Evolved gas analyses of sedimentary rocks and eolian sediment in Gale Crater, Mars: Results of the Curiosity rover’s sample analysis at Mars instrument from Yellowknife Bay to the Namib Dune


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Abstract The sample analysis at Mars instrument evolved gas analyzer (SAM-EGA) has detected evolved water, H2, SO2, H2S, NO, CO2, CO, O2, and HCl from two eolian sediments and nine sedimentary rocks from Gale Crater, Mars. These evolved gas detections indicate nitrates, organic oxchlorine phase, and sulfates are widespread with phyllosilicates and carbonates occurring in select Gale Crater materials. Coevolved CO2 (160 ± 248–2373 ± 820 μgC(CO2)/g) and CO (11 ± 3–320 ± 130 μgC(CO)/g) suggest that organic C is present in Gale Crater materials. Five samples evolved CO2 at temperatures consistent with carbonate (0.32 ± 0.05–0.70 ± 0.1 wt % CO3). Evolved NO amounts to 0.002 ± 0.007–0.06 ± 0.03 wt % NO2. Evolution of O2 suggests that oxchlorine phases (chlorate/perchlorate) (0.05 ± 0.025–1.05 ± 0.44 wt % ClO4) are present, while SO2 evolution indicates the presence of crystalline and/or poorly crystalline Fe and Mg sulfate and possibly sulfide. Evolved H2O (0.9 ± 0.3–2.5 ± 1.6 wt % H2O) is consistent with the presence of adsorbed water, hydrated salts, interlayer/structural water from phyllosilicates, and possible inclusion water in mineral/amorphous phases. Evolved H2 and H2S suggest that reduced phases occur despite the presence of oxidized phases (nitrates, oxchlorine, sulfate, and carbonate). SAM results coupled with CheMin mineralogical and Alpha-Particle X-ray Spectrometer elemental analyses indicate that Gale Crater sedimentary rocks have experienced a complex authigenic/diagenetic history involving fluids with varying pH, redox, and salt composition. The inferred geochemical conditions were favorable for microbial habitability and if life ever existed, there was likely sufficient organic C to support a small microbial population.

1. Introduction

The Mars Science Laboratory (MSL) Curiosity rover and its payload of science instruments have traversed ~12 km for 1237 Martian solar days (sols) spanning more than 100 m of sedimentary stratigraphy. Along the way, Curiosity has evaluated the sedimentology, geomorphology, geochemistry, and mineralogy of regolith fines, eolian sediments, and igneous clasts in the sedimentary rocks of Gale Crater [e.g., Meslin et al., 2013; Minitti et al., 2013; Williams et al., 2013; McLennan et al., 2014; Ming et al., 2014; Schmidt et al., 2014; Vaniman et al., 2014; Cousin et al., 2015; Grotzinger et al., 2015] (Figures 1 and 2). Gale Crater is ~155 km wide with an ~5 km high layered central mound (Aeolis Mons; informally known as Mount Sharp) [Malin and Edgett, 2000; Grotzinger et al., 2015]. Orbiting infrared analyses suggest that the lower slopes (<1 km) of Mount Sharp consist of strata that contain variable amounts of hematite, phyllosilicates, and sulfate [Milliken et al., 2010]. The remaining upper several kilometers of Mount Sharp strata do not present infrared signatures that can be interpreted in terms of mineral components. Spectral evidence for minerals on upper Mount Sharp in some cases may be obscured by dust [Milliken et al., 2010]. Limitations in spatial...
coverage and resolution, dust cover, and the types and abundances of minerals that can be detected by a given orbital observing technique mean that there is an incomplete understanding of the compositional variations within Mount Sharp strata. Results from MSL investigations are intended to enhance our understanding of the composition of Mount Sharp and to search for records of past or evidence of present geochemical and environmental conditions that could have been suitable for microbial activity [Grotzinger et al., 2014, 2015].

Current estimates place Gale Crater formation at ~3.8 to 3.6 billion years ago with deposition of sedimentary material occurring until ~3.3 to 3.1 billion years ago [e.g., Thomson et al., 2011; Schwenzer et al., 2012; Grotzinger et al., 2015]. The sedimentary rock strata encountered along the rover’s traverse are interpreted to record interfingered fluvial, deltaic, and lacustrine environments. This indicates that Gale Crater had intermittent bodies of standing water that could have lasted from 100 to 10,000 years on the surface [Grotzinger et al., 2014, 2015].

The sample analysis at Mars (SAM) instrument has been essential in providing a fundamental understanding of the volatile-bearing phases including organic materials in Gale Crater sediments. The SAM analyses of the scooped eolian drift material (Rocknest-RN) and two drilled lacustrine mudstone samples of the Sheepbed member at the base of the Bradbury group (Figure 2; John Klein-JK and Cumberland-CB) have resulted in reporting the detection of evolved water, $\text{SO}_2$, $\text{H}_2\text{~S}$, $\text{CO}_2$, NO, $O_2$, HCl, and organics [Glavin et al., 2013; Leshin et al., 2013; Archer et al., 2014; Ming et al., 2014; McAdam et al., 2014; Freissinet et al., 2015; Stern et al., 2015, 2017]. SAM results demonstrated that water was likely adsorbed to particle surfaces and also associated with amorphous phases, phyllosilicates, and hydrated salts [Leshin et al., 2013; Ming et al., 2014].

**Figure 1.** Regional geologic map, on Mars Reconnaissance Orbiter (MRO) High Resolution Imaging Experiment (HiRISE) camera image mosaic base map, showing the Curiosity rover traverse (white line) and sampling sites. (a) Gale Crater topography; the white box outlines the area shown in Figure 1b. (b) Geologic map showing the principal stratigraphic units exposed at Aeolis Palus and the foothills of Aeolus Mons (Mount Sharp) and the sample sites (red dots) with individual sample abbreviations (in parentheses) along the rover traverse where SAM analyses were conducted through sol 1237. (c) Topographic profile of the rover traverse.
Iron sulfates, amorphous S phases, and possible oxidation of iron sulfides are likely sources of SO₂ [McAdam et al., 2014]. Oxygen and HCl releases are consistent with the presence of perchlorate and chlorate phases [Glavin et al., 2013; Ming et al., 2014; Sutter et al., 2016]. Organics, carbonates, and along with some contributions from instrument background are possible sources of evolved CO₂ [Glavin et al., 2013; Leshin et al., 2013; Ming et al., 2014; Freissinet et al., 2015]. Another significant discovery was the first in situ detection of nitrate on Mars in the Gale Crater sediments [Stern et al., 2015]. Nitrate has been detected in Martian meteorites [Kounaves et al., 2014]; however, the SAM detection confirms the presence of nitrate on Mars. This discovery in sedimentary rocks and eolian sediments indicates nitrogen (N) was present for potential microbial activity on Mars.

Results from the SAM instrument have indicated that Gale Crater sediments consist of oxidized (sulfate, nitrate, and oxychlorine) and reduced (sulphides and organics) phases along with minerals that formed in acidic (e.g., Fe sulfate) and alkaline (e.g., phyllosilicate and carbonate) conditions [e.g., Leshin et al., 2013; McAdam et al., 2014; McLennan et al., 2014; Ming et al., 2014; Rampe et al., 2017]. The CheMin X-ray diffraction instrument has verified the SAM detections of phyllosilicates, iron sulfides, and iron sulfate (e.g., jarosite) [Vaniman et al., 2014; Hurowitz et al., 2017; Rampe et al., 2017]. These results indicate that sedimentary rocks in Gale Crater are composed of complex mineral mixtures that reflect varying geochemical conditions at likely different times, represented by mineral inventories at different stratigraphic levels. These mineral mixtures suggest that past geochemical conditions may have been favorable for microbial life [e.g., Grotzinger et al., 2014]. Some mineral assemblages are out of equilibrium, for example, at the base of Murray formation at Pahrump hills where iron sulfate minerals coexist with phyllosilicate minerals [Hurowitz et al., 2017; Rampe et al., 2017], which demonstrates that these samples may preserve evidence of multiple aqueous episodes.

The goal of this work is to provide a comprehensive summary of all major evolved gases detected in samples analyzed by the SAM evolved gas analyzer (SAM-EGA). The SAM-EGA data of RN, JK, and CB samples have been previously reported [e.g., Leshin et al., 2013; Ming et al., 2014] and are included here with the eight additional sampling sites to provide a complete summary of results. The objectives of this work are to (1) provide a comprehensive characterization of the evolved gas (H₂O, SO₂, H₂S, CO₂, CO, NO, O₂, HCl, and H₂) release from samples analyzed by SAM since landing (sol 0) to the analysis of eolian dune material at Gobabeb (GB) (sol 1237), (2) place constraints on the mineralogy or composition of volatile-bearing phases based on

Figure 2. Stratigraphic column describing the sedimentary facies from Yellowknife Bay to Stimson formation. The Bradbury (Aeolis Palus) group and the Murray formation contact is composed of inter fingering facies. Stimson formation unconformably overlies Murray formation. The Rocknest (RN) and Gobabeb (GB) are loose, unconsolidated modern eolian sediments that overlie the Yellowknife Bay and Murray formations, respectively. See Grotzinger et al. [2015] for detailed stratigraphic discussion.
2. Materials and Methods

2.1. Sample Analysis at Mars (SAM) Instrument

The sample analysis at Mars (SAM) instrument analyzed evolved gases from Gale sediments with three instruments: the quadrupole mass spectrometer (MS), gas chromatograph (GC), and tunable laser spectrometer (TLS) [Mahaffy et al., 2012]. Before sample acquisition, a SAM quartz sample cup was preconditioned to 870°C (5 min), and gas transfer lines and manifolds were preconditioned to 135°C (150 min) in order to minimize background contributions to the SAM analysis. Single or triple portions of scooped or drilled materials were transferred from Curiosity’s sample processing system into quartz sample cups which are then sealed in a SAM oven. Samples were heated (35°C min⁻¹) to ~870°C where helium carrier gas flows (~0.8 cm³ min⁻¹ STP; 25 mbar) up through the porous quartz frit of the cup and flushes the evolved volatiles from the sample to the MS. As the sample was heated, gases (e.g., H₂O, CO₂, O₂, and SO₂) evolve from volatile-bearing phases (e.g., minerals and organics) at characteristic temperatures were used to identify the presence of organic or particular mineral phases. The SAM plumbing is configured to allow the MS to sample a portion of evolved gases over the entire sample heating range. After the evolved gas were sample by the MS, gases flowed downstream and were then either sampled by the hydrocarbon trap for GCMS analysis or the TLS or exhausted to the Martian atmosphere.

Hydrocarbon detections using GCMS and isotopic analysis of water and CO₂ using the TLS are discussed elsewhere [Glavin et al., 2013; Leshin et al., 2013; Ming et al., 2014; Freissinet et al., 2015; Mahaffy et al., 2015]. This work focuses on evaluation of H₂O (m/z 20), H₂ (m/z 2), SO₂ (m/z 64), H₂S (m/z 34) CO₂ (m/z 44), CO (m/z 28), NO (m/z 30), O₂ (m/z 32), and HCl (m/z 36) evolved gases as analyzed by only the direct MS pathway and is referred to as the SAM-EGA mode. The isotopologue of water, m/z 20, is presented instead of the m/z 18 signal, which was saturated on every analysis because of the extreme sensitivity of the MS. Procedures and methods utilized for quantifying total evolved gas releases (e.g., μmol or nmol) and their associated errors are described in detail elsewhere [Archer et al., 2014]. Evolved gas totals were then used with estimated sample mass deliveries to SAM to calculate evolved gas contents (e.g., wt % or μg/g). Evolved gas content errors are based on propagation of errors derived from sample mass delivery and evolved gas totals.

2.2. SAM-EGA Signal Processing

All evolved gas plots were deadtime and background corrected, while many of the evolved gas plots were further processed to account for MS signal saturation and/or the presence of isotopologue fragments created in the MS. Deadtime correction refers to a process that accounts for lost counts during high count rates. Detailed description of deadtime correction procedure was outlined by Franz et al. [2014]. The SAM-EGA analysis of calcite (CaCO₃) and melanterite (FeSO₄·7H₂O) standards using the SAM instrument before integration into the rover allowed for correction factors to be derived that were applied to m/z 28, m/z 32, m/z 34, m/z 44, and m/z 64. These two standards in several cases were used as proxies for the isotopic ratios in the signal-saturated Mars samples that were attributed to CO₂ and SO₂ evolving phases that differed from the CO₂ and SO₂ evolving standard phases. This could provide inaccurate isotopic measurements which is not necessarily critical when correcting figures for presentation purposes but is important when reporting evolved gas abundances discussed in this work. Derivation of error associated evolved gas abundance determinations as a result of using these two standards is beyond the scope of this work and can be found elsewhere [Archer et al., 2014].

Many samples experienced signal loss of the most intense portions of the m/z 44 (CO₂) and m/z 64 (SO₂) peaks as a result of signal saturation of the MS detector. The missing portions of these peaks were replaced by corrected isotopologue m/z 45 (CO₂) and m/z 66 (SO₂) plots, respectively. The missing portions of the m/z 44 signal was replaced by a correction to the m/z 45 plot from the Gale sample based on m/z 44/m/z 45 ratio calculated from the calcite standard. Subscripts c and uc indicate corrected and uncorrected...
SAM-EGA plots of Gale samples, respectively, and mineral subscript refers to mineral standard analyzed in SAM before flight.

\[ m/z 44_c = m/z 45_{\text{Gale}} \times (m/z 44 / m/z 45)^{\text{calcite}}. \]

Similar to m/z 44, when m/z 64 was saturated, the missing portion of the m/z 64 peak was derived from m/z 66:

\[ m/z 64_c = m/z 66_{\text{Gale}} \times (m/z 64 / m/z 66)^{\text{melanterite}}. \]

Signal loss was not always complete for m/z 44 or m/z 64 plots, and only the missing (i.e., saturated signal) portions of the m/z 44 or m/z 64 peaks were replaced with corrections from m/z 45 and m/z 66, respectively.

The m/z 28 signal was composed of evolved CO from the sample and the CO isotopologue fragment of CO₂ created in the MS during analyses. The CO isotopologue fragment contribution was based on m/z 28//m/z 44 ratio calculated from calcite standard run in SAM where no CO producing phase was present. The CO isotopologue fragment contribution was subtracted from the uncorrected m/s 28uc plot to obtain the corrected m/s 28c plot:

\[ m/z 28_c = m/z 28_{\text{uc}} - m/z 44_c \times (m/z 28 / m/z 44)^{\text{calcite}}. \]

All low-temperature (<500°C) m/z 32 was assumed to be derived from oxychlorine thermal decomposition as there were no significant SO₂ (m/z 64) detections at those temperatures. All m/z 32 detections above 500°C had contributions from SO₂ (32S) that were subtracted out using m/z 32//m/z 64 ratio determined from the melanterite standard.

\[ m/z 32_c = m/z 32_{\text{uc}} - m/z 64_c \times (m/z 32 / m/z 64)^{\text{melanterite}}. \]

The O₂ (18O16O) and SO₂ (34S) isotopologue fragments must be subtracted from m/z 34 to acquire m/z 34 due to H₂S. The presence of high concentration of oxychlorine phase indicated that O₂ was contributing to all low-temperature (<500°C) m/z 34 and that if H₂S was present below 500°C it would be difficult to distinguish it from 18O16O from oxychlorine. The m/z 34 peak that coincided with O₂ below 500°C in the Gale sample was used to calculate m/z 34//m/z 32 ratio of the Gale sample. Contributions to m/z 34 from SO₂ (34S) were also subtracted from the m/z 34 plot using the m/z 34//m/z 64 ratio calculated from the melanterite standard.

\[ m/z 34_c = m/z 34_{\text{uc}} - \left[ m/z 32_c \times (m/z 34 / m/z 32)^{\text{Gale}} \right] - \left[ m/z 64_c \times (m/z 34 / m/z 64)^{\text{melanterite}} \right]. \]

### 2.3. MTBSTFA Contributions to CO₂ and NO Detections

The derivatization agent, N-methyl-N-(tert-butyldimethylsilyl)trifluoroacetamide (MTBSTFA), that was brought for wet chemistry SAM-GCMS analysis was determined to have leaked from one of the sealed cups in SAM and likely contributed to SAM-EGA detections of CO₂ and NO [Glavin et al., 2013; Freissinet et al., 2015; Stern et al., 2015]. The worst-case scenario contributions of MTBSTFA-C to SAM-CO₂ detections were proposed to be 900 nmol C for each SAM-EGA analysis [Glavin et al., 2013]. The detection of MTBSTFA breakdown byproducts used to estimate the level of MTBSTFA that leaked into SAM indicates that MTBSTFA did not undergo complete combustion to CO₂ and thus could cause this estimate to be lower [Glavin et al., 2013]. Nevertheless, 900 nmol CO₂-C from MTBSTFA for each analysis whether it was a single or triple portion was used as worst-case scenario when discussing CO₂-C contents. The impact this level of MTBSTFA-CO₂ has on total CO₂ contents will be discussed in section 4.6.

The MTBSTFA contribution to the NO detected by SAM-EGA was determined by the background correction method by Stern et al. [2015]. Briefly, MTBSTFA decomposes into nitrogen-free ion fragments that were detected by SAM-EGA which include tert-butyldimethylsilanol (monosilylated H₂O, MSW) (m/z 75), 1,3-bis(1,1-dimethylethyl)-1,3,3-tetramethyldisiloxane (bisilylated H₂O, BSW) (m/z 147), tert-butyldimethylfluorosilane (TBDMS-F) (m/z 134), 2-methylpropene (C₄H₈) (m/z 41), and a strong contribution at m/z 15, either CH₄ or methane ions ([Stern et al., 2015], supporting information). The detection of these fragment masses in the sample of interest indicates how much MTBSTFA decomposed. Each MTBSTFA
molecule has 1 nitrogen, and thus, the amount of nitrogen presumably evolved as NO when MTBSTFA decomposed can be calculated as follows:

\[
\text{MTBSTFA - nitrogen} = 2x\text{BSW} + \text{MSW} + \text{TBDMS} - F + C_4H_8 + 1/5xCH_4
\]

Corrected - nitrogen = Total nitrogen detected by SAM - EGA - MTBSTFA - nitrogen

2.4. O\textsubscript{2} Corrections From Nitrate Decomposition.

Evolved oxygen used to calculate the amount of oxychlorine present must account for contributions from nitrate thermal decomposition which also evolved O\textsubscript{2} [e.g., Ettarh and Galwey, 1996]. For example,

\[
2\text{Ca(NO}_3\text{)}_2 \rightarrow 2\text{CaO} + 4\text{NO} + 3\text{O}_2.
\]

Oxygen contribution from nitrate was calculated by multiplying MTBSTFA-corrected nitrate amount by 0.75 and subsequently subtracting this nitrate oxygen from total oxygen evolved at temperatures consistent with oxychlorine thermal decomposition. Evolved oxygen above 600°C from sulfate decomposition were not corrected for nitrate oxygen as these temperatures were too high to have contributions from nitrate thermal decomposition. The relatively low amounts of Gale nitrate resulted in between 1 and 8% of the evolved O\textsubscript{2} to be attributed to nitrate thermal decomposition.

2.5. Sampling Procedures

Eleven sampling sites as of sol 1237 (29 January 2016) have been examined by SAM. One sample was scooped from an eolian drift deposit dubbed Rocknest (RN), while two eolian dune samples of differing particle size distributions were scooped from Gobabeb (GB1 and GB2) (Figures 3 and 4). The other nine samples were acquired from sedimentary rocks using a drill and are listed in order of acquisition: John Klein (JK), Cumberland (CB), Windjana (WJ), Confidence Hills (CH), Mojave 2 (MJ), Telegraph Peak (TP), Buckskin (BK), Big Sky (BS), and Greenhorn (GH) (Figures 5–9). The RN, JK, CB, WJ, and CH samples included a blank
Figure 4. (a) Curiosity rover, for scale, at the Gobabeb sample extraction site on the north side of the active eolian Namib Dune sand dune in the Bagnold dune field. This is a radiometrically corrected, color-adjusted mosaic of MAHLI images acquired on sol 1228. White arrow points to location of scoop 1 (NASA/JPL-Caltech/MSSS product PIA20316). (b) Gobabeb Scoop 1 site before the sample was extracted. Illuminated by sunlight from the left, an eolian sand ripple crest runs up the center of this figure, a portion of MAHLI focus merge product 1224MH0001700000403204R00 created on board the instrument from eight focus stack images acquired on sol 1223.

Figure 5. (a) The John Klein (JK) and Cumberland (CB) sampling sites in the Sheepbed member of Yellowknife Bay formation with view to the southwest (Mastcam-34 white balanced mosaic mosaic from sequence mcam00818, sol 137). JK and CB are ~2.75 m apart. (b) The JK sample extraction drill hole (center) created on sol 182 with shallower hole (left) that was drilled on sol 180 to interrogate the rock properties before commitment to drill a full-depth hole (Portion of MAHLI image 0182MH0002300010102629C00) (c) The CB sample extraction drill hole created on sol 279 (Portion of MAHLI image 0279MH0002680020103055C00). Drill holes are ~1.6 cm wide.
SAM-EGA analysis of an empty cup prior to the sample run. Only the H$_2$ blank analysis for RN is not reported as this mass (m/z 2) nor its isotopologue (m/z 3) was analyzed by SAM. The GH blank analysis was conducted between the GH1 and GH2 analyses. The cup used for each blank analysis was always used for the subsequent sample analysis by SAM. All drilled samples were acquired with a drill that produced a 1.6 cm diameter by ~6 to 6.5 cm deep hole. The drill design causes the top 1.5 cm of drilled material to not be included in the sample and is instead extruded as cuttings onto the adjacent rock surface [Anderson et al., 2012]. The extruded cuttings are referred to as drill tailings. All samples except one were passed through a 150 μm sieve in the rover’s sample processing and handling system, which has the capability of delivering small sample portions to the analytical instruments [Anderson et al., 2012]. The exception is the GB2 sample which consisted of a scooped 150 μm to 1 mm fraction. The scooped RN sample was a single portion amounting to 50 mg ± 8 (2σ) whereas the drilled single portion samples of JK, CB, CH, MJ, and TP amounted to 45 mg ± 18 (2σ). Triple portions [135 mg ± 31(2σ)] of drilled WJ, BK, BS, GH, and scooped GB1 were delivered to enhance organic detection. The amount of scooped coarser grained (150 μm to 1 mm) GB2 sample delivered to SAM was unknown because sample delivery models used to calculate delivery amount were based on finer grained components (<150 μm) and sample delivery models for the coarse-grained components have yet to be developed.

Eight samples were only analyzed once but four samples (RN, JK, CB, and GH) analyzed multiple times to collect TLS and GCMS data over different temperature ranges and expose samples to prerun heat treatments to minimize low-temperature background organic detections (Table 1) [e.g., Leshin et al., 2013; Freissinet et al., 2015]. The SAM EGA results from all the RN subsamples are similar, and thus, the RN subsample 4 profile was presented for brevity. Four and seven subsamples of the JK and CB drilled material,
respectively, were analyzed by SAM. Despite preconditioning all sample cups before analyses to 870°C, select JK and CB subsamples underwent additional heat treatments to minimize organic background levels \cite{Ming2014,Freissinet2015}. The JK1, JK2, and JK3 subsamples underwent heat treatments to 320°C. The CB5, CB6, and CB7 empty cups were heated to 250°C allowed to cool. The CB6 and CB7 subsamples were loaded into the cups and heated again to 250°C for 27 min for before continuing the heating to 870°C (Table 1). The CB6 and CB7 subsamples were not presented because heating to these temperatures were shown to affect SAM H₂O, CO₂, CO, and O₂ low-temperature evolutions. The CB1, CB2, and CB3 were analyzed similarly without cup reheating or sample preheating. Several CB1 evolved gas abundances were much less than CB2, CB3, and CB5 which may have been attributed to less sample being dumped into the cup relative to the other subsamples; thus this subsample will not be discussed. The evolved gas release profiles for CB2, CB3, and CB5 are similar \cite{Ming2014}.

Figure 7. (a) The Pahrump Hills member sampling site for Confidence Hills (CH), Mojave 2 (MJ), and Telegraph Peak (TP). MJ is 10 m from and 1 m above CH, while TP is ~6 m above and ~39 m from MJ \cite{Majet135}. (b) The CH drill hole (center), created on on sol 759, and preceding shallow rock properties interrogation hole (top left), drilled on sol 756; \cite{MAHLI image 0759MH000397001030030C00, sol 759}. (c) Brushed MJ material showing crystal forms of lenticular morphology of about 1–2 mm length (MAHLI focus merge product 0880MH0004620000302350R00, sol 880). (d) The MJ drill hole created on sol 882 (MAHLI-0882MH0003970010302481C00, sol 882). (e) The TP drill hole created on sol 908 (MAHLI-0908MH0003970010302871C00, sol 908). Drill hole diameters are ~1.6 cm.
et al., 2014]; thus, only CB2 profiles are presented. JK4 is presented in this work as it is the only JK subsample to not have undergone any preheating before SAM-EGA analysis. Two GH subsamples were analyzed because the evolved gases from overlapping temperature ranges were examined by TLS and GCMS in GH1 and GH2, respectively. Furthermore, the GH1 subsample was deposited over the residue of the previously analyzed BS sample in the same cup, while the GH2 subsample was placed in an empty cup. The GH1 and GH2 have similar profile releases for most gases though there are some differences in release temperatures between the two samples. The reasons for the differences in temperature of gas releases between the two samples is not clear and may be attributed to GH1 and BS residue sample interaction. The GH2 analysis is presented to maintain consistency with other samples that were analyzed in unused cups.

2.6. Alpha-Particle X-Ray Spectrometer Sample Analysis

Aliquots of all samples analyzed by SAM were also analyzed by Alpha-Particle X-ray Spectrometer (APXS) for total chemistry; however, only total sulfur and total chlorine are presented, because these are the only elements that can be directly compared to SAM data, namely, evolved SO$_2$ and HCl, respectively. Details of the APXS method, the instrument, and operation are explained elsewhere [Gellert et al., 2006, 2015; Campbell et al., 2012; Schmidt et al., 2014; Berger et al., 2016; Thompson et al., 2016a]. Briefly, the scooped or drilled material was acquired by the rover’s sample acquisition, processing, and handling (SA/SPaH) system which then passed portions of the sample through a sieve to CheMin and SAM for analysis [Anderson et al., 2012]. The remaining sieved material of each sample was then dumped onto the Martian surface at a later date and subsequently analyzed by APXS. The sieved material was dumped in an amount that exceeded the APXS field of view which avoided analyzing chemistry of the underlying material. This “postsieve” sample APXS analysis was then compared to the SAM data (supporting

Figure 8. (a) The Buckskin sampling site of the Murray formation in the Marias Pass region. Red arrow marks approximate BK acquisition location behind rock. Sampling location to plateau top at top left is ~1 m (Portion of a Mastcam-34 white balanced mosaic from sequence mcam04375, sol 990). (b) The geological surroundings of the BK drill hole (Portion of Mastcam-34 white balanced image 1062ML0046720020306187E01, sol 1062). (c) The BK drill hole (Portion of MAHLI image-1060MH0003970010400402C00, sol 1060). Drill hole diameter is ~1.6 cm.
information Tables S3 and S9). Mars Hand Lens Imager (MAHLI) images of the postsieve dump sample were acquired to verify adequate coverage of underlying surface. Furthermore, APXS analyses of sample drill tailings not acquired by the SA/SPaH and presieve sample acquired by SA/SPaH were used to assess if the postsieve APXS analysis was successfully taken and if it represented the best APXS data set for comparison with SAM.

The APXS determines the composition of the sample using X-ray spectroscopy by excitation with high energetic alpha-particles and X-rays from the internal 244Cm sources. The method resembles a combination of the terrestrial standard methods of particle induced X-ray emission and X-ray fluorescence. Sulfur and chlorine are unambiguously identified by their characteristic X-ray energy and quantified with small statistical uncertainty down to a few 1000 ppm abundance. The signal for these two elements stems from about 5 to 10 μm within the sample. The Cl and S accuracy determined during calibration was 30% and 15% absolute APXS error, respectively. The underlying reason for the accuracy errors is that the APXS calibration assumes a homogeneous sample for the correction of self-absorption within the sample. The Cl and S unfortunately occur within heterogeneous natural samples that can cause absorption effects of the S and Cl X-ray signal which can skew abundance determinations.

2.7. Image Processing for Figures

The images (Figures 1 and 3–9) used in this report to document Curiosity’s sample extraction sites were acquired by the Mars Reconnaissance Orbiter (MRO) High Resolution Imaging Experiment (HiRISE) camera [McEwen et al., 2007], the MSL Mast Cameras (Mastcams) [Malin et al., 2010], the MSL robotic arm-mounted Mars Hand Lens Imager (MAHLI) [Edgett et al., 2012], and the MSL front Hazard cameras (Hazcams) [Maki et al., 2012]. The figures include cropped portions of singular Mastcam, MAHLI, and Hazcam images; cropped portions of mosaics consist of multiple Mastcam, MAHLI, and HiRISE images; and MAHLI focus merge products created on board the instrument. The images have been processed to enhance the visibility of the key illustration details via broadening image dynamic range and contrast sharpening. Some of the images and mosaics, as indicated in figure captions, have been radiometrically corrected and white balanced. White balanced color shows what the geologic materials would look like if they were in sunlight on Earth; this processing can sometimes cause the sky to appear blue or white. Other images retain their natural color as they appear under Martian illumination conditions.
3. Geologic Context

3.1. Eolian Sediment Samples Analyzed by SAM

3.1.1. Rocknest

The Curiosity rover landed in northern Gale Crater on Aeolis Palus on an exposure of the Bradbury group fluviatile conglomerate rocks which are the oldest strata exposed in this part of Gale Crater (Figure 1) [Williams et al., 2013; Vasavada et al., 2014]. The Bradbury group underlies the Mount Sharp group and is exposed in the crater plains along the north side of Mount Sharp (Figure 2). The Curiosity rover drove 400 m east-southeast of the landing site to the first SAM sampling stop at the RN eolian drift material (4518 m elevation). Sediments from the RN eolian drift deposit were sampled in five different locations within the deposit (Figure 3) but only scoop five sampled on sol 93 was examined by SAM. The Rocknest material consists of unconsolidated sand and dust [Blake et al., 2013; Minitti et al., 2013] that overlies rock of the Glenelg member of the Yellowknife Bay formation (Figure 3). The RN material likely consists of a mixture of local, regional, and global sources that yield a bulk basaltic chemistry that is similar to eolian material at other landing sites [Blake et al., 2013; Minitti et al., 2013].

The bulk Rocknest sediment possesses a basaltic mineralogy consisting of plagioclase feldspar, olivine, pyroxenes, ilmenite and magnetite and a secondary mineralogy that includes calcium sulfates and hematite [Bish et al., 2013; Blake et al., 2013]. A broad hump in the X-ray diffraction pattern is consistent with the

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aBold samples are presented in this work. See text for descriptions of sample and cup pretreatment procedures.
presence of X-ray amorphous material that may consist of basaltic glass, allophane (e.g., hisingerite) poorly crystalline sulfate, and/or nanophasic Fe oxides (e.g., ferrithiodyrite) [Bish et al., 2013].

3.1.2. Gobabeb 1 and 2

The Gobabeb sand samples were scooped from the lower stoss slope of a dune informally named Namib Dune which is part of the active informally named Bagnold Dune Field (Figure 4) [Silvestro et al., 2013]. The Gobabeb sediment unconformably overlies Murray formation mudstone (~4424 m). Examination of the sieved <150 μm fraction of the sediment indicates the GB1 samples contains grains that are mostly resolvable by MAHLI (Figure 4b). The <150 μm RN material by comparison contains at least 40% material that is unresolvable by MAHLI [Minitti et al., 2013]. The overall total chemistry of the GB samples is similar to RN except that the GB has a lower S/Cl ratio than RN [Ehlmann et al., 2017]. The overall mineralogical abundance of GB is similar to RN and consists of plagioclase feldspar, olivine, augite, pigeonite, and magnetite [Achilles et al., 2017].

3.2. Bedrock Samples Analyzed by SAM

3.2.1. John Klein and Cumberland (Bradbury Group-Yellowknife Bay Formation-Sheepbed Member-Mudstone)

The JK and CB sampling sites in the Sheepbed mudstone occur near the topographically and stratigraphically lowest point along the MSL rover traverse (~4520 m) [Grotzinger et al., 2014, 2015] and are located ~444 m east of the Bradbury landing site and ~100 m northeast of RN (Figures 1, 2, and 5). The JK and CB samples were drilled (sols 180 and 279, respectively) and extracted from the Sheepbed lacustrine mudstone. The two samples are separated by 2.75 m and 10 cm, horizontally and vertically, respectively [Ming et al., 2014] (Figure 5a).

The JK and CB materials possess a basaltic mineralogy similar to RN along with calcium sulfate, iron oxide/hydroxides, and high X-ray amorphous content (28–31 wt %) that consists mostly of Si, Fe, Ca, and S [Vaniman et al., 2014]. The JK drill hole wall showed light-toned millimeter-sized veins that were high in Ca, S, and H, which was determined to be consistent with bassanite [Rapin et al., 2016]. The CB drill hole wall had far fewer veins than JK [Vaniman et al., 2014]. Unlike RN, iron sulfides and smectite (saponite) were detected in these samples. The presence of smectite and magnetite along with the low abundance of olivine in JK and CB relative to RN suggests that these secondary phases formed under near neutral pH conditions in the Sheepbed mudstone [McLennan et al., 2014; Vaniman et al., 2014; Bristow et al., 2015]. Both of these drilled materials are grey in color highlighting the reduced nature of these materials (Figures 5b and 5c) [Ming et al., 2014].

3.2.2. Windjana (Bradbury Group-Kimberley Formation-Dillinger Member-Sandstone)

The Windjana (WJ) drilled sample was extracted from a sandstone outcrop that occurs ~4 km southwest and ~40 m stratigraphically higher than the Sheepbed mudstone samples (CB and JK) (Figures 1 and 2). The WJ sample was acquired (sol 621) from the Dillinger member of the Kimberley formation that is dominated by fluvial conglomerate and deltaic sandstone (Figure 6) [Treiman et al., 2016]. The Windjana material consists of a crossstratified sandstone of likely eolian origin that reflects reworking of fluvial sand [Grotzinger et al., 2015; Treiman et al., 2016]. The WJ sample has a potassium-rich basaltic mineralogy as indicated by the high concentration of K-feldspar (~22 wt %) [Thompson et al., 2016a; Treiman et al., 2016]. This potassium-rich feldspar mixed with basaltic minerals indicates that the WJ material is derived from multiple parent rock sources. The Windjana sample also possesses high pyroxene and magnetite contents along with a 2:1 layer type phyllosilicate and X-ray amorphous material [Downs and MSL Science Team, 2015; Treiman et al., 2016]. Postdepositional diageneric processes are likely responsible for cementing the WJ with magnetite, 2:1 layer type phyllosilicate, and Fe-rich X-ray amorphous material of ferrithiodyrite composition [Treiman et al., 2016].

3.2.3. Confidence Hills, Mojave, and Telegraph Peak (Mount Sharp Group-Murray Formation-Pahrump Hills Outcrop-Mudstone)

The CH, MJ, and TP materials were the first drilled (sols 759, 881, 908, respectively) samples of the Murray formation (basal layer of the Mount Sharp group) (Figure 7). The Murray formation forms an interfingering contact with the Bradbury group (Figure 2), and the samples are located ~2.2 km to the southwest of WJ and ~20 m higher stratigraphically [Stack et al., 2015; Grotzinger et al., 2015] (Figures 1 and 2). The Murray formation is dominated by finely laminated mudstones interpreted to have been deposited in a lacustrine environment in the proximity of a fluvial-deltaic depositional setting [Grotzinger et al., 2015]. The CH site (~4461 m) is approximately 1 m stratigraphically below and 10 m from MJ (~4459 m), while TP (~4453)
The Pahrump Hills mineralogy as determined by CheMin suggested a complex geochemical history where minerals stable under acidic (jarosite), alkaline (apatite), oxidizing (sulfate), and reducing (magnetite) conditions are present [Rampe et al., 2017]. Confidence Hills has the highest hematite concentration of all samples reported here, and this was demonstrated by CH having the reddest coloration of any of the drilled sediments (Figure 7b). The presence of jarosite and high hematite contents in CH and MJ suggests that these materials may have been exposed to acidic-oxidizing conditions after deposition. Greater magnetite contents in TP, however, suggested that more reducing conditions may have prevailed in TP. Furthermore, the occurrence of apatite in CH, MJ, and TP, which is not stable under acidic pH conditions, is difficult to reconcile suggesting again that Gale Crater sediments possess a complex geochemical history. TP also contained abundant cristobalite, marking the first detection of crystalline silica in the mission. The Pahrump Hills materials possessed a high X-ray amorphous concentration (up to 53 wt %). A 2:1 layer type phyllosilicate was also detected, but similar to WJ, the phyllosilicate concentrations were lower than what was detected in JK and CB [Rampe et al., 2017].

**3.2.5. Buckskin (Mount Sharp Group-Murray Formation-Marias Pass Outcrop-Mudstone)**

The BK sample was drilled (sol 1060) in the Murray formation about 6 m stratigraphically above and ~600 m to the southwest of TP, in the Marias Pass outcrop (Figures 1, 2, and 8). The BK sample came from a finely laminated mudstone [Morris et al., 2016]. The BK sample was drilled because it exhibited high Si (73.7 wt % SiO₂) and Ti (1.6 wt % TiO₂) contents [Morris et al., 2016]. The Buckskin mudstone contains abundant SiO₂-enriched X-ray amorphous material (~60 wt %) and has no detectable phyllosilicates [Morris et al., 2016]. Other phases present include tridymite (~14 wt %), a high-temperature SiO₂ polymorph, magnetite, minor cristobalite, and anhydrite [Morris et al., 2016]. Much of the silica polymorphs in the Marias Pass region are likely detrital from a silicic source region [Morris et al., 2016], although some of the amorphous silica could have resulted from post-depositional acidic weathering of the Buckskin materials, causing passive enrichment in Si and Ti [Morris et al., 2016; Rampe et al., 2017; Schmidt et al., 2016; Thompson et al., 2016b]. Alternatively, the amorphous Si may have been a result of postdepositional additions of Si in alkaline pH lake conditions [Hurowitz et al., 2017].

**3.2.5. Big Sky and Greenhorn (Siccar Point Group-Stimson Formation-Bridger Basin Outcrop-Sandstone)**

The Big Sky and Greenhorn samples were acquired (sols 1120 and 1137, respectively) from the dominant crossbedded eolian sandstone facies of the Stimson formation in the Bridger Basin outcrop area (Figures 1, 2, and 9). The Stimson formation lies unconformably over the Murray formation (Figure 2) [Fraeman et al., 2016]. The BS and GH sampling site are ~250 m to the south of and ~8 m higher stratigraphically than Buckskin. The dark gray Big Sky sample was assumed to be representative of typical unaltered Stimson eolian sandstone with a total chemistry similar to average Mars crustal composition [Yen et al., 2017]. The Big Sky mineralogy is consistent with a basaltic origin, but in contrast to Rocknest eolian sediments, the Big Sky sample has substantially more magnetite and no olivine [Yen et al., 2017]. The Greenhorn sample extraction site is ~1.9 m to the southwest of Big Sky and occurs in a lighter gray to white band ~75 cm wide associated with a fracture that crosscuts the otherwise dark gray crossbedded sandstone represented by Big Sky (Figure 9). The Greenhorn sample is interpreted to be chemically altered compared to BS as indicated by its lower Mg, Al, Fe, Mn, Ni, and Zn suggesting that Greenhorn has been leached by fluids that passed through the fracture and adjacent pores [Yen et al., 2017]. The Greenhorn sample has higher Si and S than Big Sky, indicating that it underwent a complicated alteration history involving loss and gain of constituents. Greenhorn consists of less feldspar and pyroxene and more anhydrite and X-ray amorphous material than Big Sky [Yen et al., 2017].

**4. Results/Discussion**

### 4.1. H₂O

All samples evolved water (0.9 ± 0.3 to 2.5 ± 1.6 wt %) during SAM-EGA analysis (Figure 10 and Table S1). Water evolved less than ~200°C could be the release of adsorbed H₂O [e.g., Tan et al., 1986]. Water
evolved between 100 and 300°C may have contributions from interlayer water [Borchardt, 1989] derived from smectite as in CB and JK [Vaniman et al., 2014], and 2:1 phyllosilicates in WJ, CH, and MJ [Treiman et al., 2016; Rampe et al., 2016]. Hydrated salts, such as bassanite in JK, CB, and WJ [Vaniman et al., 2014; Treiman et al., 2016] and oxychlorine salts, likely occur in all samples based on SAM observations (see below), and these could also contribute water released below 350°C. Dehydroxylation of small amounts of akaganeite detected by CheMin in JK, CB, and possibly WJ could also release some low-temperature H₂O [Ming et al., 2014; Vaniman et al., 2014; Treiman et al., 2016]. Finally, X-ray amorphous phases, detected by CheMin in all samples [Bish et al., 2013; Vaniman et al., 2014; Treiman et al., 2016; Achilles et al., 2017; Morris et al., 2016; Rampe et al., 2017; Yen et al., 2017], such as poorly crystalline aluminosilicate (e.g., allophane) or amorphous silicate (e.g., opal-A), and/or nanophase oxides/oxyhydroxides (e.g., ferrihydrite), could evolve H₂O in the <~450°C range.

Dehydroxylation may have yielded the evolved water above 450°C. The shoulders or peaks near 450–500°C in the CH, MJ, and TP traces can be attributed to dehydroxylation of H₂O from jarosite. The BK sample has a similar water feature at this temperature (Figure 10), but BK had no detectable jarosite in the CheMin data, suggesting either that jarosite is present but is below CheMin detection limits or the water is derived from some other source. Water evolved between 650 and 800°C from CB, JK, WJ, and MJ is interpreted to have been derived from the dehydroxylation of the phyllosilicates detected by CheMin [Ming et al., 2014; Vaniman et al., 2014; Treiman et al., 2016; Rampe et al., 2017].
The wide temperature water release \( > -500^\circ \text{C} \) in BK, BS, and GH was attributed to inclusion water in the X-ray amorphous phases detected in these materials. The X-ray diffraction amorphous features in BK and GH suggest a more silica rich amorphous fraction than other samples, which exhibits X-ray diffraction features more consistent with a basaltic glass [Bish et al., 2013; Vaniman et al., 2014; Treiman et al., 2016; Achilles et al., 2017; Morris et al., 2016; Rampe et al., 2017; Yen et al., 2017]. Inclusion water in basaltic and rhyolitic (Si-rich) glass phases can evolve above 500°C [Westrich, 1987; Pineau and Javoy, 1994; McAdam et al., 2016].

4.2. H₂

Hydrogen (H₂) was evolved over a wide temperature range from all samples (Figure 11). Rocknest, JK, CB, BS, GH, GB1, and GB2 evolved H₂ mostly above 400°C. Windjana through BK evolved most of their H₂ between \(-100^\circ \text{C}\) and \(-300\) to 700°C. GB1 also had a low contribution of H₂ in this temperature range. This low- to middle-temperature H₂ in WJ to BK is likely related to water fragmentation in the mass spectrometer because these H₂ releases tend to follow evolution of water (Figure 10a). This low-temperature H₂ attributed to water fragmentation in the mass spectrometer was also present in JK and CB. This low-temperature H₂ is not apparent in JK and CB because it is a minor release compared to the higher-temperature JK and CB H₂ releases in Figure 11a and is difficult to resolve due to the large offset in the y axis compared to the WJ and BK curves.

Possible sources of these H₂ releases are serpentinization-type reactions, inclusion H₂ (MJ and BK only) and/or refractory organic carbon. Serpentinization-type reactions involving Fe(II)-containing phases (e.g., iron sulfides, olivine, and magnetite) and evolved water during SAM analysis could produce H₂ [e.g., Sleep et al., 2004; McColloM and Bach, 2009; Mayhew et al., 2013]. The subtle yet sharp H₂ releases at 590 and 650°C in MJ and BK, respectively (Figure 11a), could also be attributed to inclusion H₂ in a mineral phase. Refractory carbon phases could also contribute to H₂ evolution. Hydrogen gas evolution from refractory carbon phases has been documented to occur between 350 and 850°C [e.g., Campbell et al., 1980]. Refractory magmatic carbon phases have been detected in Martian meteorites [Leshin et al., 1996; Grady et al., 2004; Steele et al., 2012; Agee et al., 2013], suggesting that these carbon phases could contribute to the evolved H₂ in the Gale samples.

Evolved H₂ observed above 800°C in the RN and WJ to GB samples (Figure 11a) was attributed to an instrument effect. Examination of empty cup runs including the CB blank in oven 1 indicates a weak broad release of H₂ beginning at 300 to 700°C (Figure 11a). The greater y axis scale for the CB blank relative to the subsequent blanks does not permit the visualization of the broad H₂ release in the CB blank. This broad high-temperature release during the empty cup runs could, for reasons not yet known, be magnified in the presence of a sample. If this was the case, why this \( > 800^\circ \text{C} \) release was not observed in JK and CB is not clear, though this \( > 800^\circ \text{C} \) H₂ release was observed from RN (Figure 11a)? The SAM instrument contains two nickel alloy (Inconel 693) ovens and analyses on the SAM Testbed which uses nickel alloy ovens similar to the SAM flight ovens also exhibit these trends of high-temperature H₂ (data not shown). Analyses of the same samples in other non-Inconel containing ovens, however, often do not have these \( > 800^\circ \text{C} \) H₂ releases. This suggests an instrument effect that could be related to high-temperature Ni-alloy reactions with residual water (similar to serpentinization) to form H₂.

The source of the variations in H₂ abundances and H₂ evolution temperatures between samples is complex (Figure 11 and Table S2). These variations were likely related to water evolution temperature from the sample, the mineralogy of the Fe²⁺ phases that can promote serpentinization reactions in the oven, and type of carbon phase present. Other factors affecting H₂ evolution can be attributed to other evolved gases that may react with water and H₂ to inhibit or enhance the H₂ detection. Laboratory investigations using the appropriate analog materials and oven conditions are required to more fully understand the H₂ release phenomena at all temperatures.

4.3. SO₂

All samples evolved SO₂ mostly above 500°C with minor SO₂ evolution below 500°C in CB and GB2 (Figure 12a). Minor SO₂ evolutions below 500°C were consistent with oxidative sulfide decomposition directly to SO₂ [e.g., Pelovski and Petkova, 1999; Ming et al., 2014]. CheMin analyses showed \(~1 \text{ wt } \%\) pyrrhotite (FeS) in CB, while sulfides in GB2 were not detected and were presumed to occur at concentrations below CheMin detection.
limits. The \( \text{O}_2 \) (Figure 17a) for sulfide oxidation was derived from oxychlorine decomposition that coincides with the \(<500^\circ\text{C} \) \( \text{SO}_2 \) evolutions [Ming et al., 2014]. The evolution of \( \text{SO}_2 \) above \( 500^\circ\text{C} \) was consistent with the presence of \( \text{Fe} \) and/or \( \text{Mg} \) sulfate ranging from crystalline (e.g., jarosite) to amorphous sulfate as well as adsorbed sulfate forms [McAdam et al., 2014]. The shapes of the major \( \text{SO}_2 \) release profiles above \( 500^\circ\text{C} \) vary between samples, but several samples have two or more “peaks” within the broad temperature release (e.g., RN, CB, GH, and MJ), from \(~500 \) to \( 700^\circ\text{C} \) and \( >700^\circ\text{C} \) (Figure 12a). The lower temperature peaks were consistent with \( \text{Fe} \) sulfate phases that typically decompose between \( 500 \) and \( 700^\circ\text{C} \) [e.g., McAdam et al., 2014]. Magnesium sulfate, if hydrated, will dehydrate below \( 500^\circ\text{C} \) and then decompose at temperatures higher than those reached by SAM analyses [Scheidema and Taskinen, 2011]. However, when mixed with other phases, the thermal decomposition temperature of \( \text{Mg} \) sulfate is known to drop to within the SAM temperature range and likely dominates \( \text{SO}_2 \) releases above \( 700^\circ\text{C} \) [Mu and Perlmutter, 1981a; McAdam et al., 2016].

The presence of multiple peaks at select temperatures within the \( \text{Fe} \) sulfate or \( \text{Mg} \) sulfate temperature range (e.g., BS, BK, MJ, and GB2, Figure 12a) has the potential to be utilized to further identify specific \( \text{Fe} \) sulfate and \( \text{Mg} \) sulfate species [e.g., McAdam et al., 2014]. The identification of specific \( \text{Fe} \) sulfate or \( \text{Mg} \) sulfate species from the evolved \( \text{SO}_2 \) detections will require extensive laboratory analog analyses that is beyond the scope of this work.

When crystalline sulfate species (e.g., jarosite detected by CheMin in CH, MJ, and TP) cannot account for all the evolved \( \text{SO}_2 \), then other sources of \( \text{SO}_2 \) are possible, including X-ray amorphous materials (e.g., amorphous sulfates), adsorbed sulfates, \( \text{S} \) phase inclusions in minerals or glass, oxidation of sulfide phases, or trace crystalline sulfur phases present below CheMin detection limits. X-ray amorphous materials (30–60 wt %)

Figure 11. (a) Hydrogen (\( \text{H}_2 \), m/z 2) \( \text{(c/s)} \) versus temperature as detected by the SAM-EGA. Y axis scale is \( 10^5 \) c/s except where noted. Dotted plots refer to empty cup runs that were run before sample analysis. (b) Total evolved \( \text{H}_2 \) content. Far right bracketed notation refers to windblown eolian material (E) and sedimentary rock formations (Figure 2) Yellowknife Bay (YB), Kimberly (K), Murray (M), and Stimson (ST).
containing S (reported as 2–12 wt % SO3) are present and could contribute significantly to SAM-EGA S detections [e.g., Blake et al., 2013; Vaniman et al., 2014; Morris et al., 2016; Treiman et al., 2016; Rampe et al., 2017; Yen et al., 2017]. Sulfate adsorbed onto iron (hydr) oxides is commonly observed in terrestrial environments [e.g., Aylmore et al., 1967; Parfit and Smart, 1978; Rampe et al., 2016] and would not be detected by XRD if such sulfates were present in Gale Crater. The CheMin detection of pyrrhotite (FeS) in JK and CB indicate that the >500°C SO2 could result from iron sulfate derived from sulfide oxidation by O2 evolving from (per) chlorate at lower temperatures or oxidation of any sulfide remaining at higher temperatures as a result of incomplete oxidation below 500°C [McAdam et al., 2014].

Other possible phases that could contribute to evolved SO3 include Al sulfates or organo S-bearing phases [e.g., Mu and Perlmutter, 1981a; McAdam et al., 2014]. CheMin and APXS data sets did not provide compelling mineralogical or geochemical evidence for the presence of Al sulfates though Al sulfates do form in acidic environments similar to Fe sulfates [Lindsay, 1979] and could be present at low concentrations. The oxidation of sulfur-bearing organics from O2 derived oxychlorine phases could also contribute to SO3 releases [McAdam et al., 2014]. If all the evolved CO and CO2 in WJ (~2373 μgC/g) was derived from indigenous carbon sources, up to 200 μg S/g could accompany this carbon and thus could contribute a small fraction to SAM-S (<0.05 wt % SO3) (assumes 12.3 C/S wt % ratio from Murchison meteorite analyses) [Okumura and Mimura, 2011].

The presence of Ca sulfate in most samples (RN, JK, CB, BS, GH, and GB) [Bish et al., 2013; Vaniman et al., 2014; Morris et al., 2016; Yen et al., 2017] caused the amounts of S detected by SAM (SAM-S) to be less than the amounts of S detected by the APXS (APXS-S) (Figure 12b and Table S3). This is because all Ca sulfates decomposes at temperatures beyond the SAM range and thus cannot be detected by SAM [e.g., Mu and Perlmutter,
When SAM-S and APXS-S contents were similar, Ca sulfate was either not detected by CheMin, or its detection was questionable (e.g., WJ, CH, MJ, and TP) [Rampe et al., 2017; Treiman et al., 2016]. For the WJ, CH, MJ, and TP samples, SAM-EGA is capturing all the APXS detected S (Figure 12b) which was determined to be mostly associated with Fe and/or Mg [Treiman et al., 2016; Rampe et al., 2017].

4.4. H$_2$S

The majority of H$_2$S was evolved at temperatures greater than 500°C (Figure 13a), though a gradual rise in H$_2$S evolution began at ~150°C in many samples (e.g., CH, MJ, and TP) [Rampe et al., 2017; Treiman et al., 2016]. For the WJ, CH, MJ, and TP samples, SAM-EGA is capturing all the APXS detected S (Figure 12b) which was determined to be mostly associated with Fe and/or Mg [Treiman et al., 2016; Rampe et al., 2017].

The source of H$_2$S could be derived from H$_2$O and/or HCl reactions with sulfides in the sample (reactions (1) and (2)) [e.g., Uno, 1951; Ingraham et al., 1972].

$$\text{FeS} + \text{H}_2\text{O} \rightarrow \text{H}_2\text{S} + \text{FeO}$$

$$\text{FeS} + 2\text{HCl} \rightarrow \text{H}_2\text{S} + \text{FeCl}_2.$$  

Releases of H$_2$O and HCl occurred at similar temperatures as the H$_2$S releases especially at high temperature (Figures 10a and 18a), suggesting hydrogen from HCl or H$_2$O could be reacting with sulfide-bearing phases that may not have been completely oxidized or decomposed at lower temperatures. Pyrrhotite was detected in JK and CB and thus could react with water or HCl to evolve H$_2$S [Ming et al., 2014]. Sulfide minerals were not detected in the other samples, but these sulfides could occur below CheMin detection limits or occur in the X-ray amorphous phase(s). Basaltic and silicic glasses can contain reduced sulfur [e.g., Mathez, 1976; Paris

Figure 13. (a) Hydrogen sulfide (m/z 34) (c/s) versus temperature as detected by the SAM-EGA. Dotted plots refer to empty cup runs that were run before sample analysis. Y axis scale is $x10^4$ c/s except where noted. (b) Evolved hydrogen sulfide (H$_2$S) content. Far right bracketed notation refers to windblown eolian material (E) and sedimentary rock formations (Figure 2) Yellowknife Bay (YB), Kimberly (K), Murray (M), and Stimson (ST).
et al., 2001] that might contribute to reactions that evolve H2S. Other possible sulfides sources include nanophase sulfides or organic S-bearing phases that can occur in meteoritic materials [Dai and Bradley, 2001; Sephton, 2002; Okumura and Mimura, 2011] which are expected to be incorporated in Martian surface material [e.g., Flynn, 1996]. Evolved gas reactions between H2 and SO2 are another possible source of H2S (reaction (3)) [Arutyunov et al., 1991; Binns and Marshall, 1991; McAdam et al., 2014].

\[
\text{SO}_2 + \text{H}_2 \rightarrow \text{H}_2\text{S} + \text{O}_2.
\] (4)

The bulk of H2S is released at similar temperatures, whereas most of the SO2 and H2 evolved together in several samples (e.g., JK, CB, BS, and GH) (Figures 11a and 12a) indicating that reaction (3) could also be occurring in those samples.

The "flat" gap between 400 and 550°C in BS, GB1, and GB2 was an artifact caused by the correction for the oxychlorine $^{18}\text{O}^{16}\text{O}$ isotopologue (Figure 13a). Contributions to m/z 34 from oxychlorine $^{18}\text{O}^{16}\text{O}$ isotopologue and sulfate $^{34}\text{S}$ isotopologue were subtracted from all m/z 34 plots as discussed in section 2.2. This feature was observed in these three samples because O2 was observed at higher temperatures and at higher intensity than most other samples. Furthermore, m/z 34 attributed to H2S achieved a higher intensity than most samples below and above the temperatures where the correction was applied. These factors all conspired to result in the effect of "cutting" into the H2S evolution. Subtracting out less O2 ($^{18}\text{O}^{16}\text{O}$) would result in peak in the gap that is clearly attributed to decomposition of oxychlorine.

The complexity of reactions between evolved gases and solid phases during SAM-EGA demonstrates that laboratory analysis of analog materials will be required to determine the most likely sources of H2S in the Gale Crater samples. For example, the JK and CB samples have the only CheMin detections of a mineral iron sulfide (pyrrhotite), yet they have the some of the lowest H2S content along with WJ (Figure 13b and Table S4). The other samples could have higher X-ray amorphous sulfide content that is undetectable by CheMin. The sulfides in CB could have undergone more oxidation due to greater O2 evolution than the other samples (Figure 17b), thus obscuring the H2S content that otherwise would have been evolved from CB.

4.5. NO (m/z 30).

SAM detected nitrates (0.002 ± 0.005 to 0.06 ± 0.03 wt %) in all sediments as indicated by evolved NO (Figure 14a). Multiple peak NO releases suggest that multiple cation species may be associated with nitrates in the Martian sediments. Fe-bearing nitrates tend to evolve NO at lower temperatures followed by Mg-, Ca-, Na-, and K-bearing nitrates at correspondingly higher temperatures [e.g., Gordon and Campbell, 1955; Mu and Perlmutter, 1982; Ettrah and Galwey, 1996; Stern et al., 2015]. However, the Fe-bearing phases like hematite, magnetite, and nanophase Fe oxides and oxychlorine phases present in the Gale materials have the potential to lower nitrate thermal decomposition temperatures relative to nitrates in the absence of such phases [Hoshino et al., 1981; Navarro-González et al., 2013]. Thermal shock processes involving impact events or lightning acting on atmospheric N2 have been invoked to be the main sources of nitrates on Mars [e.g., Mancinelli, 1996; Segura and Navarro-González, 2005; Manning et al., 2008].

Despite the calculated instrument background contributions to NO in the Gale samples, SAM-EGA detected sufficient NO to indicate the presence of nitrate (Figure 14b and Table S5). The only cases where calculated background contributions could account for most of the evolved NO are in the CH and WJ samples. The CH had nearly as much background NO detected as JK but because CH had much less total NO detected, most of its detected NO could be attributed to background sources (Figure 14b). The WJ sample had much less total NO detected and despite its relatively lower background NO, corrected NO contributions resulted in the second lowest nitrate level.

4.6. CO2

Carbon dioxide evolved over most of the SAM temperature range with candidate sources that include atmospherically adsorbed CO2, organic carbon, carbonate, CO2 inclusions in mineral or glass phases, and SAM background (Figure 15) [Macpherson and Mattey, 1994; Grady et al., 2004; Cannon et al., 2012; Steele et al., 2012; Glavin et al., 2013; Leshin et al., 2013; Ming et al., 2014; Eigenbrode et al., 2014; Freissinet et al., 2015]. The presence of adsorbed CO2 onto Martian regolith has long been proposed [e.g., Fanale and Cannon,
1974; Zent and Quinn, 1995], and it is likely that any adsorbed CO₂ present on Gale sediments could contribute to CO₂ evolved below 250°C [Hochstrasser and Antonini, 1972; Jänchen et al., 2009].

The possible types of organic C that could contribute to Gale Crater CO₂ include simple organic carbon (<350°C), refractory macromolecular carbon (300–600°C), trapped organics (>600°C), and magmatic carbon (>600°C) [e.g., Campbell et al., 1980; Espitalié et al., 1984; Grady et al., 2002, 2004; Steele et al., 2012; Francois et al., 2016]. Combustion of organics can explain some of the CO₂ releases. Crosshatched bars underneath each CO₂ release in Figure 15a correspond to temperatures of O₂ releases, while tick marks indicate position of O₂ peaks derived from the thermal decomposition of oxychlorine phases. This suggests that O₂ releases coinciding with CO₂ releases is consistent with combustion as a contributor to CO₂. The CB samples evolved chlorohydrocarbons that were detected by GCMS in the 150 to 350°C temperature range [Freissinet et al., 2015] indicating organic combustion likely contributed to CO₂ in CB. The JK sample could represent an extreme case where all the O₂ was consumed by combustion resulting in a dip of the O₂ release (dotted tick mark) between the two O₂ peaks (solid tick marks) [Ming et al., 2014] (Figures 15a and 17a). The O₂ dip (dotted vertical marker in Figure 15a) coincides with the main CO₂ release in JK.

Decarboxylation of organic compounds is another potential source of CO₂ over a broad temperature range (150 to 800°C) [Mu and Perlmutter, 1981b; Dollimore, 1987; Okumura and Mimura, 2011; Eigenbrode et al., 2014]. Oxygen-bearing carbonyl and carboxyl groups in refractory organic material have been detected in Tissint Martian meteorite [Steele et al., 2014]. Furthermore, oxygen-bearing organics are commonly found in carbonaceous chondrites [Okumura and Mimura, 2011], which through meteoritic infall, are expected to occur on the Martian surface [Flynn, 1996]. Alternatively, organics of either Martian or exogenous sources could have been oxidized to carboxylic acids [e.g., mellitic acid (RCOOH), acetate (CH₃CO₂⁻), and oxalates

Figure 14. (a) Evolved NO (m/z 30) (c/s) versus temperature as detected by the SAM-EGA. Dotted plots refer to empty cup runs that were run before sample analysis. Y axis scale is 10⁵ c/s except where noted. (b) Nitrate content (wt % NO₃⁻) calculated from evolved NO. Corrected contributions of nitrate from sample are in blue with estimated contribution from background in red. Far right bracketed notation refers to windblown eolian material (E) and sedimentary rock formations (Figure 2) Yellowknife Bay (YB), Kimberly (K), Murray (M), and Stimson (ST).
Carbon dioxide releases above 600°C were consistent with decomposition of organics trapped in minerals [e.g., Aubrey et al., 2006; Bowden and Parnell, 2007]. Coprecipitated phthalic acid with Mg sulfate was demonstrated to be protected from combustion by oxychlorine derived O₂ below 600°C [Francois et al., 2016]. Carbon dioxide evolution only occurred when Mg sulfate decomposed at ~700°C, and the phthalic acid subsequently combusted. CO₂ releases associated with natural jarosite thermal decomposition have also been demonstrated [Lewis et al., 2015]. The presence of Fe and/or Mg sulfate in most samples suggests that any organics trapped within Martian sulfate minerals, if present, could have contributed to evolved CO₂ at these high temperatures.

Combustion of magmatic carbon (MC) was another potential source of low-level CO₂ above 600°C [Pineau and Javoy, 1994; Grady et al., 2004; Steele et al., 2012] in CH, MJ, BK, BS, and GH. Magmatic carbon refers to carbon that occurs in inclusions or vesicles of mineral or glass phases and along grain surfaces [Grady et al., 2004]. Magmatic carbon can occur as graphite, carbynes, polycyclic aromatic hydrocarbons, and/or “amorphous” carbon [Mathez and Delaney, 1981]. Magmatic carbon has been detected in several Martian meteorites through pyrolysis and combustion analyses above 600°C and by Raman imaging [Leshin et al., 1996; Grady et al., 2004; Steele et al., 2007, 2012, 2016; Agee et al., 2013]. The CH, MJ, and BK samples had additional O₂ releases at these temperatures due to sulfate decomposition (see section 4.8 below) indicating that...
The combustion of magmatic carbon caused by O$_2$ releases from sulfate decomposition in all samples could be possible. The nondetection of sulfate-derived O$_2$ in other samples does not necessarily mean that O$_2$ was not released. The possibility exists that undetected O$_2$ was released but was consumed by other processes such as oxidation of Fe$^{2+}$ mineral phases (e.g., magnetite and pyroxenes) and/or organic combustion.

High-temperature (>600°C) evolved CO$_2$ in CH, MJ, BK, BS, and GH (Figure 15a) could also have contributions from CO$_2$ inclusions within mineral/glass phases [e.g., Macpherson et al., 1999; Manning et al., 2013]. The presence of 15–60 wt % amorphous material in the Gale Crater sediments of [e.g., Bish et al., 2013; Vaniman et al., 2014; Morris et al., 2016] suggests that some of the high-temperature CO$_2$ could be derived from this amorphous material.

Carbonates are potential sources of CO$_2$ evolved from 450 to 800°C [Leshin et al., 1996; Sutter et al., 2012; Cannon et al., 2012; Lesin et al., 2013; Ming et al., 2014]. The temperatures of the highest Gale CO$_2$ release peaks are consistent with Fe-rich carbonate for RN and CH, Mg-rich carbonate for TP, and Fe-Mg carbonate for GH and GB. The amounts of carbonates if present are low (<0.7 wt %) and below CheMin detection limits.

Reactions of evolved HCl with carbonates present in the sample could also have contributed to evolved CO$_2$ below 450°C [Cannon et al., 2012]. Laboratory analysis of evolved HCl from thermally decomposing Mg-perchlorate has been shown to react with carbonates to evolve CO$_2$ below temperatures expected for thermal decomposition of carbonate. Evolved HCl observed between 200 and 500°C in CB, CH, TP, and BS (Figure 18) could have reacted with carbonate to contribute observed CO$_2$ in these samples. The coinciding evolution of CO with CO$_2$ (Figures 15 and 16) in these samples, however, suggests that organic C sources supports organic C sources of CO$_2$ in this temperature range as well. Nevertheless, evolved HCl reactions...
with Gale Crater carbonates should be considered as potential contributors to evolved CO$_2$ at temperatures below 450°C.

Contributions of background CO$_2$ from the derivatization reagent MTBSTFA are also likely contributors to the evolved CO$_2$ at temperatures <600°C. When compared to the total evolved CO$_2$, the amount of background CO$_2$ indicated in orange in Figure 15b could not account for all of the evolved CO$_2$. Total CO$_2$ evolved from all samples except TP indicated that background CO$_2$ in the worst case would consist of less than 36% of the evolved CO$_2$ (Figure 15b). The TP sample evolved so little CO$_2$ that background CO$_2$ had the potential to contribute at least 60% evolved CO$_2$ to the TP release (Figure 15b). The 900 nmol background CO$_2$ estimate was applied to all samples regardless if they were single or triple portions [Glavin et al., 2013]. Thus, the WJ, BK, BS, and GH have lower contributions to background relative to the other samples because these triple portion samples when normalized for comparison to single portions had less background CO$_2$ than the other single portion samples.

Most CO$_2$ (except in GB) was evolved below 450°C, which was consistent with CO$_2$ derived from organics. Excluding the possible background sources, the organic sources could be indigenous to Mars from igneous, hydrothermal, atmospheric or biological sources, or exogenous from interplanetary dust particles, cometary, or meteoritic infall [e.g., Flynn, 1996; Jull et al., 2000; Yen et al., 2006; Heinrich et al., 2007; Steege et al., 2007; Leshin et al., 2013; Ming et al., 2014; Freissinet et al., 2015]. Carbonates (RN, TP, and GH) and MMC (CH, MJ, BS, and GH) are possible higher-temperature sources of CO$_2$. The GB sample is different than all other samples in that most CO$_2$ was evolved above 450°C suggesting GB CO$_2$ was derived from carbonate, although organics cannot be excluded.

Figure 17. (a) Evolved O$_2$ (m/z 32) (c/s) versus temperature as detected by SAM-EGA. Dotted plots refer to empty cup runs that were run before sample analysis. Possible sources of O$_2$ are indicated in brackets. Y axis scale is 10$^6$ c/s except where noted. (b) Evolved O$_2$ content below 600°C calculated as perchlorate (ClO$_4$) and calculated as O$_2$ above 600°C (>600°C–O$_2$). Far right bracketed notation refers to windblown eolian material (E) and sedimentary rock formations (Figure 2) Yellowknife Bay (YB), Kimberly (K), Murray (M), and Stimson (ST).
The CO₂ carbon contents (160 ± 248–2373 ± 820 μgC(CO₂)/g; corrected for background CO₂) vary over a wide range (Figure 11b and Table S6). The highest CO₂ carbon contents occurred in the eolian RN and GB sediments and the JK mudstone and WJ sandstone, whereas the lowest contents were found in the TP, BK, and GH samples (Figure 15b). There appears to be a general trend of decreasing CO₂ carbon content with increasing elevation beginning with WJ (Figure 15b).

4.7. CO

Carbon monoxide was evolved from all samples over a wide temperature range (Figure 16). The majority of the CO evolved between 150 and 575°C and was mostly coincident with at least one of the CO₂ release peaks. The CO carbon content associated with CO₂ was always less than CO₂ (Figures 15b and 16b and Tables S6 and S7). Lower CO release intensities were evolved between 575 and 750°C in all samples except CB and JK where no major CO releases were observed at those temperatures.

The evolution of the CO was consistent with the presence of incomplete combustion of organics and/or oxygen-bearing organics in the Gale Crater materials. The oxygen-bearing organics as discussed for CO₂ can be derived from initially deposited oxygen-bearing organics [e.g., Okumura and Mimura, 2011; Steele et al., 2014] or organics (e.g., hydrocarbons) that oxidized to carboxylic acids from exposure to oxidizing conditions on the Martian surface [Benner et al., 2000]. Laboratory analysis of oxygen-bearing organics from thermal decomposition of carbonaceous chondrite and carboxylic acids demonstrates that the main CO and CO₂ releases can occur at the temperature ranges observed by SAM [e.g., Dollimore, 1987; Okumura and Mimura, 2011; Eigenbrode et al., 2014].

Evidence of background CO evolution was present in the blanks above 400°C and especially above 820°C (Figure 16a). Background CO releases above 400°C was observed in the GH blank and could be considered significant for GH and possibly WJ because these samples released so little overall CO. Comparison of the WJ and GH blank intensities to other sample intensities illustrates that background CO was not as significant contributor to overall CO releases. The consistency of background CO peaks above 820°C in the blanks suggests that CO peaks with similar intensities (e.g., GB, TP, and BS) (Figure 16a) could have significant contributions from background above 820°C. The source of background CO below 600°C could have contributions from MTBSTFA. The CO background sources above 600°C are unknown, and laboratory analog work will be required to understand this source.

The CO carbon content (11 ± 3 to 320 ± 130 μgC(CO)/g) was variable throughout Gale Crater (Figure 16b and Table S7). The CO carbon content unlike CO₂ was low in RN and GB compared to the much higher CO carbon content in CH. Similar to CO₂, the CO carbon content decreased with elevation beginning with CH (Figure 16b).

4.8. O₂

Oxygen was evolved from all samples suggested the presence of oxychlorine phases like chlorate or perchlorate (0.05 ± 0.025 to 1.05 ± 0.44 wt % as ClO₄⁻) (Figure 17 and Table S8). However, the O₂ peak temperatures were lower than what is typical of pure reagent grade Ca, Mg, Na, and K perchlorate and higher than what is expected for iron perchlorate [Glavin et al., 2013]. Alternatively, pure Mg, Ca, or Na chlorate have O₂ release temperatures that occur coincident with the O₂ releases from most of the Gale samples with the exception of the CB and low-temperature JK peaks [Sutter et al., 2014; Clark et al., 2016]. Iron phases detected in Gale Crater sediments could catalyze the perchlorate and/or chlorate thermal decomposition to lower temperatures; analyses of these mixtures have resulted in O₂ release temperatures consistent with those detected by SAM [Rudloff and Freeman, 1970; Sutter et al., 2015; Clark et al., 2016].

Thermal decomposition of sulfate was the likely source of the second high-temperature O₂ evolution in CH, MJ, and BK (Figure 17a). Unlike the other Gale sediments, the CH, MJ, and BK samples have two distinctive O₂ releases. The lower temperature (<600°C) O₂ releases in CH, MJ, and BK are attributed to an oxychlorine phase(s). The higher-temperature (>600°C) O₂ releases in CH, MJ, and BK were not attributed to an oxychlorine phase as this O₂ release temperature was typically too high for perchlorate and chlorate thermal decomposition. Although the high-temperature O₂ release profile does not correspond precisely with the SO₂ release profile, the high-temperature O₂ peak does correspond with at least one of the SO₂ peaks (Figures 12a and 17a). This suggests that sulfate decomposition described above could have supplied the
high-temperature O\textsubscript{2}. Magnesium sulfate and Fe sulfate (e.g., jarosite), which have been detected in the Pahrump rocks, commonly evolve O\textsubscript{2} upon thermal decomposition [e.g., Gallagher et al., 1970; Scheidema and Taskinen, 2011; Lewis et al., 2015].

Dehydrated jarosite $2K\text{Fe(SO}_4\text{)}_3\rightarrow3K_2\text{SO}_4+Fe_2O_3+3\text{SO}_2+1.5\text{O}_2$. \hspace{1cm} (5)

Why the high-temperature O\textsubscript{2} release profile in CH, MJ, and BK does not follow the SO\textsubscript{2} release profile with same number of peaks or why high-temperature O\textsubscript{2} releases were not observed in the other samples was not clear. Complex reactions during SAM-EGA analysis between sulfate evolved O\textsubscript{2} and other mineral phases or evolved gases present in some Gale materials and not in others could inhibit O\textsubscript{2} detection. More work involving SAM-EGA analog analyses will be required to understand this inconsistency of high-temperature O\textsubscript{2} evolution.

The presence of multiple O\textsubscript{2} peaks between 250 and 500°C in BS and GH suggests that these sediments could possess chlorates and perchlorates or multiple cation phases of perchlorate and/or chlorate (Figure 17a). The cation present in the oxychlorine phase affects the decomposition temperature where perchlorate and chlorate decomposition temperatures tend to follow the order Fe perchlorate $<$ Mg chlorate $<$ Ca chlorate $<$ Mg perchlorate $<$ Ca perchlorate $\sim$ Na chlorate $<$ Na perchlorate $<$ K chlorate $<$ K perchlorate [e.g., Glavin et al., 2013; Sutter et al., 2015]. For example, one possible scenario for the three O\textsubscript{2} peaks in GH sediment was that GH possesses Mg chlorate, Mg perchlorate, and Ca perchlorate, all of which could yield three distinctive O\textsubscript{2} peaks in GH.
The content distribution of oxychlorine phases could be affected by one or more factors. The oxychlorine contents (as ClO\textsubscript{4}\textsuperscript{-}) mostly range from 0.05 to 0.38 wt % with the CB material having the significantly highest content (1.05 wt % ClO\textsubscript{4}\textsuperscript{-}, Figure 17b and Table S8). The source of all the sediments could simply have variation in oxychlorine content that causes the observed distribution. Postdepositional solutions enriched or depleted in soluble salts could raise or lower the soluble salt contents, respectively, of the Gale materials. Other post depositional oxidizing or reducing processes could also affect the oxychlorine contents [e.g., Catling et al., 2010; Quinn et al., 2013; Wilson et al., 2016]. More detailed analysis of the mineralogy and geochemistry coupled with laboratory analysis will be required to understand the nature and origins of the oxychlorine distribution in Gale Crater.

4.9. Hydrochloric Acid (HCl)

Hydrochloric acid was evolved from all Gale materials (0.004 ± 0.005 to 0.1 ± 0.08 wt % Cl\textsubscript{HCl}) (Figure 18 and Table S9). All HCl evolved at temperatures greater than the initially evolved O\textsubscript{2} with the exception of the CB and BS samples. Hatched rectangles in Figure 18a indicate the temperature range of O\textsubscript{2} releases. The CB and BS samples evolved HCl with O\textsubscript{2} but also possessed a high-temperature (>600°C) HCl peak that was not associated with any O\textsubscript{2} release (Figures 17a and 18a).

The evolution of HCl with O\textsubscript{2} in the CB and BS samples suggests the presence of Mg perchlorate and/or Mg chlorate phases. Only the Fe and Mg oxychlorine species chlorate and perchlorate are known to release HCl. All other oxychlorine phases typically form corresponding chloride phases (e.g., CaCl\textsubscript{2}, NaCl, and KCl) with no chlorate phases. Only the Fe and Mg oxychlorine species chlorate and perchlorate are known to release HCl.

The evolution of high-temperature HCl not associated with oxychlorine decomposition (Figure 18a) was consistent with water vapor reacting with vapor from melting chloride salts [e.g., Fraisslera et al., 2009].

\[
\text{CaCl}_2(g) + \text{H}_2\text{O}(g) \rightarrow 2\text{HCl} + \text{CaO}. \tag{6}
\]

The source of the chloride salts in the Gale materials was consistent with thermal decomposition products associated with Ca, Na, and/or K oxychlorine phases that decompose to chlorides [Marvin and Woolaver, 1945; Markowitz, 1963; Migdal-Mikuli and Hetmanický, 2008; Cannon et al., 2012]. The temperature of O\textsubscript{2} and HCl evolution was likely too high to be Fe perchlorate or Fe chlorate [e.g., Glavin et al., 2013]. This suggests that Mg perchlorate or Mg chlorate are the best candidates for CB and BS.

The Cl content derived from evolved HCl (Cl\textsubscript{HCl}) did not always agree with the Cl content calculated from the amount of measured SAM-O\textsubscript{2} (Cl\textsubscript{Calc}) assuming that all O\textsubscript{2} was derived from perchlorate (Figure 18b). The RN, JK, and CB samples had lower Cl\textsubscript{HCl} than Cl\textsubscript{Calc}, whereas the WJ CH, MJ, TP, BK, BS, GH, and GB materials had, within error, similar amounts of Cl\textsubscript{HCl} and Cl\textsubscript{Calc}. This suggests that the chloride salts produced from oxychlorine decomposition in the RN, JK, and CB samples did not completely react with water to yield HCl. The cause for this discrepancy is not known, and analog laboratory work will be required to understand this behavior.

The Cl\textsubscript{Calc} was always lower than total CI (Cl\textsubscript{Total}) measured by APXS (Figure 18b and Table S9), indicating that there was a CI component that does not occur as oxychlorine phases. Chlorine in Martian meteorites has been found to be associated with apatite, amphiboles, NaCl, biotite, scapolite, and glass [Williams et al., 2016, and references therein]. Apatite which was detected by CheMin in WJ, CH, MJ, and TP (0.8–2 wt %) cannot make up for the missing CI because apatite provides only a fraction (<17%) of total CI in those samples (assumes apatite CI content detected in Mars meteorites, e.g., Ca\textsubscript{3}(PO\textsubscript{4})\textsubscript{3}(Cl\textsubscript{0.5}, F\textsubscript{0.3}, OH\textsubscript{0.2}) [Filiberto and Treiman, 2009]). Amphiboles or scapolite are another a source of CI in Martian meteorites [e.g., Filiberto and Treiman, 2009; Filiberto et al., 2014], yet no amphiboles or scapolite have been detected by CheMin. Phyllosilicates have been detected in Gale sedimentary rock [Vaniman et al., 2014; Treiman et al., 2016; Rampe et al., 2017]; however, no evidence of biotite has been detected. This leaves the remaining CI component to reside in some glassy/amorphous component and/or in a crystalline phase that was below the
CheMin detection limits. If the remaining Cl component was crystalline NaCl or CaCl₂, it would amount to <1 wt % NaCl or CaCl₂, which was below CheMin detection limits.

5. Implications

5.1. Introduction

The SAM-EGA detections provide a substantial advancement in understanding the geochemical and mineralogical composition of the sediments and sedimentary rocks sampled in Gale Crater. The SAM-EGA analyses have corroborated detections of phyllosilicates and jarosite by CheMin and amorphous sulfate by CheMin/APXS and provided insight into the phases that bear water as detected by the DAN instrument [Mitrofanov et al., 2014]. SAM-EGA analyses have also been critical in detecting phases such as nitrate, oxychlorine, and organic/inorganic carbon that were not detected by other MSL instrumentation. The following section evaluates RN and GB eolian sediments, Sheepbed mudstone, Windjana sandstone, Murray mudstone, and the Stimson sandstone materials with a focus on highlighting notable evolved gas detections specific to SAM-EGA that assist in adding further understanding to the compositions and origins of Gale Crater sediments and sedimentary rock.

5.2. Rocknest/Gobabeb Eolian Sediments

Notable characteristics of RN and GB include relatively high CO₂ and NO releases and evidence of organic carbon and carbonate. There was also evidence for a grain size dependence of water type, SO₂ evolution temperatures, and possible carbon speciation.

The eolian RN and GB samples evolved CO₂ at temperatures consistent with the presence of organic C (<450°C) and carbonate (450–650°C) (Figure 15a). The GB sample is dominated by coarser grains (>100 μm) [Ehlmann et al., 2017] and released most CO₂ in the 450–650°C CO₂ range consistent with the presence of carbonate (~0.70 ± 0.1 wt % CO₃) though organic C cannot be completely excluded. Lower temperature (<450°C) CO₂ indicated that some organic C could be present in GB. The RN sample had significant fine-grained component (~40–60% <100 μm) [Minitti et al., 2013] which evolved most CO₂ at temperatures consistent with organic C, whereas ~30% of its CO₂ is consistent with carbonate (~0.32 ± 0.05 wt % CO₃) [Leshin et al., 2013]. These results demonstrate a possible grain size dependence of carbon species in eolian sediment, where organic C and carbonate may mostly reside in fine-grained (<100 μm) and coarser-grained (>100 μm) eolian sediments, respectively.

The RN and GB eolian surface materials have some of the highest nitrate and CO₂ carbon contents measured to date (Figures 14b and 15b and Tables S5 and S6), which could indicate that other sources with high nitrate and carbon contents exist in Gale Crater. The coarser grained components (e.g., >100 μm) of RN and GB suggest that carbon and nitrate were mostly acquired from the erosion of local and regional sources. The amount of eolian dust (<10 μm) [Morris et al., 2006, and references therein] in RN is not known but a fine-grained component (~<100 μm) (~40–60%; [Minitti et al., 2013]) is present indicating that RN could consist of some globally mixed nitrate- and carbon-bearing dust with similarly high C and N.

The overall water release profile for RN and GB differed, which is attributed to grain size distribution differences between the two samples. Rocknest released a greater amount of water below 200°C than GB (Figure 10a). The GB sediment material is dominated by >100 μm grain sizes [Ehlmann et al., 2017], whereas RN has a substantial <100 μm size fraction [Minitti et al., 2013]. The greater amount of fine-grained material in RN likely has a greater surface area that can retain more adsorbed water than the coarser grained GB material.

Grain size effects appeared to have an effect on SO₂ releases between the GB1 (<150 μm) and the GB2 (150 μm–1 mm) samples. The GB1 SO₂ release temperatures were consistent with SO₂ being derived from Mg sulfates, while the SO₂ release temperatures in the larger-grained GB2 were more consistent with releases from Fe sulfate phases (Figure 12a). The Murray material near to the GB samples contained Mg sulfate concretions that were also detected at Pahrump which may erode to <150 μm particle sizes and make contributions to the GB materials. CheMin detections of jarosite in the Pahrump material of the Murray formation may be the source of Fe sulfates indicated by the SAM-EGA for the larger particle GB2 sample.
5.3. Sheepbed Member-Yellowknife Bay Formation

The JK and CB mudstones had the highest evolved H$_2$ content which is consistent with the hypothesis that these lacustrine sediments were deposited under reducing conditions [Grotzinger et al., 2014; Ming et al., 2014]. The SAM-GCMS detection of organics and sulfur containing organics in CB [Freissinet et al., 2015] and the CheMin detection of Fe sulfide (pyrrhotite) only in JK and CB [Vaniman et al., 2014] highlight the reduced nature of the Sheepbed rocks. Hydrogen could be evolved from the thermal decomposition of hydrocarbons [Campbell et al., 1980] or through reactions of evolved water and Fe sulfides:

$$4\text{FeS} + 2\text{H}_2\text{O} \rightarrow \text{Fe}_3\text{S}_4 + \text{Fe(OH)}_2 + \text{H}_2.$$  (7)

Rickard and Luther, 2007].

H$_2$ was evolved from all the other samples, albeit at lower contents, even though CheMin detected no FeS (Figure 11b and Table S2). Other Fe$^{2+}$-bearing phases are present in most of the other Gale Crater rocks and eolian sediments (e.g., olivine, magnetite, and pyroxene) that could react with water to form H$_2$ during SAM-EGA analysis [e.g., Mayhew et al., 2013]. However, FeS thermally decomposes within the SAM temperature range allowing for more complete reactions with water to form H$_2$ [Yan et al., 2008]. Only a fraction of Fe(II) silicates reacted with 50–100°C water to produce H$_2$ in laboratory experiments lasting up to 100 d [Mayhew et al., 2013]. This suggests greater H$_2$ production in Gale materials possessing FeS than Gale materials without FeS. The highest H$_2$ production from the Sheepbed JK and CB mudstones demonstrates the significantly reduced nature of these rocks and that despite possessing oxidizing phases (e.g., sulfate, nitrate, and oxychlorine), the reduced nature is preserved.

While the CB sample has reduced phases (Fe sulfides), the CB rock also has the highest oxidized nitrate and oxychlorine contents relative to all the other Gale Crater materials examined to date. These oxidized salts probably did not codeposit with the iron sulfide in the JK and CB sediments. Subsequent to lacustrine deposition, fluids, or evaporative conditions concentrated these oxidized salts in CB. The lower nitrate and oxychlorine contents in JK could be the result of being leached by postdepositional fluids high in Ca sulfate that yielded the Ca sulfate veins (~1 mm) which crosscut the JK but not the CB drill hole [Vaniman et al., 2014].

5.4. Kimberley Formation

SAM-EGA analysis demonstrated that the WJ sandstone had the highest CO$_2$ carbon content and one of the lowest nitrate contents of all the samples (Figures 14b and 15b and Tables S5 and S6). Windjana, unlike the previously analyzed RN, JK, and CB, was characterized by having equal amounts of sulfur detected by SAM-EGA and the APXS (Figure 12b and Table S3).

Windjana has the highest CO$_2$ carbon content (2373 ± 820 μgC(CO$_2$)/g) with most CO$_2$ being evolved at temperatures (<450°C) that are consistent with organic C. A portion of the CO$_2$ below 250°C could be attributed to adsorbed CO$_2$. Nevertheless, if half the detected CO$_2$ was attributed to adsorbed CO$_2$, there was still a substantial amount of evolved CO$_2$ that could be attributed to organic C. The relatively less altered nature of WJ could promote retention of organic C, especially if organic C is dominated by water-soluble organic carbon such as the carboxylic acids that are predicted to exist on the Martian surface [Benner et al., 2000]. Limited postdepositional aqueous processes would allow for retention of water-soluble organic carbon in WJ. Reduced organic C was not detected by SAM-EGA above background levels or by SAM-GCMS in WJ, suggesting that organic C represented by evolved CO and CO$_2$ could be oxidized to water-soluble organic carbon. Alternatively, the WJ source materials (parent rocks) might have had inherently high organic C relative to the other materials thus far analyzed by SAM-EGA.

Windjana has the lowest nitrate concentration in the Gale materials (Figure 14b and Table S5) analyzed by SAM suggesting that the WJ sandstone was either leached of initially deposited nitrate or that it was exposed to solutions with much less nitrate than the other Gale materials. If WJ was leached of its nitrate, then this event preceded any deposition of soluble organic C described above. The WJ O$_2$ and Cl contents were not the lowest of all the Gale Crater materials examined by SAM-EGA, suggesting soluble oxychlorine and chloride were deposited after nitrate was leached.

The nearly equivalent WJ SAM-S and APXS-S contents indicates that WJ has a low Ca sulfate content. This suggests that the WJ was not exposed to depositional or postdepositional fluids enriched in Ca sulfate.
Other Gale Crater samples (e.g., JK, CB, BK, BS, and GH) that have SAM-S contents less than APXS-S contents (Figure 12b and Table S3) were the result of having greater Ca sulfate inputs than what occurred in WJ. This is supported by the CheMin/APXS analysis that indicated SAM-S is dominated by X-ray amorphous Mg-SO$_4$-bearing phases [Treiman et al., 2016].

The source of this amorphous Mg-SO$_4$ is consistent with dehydration of hydrated crystalline Mg sulfate phases (e.g., starkeyite and hexahydrite). Laboratory analog studies have demonstrated that dehydration of crystalline-hydrated Mg sulfates leads to amorphous sulfate phases [Chiper and Vaniman, 2007]. The amorphous Mg-SO$_4$ in WJ is consistent with an initially deposited, highly hydrated crystalline Mg sulfate that subsequently dehydrated to an amorphous phase. Kieserite has been shown to be stable under vacuum conditions and is not expected to dehydrate. The geochemical conditions (e.g., ionic activities, pH, and temperature) required to form these higher hydration state Mg sulfates differs from lower hydration state (e.g., kieserite) [e.g., Doner and Lynn, 1989]. Thus, the geochemical conditions in the Windjana sandstones were likely not favorable for kieserite formation but may have been more favorable for other higher-hydrated Mg sulfate formation. Amorphous Fe sulfates that may be present in other samples (e.g., RN, JK, CB, and BK) suggest that similar processes could have operated where Fe sulfates with high hydration states (e.g., copiapite and melanterite) were initially deposited that subsequently dehydrated to yield amorphous Fe sulfates [e.g., Wang and Zhou, 2014; Sklute et al., 2015; Rogers et al., 2016]. The geochemical conditions for highly hydrated Fe sulfates formation differ from those of lower hydration state jarosite [e.g., Spencer, 2000; Jambor et al., 2000], which may provide clues to past geochemical conditions where only amorphous Fe sulfate is found and not jarosite.

5.5. Murray Formation

The Murray formation (CH, MJ, TP, and BK) mudstones were deposited in a lacustrine environment and the composition of these mudstones record a complex pH and redox geochemical history [Hurowitz et al., 2017; Rampe et al., 2017]. This record of environmental change is indicated by the presence of oxidizing (hematite), reducing (magnetite), acid (jarosite), and alkaline (apatite and phyllosilicate) mineralogies at different stratigraphic positions. The SAM-EGA analyses provided additional evidence of this complicated geochemical history as indicated by presence of oxidizing (nitrate, sulfate, and oxychlorine), and alkaline (carbonate) phases, along with evidence of organics.

Some of the lowest CO$_2$ yet highest CO carbon contents occur in Murray samples (Figures 15b and 16b and Tables S6 and S7), with evolution temperatures that occur mostly below 450°C, consistent with presence of organic C. No organic C fragments were detected below 450°C by SAM-EGA or SAM-GCMS above background levels suggesting that most organics are oxygen bearing, consistent with the observations of oxygen-bearing organic C in the Tissint Martian meteorite [Steele et al., 2014]. The CO$_2$ and CO contents decreased with increasing elevation (Figures 15b and 16b). This carbon content trend with elevation could simply reflect the organic C content of the initially depositing material. Though the possibility exists that prior to nitrate and oxychlorine additions the water-soluble organics were mobilized to greater depths in the Murray via leaching processes.

The BK sample has a higher nitrate content than the other Murray materials, which is attributed to the addition of a nitrate from a different source region than those that contributed sediments to the other Murray rock samples. The NO intensity and release profile between 200 and 450°C in BK are similar to CH, MJ, and TP NO releases (Figure 14a) indicating the presence of a nitrate derived from a similar source. The NO peak at 521°C in BK does not exist in the other samples and is consistent with the addition of nitrate from a different source. Nitrate was initially deposited in CH, MJ, TP, and BK from a similar source, but additional fluids containing nitrate from a different source subsequently affected BK and not the other lower Murray rocks.

The BK sample also possesses more oxychlorine than the other Murray rocks (Figure 17b and Table S8). There are variations in oxychlorine-O$_2$ peak temperatures between the BK and the other Murray materials but not as much as with the NO peak temperatures. Nevertheless, it appears that along with nitrate, BK was exposed to more oxychlorine than the other Murray materials.

Diagenetic complexity in the Murray formation is indicated by evidence of both acidic and alkaline fluids. A high-temperature CO$_2$ peak (~600°C) consistent with Mg-bearing carbonate was detected in TP and an ~800°C water peak consistent with a Mg and/or Al smectite was detected in MJ. Jarosite, a mineral that forms under acidic (~pH 2–4) conditions, was also detected in TP and MJ [Rampe et al., 2017]. CheMin detected ~4
from 6.2 × 10³ to 2.2 m below the sea (SPG) region supported by organic C in sea sediments given the detected C contents was estimated based on comparisons with observed cell densities. Nevertheless, the size of microbial populations that could potentially have been supported in Gale Crater lake alone cannot necessarily be used to determine if life was possible or that if life ever existed on Mars.

suggest that heterotrophic microbiology could have been possible on ancient Mars. Results from this work

5.6. Stimson Formation

Notable characteristics of the Stimson materials are that they have low water and CO₂ carbon contents, have been exposed to alteration fluids that varied in pH, and have undergone a complex geochemical history characterized by elemental gains and losses [Yen et al., 2017]. The low water content may be due to less adsorbed H₂O because of the coarser particles with low surface area as well as lower levels of hydrated mineralogy (e.g., phyllosilicates and hydrated salts). The nondetection of phyllosilicates and jarosite suggests that most water above 500°C could be associated with hydroxylated amorphous Al/Si phases or water inclusions trapped in amorphous and/or mineral phases.

The alteration of GH resulted in lower nitrate, oxychlorine, and organic C (<450°C, CO₂) in GH relative to BS (Figures 14b, 15b, and 17b and Tables S5, S6, and S8). Assuming that BS is an unaltered baseline material, organic C was lost from GH as indicated by the intensity decrease of the 345°C CO₂ peak in GH relative to BS (Figure 15a). Both CO peak intensities in BS also drop dramatically relative to GH (Figure 16a). The losses are interpreted to have been caused by leaching of these soluble salts and organic C from the rock. These losses are consistent with the alteration in GH that also resulted in feldspar and pyroxene loss [Yen et al., 2017].

Although GH is characterized by chemical losses [e.g., Frydenvang et al., 2016; Yen et al., 2017], GH has gained sulfate. The SAM-S contents are lower than APXS-S contents in both BS and GH demonstrating that Ca sulfate-bearing fluids have infiltrated both BS and GH (Figure 12b). However, a SO₂ peak at 815°C in GH that is not present in BS (Figure 12a) likely results from addition of Mg sulfate to GH. Furthermore, the SAM-S/ APXS-S content ratio is higher in GH (5.4) than BS (4.8) which is consistent with GH having more Ca sulfate than BS (Figure 12b). This is also supported by GH (7.8 ± 0.08 wt % Ca) having greater Ca content than BS (6.12 ± 0.07 wt % Ca) and GH having more anhydrite than BS [Yen et al., 2017]. These results indicate that although sulfate solutions have interacted with BS, the geochemical alteration that led to GH formation also resulted in greater additions of Mg and Ca sulfate than in BS.

Along with sulfate additions, carbonate appears to have also been added to GH. The GH sample has a CO₂ release (~560°C) consistent with carbonate that is not present in BS. The level of carbonate in GH is below CheMin detection limits. The extensive alteration of GH suggests that the acid fluids were involved; however, the detection of carbonate indicates that a separate and likely later event consisting of alkaline fluids has also affected the GH material.

5.7. Implications for Habitability

The SAM instrument detections of nitrate N and organic C indigenous to Gale Crater sedimentary rocks suggest that heterotrophic microbiology could have been possible on ancient Mars. Results from this work alone cannot necessarily be used to determine if life was possible or that if life ever existed on Mars. Nevertheless, the size of microbial populations that could potentially have been supported in Gale Crater lake sediments given the detected C contents was estimated based on comparisons with observed cell densities supported by organic C in seafloor clay-dominated sediments from middle portions of the South Pacific Gyre (SPG) region [D’Hondt et al., 2010, 2015]. The SPG noncalcareous sediments at Integrated Ocean Drilling Program Sites U1366 and U1369 from 0.2 to 2.2 m below the seafloor possess organic C contents of 600–2000 μgC/g. At these subseafloor horizons, microbial communities achieved cell densities that ranged from 6.2 × 10³–4.8 × 10⁵ cells/g sediment. The Gale Crater materials possess similar organic C contents (up to 2384 μg C/g; combined C(CO₂) and C(CO)) as these SPG sediments, supporting the notion that the extremely oligotrophic SPG microbial environment provides a useful approximation of the size of heterotrophic cell densities that could have been supported in Gale Crater, provided the organic C was bioavailable.

However, factors such as radiation, hydration, and temperature fluctuations could have been significant barriers to habitability on ancient Mars. Additionally, Gale nitrate N contents (10–100 μg N/g) suggest that unlike the SPG sediments, any microbial population at Gale Crater would have likely been nitrogen limited. Due to these differences with the SPG environment along with the numerous assumptions required to extrapolate from Earth to Mars, the SPG cell densities (10³–10⁵ cells/g sediment) should be considered primarily as
a useful upper bound on what could have been possible in the putative ancient habitable environment in Gale Crater. The intent here is to demonstrate that if all necessary environmental and other essential geochemical conditions (e.g., pH, redox, and abundance other essential nutrients) were favorable for life, there was sufficient organic C to support a small microbial population similar to what is observed in the most oligotrophic environments on Earth.

6. Conclusions

Noteworthy findings specific to results of the SAM investigation are the detections of nitrate, organic C, oxochlorine phases, carbonate, and Mg sulfate. The RN, GB, GH samples possessed evolved CO₂ results consistent with carbonate indicating those samples may have been exposed to episodes of alkaline pH solutions. Oxochlorine and Mg sulfate detections in all samples, although not necessarily linked, coupled with detections made elsewhere on Mars [e.g., Gendrin et al., 2005; Murchie et al., 2009; Hecht et al., 2009; Øjha et al., 2015] demonstrate the widespread occurrence of oxochlorine and Mg sulfate phases across Mars. Water evolutions were complicated and likely derived from multiple sources including adsorbed water, interlayer and structural water from phyllosilicates, hydrated salts, jarosite hydroxyls, and mineral/glassy inclusion water. The SAM results when coupled with CheMin mineralogical and APXS geochemical analyses indicate that the sampled Gale Crater sedimentary rocks were exposed to a complicated authigenic and diagenetic history that involved temporal (spatial?) variations in pH, redox chemistry, and salt concentration. All samples possessed oxidized phases (e.g., nitrate, oxochlorine, and sulfate), suggesting that all sediments were exposed to oxidizing conditions. However, the presence of iron sulfides [Vaniman et al., 2014] and reduced hydrocarbons [Freissinet et al., 2015] in the Sheepbed mudstone suggests that these sediments may have initially been deposited in a geochemically more reducing environment that allowed for the preservation of these reduced phases. The evolution of H₂ and H₂S from all samples also indicates that all Gale samples still possess some level of reducing potential despite the presence of oxidized phases. Organic C and nitrate detections in Gale indicate that two very important constituents for microbiology were present in Gale Crater. Although nitrogen may have been limiting, the presence of organic C suggests that heterotrophic microbiology could have been possible on ancient Mars.

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