Volatile, Isotope, and Organic Analysis of Martian Fines with the Mars Curiosity Rover

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Samples from the Rocknest aeolian deposit were heated to ~835°C under helium flow and evolved gases analyzed by Curiosity's Sample Analysis at Mars instrument suite. H₂O, SO₂, CO₂, and O₂ were the major gases released. Water abundance (1.5 to 3 weight percent) and release temperature suggest that H₂O is bound within an amorphous component of the sample. Decomposition of fine-grained Fe or Mg carbonate is the likely source of much of the evolved CO₂. Evolved O₂ is coincident with the release of Cl, suggesting that oxygen is produced from thermal decomposition of an oxychloride compound. Elevated δD values are consistent with recent atmospheric exchange. Carbon isotopes indicate multiple carbon sources in the fines. Several simple organic compounds were detected, but they are not definitively martian in origin.

The exchange of materials between a planet's interior, surface, and atmosphere drives the composition of mineral and chemical constituents that can create habitable environments on the terrestrial planets. Surface deposits, including

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provide information about the average composition of the martian crust (6).

The Sample Analysis at Mars (SAM) instrument suite onboard the Mars Science Laboratory (MSL) rover Curiosity provides diverse analytical capabilities for exploring martian materials, including volatile and isotopic compositions, and a search for organic compounds, whether of abiotic or biological origin (7). Traces of organic compounds have been found in martian meteorites (8–12), but previous landed missions, most notably Viking, did not find definitive evidence of martian organic material (13).

Curiosity's first sampling campaign took place at Rocknest, an aeolian sand shadow. The rover ingested fine-grained Rocknest material into its two analytical instruments: Chemistry and Mineralogy (CheMin), for x-ray diffraction, and SAM, for analysis of volatiles. Both SAM and CheMin sampled portions from scooped materials that were sieved to contain grain sizes <150 µm. Mineralogical and chemical results summarized in a companion paper (14) indicate bulk composition similar to martian fines analyzed by previous missions. Plagioclase, olivine, augite, pigeonite, and minor magnetite are the major igneous minerals (15). Minor anhydrite and hematite are the only nonigneous minerals detected. Along with these crystalline phases, the chemical and mineralogical analyses indicate that almost half of the <150-µm fraction comprises amorphous material (14). SAM performs evolved gas analysis (EGA) with the quadrupole mass spectrometer (QMS) and isotope measurements of evolved gases using both the QMS and the tunable laser spectrometer (TLS), the latter being sensitive to isotopes of CO₂ and H₂O. Organic analyses can be performed with the QMS alone or when it is coupled to the gas chromatograph (GC). SAM analyzed four separate portions from the fifth scooped sample at Rocknest

Table 1. Experiment parameters for four analyses of Rocknest fines. All evolved gases were analyzed by the QMS; temperature (*T*) range of gases that were then sent to the GC and TLS are shown.

Rocknest run	Sol (mission day)	Sample T range of gas sent to GC (°C)	Sample T range of gas sent to TLS (°C)	Rationale
Run 1	93	146–533	547–702*	GC: Low-T organics TLS: Predicted T for thermal decomposition of carbonates
Run 2	96	98–425	440–601	GC: Low-T organics below SO ₂ evolution T TLS: Target CO ₂ from suspected carbonate peak
Run 3	99	533–822	234–425	GC: High-T organics TLS: Low-T CO_2 and H_2O evolution
Run 4	117	251–289	350–443	GC: Narrow T cut for organics below O ₂ evolution T TLS: Narrow T cut targeting suspected carbonate

*Due to the low volume of gas released by Rocknest in this temperature range, isotope data were not obtained for this run.

(see Table 1 and Materials and Methods). The exact mass of each Rocknest portion delivered to SAM is not measured by Curiosity, but tests on Earth are consistent with 50 ± 8 mg per portion (*16*).

Results and Discussion

Volatile Release

The volatile compounds observed in EGA typically reflect a combination of processes including desorption of trapped volatiles, mineral thermal decomposition, and chemical reaction during heating of the samples (17, 18). Pure minerals and chemicals produce volatile products at predictable temperatures; however, in natural mixtures, these temperatures can be strongly shifted by physical characteristics of the samples (e.g., grain size) and by interactions between mineral and chemical components (17).

All four Rocknest analyses yielded H_2O , SO_2 , CO_2 , and O_2 , in descending order of average abundance (Fig. 1 and Table 2). H_2O , CO_2 , and O_2 abundances are relatively consistent from run to run and track each other within experimental uncertainty, whereas SO_2 abundance is variable from run to run. Repeated observation of H_2O , CO_2 , and O_2 gas abundances with similar values suggests that differences in sample mass cannot explain the heterogeneity in SO_2 abundance, and thus the variability must be due to variation in the abundance of S-bearing minerals in different portions.

The H₂O observed in Rocknest EGA comprises a broad peak centered at ~300°C. Abundance estimates are ~1.5 to 3 weight percent (wt %) H₂O in the <150-µm fraction. The peak temperature and breadth of the H₂O release is most consistent with bound H₂O in amorphous phases. Specifically, adsorbed H₂O, H₂O bound to amorphous phases (e.g., amorphous aluminosilicate materials, nanophase ferric oxides and oxyhydroxides), interlayer H2O from phyllosilicates, dehydration of several salts, and dehydration of ferric oxyhydroxides could have contributed to the lower-temperature H₂O release (Fig. 2). Higher-temperature H₂O could result from more tightly bound structural H2O and/or OH in minor minerals present below the CheMin detection limit, as well as H2O occluded in minerals and glasses. However, if the water detected was released from a single host mineral, CheMin should have detected that host mineral. The lack of observed hydrous crystalline phases in the <150-µm fraction (15) implies that H₂O/OH is derived from the amorphous component. H₂O concentrations in the amorphous component are estimated to be 3 to 6 wt % H₂O.

Unlike the situation for H_2O , calculated abundances of carbonate inferred from CO_2 released, sulfate minerals from SO_2 , and oxychloride compounds (e.g., chlorate or perchlorate) from O_2 would all be at or below the detection limits of CheMin, affirming the complementarity of SAM and CheMin on Curiosity. The data do not allow specific determination of whether host materials for these evolved gases exist as crystalline phases at abundances less than the 1 to 2% detectable by CheMin, or whether these volatiles are also hosted in amorphous materials in the <150-µm fraction. However, the release temperatures of the gases suggest fine-grained and/or poorly crystalline materials as the hosts, as discussed below.

The CO₂ released from all four Rocknest runs comprises two major peaks, at ~400° and ~510°C, and a lower-temperature shoulder, which can be fit as two discrete releases at ~225° and ~295°C (Fig. 3). The two major CO₂ peaks together comprise >70% of the CO₂ released. The highest-temperature CO₂ release is consistent with the thermal decomposition of siderite (*19*). If this peak is due entirely to siderite decomposition, it would imply ~1 wt % siderite in the Rocknest <150-µm fraction. A second possibility is that this release evolved from the thermal decompo-

sition of nanophase magnesite, because nanophase carbonates decompose at temperatures at least 100°C lower than 2- to 50-µm-sized particles (17, 20). Calcite is not a likely candidate because its decomposition begins at 685°C, a temperature substantially higher than that of the vast majority of CO2 released from the Rocknest <150-µm fraction. A third possibility is that the two major CO2 peaks correspond to CO2 chemically evolved from two mineral phases, such as siderite and magnesite, by reaction with HCl (18), which is observed in the Rocknest EGA (Fig. 1B), likely from decomposition of a perchlorate salt (see below). Most likely, all three factors (grain size, mineralogy, and reaction with HCl) contribute to the two major CO2 peaks.

The concurrent evolution of CO_2 and O_2 from Rocknest suggests that organic carbon (i.e., C con-



tained in molecules having C, H, O, N, and/or S) oxidized within SAM is another potential CO_2 source. Such reduced carbon might be indigenous to Mars, delivered from space in the form of interplanetary dust particles and micrometeorites, or part of the instrument background. Molecular fragments from a reagent carried to Mars for use in a SAM wet chemistry experiment, MTBSTFA (*N*-methyl-*N*-tert-butyldimethylsilyl-trifluoroacetamide), have been identified in both empty-cup blank and Rocknest runs. A small fraction of CO_2 (<10% of the total CO_2 observed) from combustion of these organics is suggested by the amount of the most abundant MTBSTFA-related products, mono- and bi-silylated H₂O (*tert*-butyldimethylsilanol and 1,3-

bis(1,1-dimethylethyl)-1,1,3,3-tetramethyldisiloxane, respectively). These sources are discussed below in conjunction with δ^{13} C measurements and organic molecular analyses.

Although the intensity and shape of traces attributable to SO₂ vary between the Rocknest samples, overall, the EGA traces indicate that SO₂ evolves from ~450° to 800°C. Two main peaks are observed, at ~500° to 550°C and ~700° to 750°C (Fig. 1). Possible sources of the evolved SO₂ include the thermal decomposition of sulfates and/or sulfites, oxidation of sulfides, and S adsorbed onto particle surfaces, which can persist to relatively high temperatures (21). Laboratory EGA under SAMlike conditions shows that iron sulfates produce

Table 2. Abundance of major species released upon heating of Rocknest as measured with the SAM QMS. Errors reported for molar abundances are the 2σ SD from the mean of calculations done with different m/z values for the same species. Weight % values were calculated with an estimated sample mass of 50 ± 8 mg (2 σ), with errors propagated including the uncertainty in molar abundance (14).

	M	olar abundances (µmo	l)	
	Run 1	Run 2	Run 3	Run 4
CO ₂	8.3 ± 2.0	10.8 \pm 2.6	10.1 \pm 2.4	10.4 \pm 2.5
50 ₂	$\textbf{2.9} \pm \textbf{0.2}$	$\textbf{13.7} \pm \textbf{1.9}$	21.7 ± 2.9	$\textbf{10.5}~\pm~\textbf{1.4}$
H ₂ O	$\textbf{43.3} \pm \textbf{10.7}$	66.5 ± 16.2	54.5 ± 9.9	55.9 ± 11.9
0 ₂	$\textbf{3.0} \pm \textbf{0.4}$	$\textbf{5.1} \pm \textbf{0.6}$	$\textbf{3.7} \pm \textbf{0.4}$	$\textbf{3.7}~\pm~\textbf{0.5}$
		Sample weight %		
	Run 1	Run 2	Run 3	Run 4
CO2	0.7 ± 0.2	$\textbf{1.0} \pm \textbf{0.3}$	0.9 ± 0.3	$\textbf{0.9} \pm \textbf{0.3}$
SO3 equiv.	$\textbf{0.5}~\pm~\textbf{0.1}$	$\textbf{2.2} \pm \textbf{0.5}$	$\textbf{3.5} \pm \textbf{0.7}$	$\textbf{1.7} \pm \textbf{0.3}$
H ₂ O	$\textbf{1.6} \pm \textbf{0.5}$	$\textbf{2.4} \pm \textbf{0.7}$	$\textbf{2.0} \pm \textbf{0.5}$	$\textbf{2.0} \pm \textbf{0.5}$
ClO4 equiv.	$\textbf{0.3}\pm\textbf{0.1}$	$\textbf{0.5}\pm\textbf{0.1}$	$\textbf{0.4}~\pm~\textbf{0.1}$	$\textbf{0.4} \pm \textbf{0.1}$



Fig. 2. Water release from Rocknest compared to laboratory measurements of mineral breakdown. Water release versus temperature for Rocknest <150- μ m fraction measured by the SAM QMS. Arrows indicate temperatures of water-release peaks determined by laboratory analysis for select hydrous minerals phases under conditions similar to that in SAM (17).

SO₂ at temperatures consistent with Rocknest observations. Mg- and Ca-sulfates, including the anhydrite observed in Rocknest <150-µm fraction by CheMin (15), have SO₂ evolution temperatures too high to explain the observed SO₂. The hightemperature tail of O₂ peak at ~460°C is coincident with the initial rise of SO₂. This observation and SAM EGA detections of small amounts of H₂S, OCS, and CS₂ evolved at temperatures close to the higher-temperature SO₂ release (Fig. 1) support the hypothesis that oxidative reactions of reduced sulfur phases during heating also contributed to the evolved SO₂.

The onset of release of O_2 correlates with the release of chlorinated hydrocarbons (Fig. 1), suggesting that an oxychloride compound, such as a chlorate or perchlorate, is the source of the oxygen and chlorinated volatiles. Laboratory evaluation of various perchlorates and chlorates has not identified an unequivocal match to the SAM Rocknest data, but Ca-perchlorate provides the most reasonable match, with Fe- and Mg-bearing perchlorate, various chlorates, and mixtures with other minerals that may affect decomposition temperatures (22–24) as other possibilities.

The likely detection of an oxychloride compound by SAM is consistent with perchlorate observed in samples analyzed by the Wet Chemistry Laboratory (WCL) and the Thermal and Evolved Gas Analyzer (TEGA) instrument on the Phoenix lander (25), which observed a similar O_2 release during analysis of a soil sample. On the basis of WCL results, Phoenix soils were calculated to contain 0.4 to 0.6 wt % ClO_4^- (25). If all of the oxygen detected by SAM resulted from perchlorate decomposition, the estimated ClO₄⁻ abundance in the Rocknest <150-µm fraction (Table 2) would be comparable to the abundances observed by Phoenix. This abundance does not account for all of the chlorine detected by Curiosity's Alpha Particle X-ray Spectrometer (APXS) (14), implying the presence of other chlorinebearing species at Rocknest.

Chlorine has been detected in every soil ever analyzed on Mars-in situ at the equatorial and mid-latitude sites of the two Viking landers (2)and from equator to mid-latitude by remote sensing from Mars Odyssey spacecraft (26). The process of perchlorate formation is believed to start with the oxidation of chlorine in gas-phase reactions in the atmosphere (27), various chlorine oxides produced by energetic electrons from galactic cosmic-ray interaction with the surface ice (28), heterogeneous mineral-catalyzed photooxidation of surface chlorides (29), or on airborne dust. The global presence of chlorine, and the detection of perchlorate in fines at two very different locations (Phoenix and Curiosity landing sites), support the hypothesis that perchlorates are globally distributed in the regolith of Mars. Perchlorates can be a sensitive marker of past climate and a potential terminal electron acceptor for martian biota. They may also form liquid brines under current martian conditions and contribute to the oxidation and transformation of martian

organic matter when exposed to ionizing radiation at or near the surface or during analytical processing. Thus, a widespread presence of perchlorate salts, spatially and temporally, would have an important bearing on understanding habitability, organic matter preservation potential, and organic biosignature detection on Mars.

Isotopes

The results of the TLS isotopic analyses at Rocknest are summarized in Table 3. The strategy for the different temperature ranges of evolved gas sent to the TLS was developed with the EGA data to iteratively design experiments that selectively focused on various gas releases. For example, run 3 captured the bulk of the H₂O peak, and run 4 focused on the first of the two large CO₂ peaks. The EGA data were also used to constrain the isotopic composition of C in CO₂ and S in SO₂.

Hydrogen in all Rocknest samples is highly enriched in deuterium compared to terrestrial materials (Fig. 4), with δD values ranging from ~+3900 to +7000 per mil (‰). Run 3 should be most representative of the "bulk" of the water in Rocknest, with a value of ~+7000‰. However, significant variation in the δD value with temperature is observed, with the lower-temperature cut having the highest δD value and the highesttemperature cut having the lowest.

The δD values measured in the Rocknest <150-µm fraction are consistent with the SAM TLS measurements of water in the martian atmosphere taken before Rocknest, which show a δD value of +5000 ± 1000‰ (*30*). In addition, the Rocknest δD values are within the range of values observed by remote-sensing analysis of the martian atmosphere (*31*), where telescopic measurements from Earth have previously suggested a reservoir enriched in D by a factor of ~5 over terrestrial values. The D-enriched values in a martian soil are also consistent with D-enriched H₂O observed in both bulk (*32*) and single grains (*33*) in martian meteorites.

The close match between the δD values from H₂O in both atmospheric gas and Rocknest suggests that the H2O-rich phases in the amorphous material were formed either in direct contact with the atmosphere or through interaction with volatiles derived from it. The variation of δD value with temperature may either record longterm variation of D/H through time or represent seasonal variations reflecting changes in the water cycle. It is likely that the water evolved at the lowest temperatures represents water in active exchange with the present atmosphere, whereas the higher-temperature releases could represent water from a more ancient time. Telescopic measurements suggest that there could be large variations in atmospheric δD value with water content of the atmosphere and season (31), and such variations may be reflected in the Rocknest results.

Like hydrogen in $\rm H_2O,\ ^{13}C\text{-enriched}\ \rm CO_2$ has also been observed in the atmosphere at Gale

crater with SAM TLS (30) and QMS (34), with an average δ^{13} C value measured to date of ~+46‰. Unlike hydrogen, however, the CO2-bearing phases in Rocknest soil do not fully reflect this ¹³Cenriched atmospheric value. Rather, δ^{13} C values of CO₂ evolved from Rocknest and analyzed by TLS range from -6 to +20% (Table 3), and estimates of δ^{13} C over the two major CO₂ peaks using QMS data average $\sim +18 \pm 10\%$, consistent with the TLS results. These values overlap with δ^{13} C values from both carbonates and refractory/ reduced carbon in martian meteorites (Fig. 5). Consistent with the above discussion of several possible CO2 sources in SAM analyses of Rocknest, the δ^{13} C compositions likely reflect mixing of multiple carbon sources. The concurrent evolution of CO2 and O2 from Rocknest suggests that partial combustion of reduced carbon could contribute to evolved CO₂. δ^{13} C associated with the CO₂ release between 250° and 450°C might reflect some contribution from this combusted carbon. Previous studies of martian meteorites have shown that reduced carbon is present either as an indigenous component or from exogenous meteoritic input (8, 10-12).

The Rocknest δ^{13} C values suggest a hint of $^{13}\mathrm{C}$ enrichment, consistent with $\delta^{13}\mathrm{C}$ values observed in martian meteorite carbonates. Specifically, the data from run 4, which most closely capture the largest CO₂ peak, has a δ^{13} C value of $\pm 10\%$, which is similar to carbonate measured in the Nakhla meteorite (35). This value is lower than would be expected for carbonate formed from the modern atmosphere as measured by SAM TLS (30). It is possible that this CO₂ release is a mixture of carbonate-derived CO2 with a high δ^{13} C value and CO₂ depleted in ¹³C and thus does not reflect the true carbon isotopic composition of the carbonate. It is also possible that the carbonate does have low δ^{13} C values as observed in some of the martian meteorites, suggesting that the atmosphere has changed through time (36). Overall, the data support a minor amount of carbonate in martian soil derived from atmosphere interaction with only transient water (37).

The sulfur isotopic composition of SO₂ released during run 4 was determined from QMS data at a mass-to-charge ratio (m/2) of 64, 65, and 66. The Rocknest <150-µm fraction, including analyses of both of the major SO₂ evolution peaks,



Fig. 3. Deconvolution of CO₂ release from Rocknest. Rocknest run 2 CO₂ (mass 45) versus temperature (red). Gray peaks are Gaussian fits to overall CO₂ release that sum to mass 45 fit (blue line). CO_2 fractions in each of the four peaks are 0.07, 0.22, 0.41, and 0.30, respectively.

 Table 3. Isotopic composition of volatiles released upon heating of Rocknest as measured with

 the SAM TLS.
 Blank cup corrections have been applied as described in materials and methods.

Rocknest run	T range sampled (°C)	δ^{13} C in CO ₂ (‰)	δD in H ₂ O (‰)
Run 3	234–425	-6 ± 14	7010 ± 66
Run 4	350-443	20 ± 10	$4250~\pm~60$
Run 2	440-601	3 ± 9	$\textbf{3870}\pm\textbf{60}$

have $\delta^{34}S_{VCDT}$ of $0 \pm 10\%$, consistent with sulfur isotopic compositions measured in martian meteorites (38, 39).

Organic Matter

Chlorohydrocarbons comprising chloromethane (CH₃Cl), dichloromethane (CH₂Cl₂), trichloromethane (CHCl₃), and chloromethylpropene (C₄H₇Cl) were detected during SAM GC-MS analyses (Fig. 6 and Table 4). Chloromethanes detected by SAM

Fig. 4. Tunable laser spectrometer data showing hydrogen isotope enhancement in Rocknest. Section of a single spectrum (60 s integration) downloaded from Curiosity (black) for the Rocknest 3 sample run, showing large HDO line depth compared to calculated HITRAN spectrum (red) based on terrestrial SMOW water isotope ratios. The HDO line is ~4 times the depth of that predicted for SMOW, so that the D/H ratio is ~8 times that of SMOW, corresponding to a δD value of ~7000‰, as reported.

in runs 1, 2, and 4 were at ~nanomole levels and above SAM background. Run 3 produced lower abundances of chloromethanes (typically observed at <300°C) because only a high-temperature cut of evolved gases were transferred to the GC. Minor amounts of HCN, CH₃Cl, CH₂Cl₂, and CHCl₃ are also observed in SAM EGA data (Fig. 1B). The abundance of these species is more than two orders of magnitude lower than that of the most abundant volatile released—H₂O.





Fig. 5. Carbon isotopes in relevant solar system reservoirs. Carbon isotopic composition of materials from Mars (44–46), Earth (47), and carbonaceous chondrite meteorites (48) for comparison the values measured in Rocknest and the martian atmosphere (30) by the Mars Curiosity Rover.

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The abundances measured by SAM are higher than the picomole levels (up to 40 parts per billion) for chloromethane and dichloromethane previously measured by the Viking pyrolysis gas chromatography-mass spectrometry (GC-MS) instruments after heating the samples of scooped fines up to 500°C (13). Biemann et al. (13) attribute the Viking results to chlorohydrocarbons derived from cleaning solvents used on the instrument hardware, not from the martian samples themselves. Recently, Navarro-González et al. (40) suggested that these chlorohydrocarbons may have formed by oxidation of indigenous organic matter during pyrolysis of the soil in the presence of perchlorates, but Biemann and Bada (41) disagree with this conclusion.

The absence of detectable chlorohydrocarbons in the SAM blank run indicates that the chlorohydrocarbons measured at Rocknest are not directly attributable to the SAM instrument background signal. However, the associated release of chloromethanes, O2, and HCl strongly suggests that these chlorohydrocarbons are being produced within SAM by chlorination reactions involving an oxychloride compound in the Rocknest <150-µm fraction and an organic carbon precursor (23). Three sources for the organic carbon of this reaction are possible: (i) terrestrial sources within the SAM instrument or the Curiosity sample chain; (ii) exogenous carbon in the martian surface materials derived from infalling meteoritic carbon; and (iii) martian indigenous organic matter. A feasible explanation involves terrestrial carbon derived from the MTBSTFA, whose reaction products were identified in both the blank and soil EGA and GC analyses. On the basis of laboratory pyrolysis GC-MS experiments, pyrolytic reaction of martian Cl with organic carbon from MTBSTFA in SAM can explain the presence of the chloromethanes and chloromethylpropene detected by SAM. However, we cannot rule out the possibility that traces of organic carbon of either martian or exogenous origin contributed to some of the chlorohydrocarbons measured by SAM at Rocknest.

Overall, SAM analyses indicate that martian fines contain a number of materials with bound volatiles that can be released upon heating. These volatile-bearing materials are likely very finegrained and associated with the amorphous component of martian regolith. The fines could be a good source of water, CO2, and other volatiles to be leveraged by future human explorers on Mars. Isotopic compositions support an atmospheric source of the water and possibly CO2, consistent with previously proposed formation mechanisms for carbonate and perchlorate in the fines that involve interaction with the atmosphere. Although martian organic matter was not definitively detected, the presence of materials that produce substantial amounts of oxygen upon heating suggests that detection of such compounds in martian soils will be difficult with pyrolysis techniques. The fines on Mars reveal a complex history, reflecting global, regional, and local-scale processes.

Materials and Methods

SAM Instrument Overview and Operations

The SAM instrument suite supports the MSL mission and sits inside the Curiosity rover at Gale crater on Mars. The SAM instruments are a QMS, a TLS, and a six-column GC with thermal conductivity detectors (TCDs) (7). These three instruments share a solid sample- and gasprocessing system to generate complementary compositional and isotopic observations on each sample delivered by the rover's Sample Acquisition, Sample Processing and Handling (SA/SPaH) hardware to a SAM solid sample inlet tube or ingested directly through a gas inlet. Before each analysis, the oven for solid samples, gas-processing system, and instruments are purged with helium and heated to release any residual volatiles in the system to, in effect, precondition and clean SAM.

Scooped, sieved <150-µm-particle-size fraction, and portioned (<76 mm³) sediments of the Rocknest aeolian drift were heated to thermally evolve gases for processing and analysis. These volatiles are the result of the following processes often happening concurrently: (i) desorption of surface-adsorbed volatiles and anions, (ii) mineral thermal decomposition, and (iii) thermochemical reactions among chemical components (18). When organic materials are present in solid samples, they might be desorbed at low temperatures (usually below 320° C), as is the case for small individual molecules; undergo pyrolysis (i.e., thermal bond cleavage) at higher temperatures; or contribute to thermochemical reactions (at all temperatures) (42).

SAM performs EGA with the QMS and isotope measurements of evolved gases with both the QMS and the TLS, with the latter being sensitive to CO₂, water, and methane (methane detection capability was not used during Rocknest runs). Organic analyses can be performed with the QMS alone or when it is coupled to the GC. SAM heated each Rocknest sample to ~835°C at a rate of 35°C/min with a He carrier gas flow rate of ~ 0.77 standard cm³ per minute and at an oven pressure of ~25 mbar. The SAM QMS analyzed abundances of gases across the entire temperature range, while selected temperature ranges of the evolved gases in each run were sent to the TLS and GC for analysis (Table 1). For each portion ingested by SAM (called runs 1 through 4), the gases evolved across the selected range of temperatures were accumulated inside the TLS Herriott cell, where hydrogen isotopes in water and carbon isotopes in CO_2 were analyzed in the bulk gases. Evolved gases from a different selected temperature range sent to the GC were first captured on the hydrocarbon trap held at 5°C. The trap was subsequently heated to ~300°C under He flow, and the desorbed gases were sent to a GC channel (composed of an injection trap, MXT-CLP column, and TCD) suited for the analysis and separation of volatile organic compounds. The TCD and QMS provide detection and identification of the chemical molecules eluted from the GC.

Solid Sample Analysis Details

At Rocknest, fines from scoop #5 were delivered to SAM four times and placed in separate quartzglass cups. The sample in each cup was flushed with pure helium (99.999%) at ~ 25 mbar at ~0.77 standard cm³ per minute and heated at ~75°C for 15 min to release water adsorbed on mineral surfaces and minimize saturating the system with excess water. This early gas release was directly measured by the QMS. Each cup was heated in SAM oven #1 at a rate of ~35°C min⁻¹ to ~835°C, where the final temperature varied slightly due to different ambient Mars environmental conditions. All evolved gases from the 75° to 835°C range were sampled by the QMS



the Rocknest figure) were identified above measured background levels: 2, carbon dioxide; 3, sulfur dioxide; 4, hydrogen cyanide; 5, hydrogen sulfide; 6, chloromethane; 7, dichloromethane; 9, trichloromethane; 11, chloromethylpropene; 15, chlorobenzene. The following peaks are consistent with measured background levels: 1, carbon monoxide; 8, acetone; 10, acetonitrile; 12, benzene; 13, toluene; 14, *tert*-butyldimethylsilanol; 16, phenylethyne; 17, styrene; 18, 1,3-bis(1,1-dimethylethyl)-1,1,3,3-tetramethyldisiloxane; 19, trimethylsilylborate; and 20, biphenyl. See Table 4 for discussion of possible origins of each peak.

through a capillary flow restrictor. This experiment is referred to as evolved gas analysis and produced data in the form of pyrograms for individual ions defined by their m/z ratios. The main split of evolved gas was then passed through gas manifolds heated at 135°C to either the hydrocarbon trap for GC analysis or the TLS Herriot cell for isotopic and mixing-ratio measurements of H₂O, CO₂, or O₂, or vented to Mars. For each analytical run, different temperature cuts of gas were selected to go to the GC or TLS, but in no case was the gas sent to both at the same time. Temperature cuts for the GC and TLS are listed in Table 1.

Figures 2 and 3 show detailed analyses of the H₂O and CO₂ releases, respectively. The plot of H₂O release in Fig. 2 is generated with the QMS data from m/z 20, because the molecular ion for H_2O (m/z 18) is saturated in these runs. The release temperatures of various hydrous mineral phases marked on the plot are derived from laboratory measurements performed under conditions similar to those for the EGA in SAM. These were typically determined for single minerals, and mineral mixtures and grain-size effects can change these values. Nonetheless, the broad H₂O release peak is not clearly indicative of any one mineral phase. Figure 3 shows evolved CO₂ (m/z45) as a function of temperature for Rocknest run #2, for which four discrete peaks can be fit to Gaussian peak shape to model the summed CO2 release. The integrated areas for the fitted

peaks are used to quantify the contributions from each release event to the total abundance of evolved CO₂. Although it is not possible to assign definitively specific species to each of the four peaks, oxidized organics (terrestrial or martian) and several types are carbonate are discussed in the text as likely contributing to the CO₂ peaks, especially the two major peaks.

Methods for Molar Abundance Calculations

Molar abundances were primarily computed by referencing measurements on Mars to pre-launch SAM calibration runs of quantified samples of calcite (CaCO₃) and a hydrated iron sulfate (FeSO₄·4H₂O) (6). A calibration factor [counts per second (cps)/mol] was determined for the relevant m/z value in laboratory standard runs by integrating under the evolved gas curve and dividing by the number of moles evolved from the sample, assuming complete decomposition.

Under nominal SAM operating conditions, the most abundant ion of major species (e.g., m/z18 for H₂O) often saturates the detector. Given a fixed detector range, this makes the instrument more sensitive to low-abundance materials. To quantify amounts with high abundances, doubly ionized molecules, ion fragments, and isotopes were used to calculate evolved gas abundances. For example, m/z 44 (CO₂⁺) saturated the detector for most of the Rocknest runs because the amount of CO₂ evolved exceeded detector lim-

Table 4. Inorganic and organic volatile species detected by the SAM GC-MS upon heating of Rocknest and their possible origins. MTBSTFA (*N*-tert-butyldimethylsilyl-*N*-methyltrifluoroacetamide) and DMF (dimethylformamide) are both carried within SAM for future derivatization experiments. Tenax TA is a porous polymer adsorbent resin used to concentrate organic compounds on the SAM hydrocarbon traps. Those sources that are known to be terrestrial in origin are shown in italics. Compounds in bold are observed above measured background levels.

Peak no. from Fig. 6	Compound	Possible origin(s)
1	Carbon monoxide	Unknown
2	Carbon dioxide	Martian carbonates/carbon?, MTBSTFA or DMF
3	Sulfur dioxide	Martian S-bearing minerals
4	Hydrogen cyanide	MTBSTFA + perchlorate or high-T martian source?
5	Hydrogen sulfide	Product of S-bearing minerals
6	Chloromethane	MTBSTFA or martian carbon? + perchlorates
7	Dichloromethane	MTBSTFA or martian carbon? + perchlorates
8	Acetone	MTBSTFA or DMF
9	Trichloromethane	MTBSTFA or martian carbon? + perchlorates
10	Acetonitrile	MTBSTFA or DMF
11	Chloromethylpropene	MTBSTFA + perchlorates
12	Benzene	Tenax TA
13	Toluene	Tenax TA
14	tert-Butyldimethylsilanol	MTBSTFA + water
15	Chlorobenzene	$HCl + Cl_2 + Tenax TA$
16	Phenylethyne	Tenax TA
17	Styrene	Tenax TA
18	1,3-bis(1,1-dimethylethyl)- 1,1,3,3-tetramethyldisiloxane	MTBSTFA + water
19	Trimethylsilylborate	MTBSTFA + glass beads
20	Biphenyl	Tenax TA

its. To quantify the abundance of CO₂, m/z 12 (C⁺), m/z 22 (CO₂²⁺), and m/z 45 and 46 (isotopologs of CO₂) were used instead of m/z 44. The number of moles of CO₂ evolved from Rocknest samples was determined by taking an average of the total areas calculated for each m/z listed above. The error was calculated as 2σ SD from the mean. For H₂O, m/z 19 and 20 were used because m/z17 and 18 were saturated, in both Rocknest and earlier laboratory calibration runs. To calculate SO₂ abundances, m/z 66 and 50 were used (isotopologs of SO₂ and SO) because m/z 64 and 48 saturated in calibration runs.

There are two additional complications encountered when calculating H₂O abundances. First, FeSO4·4H2O begins to lose H2O as soon it is exposed to lower pressures, which is before the QMS begins monitoring gas evolution. Fortunately, there is a very distinct and repeatable H₂O release at slightly higher temperatures (~200°C) with a measurable mass loss. This H2O release was used to calibrate Rocknest data. Second, Mars has a much higher D/H ratio than Earth, which can affect peak integration values, especially for m/z 19 (HDO). We corrected for this effect when calculating the water abundances on Mars using QMS data during EGA. The differences in the δ^{18} O, δ^{13} C, and δ^{34} S isotopic values between Earth and Mars are small compared to the other uncertainties involved in these abundance calculations and, therefore, were not included.

Oxygen abundance values were calculated in a slightly different way because none of the minerals run during prelaunch testing released O_2 . However, a separate prelaunch characterization run was done with an equimolar gas mix of O_2 , CO_2 , Ar, and N_2 . These data were used to determine relative calibration factors for the major atmospheric species, as discussed in Mahaffy *et al.* (*34*). Such calibration factors yield a value for relative ionization rates for O_2 and CO_2 at equivalent abundances, which were applied to the EGA data to determine calibration factors for O_2 in cps/mol.

Isotope Data Reporting Convention

All isotope results are presented in standard delta notation (δD , $\delta^{13}C$, $\delta^{34}S$) with respect to Vienna standard mean ocean water (VSMOW) for hydrogen, Vienna Pee Dee belemnite (VPDB) for carbon, and Vienna Cañon Diablo troilite (VCDT) for sulfur. Here, $\delta(\%) = [R_{\text{meas}}/R_{\text{std}} - 1] \times 1000$, where R_{meas} is the measured isotope ratio (heavy/light), and R_{std} is the ratio of the relevant reference standard.

TLS Operational Conditions and Data Reduction

TLS is a two-channel tunable laser spectrometer that uses direct and second harmonic detection of infrared laser light absorbed after multipassing a sample cell. For the results reported here, the sample cell path length is 43 passes of a ~19.5-cm cell length, or 840 cm. TLS scans over individual rovibrational lines in two spectral regions near 2.78 μ m; one centered at 3590 cm⁻¹ for

 CO_2 isotopes, and a second centered at 3594 cm⁻¹ for both CO2 and H2O isotopes. The lines used in both regions have no discernable interferences. In the 3594-cm⁻¹ region, the CO₂ and H₂O lines used interleave across the spectrum without interference, allowing determination of isotope ratios across widely varying CO2 and H2O abundances in both atmospheric and evolved gas experiments. For carbon isotopes, the values given are the weighted means of two separate pairs of lines, one pair from each region. Further data-processing details and calibration are described in the supplementary materials accompanying Webster et al. (30). Figure 4 is a good example of a TLS flight spectrum used for isotope ratio measurement, showing the large enhancement of the HDO line over that expected (HITRAN database) for terrestrial water.

The TLS sample cell (Herriott cell) is first pumped out by using the SAM turbomolecular pump with empty cell pressures of CO₂ and H₂O that are insignificant compared to either EGA or "blank cup" runs. At some predetermined time during either the four EGA or single blank cup runs, temperature cuts of evolved gas are sent to the TLS (Table 1), where they produced Herriott cell pressures of 4 to 9 mbar of principally helium, with evolved water and carbon dioxide as minor components. Before the Rocknest EGA runs reported here, a blank cup run was conducted under the identical conditions (He flow, temperature cut, pressure, etc.) but without solid sample in the pyrolysis oven. Resulting signals (abundances) for CO₂ and H₂O were not large compared to the Rocknest abundance values, and isotope values are similar to those of the samples, but nonetheless it is appropriate to make a correction. This correction was weighted by the relative abundance of the gas of interest. Specifically, the H₂O abundances in the blank were $\sim 3\%$ of the total water measured, and the CO₂ abundances were ~5 to 10% of the total CO2 measured in Rocknest aliquots. Blank cup values for measured δ^{13} C in CO_2 was –80‰ and δD in H2O was 3880‰. The TLS measured results are therefore the combination of an underlying background (blank cup) contribution and the Rocknest sample contribution in proportions dependent on the relative abundances of water and carbon dioxide from each. Because the blank cup abundances of these gases is much smaller than those evolved from the Rocknest samples, the corrections to the measured isotope ratios are usually small. Given the small abundance of H_2O and CO_2 in the blank, the blank isotope values have relatively large uncertainties, and these are propagated through the correction calculation. The results given in Table 3 are the Rocknest sample isotope ratios after correction for the blank cup values.

QMS Isotope Value Calculations

Isotope ratios from QMS data obtained through EGA of solid samples are computed from the time-integrated signal at isotopologs for the compounds of interest. For CO₂, the ${}^{13}C/{}^{12}C$ ratio is

determined from m/z 45 and 46, with correction for oxygen contributions based on the most relevant TLS measurements of the oxygen isotopic composition in CO₂. The values of δ^{13} C given in the main text were computed on the basis of an estimated δ^{18} O value of $-28\%_0$, as determined by the TLS for CO₂ released during Rocknest 4, because the gas sample sent to TLS during this run was the most representative of the peak CO₂ release from Rocknest samples. The carbon isotopic composition of CO₂ during these runs cannot be calculated from the more typical pairs of m/z 12-13 or 44-45 because of interference from MTBSTFA background at m/z 13 and detector saturation at m/z 44.

For SO₂, the ³⁴S/³²S ratio is computed from m/z 64-66, with correction for oxygen contributions (assumed δ^{18} O of 50 ± 5‰) based on TLS measurements of the oxygen isotopic composition of both CO₂ and H₂O in the martian atmosphere. For both CO₂ and SO₂, ¹⁷O contributions were estimated from the assumed δ^{18} O on the basis of a Δ^{17} O of 0.32‰, the average for martian silicates (*43*).

GC-TCD and GC-MS Operational Conditions

The SAM GC hydrocarbon trap consists of three layers: nonporous silica beads, Tenax TA adsorbant (porous 2.6-diphenylene oxide polymer resin), and Carbonsieve G adsorbant (graphitized carbon) (7). Gases were passed through the hydrocarbon trap that had been cooled to 5°C and selectively condensed onto glass beads or adsorbed on the basis of volatility, molecular size, and chemistry. Thermal desorption at 300°C for 4 min under He (0.5 standard cm³ per minute, 0.9 bar) released analytes from the hydrocarbon trap in the opposite direction. Analytes then collected on a Tenax TA injection trap of the GC. The injection trap was then flash heated to 300°C. All four Rocknest GC analyses used the MTX-CLP column (30-m length, 0.25-mm internal diameter, 0.25-µm film thickness), which has a polydimethylsiloxane with phenyl and cyanopropyle film (7) and is designed for separating mid-molecular weight hydrocarbons. The column temperature was programmed from 50° to 220°C at 10°C min⁻¹ and He carrier gas was held at a constant column inlet pressure of 0.9 bar. Gases eluting from the GC were nondestructively detected by the TCD and then ionized by electron impact at 70 eV in the QMS source, which fragmented molecules in a predictable fashion. The OMS scanned for ions in a *m/z* range of 2 to 535 using the Smart Scanning algorithm previously described (7). The GC-TCD result is shown as a single chromatogram for all detected molecules in which the retention time (x axis) is normalized to the GC retention time. GC-MS results are given as individual ion chromatograms and mass spectra for isolated peaks, in which the mass spectra were generated with a custom dataprocessing program.

The identification of the chlorohydrocarbons in the GC-MS data was based on the retention times and comparisons of the mass fragmentation patterns to the NIST11 library. The abundances of the chlorohydrocarbon compounds ($\sim 10^{-2}$ to 1 nmol) were determined by comparing the fitted peaks in the Rocknest data to those of known amounts of hexane measured during preflight calibration runs on SAM GC, corrected for differences in ionization efficiency (*34*). These abundances are also corrected for the gas fraction sent to the hydrocarbon trap with EGA data (Table 1).

EGA, GC-TCD, and GC-MS Background Measurements

Within SAM, there are several common sources of organic molecules that are typical of similar, nonflight instruments. These include (i) wet chemistry reagents used for derivatization (i.e., MTBSTFA and DMF) and thermochemolysis [i.e., tetramethylammonium hydroxide in methanol (TMAH)] and their breakdown products; (ii) the Tenax TA, a porous, 2.6-diphenylene oxide polymer resin adsorbant that slowly degrades with use into benzene, toluene, tropylium ion, biphenyl, and other single-aromatic ring structures; and (iii) the polymer films inside the capillary columns that promote selective molecular separation. The molecules from these sources are known and make up what is referred to as instrument background signal, which is expected to change over the course of the SAM instrument's lifetime and is continuously monitored by analysis of empty sample cup or "blanks." Contributions of aromatics from the Tenax TA traps to TCD and QMS (post-GC) data have been determined on the basis of their absence in EGA data. The blank analysis run before Rocknest fines was the first analysis performed since prelaunch testing, and it showed the presence of DMF, MTBSTFA products, Tenax TA products, and terrestrial water (Table 4). TLS measurements of D/H in water indicated that further heating and purging of the system largely eliminated the terrestrial water.

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Supplementary Materials

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Volatile, Isotope, and Organic Analysis of Martian Fines with the Mars Curiosity Rover

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