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In situ detection of boron by ChemCam on Mars

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Abstract We report the first in situ detection of boron on Mars. Boron has been detected in Gale crater at levels <0.05 wt % B by the NASA Curiosity rover ChemCam instrument in calcium-sulfate-filled fractures, which formed in a late-stage groundwater circulating mainly in phyllosilicate-rich bedrock interpreted as lacustrine in origin. We consider two main groundwater-driven hypotheses to explain the presence of boron in the veins: leaching of borates out of bedrock or the redistribution of borate by dissolution of borate-bearing evaporite deposits. Our results suggest that an evaporation mechanism is most likely, implying that Gale groundwaters were mildly alkaline. On Earth, boron may be a necessary component for the origin of life; on Mars, its presence suggests that subsurface groundwater conditions could have supported prebiotic chemical reactions if organicies were also present and provides additional support for the past habitability of Gale crater.

1. Introduction

Boron is an intriguing element for geology and astrobiology. On Earth, boron is a highly soluble element found concentrated in phyllosilicate-bearing sediments and evaporite deposits, making boron useful for understanding surface and subsurface aqueous processes [Goldschmidt and Peters, 1932; Fredrickson and Reynolds, 1960; Hunt et al., 1966; Lerman, 1966; Couch and Grim, 1968; Harder, 1970; Perry, 1972] and in reconstructing fluid temperature and pH [Spivack et al., 1987; You et al., 1996; Goldberg, 1997; Williams et al., 2001]. Borate anions (BO$_4^{-}$) may also be necessary for the origin of life [Scorei, 2012]. Borate anions have been shown to slow the decomposition of ribose in solution, an important prebiotic compound and component of ribonucleic acid (RNA), and mediate ribose formation [Kim et al., 2011] by the formose reaction [Boutierow, 1861] by forming ribose-borate complexes [Ricardo et al., 2004; Scorei, 2012; Furukawa et al., 2013].

Boron and borates are potentially detectable on Mars by instruments deployed on the surface, including ChemCam [Wiens et al., 2012; Maurice et al., 2012] and CheMin [Blake et al., 2012] on Curiosity, or by orbital multispectral infrared remote sensing instruments [Cloutis et al., 2016]. Boron has also been detected in Martian meteorites (Table 1). Although Martian evaporite deposits have been detected by Mars Global Surveyor TES and Mars Odyssey THEMIS instruments [e.g., Osterloo et al., 2008; 2010; Hynek et al., 2015], multispectral imagers have not detected borates remotely as of this publication [Cloutis et al., 2016]. CheMin, the X-ray diffraction instrument on Curiosity, is capable of detecting borate minerals but has not detected them in any drilled or scooped samples. Thus, this detection of boron by ChemCam represents the first time boron has been observed in situ on Mars.
Here we show that boron is enriched relative to igneous Martian materials (Table 1) and found in several calcium-sulfate-filled fractures (henceforth called “veins,” described by Schwenzer et al. [2016], Rapin et al. [2016], and Nachon et al. [2014, 2017]) hosted by the “Yellowknife Bay” formation, the lacustrine “Murray” formation mudstone, and the unconformably overlying aeolian “Stimson” formation sandstone units (described in Grotzinger et al. [2014, 2015], Fedo et al. [2017], and Banham et al. [2017]).

### 2. Methodology

Boron observations were made using the ChemCam instrument suite: a laser-induced breakdown spectroscopy (LIBS) instrument and a remote microimager (RMI). LIBS provides elemental emission spectra for 350–550 μm sized observation points on rocks up to a 7 m standoff distance [Wiens et al., 2012; Maurice et al., 2012]. The RMI records high-resolution images of target points before and after LIBS observations, allowing visual identification of and context for each observation point [Le Mouélic et al., 2015]. Mastcam images are also recorded for each ChemCam target using the 100 mm focal length camera (Right Mastcam) [Malin et al., 2010] to provide geologic context and color images.

Boron is observed in LIBS data via neutral atomic emission lines at 249.75 and 249.84 nm (Figure S1a) [Sansonetti and Martin, 2005]. Peak areas are obtained by fitting a Voigt lineshape to the LIBS spectra [cf. Rapin et al., 2016]. A preliminary calibration curve is created from standards, yielding the relative abundance of boron (see supporting information). Because the majority of ChemCam targets in Gale contain ~18 wt % FeOT [Mangold et al., 2017], interference from a Fe II emission line at 249.96 nm (Figure S2) limits boron detection in Mars data to targets with ≤10 wt % FeOT. Laboratory LIBS spectra on standards show that B peaks in samples with ~0.05 wt % B abundance are visible in samples containing ~11 wt % FeOT, and the detection limit is ~100 ppm B in Fe-free standards. The detection limit is defined here as the composition at which the boron peak is visible above background noise plus the average area of any interfering lines (Figure S2). Most low Fe observations correspond to CaSO₄-filled veins or high silica targets [e.g., Morris et al., 2016; Frydenvang et al., 2017]. In vein targets, the main interfering line is Ca III (249.84 nm) rather than Fe. The statistical significance of boron detections on Mars are calculated from interfering line average peak area and standard deviation are.

### 3. Results

Boron is observed in 43 vein targets (45 2σ detections and 30 3σ detections totaling 98 observation points) (Figure 1, Table S1, Figure S1). We estimate a detection limit of ~100 ppm B in LIBS spectra and boron abundances in veins up to 500 ppm B based on preliminary laboratory studies of boron-bearing standards (see supporting information). Boron was detected in five calcium sulfate vein targets in the Yellowknife Bay Group “Sheepbed” member mudstone, “Point Lake,” and “Shaler” sandstone (Figure 2, inset A) and in 38 veins in the Mt. Sharp Group Murray mudstone and Stimson sandstone (Figure 2). Although veins were targeted on rare occasions during the traverse through the Bradbury Group after visiting Yellowknife Bay, no boron was detected there (Figure 2). The first observation of boron coincided with the first ever vein observation by ChemCam on Martian day (sol) 113 of the mission [Nachon et al., 2014], but the majority of detections

| Table 1. Literature Summary of Boron Content in Bulk Silicate Mars and Martian Meteorites |
|--------------------------|-----------------|-----------------|
|                         | ppm B References |
| **Bulk Silicate Mars** | 0.6             | Stephenson et al. [2013] |
| **Meteorite(s)**        | Associated Phase |
| MIL 090030              | olivine          | 0.03             | Stephenson et al. [2013] |
| Nakhla, Lafayette, MIL 090030 | pyroxene     | 0.1–1           | Stephenson et al. [2013], and Spivak-Birndorf et al. [2008b, 2008c] |
| **Nakhla**              | whole rock       | 4.6             | Loders [1998] |
| **Nakhla**              | melt inclusion   | 5               | Spivak-Birndorf et al. [2008c] |
| **Nakhla**              | mesostasis       | 4–7             | Spivak-Birndorf et al. [2008c] |
| Nakhla and Lafayette    | iddingsite      | 7–18            | Spivak-Birndorf et al. [2008a, 2008b] |
| MIL 090030              | alteration vein  | 154–166         | Stephenson et al. [2013] |
| Chassigny               | whole rock       | 6.3             | Loders [1998] |
have been made since sol 1350 in Upper Murray veins, at elevations above ~4435 m (Figure 2, inset B), and are colocated with clay-rich Murray bedrock [Rampe et al., 2017a].

Boron has been detected in a variety of calcium sulfate vein morphologies and settings (Table S1). Boron does not preferentially appear in any one vein morphology. Vein targets in Yellowknife Bay include the ubiquitous thin (<2 cm) veins, veins with nodular textures, small nodules or “bowls,” polygonal textures, and “protruding” veins (e.g., Figure 1b) [Nachon et al., 2014]. Subsequent investigations of veins in the Murray and Stimson bedrock units [Nachon et al., 2017] found complex and large veins (>2 cm wide) in the “Pahrump Hills” region. The only detection of boron in Pahrump Hills was the “Indianola” target, which is a light-toned part of a large vein in the “Garden City” outcrop. Other targets found after “Naukluft Plateau” include “horizontal veins” that are approximately parallel to mudstone laminations as seen directly above the “Cadillac_Mountain” target (Figure 1c).

4. Discussion

All boron detections in Gale crater are in veins. However, the lack of detections in bedrock targets is very likely a consequence of observation bias: since Fe and B emission lines interfere with each other in the LIBS spectra, boron detection is limited to targets with ≤10 wt % FeOT, which represents unusually low Fe for Gale bedrock. Nevertheless, the vein geological context points to emplacement of boron by water. Boron is very easily oxidized; boron is always a borate in conditions where water is stable and in past and present Mars surface oxygen fugacity conditions, which implies that the boron was a borate when emplaced and remains a borate at present [Ingr ri, 1963; Schubert, 2015]. We suggest that the boron was at least partly emplaced by a late-stage groundwater and was likely redistributed from a primary source [cf. Schwenzer et al., 2016]. Although the iron
content of typical bedrock targets on Mars precludes reliable measurement of boron, boron may indeed be present in these materials.

4.1. Models for Boron Enrichment in Veins

There are several possible mechanisms of boron enrichments in veins, and each has intriguing implications for Mars. Both models considered suggest that boron (a) weathered out of primary igneous crust in the Gale crater watershed [Ehlmann and Buz, 2015], (b) was dissolved in lake water, and (c) was dissolved in the late-stage groundwater system. However, the different possible mechanisms of initial boron concentration from lake waters have important consequences for fluid pH and the resultant borate mineralogy, since possible borate mineral precipitates are dependent on fluid temperature, pH, and composition [Hunt et al., 1966; Garrett, 1998; Schubert, 2015].

The temperature of the vein-emplacing fluids may be constrained using a number of observations. First, calcium sulfate veins in Gale are predominantly composed of bassanite (CaSO₄·H₂O) [Rapin et al., 2016],
which likely forms from the dehydration of gypsum due to hyperarid conditions. This interpretation is consistent with mineralogical results from CheMin [Vaniman et al., 2017]. The presence of gypsum indicates that the groundwater was <60°C [Rapin et al., 2016]. Other studies of sulfate veins and phyllosilicates in the crater suggest an upper limit of ~50°C [Vaniman et al., 2014; Schwenzer et al., 2016; Nachon et al., 2014, 2017]. Thus, the majority of groundwaters in Gale were likely low temperature.

Fluid pH is the main driver of borate-rock interaction, but pH of Gale groundwater is not well constrained. While most evidence suggests that Gale was host to neutral to alkaline groundwater [e.g., Frydenvang et al., 2017; Hurowitz et al., 2017], some observations suggest that acidic groundwater (pH <6) may have interacted with the bedrock [Yen et al., 2017; Haur stout et al., 2017; Rampe et al., 2017b]. Borate species form alkaline solutions with water, providing more evidence that Martian groundwater was alkaline. Although we have no constraints on the concentration of the other ions in solution that affect groundwater pH, when in solution with pure water, borate acts as a buffer with pH ~9. Thus, ~100 ppm of boron in solution will tend to make the fluid alkaline, preventing pH shifts outside of pH 8–10. Higher dissolved borate concentrations increase the effectiveness of the buffer. At elevated pH, borates can be adsorbed to 2:1 phyllosilicates and the concentration of borates in clays has been used as a proxy for terrestrial paleosalinity [Fredrickson and Reynolds, 1960; Lerman, 1966; Harder, 1970; Perry, 1972]. The borate anion is adsorbed by positively charged sites on the surfaces of 2:1 phyllosilicates that form in alkaline fluids. The optimal pH for uptake of borates by clays depends on the type of clay and fluid pH, with a pH range for terrestrial clays of ~8.5–10 [Couch and Grim, 1968; Keren and Mezuman, 1981; Keren and O’Connor, 1982; Karahan et al., 2006] at low temperatures <120 °C [You et al., 1996]. Additionally, borates can adsorb to amorphous materials in soils in similar pH conditions to clays, depending on the oxide [Goldberg, 1997]. While the degree of borate adsorption to Fe- and Mg-rich 2:1 phyllosilicates and its relationship with pH is not well constrained for Martian conditions, studies of common terrestrial soil uptake of B [Goldberg, 1997] can be used to estimate that ~100–1000 ppm of B could be adsorbed by clays and amorphous materials in favorable conditions.

With these constraints in mind, we consider two models to explain the observed boron in veins. The first model involves initial adsorption of boron by phyllosilicates and amorphous materials during their deposition in the lake, followed by later groundwater interaction with bedrock to extract the boron before it was coprecipitated with Ca-sulfates in veins. The second model involves initial enrichment of boron in evaporite layers in the lake, followed by later groundwater interaction with bedrock to extract the boron before it was coprecipitated in fractures. In the first model, borates dissolved in Gale lake water adsorbed on to phyllosilicates and amorphous materials as the lacustrine bedrock was deposited. Later, borate ions may have been mobilized by groundwater and re-emplaced in veins. This model is supported by the observation that the lacustrine Murray and Yellowknife Bay units that host the highest abundance of boron-bearing veins are rich in 2:1 phyllosilicates and amorphous phases [Rampe et al., 2017a; Bristov et al., 2017]. Hence, postlithification interactions of groundwater with these clay-amorphous-rich deposits could lead to boron being desorbed from the bedrock and then precipitated in veins. Groundwater with a low pH or with constituents that have a stronger affinity than borates to clay or amorphous materials could release borates back into solution and lead to borate enrichments in fracture filling materials. Yellowknife Bay pore fluids may have reached pH ~12 [Bridges et al., 2015], which could release any boron from or inhibit boron adsorption to clay and amorphous materials. If the borate is adsorbed by the bedrock in water with pH of 9, then fluids of pH 6–7 would be sufficient to free the majority of the borate anions trapped in the bedrock.

Although boron adsorption on to clays and amorphous materials is plausible, many observations suggest that this first model is unlikely because of the likely fluid pH ranges that were present in Gale Lake. Lake waters were near neutral to alkaline pH [Vaniman et al., 2014; Hurowitz et al., 2017]; optimal adsorption of boron to clays and amorphous materials occur at higher alkalinitities (pH 8.5–10), and boron adsorption to these materials at pH 6–7 is not favored. Borate concentration in the lake was likely low during deposition of clay- and amorphous-bearing mudstone because it is only present in extremely low concentrations in the crust (Table 1). Thus, boron concentration requires both extensive weathering in the Gale watershed region and subsequent concentration by evaporation of water, e.g., in a closed drainage basin. In a deep near-neutral pH lake [Hurowitz et al., 2017], conditions would not have been conducive to adsorption of boron to clay or amorphous materials.
In the second model, borates previously deposited in evaporite layer deposits were remobilized by a subsequent wet period or circulation of basinal fluids after burial. This boron-bearing groundwater could then have precipitated borates in evaporite veins in lower strata. This scenario is supported by the fact that borates are highly soluble in water and in very arid environments on Earth are typically found in evaporites.

Although a primary source of boron-bearing evaporates has yet to be discovered, remote sensing studies suggest that sulfate/phyllosilicate layers above the rover’s current location in the Murray formation could represent the source evaporitic layer [Milliken et al., 2010; Baldridge et al., 2009; Grotzinger et al., 2015; Fraeman et al., 2016]. During the transition to a drier Mars climate, the Gale crater basin could have been analogous to saline playa lakes in Western Australia, where acidic and alkaline groundwater interact, leading to the coeval deposition of clays and sulfates [Baldridge et al., 2009]. Alternatively, Gale could have hosted a perennial lake system such as those in Death Valley California, where layers of borates and other evaporites build up between mudstone deposits after multiple wet-dry cycles [Eugster and Hardie, 1978; Lowenstein et al., 1999; Tanner, 2002]. Later groundwater interaction with these evaporite layers could dissolve some or all of the salts, transporting them into underlying layers of mudstone through fractures [Schwenzer et al., 2016]. This hypothesis is supported by the observation of possible desiccation features in the Upper Murray [Stein et al., 2017]. More broadly, evaporite deposits have been remotely detected on Mars throughout the Southern Highlands [Osterloo et al., 2008; Hynek et al., 2015; Ehmann and Buz, 2015] and evaporite phases including halite are common in nakhlite martian meteorites [Gooding et al., 1991; Bridges and Grady, 2000; Hicks et al., 2014]. Boron has recently been observed in the Martian meteorite Nakhla in association with halite and carbonaceous material [Thomas-Kept, 2015]. ChemCam has observed chlorine and high sodium that may be interpreted as possible halite or sodium perchlorate in the same vicinity as boron-bearing vein targets [Thomas et al., 2017]. This model may be tested by searching for evaporate deposits as the rover continues its traverse up the Mt. Sharp stratigraphy.

Based on the chemical and stratigraphic evidence presented above, we find the second scenario the more plausible of the two models for B emplacement into veins (Figure 3). The initial deposition of B in evaporites required drying the lake, lowering the water table, remobilizing soluble elements (primarily Ca, Mg, and Na), and precipitating them in evaporite veins lower in the stratigraphy (Figure 3). Deep lake water conditions prevented borate adsorption onto phyllosilicates or amorphous materials (Figure 3a), thus leading to formation of boron-bearing evaporites as Gale lake dried (Figure 3b). Gale then became a predominately closed-drainage basin, forming a perennial lake system; following terrestrial models, layers of mudstone and evaporites would alternate in the bedrock (Figure 3c) [Eugster and Hardie, 1978]. In this perennial lake stage, the shallow lake would have higher borate concentrations and borates could then be adsorbed by lakebed clay and amorphous materials. Later, Gale underwent multiple episodes of eolian burial and exhumation [Grotzinger et al., 2015; Banham et al., 2017; Frydenvag et al., 2017] and burial overpressure plus groundwater intrusion caused fractures to form within the bedrock [Watkins et al., 2017]. Groundwater events then dissolved some of the evaporite deposits (Figure 3d), and the observed evaporite minerals (Ca sulfates) were then coprecipitated with borates in veins throughout the lower stratigraphic layers. The groundwater was enriched in borate from dissolution of the evaporites, implying that the water was alkaline, and borate would be adsorbed to host-rock clay and amorphous components as the boron-rich groundwater circulated through lower stratigraphic layers. The Murray unit then experienced erosion and exhumation before the deposition of the overlying aeolian Stimson sandstone, which itself became lithified and fractured [Grotzinger et al., 2015; Frydenvag et al., 2017; Banham et al., 2017; Watkins et al., 2017]. Fractures in Stimson then also became filled with calcium sulfate and borate (Figure 3e). The groundwater system likely shut down sometime after the deposition of sulfate into the fractures (Figure 3e) but before the erosion of Gale sediments to its present surface (Figure 3f).

ChemCam cannot directly measure mineralogy, but the possible mineralogies of borate-bearing materials in veins may be inferred for a Ca-rich, 0–60°C, and alkaline fluid (for other conditions, see supporting information). The borate anion may isomorphically replace sulfate in the Ca-sulfate minerals in the veins during burial [e.g., Orti and Alonso, 2000], or borates are present as their own phase that coprecipitated with Ca sulfates as Ca borates and Ca sulfate stability field conditions significantly overlap [Crowley, 1996].
4.2. Implications for Astrobiology

The presence of boron on Mars opens up new possibilities for habitability because of the important role borate may have played in prebiotic chemistry on early Earth [Scorei, 2012]. Borates stabilize ribose, the simple sugar that forms the backbone of RNA with phosphate [Ricardo et al., 2004; Furukawa et al., 2013]. Without borates, ribose quickly decomposes in water [Larralde et al., 1995]. Borates may thus have been a necessary bridge from abiotically produced organic molecules to RNA-based protolife on Earth [Scorei, 2012]. Although other methods are proposed for producing RNA [Ruiz-Mirazo et al., 2014], this work nevertheless shows that borate is available in an ancient planetary setting. On Mars, we have shown borate was present in a long-lived hydrologic system, suggesting that important prebiotic chemical reactions could plausibly have occurred in the groundwater, if organics were also available. Thus, the discovery of boron in Gale crater opens up intriguing questions about whether life could have arisen on Mars.

The timing of the initial concentration of borates in sediments provides key bounds for when life could have formed on Mars. It is unknown if borate evaporite minerals were present on the Archean Earth [Grew et al., 2011]. Borates stabilize ribose, the simple sugar that forms the backbone of RNA with phosphate [Ricardo et al., 2004; Furukawa et al., 2013]. Without borates, ribose quickly decomposes in water [Larralde et al., 1995]. Borates may thus have been a necessary bridge from abiotically produced organic molecules to RNA-based protolife on Earth [Scorei, 2012]. Although other methods are proposed for producing RNA [Ruiz-Mirazo et al., 2014], this work nevertheless shows that borate is available in an ancient planetary setting. On Mars, we have shown borate was present in a long-lived hydrologic system, suggesting that important prebiotic chemical reactions could plausibly have occurred in the groundwater, if organics were also available. Thus, the discovery of boron in Gale crater opens up intriguing questions about whether life could have arisen on Mars.

The timing of the initial concentration of borates in sediments provides key bounds for when life could have formed on Mars. It is unknown if borate evaporite minerals were present on the Archean Earth [Grew et al., 2011], but the borate evaporites in Gale are much older than the oldest inferred age of evaporites on Earth [Grew et al., 2011]. Additionally, there is evidence for ancient, evolved igneous rocks from the Southern Highlands in the Gale lake watershed [e.g., Cousin et al., 2017], which are required precursor rocks for boron enrichment [Grew et al., 2011]. Hence, the presence of boron enrichments in Gale suggests that boron-concentrating processes can occur early in a terrestrial planet’s history. If the primary deposition of borate was in evaporite layers, such as those that form in perennial lakes, borates would be both highly concentrated and in close proximity to clays and any organics, if present. Such wet and dry cycles occurring in perennial lakes are considered ideal for promoting prebiotic reactions on Earth [Lahav and Chang, 1976, 1982]. Additionally, evidence for boron in veins in the younger

**Figure 3.** An illustration of the hypothesized steps that lead from boron dissolved in Gale lake to the present day observation of boron in veins. Starting with a) boron dissolved in lake water, followed by b) evaporite deposition, c) groundwater infiltration, d–e) evaporite remobilization by groundwater, and redistribution of dissolved salts into fractures within the Murray mudstone and Stimson sandstone units, and ending with f) erosion to the present surface. White diagonal lines in d–f represent Ca-sulfate in fractures. Based on events described by Grotzinger et al. [2015] and Frydenvang et al. [2017].
Stimson sandstone suggests that the groundwater system in Gale crater was late-stage [Frydenvang et al., 2017]. And although the source of groundwater in Gale crater is as of yet unknown, it is clear that Gale has undergone multiple episodes of groundwater over its history with changes to its chemistry, pH, and Eh over time [Schwenzer et al., 2016; L’Harrard et al., 2017; Lanza et al., 2016; Frydenvang et al., 2017; Yin et al., 2017; Nachon et al., 2017]. This complex and changing groundwater system was not only habitable, providing geochemical and energy gradients needed to support life, but it would also have promoted prebiotic chemical reactions between borates and any organics present. If borate-organic molecules are directly observed in Gale, the early hydrologic system on Mars may shed light on how life arose on Earth while also opening up new possibilities for life on Mars. The discovery of boron on Mars makes the search for borates a high priority for Curiosity and future missions to Mars.

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