mt Sharp Group to have notably higher CIA values compared to the Bradbury Group which has led these authors to hypothesise that Murray formation mudstones have undergone more chemical weathering at their source compared to Sheepbed Member mudstones. Our ChemCam analyses of the constrained bulk rock dataset also show Murray formation CIA values to be higher than those of the Sheepbed Member (Murray formation CIA = 51 ± 6 to 55 ± 5, Sheepbed Member = 40 ± 4; Table 2). Hurowitz et al. (2017) hypothesised that the differences in APXS derived CIA values between the Sheepbed (CIA 34 ± 2) and Murray Pahrump Hills (CIA 45 ± 4) mudstone units reflect modest, short-term paleoclimatic fluctuations. According to Hurowitz et al. (2017), the Sheepbed mudstone sediments were eroded and transported during a colder, drier period compared to the Murray formation sediments, and that geochemical variation between these mudstone units is unlikely to be related to a different igneous source. The early stages of chemical weathering in basalts can liberate substantial amorphous silica from primary igneous phases which are susceptible to weathering (McLennan, 2003; Nesbitt and Wilson, 1992; McLennan et al., 2014). On this basis, Hurowitz et al. (2017) also associated the relatively elevated SiO₂ concentrations in the Murray formation with respect to Sheepbed mudstone as being caused by the greater degree of weathering, particularly as CIA and SiO₂ share a (weak) positive correlation between the two mudstone units for APXS analyses.

Fig. 10 shows calculated CIA values for our constrained ChemCam bulk rock dataset plotted against SiO₂, MgO and FeO₁₇. We also conducted SLR models of the mudstone dataset for SiO₂, MgO and FeO₁₇ against CIA (Fig. 10; Appendix C). Our results show a minor (r² = 25.5%), but statistically significant (p < 0.05) correlation between CIA and SiO₂ (Fig. 10A). This slight correlation suggests that chemical weathering may have somewhat increased SiO₂ concentrations between the Murray and Sheepbed mudstones (the model suggests an increase in 2.6 wt% SiO₂ for every 10% increase in CIA values) prior to Murray formation deposition, but in general the variation in SiO₂ concentrations cannot be explained by CIA values alone. In addition, the regression models show that neither MgO nor FeO₁₇ present a significant correlation with CIA values (Fig. 10B, C). The FeO₁₇ regression model does not show a statistically significant correlation with CIA, having high p-values (p = 0.67; Appendix Table C.3). On the other hand, modelled MgO concentrations do show a very weak negative correlation with CIA (r² = 11.1%), for a p-value < 0.01 denoting statistical significance (Appendix Table C.2), though this still leaves most of the MgO variation between the mudstone units unaccounted for. Hence, chemical weathering does not appear to fully explain chemical differences between the Sheepbed and Murray mudstones. According to these models, the increased SiO₂, and decreased MgO present in the Mt Sharp Group is likely a factor of different source region geochemistry, and not sufficiently explained by different degrees of chemical weathering. Furthermore, the strong association of Na₂O with Al₂O₃ in both stratigraphic groups according to MLR and SLR models (Appendix B) suggests that feldspar is still the dominant control on this major oxide across Gale samples.

The slight elevated CIA values in the Murray formation mudstones must therefore be related to the lower CaO, and higher Al₂O₃ contents compared to Sheepbed. A decrease in CaO was also noted by Mangold et al. (2018) to be correlated with CIA within the Mt. Sharp Group members that were studied in that work. In the Sheepbed mudstones, CaO and Al₂O₃ have a Pearson correlation coefficient indicative of a strong correlation (r = 0.60; Appendix B), but these elements are poorly correlated in the Murray formation (r = 0.23). These correlation coefficients suggest that in the Sheepbed mudstone, CaO is present within Al-bearing mineral phases such as plagioclase feldspar and pigeonite – which is supported by CheMin derived mineral compositions (Morrison et al., 2018). Although Murray formation pigeonite determined by CheMin possesses little or no CaO, CaO was identified within the plagioclase feldspar component (Morrison et al., 2018). Hence, dissolution of plagioclase feldspar by chemical weathering at the source region or during transport may have liberated some Ca²⁺ and Al³⁺ cations into solution. The most common secondary minerals to form from the chemical weathering of basaltic minerals are smectites and kaolinites, both of which are associated with Al³⁺, but neither smectites nor kaolinites incorporate significant (>3.5 wt%; Deer et al., 1992) Ca²⁺ into their crystal structure (Deer et al., 1992; Nesbitt and Wilson, 1992). This would result in some Ca²⁺ remaining in solution and being transported away from the system. As plagioclase feldspar is more resilient to chemical weathering than olivine and volcanic glass (Nesbitt and Wilson, 1992), it is likely that these will have also experienced some chemical weathering, liberating Mg and Fe ions into solution that could bond with Al to form Mg₂⁻ and Fe-bearing smectites.

In addition to the relatively low CaO content in the Murray mudstones potentially relating to chemical
weathering, the apparent lack of substantial CaO in Murray pigeonites may also be a product of relatively low CaO at the source. If the source of the Murray mudstones was richer in felsic material through a more evolved magmatic composition (indicated by the pigeonite’s intermediate Fe/Mg ratio; Morrison et al., 2018), this would generate naturally higher CIA values than a more mafic source. In this scenario, these naturally higher CIA values may have been accentuated by minor chemical weathering, but not enough to drastically alter or mask source region characteristics. As the CIA values only show a weak correlation with SiO₂ and MgO, no correlation with FeO₇, and average Murray CIA values are not situated considerably above 50—the CIA value of pure feldspar—we conclude that the degree of chemical weathering of Murray formation source material was not significant enough to mask source region characteristics in our constrained bulk rock dataset. Therefore, we favour that geochemical variation between the Sheepbed Member and Murray formation mudstones is more likely a factor of different source regions.

4.2. Variation of source region characteristics with grain size

4.2.1. Endmembers identified in the coarse-grained (>1 mm) units

Coarse-grained targets (Fig. 2; Le Deit et al., 2016; Mangold et al., 2016; Siebach et al., 2017) were identified within the Darwin Outcrop (sols 392–401), on the Kimberley to Darwin traverse (sols 401–574), at the Kimberley Waypoint (Point Coulomb, Jun Jun target and Liga Members; sols 574–632), and at the base of the Mt. Sharp Group at Pahrump Hills (Bald Mountain 1 and 2; sols 775–778). Bulk ChemCam observation point analyses for these pebbly units plot within the basalt to trachybasalt igneous float and clast density contours, and constitute the majority of point analyses in the Bradbury Group’s high aluminium sub-focus (Figs. 11A, 12A). Total-alkali versus silica (TAS) plots are commonly applied to terrestrial magmatic rocks in order to distinguish the extent of fractionation that a magma has undergone (Le Bas et al., 1986). On Earth, geochemical variation of coarse sediments (grain size greater than 1 mm) is largely inherited from source rock
characteristics (Tolosana-Delgado and von Eynatten, 2009). At Gale crater, the number of igneous float and clasts identified within coarse-grained units and targets (Mangold et al., 2016; Edwards et al., 2017) suggests that this is also true for Gale’s conglomerates. Due to the lack of large scale chemical remobilisation in the Bradbury Group and presence of primary igneous minerals (Fig. 9), this plot reveals likely detrital source compositions and geochemical differences between the groups relating to the alkali contents (Mangold et al., 2017b), especially when compared to the Gale crater igneous float and clasts. The distribution of igneous float rock and Bradbury Group density contour focal compositions are consistent with smoothed estimates of bulk rock compositions. The foci of the compositional contours for the Bradbury group and igneous float rocks are not coincident with the most abundant primary igneous minerals. Therefore, similar to the results of Cousin et al. (2017), this indicates that a mixture of minerals is analysed by ChemCam at most observation points, including for relatively coarse mineral grains.

Darwin-type conglomerate observation point analyses (Darwin and Darwin to Kimberley traverse analyses) are concentrated within the high-Al$_2$O$_3$ igneous float and Bradbury sub-focus (Fig. 11A), which make it likely that the coarse-sand matrix and small pebbles of Darwin-type conglomerates are derived from the trachybasalt parent rock. Meanwhile, the Kimberley-type conglomerates are relatively more enriched in K$_2$O than Darwin-type conglomerates (Figs. 11D and 12A) suggesting that a potassium-rich endmember has contributed to this stratigraphic unit (Le Deit et al., 2016). The potassium enrichment correlates with the presence of sanidine detected at high concentrations (~20%) in the Windjana drill hole within the Dillinger Member sandstone linked to a potassic-igneous source (Treiman et al., 2016). The JumJum conglomerate target (sol 550) representative of the lowermost Point Coulomb Member in the Kimberley formation also has elevated K$_2$O/Na$_2$O ratios. Hence, during the period of deposition in the sedimentary units leading up to Kimberley, this trachytic endmember grew in significance as a Gale source region (Le Deit et al., 2016).

Aside from source region variation of the sediments in Gale, an alternative hypothesis relating to Martian mineral sorting regimes could explain why both the igneous and high-aluminium Bradbury Group density contours are co-located between the alkali and plagioclase feldspar compositions (Figs. 11A, 12A). Feldspars have been noted to be more resilient to physical weathering than mafic minerals (Nesbitt and Young, 1996; Fedo et al., 2015), and hence are more concentrated in coarse-grained fluvial sediments.

![Fig. 11. ChemCam observation point analyses for Gale crater stratigraphic units and targets shown on Al$_2$O$_3$ versus MgO major oxide diagrams and total alkali-silica diagrams. Plots are displayed in accordance to the grain size fraction represented (left column is coarse (>1 mm), middle column sand grain (0.062–1 mm) and the right column is mud grain size (<0.062 mm)). Igneous group contours are derived from Gale crater igneous float rocks and clasts data (Mangold et al., 2016; Edwards et al., 2017) and olivine fractionation line determined by Edwards et al. (2017), CheMin derived Martian mineral compositions for the Rocknest (olivine, plagioclase, augite) and Windjana (alkali feldspar) samples are taken from Morrison et al. (2018), Adirondack Class basalt compositions (Gellert et al., 2006).](image-url)
eroded from basaltic provenances (Siebach et al., 2017). The scatter of these observation point analyses towards modelled Gale feldspar compositions could suggest that the trachybasalt source signature identified in coarse sediments may be a result of the concentration of feldspar resulting from this Martian fluvial mineral sorting regime. However, as most conglomerate targets are poorly sorted, and have therefore been derived from multiple source rocks (Grotzinger et al., 2014, 2015), we favour the initial interpretation that the felsic nature of the coarse-grained units are considered to be representative of input from the trachybasalt source composition described by Edwards et al. (2017), and a potassic source described by Treiman et al. (2016) in the Kimberley formation Liga conglomerate Member.

4.2.2. Endmembers identified in the sandstone units (0.062–1 mm)

Fluvial sandstones are present within the Glenelg and Gillespie Lake Members of the Yellowknife Bay formation (Grotzinger et al., 2014; Anderson et al., 2015) as well as the Kimberley formation’s Square Top, Dillinger, and Mt. Remarkable Members (Grotzinger et al., 2015; Treiman et al., 2016). On Earth, sandstones and shales have been related to source regions on the basis of their SiO₂/Al₂O₃ and Na₂O/K₂O ratios (Pettijohn et al., 1972; Roser and Korsch, 1986). They are also classified according to their relative abundances of quartz, feldspar and lithic fragments as these mineral populations are abundant in terrestrial sediments (Pettijohn et al., 1972; Dickinson, 1988; Weltje, 2012). On Mars, quartz has not been detected with CheMin at concentrations >1% (Fig. 9; Vaniman et al., 2014; Treiman et al., 2016; Rampe et al., 2017a; Yen et al., 2017), therefore the SiO₂/Al₂O₃ ratios employed by most terrestrial sediment geochemical classification schemes (Pettijohn et al., 1972; Herron, 1988) cannot distinguish key mixing components on Mars. Instead, due to the abundance of mafic minerals (especially clinopyroxene) detected in the CheMin analyses of drilled samples, we use the log-ratio Al₂O₃/(MgO + FeO T) to distinguish between these and the major felsic-felsic contributions in sandstone and fine-grained units (Fig. 12D–F). The log-ratio SiO₂/K₂O (Fig. 12E–D) and K₂O/Na₂O (Fig. 12A–C and 13, plotted against SiO₂ to differentiate between possible igneous rock sources) are employed to distinguish volcanic lithic-fragments and feldspars from evolved (K-rich feldspar) sources (Grotzinger et al., 2014, 2015) the mafic-felsic mineral sorting regime outlined in Section 4.2 and described by Siebach et al. (2017) is also unlikely to have had a significant effect on sandstone compositions. Hence, this scatter may relate to sediment contribution from the trachybasalt endmember identified in the conglomerate units. However, as there is grain size variation within the units themselves, it is not clear to what extent the high proportion of plagioclase feldspar in this unit has been concentrated in the coarser fraction as a result of the mineral sorting regime (Siebach et al., 2017). Therefore, it is impossible to discern whether these feldspar-rich sediments were related to the trachybasalt source type or feldspar from the low-Al₂O₃, basaltic source that was concentrated in the coarser beds.

A clear difference can be seen between the Yellowknife Bay and Kimberley formation sandstones (Square Top, Dillinger and Mt. Remarkable Members) based on potassium-enrichment (Figs. 11E and 12B). The upper Dillinger and Mt. Remarkable Members are particularly distinct, forming their own cluster. As the CheMin analysis was conducted within the Dillinger Member, it can be assumed that the less K₂O-rich Square Top Member of the Kimberley formation contains a lower abundance of sandine. The Glenelg and Gillespie Lake Members also possess higher K₂O/Na₂O ratios (cf. Anderson et al., 2015) than the sodic Darwin and Hummocky plains conglomerates. If this is attributed to relative proportions of potassic and sodic feldspars this suggests that these coarse sandstone units also contain some sandine mineral grains. Therefore, the hypothesised potassic-igneous endmember has had an influence on Gale’s sediment mineral assemblage throughout its depositional history. But, both due to a change in source region contribution, or increased sandstone maturity (Grotzinger et al., 2015; Treiman et al., 2016), this endmember is only seen to significantly contribute to the later Dillinger and Mt. Remarkable stratigraphic units.

In summary, the sandstone units within Gale crater are largely derived from a subalkaline basalt source, similar to the Adirondack Class basalts of Gusev crater, with a minor contribution from a potassic volcanic source (Treiman et al., 2016). Sediment contribution from the potassic volcanic source appears to increase from the Darwin waypoint to the Kimberley formation (Le Deit et al., 2016) where it is seen to strongly influence the geochemical composition of the Dillinger and Mt Remarkable Members. The trachybasalt endmember described by Edwards et al. (2017) may have also contributed to the sedimentary record within this
However, it is not clear whether the increased abundance of feldspars present within these units are the result of the trachybasalt source region, or grain size variation within units.

4.2.3. Endmembers within the mudstone (<0.062 mm) units

The Sheepbed Member and Murray formation constitute Gale crater’s mudstones (Grotzinger et al., 2014; Grotzinger et al., 2015). The vast majority of the Mt. Sharp Group’s lower Murray formation possesses a relatively uniform composition throughout the East and West Naukluft, and Murray Buttes localities (Figs. 11 and 12). This general chemistry is more enriched in Al₂O₃, K₂O, SiO₂, and depleted in CaO, MgO, and Na₂O than the Sheepbed mudstone in the Bradbury Group (which plot within the mafic Bradbury contour focus). The very fine grain size of the Sheepbed mudstone and Murray formation results in more homogeneous compositions observed by ChemCam. Due to the removal of alteration features and targets potentially exposed to open system weathering (see methods Section 2.3, and Sections 3.2 and 4.1; Figs. 7–10), the observed geochemical variation between the Bradbury and Mt. Sharp Group (lower Murray) mudstone units must be representative of either lacustrine mineral sorting processes (Section 4.2.3.1), or different endmember mixing proportions (Section 4.2.3.2).

4.2.3.1. The effects of mineral sorting processes within the Gale lake on sediment geochemistry. A possible explanation for the observed geochemical differences between stratigraphic groups could relate to transport fractionation processes in a lake body. Supposing no change in sediment source region and assuming a homogeneous grain size of input materials, distal lake deposits will possess higher Al₂O₃ and alkali (especially Na₂O) concentrations due to greater, low density felsic and possibly clay-rich mineral abundances (Das et al., 2006; Jin et al., 2006; Babechuk et al., 2014). For Gale crater’s Sheepbed mudstone, bed thicknesses and close association with overlying fluvial sandstone have been interpreted as a depositional environment that was relatively rapid, close to the shoreline and sediment source (McLennan et al., 2014; Hurowitz et al., 2017). It has also been argued by Hurowitz et al. (2017) that hematite-bearing sections of the Murray formation such as the Pahrump Hills were deposited in shallow water potentially near the shoreline due to high abundances of hematite and Mn-oxide pertaining to an oxidising environment. Trough cross-bedding and interstratified siltstone and fine sandstone near Pahrump also support deposition in shallow waters and/or proximity to the lake shoreline (Hurowitz et al., 2017). Therefore, if both the Sheepbed Member and Pahrump Hills Murray formation mudstones are deposited at near-shore locations in the paleolake, their differences in mineralogy and chemistry must reflect a change in source regions. The Pahrump Hills mudstones located near the interfingering boundary with the Bradbury Group are more Na₂O- and K₂O-rich, and possess greater abundances of plagioclase feldspar than the Sheepbed
mudstone (Figs. 9 and 12F) which is not what we would expect based on mineral density and sediment sorting alone.

In contrast, Rampe et al. (2017a) noted that the observed secondary mineral assemblages in Pahrump Hills that Hurowitz et al. (2017) attributes to a shallow lake environment could also be produced through diagenesis in short-lived, oxidizing ground waters. However, the morphological evidence for shallower lake conditions for the Mt Sharp Group near Pahrump is significant (thicker laminations, interstratified, cross-bedded, fine sandstones; Grotzinger et al., 2015; Hurowitz et al., 2017). Thus differing source regions between the Murray formation and Sheepbed Member mudstones are the most plausible explanation for their compositional differences.

4.2.3.2. Endmembers and source region contributions. In this section we build on the model above that chemical weathering in source materials, diagenesis and transport processes have not erased the source rock compositions of the mudstones. The Sheepbed Member consistently plots within the dominant, Mg-rich local composition of the Bradbury Group (Figs. 11C, F, 12C) and is similar in composition to the Adirondack Class basalts and subalkaline basalt endmember determined by Edwards et al. (2017) from Gale crater’s igneous pebbles and float (Fig. 11C). Therefore, the Sheepbed mudstone unequivocally relates to the dominant, subalkaline basalt source region that has also contributed sediment to the Bradbury Group’s sandstones and conglomerates. In contrast, the Mt Sharp Group MgO composition appears intermediate between those of the Bradbury trachybasalt and subalkaline basalt subfocal compositions (Figs. 6D and 11C) suggesting similar mixing between a mafic and felsic source. However, the increased K₂O, Al₂O₃/(MgO + FeO T) and SiO₂ content of the Murray formation compared to the Sheepbed mudstones (Figs. 11C, F, 12C, and F) requires a greater contribution of felsic minerals and silica-rich material. CheMin analyses (Fig. 9) also demonstrate this greater felsic/mafic mineral proportion.

CheMin XRD analyses show sanidine across the Murray formation (except for Mojave 2 and Oudam) with between 4.2% at the Buckskin drill hole, to 7.7% in Marimba 2 (Morris et al., 2016; Rampe et al., 2017b). These are higher abundances than those of sanidine detected in the Yellowknife Bay formation (2.9–3.4%). On average, Murray alkali feldspars are inferred to possess higher abundances than those of sanidine detected in the Yellowknife Bay formation (2.9–3.4%). On average, Murray alkali feldspars are inferred to possess higher abundances than those of sanidine detected in the Yellowknife Bay formation (2.9–3.4%). On average, Murray alkali feldspars are inferred to possess higher abundances than those of sanidine detected in the Yellowknife Bay formation (2.9–3.4%). On average, Murray alkali feldspars are inferred to possess higher abundances than those of sanidine detected in the Yellowknife Bay formation (2.9–3.4%). On average, Murray alkali feldspars are inferred to possess higher abundances than those of sanidine detected in the Yellowknife Bay formation (2.9–3.4%). On average, Murray alkali feldspars are inferred to possess higher abundances than those of sanidine detected in the Yellowknife Bay formation (2.9–3.4%). On average, Murray alkali feldspars are inferred to possess higher abundances than those of sanidine detected in the Yellowknife Bay formation (2.9–3.4%).
Alternative explanations have been proposed by others to explain the presence of the high-silica Marias Pass unit. The first suggests that it was deposited in a temporally isolated event, such as a relatively sudden, volcanic eruption of a highly evolved silicic magma (Morris et al., 2016). This could have generated a thick layer of ash that was later washed downstream into the lake (Morris et al., 2016). Once this eruption had ceased, a return to the mafic sediments deposited prior to the eruption occurred, resulting in the distinct Marias Pass stratigraphic unit within the Murray formation (Morris et al., 2016; Frydenvang et al., 2017). The second explanation for silica-enrichment in Marias Pass discusses that sediments may have been leached in place by short-lived, mildly acidic, late-stage fluids (Rampe et al., 2017a). These diagenetic fluids could have dissolved ferromagnesian minerals and created the observed trace element trends (Rampe et al., 2017a). As tridymite is strongly inferred to be detrital from a silicic source region (Morris et al., 2016), and as the abundance of magnetite and lack of hematite at this locality is more indicative of reducing settings (Hurowitz et al., 2017), we prefer a source region and deposition-related hypothesis to explain the presence of this unusual unit.

5. CONCLUSIONS

Investigation of the ChemCam database from the beginning of the mission through to sol 1482 suggests that the main influences on observed geochemical variation are changing sediment sources (see Table 3 for endmember compositions), and mineral sorting regimes once diagenetic features are removed from the database and the effects of chemical weathering have been taken into account. The dominance of primary igneous minerals, and CIA values unrelated to SiO$_2$ and MgO variations in our constrained dataset, all suggest that chemical weathering processes have not dominated the endmember compositions despite CIA values being higher in the Mt Sharp Group mudstones compared to the Sheepbed mudstone Member. Instead, bulk compositions for the stratigraphic units and groups are likely determined by varying proportions of primary igneous minerals, with isochemical weathering hypothesised to generate amorphous and phyllosilicate component compositions similar to bulk rock geochemistry. Our results also support the Martian fluvial mineral sorting regime whereby felsic minerals are more abundant in coarse sandstone fractions due to their resilience to diminution during transportation (Siebach et al., 2017), but are more likely to travel farther into a lake body because of their lower density compared to mafic minerals. As a consequence of the Murray formation and Sheepbed mudstone possessing a possibly similar depositional setting in relatively shallow water and/or close to the shoreline, the compositional differences between these units likely reflect changes in sediment source region. This change in sediment provenance is particularly notable in the Marias Pass region with the appearance of the silica-rich endmember and we propose a silica-rich, subalkaline basalt endmember to be the main source of sediments at the base of Mt Sharp.

In isolating the analysis to grain size dependent subpopulations, four previously identified endmembers and one new endmember are observed throughout Gale crater sediments in this part of the mission, which can be related to specific igneous source regions (Fig. 13). These are:

1. A subalkaline basalt endmember (Anderson et al., 2015; Siebach et al., 2017); similar in composition to the Adirondack Class basalts observed by the MER Spirit rover at Gusev. This endmember is observed throughout the Bradbury Group stratigraphy and is concentrated within the Sheepbed mudstone Member of Yellowknife Bay. Due to the similarity of this composition to Adirondack basalts it can be inferred that this is most likely a regional endmember composition and contributes significantly to Gale’s geochemical record.

2. A trachybasalt endmember (Edwards et al., 2017); likely formed during crystal-melt fractionation of the mafic, Adirondack-type basalt that constitutes endmember (1). This is largely identified in clasts within the conglomerate outcrops (especially Darwin), and is characterised by Na$_2$O and Al$_2$O$_3$ enrichment associated with an increase in plagioclase feldspars.

3. A trachyte endmember (Treiman et al., 2016); related to an increased abundance of sanidine in comparison to the more Na-rich plagioclase feldspars from endmember (2). This endmember is recognised to an extent within the Gillespie Lake and Glenelg Members of the Yellowknife Bay formation, plus the later Mt. Sharp Group mudstone. It is seen to contribute significantly to the Kimber-

### Table 3

A table of the ChemCam derived sediment endmember compositions in Gale crater from this study and Edwards et al. (2017). Propagated 1σ error is provided in brackets for the modelled endmember compositions. Exact compositions cannot be derived from bulk sediment analysis for the trachyte and rhyolite endmembers as these are based on the presence of specific minerals within complex sedimentary units that were either derived from multiple igneous source regions (i.e., Windjana sandstone, Treiman et al., 2016) or have experienced uncertain levels of aqueous alteration (i.e., the Marias Pass mudstones, Frydenvang et al., 2017).

<table>
<thead>
<tr>
<th>Endmember</th>
<th>SiO$_2$</th>
<th>TiO$_2$</th>
<th>Al$_2$O$_3$</th>
<th>Fe$_2$O$_3$</th>
<th>MgO</th>
<th>CaO</th>
<th>Na$_2$O</th>
<th>K$_2$O</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Subalkaline</td>
<td>±1.4</td>
<td>±0.2</td>
<td>±0.8</td>
<td>±1.2</td>
<td>±1.9</td>
<td>±0.9</td>
<td>±1.6</td>
<td>±1.4</td>
<td>±0.3</td>
</tr>
<tr>
<td>Basalt</td>
<td>±1.9</td>
<td>±0.2</td>
<td>±0.8</td>
<td>±1.4</td>
<td>±0.6</td>
<td>±0.6</td>
<td>±0.2</td>
<td>±0.2</td>
<td>This paper, Edwards et al. (2017)</td>
</tr>
<tr>
<td>Trachybasalt</td>
<td>±3.5</td>
<td>±0.2</td>
<td>±2.5</td>
<td>±1.9</td>
<td>±1.6</td>
<td>±1.2</td>
<td>±1.7</td>
<td>±1.3</td>
<td>±0.9</td>
</tr>
<tr>
<td>Silica-rich</td>
<td>±1.9</td>
<td>±0.2</td>
<td>±1.4</td>
<td>±1.4</td>
<td>±1.6</td>
<td>±1.4</td>
<td>±1.4</td>
<td>±0.3</td>
<td>This paper</td>
</tr>
<tr>
<td>Basalt</td>
<td>±1.9</td>
<td>±0.2</td>
<td>±0.8</td>
<td>±1.4</td>
<td>±0.6</td>
<td>±0.6</td>
<td>±0.2</td>
<td>±0.2</td>
<td>This paper</td>
</tr>
</tbody>
</table>
ley formation sandstones and conglomerates and – due to the detected sanidine being of a high order – it may be associated with erosion from a volcanic alkaline or trachytic source (Treiman et al., 2016).

4. A silica-rich (rhyolite) igneous endmember (Morris et al., 2016); identified at the Marias Pass locality within the Mt. Sharp Group, this endmember is thought to resemble the erosion of a late-stage magmatic source, or ash cloud generated from a volcanic event of an evolved magma composition, due to the predominance of high-temperature silica polymorphs.

5. A silica-rich, subalkaline basaltic endmember; proposed by this study, this endmember contributes most to the Mt. Sharp Group and is more SiO₂, Al₂O₃ and K₂O-rich than endmember (1). This endmember is seen to become more prevalent up the stratigraphic succession.

The above source compositions have strongly influenced the geochemistry and mineralogy of the sedimentary rocks which constitute the Bradbury and Mt. Sharp (Murray formation up to sol 1482) fluviolacustrine stratigraphic groups traversed by MSL. Endmembers (1) and (2) are largely associated with the distal alluvial fan deposits seen within the Bradbury Group (Grotzinger et al., 2014; Grotzinger et al., 2015) while endmembers (4) and (5) are solely recognised in the lower Murray formation. Endmember (3) is identified across the fluviolacustrine deposits, predominately in Bradbury Group fluvial sandstone and Murray mudstone, but contributes little to the Yellowknife bay formation. Instead, the Yellowknife bay formation appears to be largely derived from endmember (1) with some local influence of endmember (2).

This study shows that despite temporal and spatial influences on sediments, unique source region characteristics can still be distinguished within the Martian sedimentary record. This can be achieved through identifying chemical and mineralogical trends that contradict physical sorting processes, as well as analysing sediments according to grain size. Gale crater stratigraphy therefore shows that the ancient highland crust which contributed to its basin fill is more geochemically complex than previously anticipated for Mars. This is despite the lack of evidence for evolved crustal compositions detected from orbit which we speculate may relate to evolved rocks being volumetrically smaller relative to basalt in Mars’ crust.

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APPENDICES A–D. SUPPLEMENTARY MATERIAL

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REFERENCES


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