Contents

Frontispiece: Mars 2020 Sample Collection Map ................................................................. i
Preface ..................................................................................................................................... iii
Mars 2020 Returned Sample Science Participating Scientists ........................................ iv
Explanatory Notes for Mars 2020 Initial Reports ................................................................. v
Cite as, Revision History, Acknowledgements ................................................................. xi
M2020-120-1 WB1 ................................................................................................................ 1-1
M2020-164-2 Roubion .......................................................................................................... 2-1
M2020-190-3 Montdenier ................................................................................................. 3-1
M2020-196-4 Montagnac ................................................................................................. 4-1
M2020-262-5 Salette .......................................................................................................... 5-1
M2020-271-6 Coulettes ..................................................................................................... 6-1
M2020-295-7 Robine .......................................................................................................... 7-1
M2020-337-8 Malay ............................................................................................................ 8-1
M2020-371-9 Hahonih ........................................................................................................ 9-1
M2020-374-10 Atsah .......................................................................................................... 10-1
Preface

A central goal of the Mars 2020 mission is collection of scientifically-selected martian rock, regolith, and atmosphere samples for possible return to Earth by future missions. This is an ambitious undertaking: Perseverance carries 38 sample tubes and 5 contamination-knowledge “witness tubes” to be filled over the course of the mission. Using the rover’s instruments, high priority outcrops must be identified, and the characteristics of each acquired sample and its surroundings investigated. With such a large number of samples to acquire and with an anticipated short interval between their acquisition, it is important to record in near real-time what the Mars 2020 Science Team has learned about each sample. The Mars 2020 Initial Reports series fills this need. A sample’s Initial Report is a set of field notes recording what is known about the sample and how it fits into its surroundings. The goal is to have each report completed within three weeks of sample acquisition, long enough to have downlinked, processed, and completed an initial high-level interpretation of the data, but short enough that the sample is still fresh in the team’s memory. Just like conventional field notes, the Initial Report is not updated after completion. The Initial Report is therefore preliminary, and the interpretations and even some of the underlying measurements will be revised and described more fully in detailed follow-on publications. The goal of the Initial Reports collection is to provide a timely, routine, systematic and preliminary narrative description of all the acquired samples in a single compendium. The Initial Reports complement the Sample Dossiers - a compilation of the digital records related to each sample. The Initial Reports series facilitates intercomparison among samples already collected as well as rocks under consideration for sampling over the course of the mission. It may also prove useful for prioritization of samples for Earth return and as a quick reference guide after samples arrive on Earth.

K.A. Farley, Project Scientist

K.M. Stack, Deputy Project Scientist
Mars 2020 Returned Sample Science Participating Scientists

The Mars 2020 Science Team includes individuals selected by NASA to guide sample selection, interpretation, and documentation, including preparation of these Initial Reports. For the Prime Mission these individuals are:

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Explanatory Notes for Mars 2020 Initial Reports

**Sample Designation**

The standardized format for sample designations is M2020-sol-N name. M2020 refers to the mission that collected the sample, sol indicates the mission sol of sample coring (sealing for witness tubes), N refers to the sequential number of the sample (or witness blank) acquisition, and name is the informal name applied to that sample and derived from the quadrangle in which the core was acquired. In the case of a witness blank acquisition, name is simply “WB” followed by the sequential number of the witness blank, starting with 1 and incrementing (e.g. WB1).

The same naming scheme is used for paired samples (i.e., two cores acquired with one associated STOP list execution; see below for definition of STOP list), except sol, N, and name all refer to the second member of the pair.

In typical usage the sample name alone is sufficient to uniquely identify a specific core. Because paired samples share a STOP list data set, when describing characteristics that refer to both members of paired samples, hyphenation of sample names will provide clarity: e.g., name1-name2.

**Date of Coring (or Exposure-Activation-Sealing)**

This is the calendar date on which the corer was placed on the rock surface to begin acquisition. Ordinarily, coring is completed in one sol. In the case of a witness blank, the dates of initial exposure, activation (inner seal puncture), and sealing are indicated.

**Estimated Volume Recovered**

The rock volume \(V_{rock}\) estimate is derived from the penetration depth of the volume probe, a rod inserted into the tube after sample acquisition (in practice the tube is manipulated while the rod is fixed in place). Volume is computed from the implied length of sample multiplied by an assumed cylindrical cross section corresponding to the coring bit inner diameter \(d=13.4\) mm. The core itself has \(d=13\) mm; the 0.4 mm difference in diameter between the bit and the core is assumed to be filled by cuttings and so contributes to acquired rock volume. A full length core is typically 6 cm long, though shorter lengths can be commanded and will sometimes be all that is acquired. Note that void spaces between core fragments may exist, such that the true sample volume may be less than this estimate.

**Coring Bit Number**

*Perseverance* carries 6 rock coring bits and 2 regolith bits. The bit number used here provides insight to usage history that may be relevant for contamination and cross-contamination assessment.

**Core Orientation**

Three pieces of information are needed to orient the core in the Martian geographic frame (also known as the SITE frame) (Fig. 1). First, two angles orient the core’s pointing vector. Second, the core’s roll is noted by marking the pre-drilling WATSON image of the core face. The core orientation pointing vector is estimated as equal to the orientation of the coring drill after pre-loading on the core target.
The core pointing vector is quantified by the azimuth and hade. The azimuth is defined as the clockwise angle of the horizontal projection of the core y-axis, \( \mathbf{cy} \), from geographic north, \( \mathbf{sx} \), where the \( \mathbf{cy} \)-axis lies in the martian geographic vertical plane. The hade is defined as the angle of the core face from martian geographic horizontal (\( \mathbf{sx}-\mathbf{sy} \) plane). These angles are obtained from coring drill and rover body ancillary orientation data at the time of pre-loading of the coring drill.

The core roll is quantified by an angle \( \alpha \), defined as the clockwise angle from the WATSON y-axis (\( \mathbf{wy} \)) to \( \mathbf{cy} \) as viewed in the pre-coring WATSON images. WATSON 6-7-cm standoff images are typically used for this purpose.

**Fig. 1.** Orientation system for cores collected by the Perseverance rover. Shaded cylinder is core with cutaway view in lower right quadrant. Martian geographic east, north, and down are the SITE axes, \( \mathbf{sx} \), \( \mathbf{sy} \) and \( \mathbf{sz} \), respectively. The pointing vector, \( \mathbf{cx} \), points into outcrop and is normal to the core face and the \( \mathbf{cy} \) axis lies in the vertical plane. The azimuth is the clockwise angle from geographic north of the projection of the \( \mathbf{cy} \)-axis onto the Martian horizontal (\( \mathbf{sx}-\mathbf{sy} \) plane). The hade is the angle of \( \mathbf{cx} \) from vertical. The core roll, \( \alpha \), is defined as the clockwise angle from the WATSON pre-coring image y-axis, \( \mathbf{wy} \), to \( \mathbf{cy} \); this direction is marked with an arrow on the WATSON image. Adapted from Butler (1992) *Paleomagnetism: Magnetic Domains to Geologic Terranes*, Blackwell Scientific, 319 pp.

**Sample Tube, Seal, and Ferrule Serial Numbers**

Every sample tube carries a set of serial numbers that will allow downstream identification (Figure 2). Sample Tube serial numbers are inscribed on the side of the tube and on the top of the bearing race (left). The number on the bearing race is partially covered by the seal once it is inserted. Each tube has a unique serial number and this number should be considered the primary identifier for a sample tube. Secondary identifiers are visible in Cachecam images (Figure 2B,C), including the Seal Assembly number and the Seal Ferrule serial number. The latter is non-unique and should not be used to identify a tube.
Fig. 2. Sample tube identifiers include the Sample Tube serial number inscribed on the top of the bearing race (A) and the side of the tube (B). Looking at the end of a sealed tube with Cachecam, the Seal Assembly number (red arrow) and the Seal Ferrule serial number (black arrow) are visible.
**ACA Temperature at Time of Sealing**

Sample tubes are sealed in the Adaptive Caching Assembly (ACA). Mechanisms in this subsystem are actively heated, so sealing occurs at temperatures higher than ambient conditions around the rover. The temperature recorded here is the average of the temperatures recorded on the Sealing Station and the Sample Handling Arm end effector. These temperatures are likely warmer than the sample and the associated head space gas.

**Estimated Rover-Ambient Pressure and Temperature at Time of Sealing**

Atmospheric temperature and pressure are obtained from MEDA. When MEDA data was not acquired on the same sol as sample sealing (often the case given energy limitations), the average values of pressure and temperature are estimated from the closest sol or sols. The reported temperature is the minimum of MEDA ATS 4 and 5, the sensors located 0.84 m above the surface.

**Estimated Amount of Martian Atmosphere Headspace Gas**

The estimated amount of headspace gas in moles \( (n_h) \) is computed from the ideal gas law and assuming the rover ambient temperature \( (T) \) and pressure \( (P) \) described above.

\[
n_h = \frac{P \cdot (V_{\text{tube}} - V_{\text{rock}})}{RT}
\]

where \( R \) is the gas constant. \( V_T \) is assumed to be 12 cm\(^3\). The actual temperature of the gas upon sealing is difficult to estimate because the Adaptive Caching Assembly (ACA) is substantially warmer than rover surroundings, typically by almost 100 K. For consistency we assume rover ambient temperature in this calculation, recognizing that it is an upper limit. A lower limit would be obtained by using the reported ACA temperature in the above equation.

**Abrasion Patch Name and Depth**

To minimize possible degradation of the sample, much of the data associated with a given sample is acquired on a 5 cm diameter abrasion patch acquired within a few tens of cm of the coring site, and in the same lithology. This IR entry gives the name of that patch and its depth relative to the highest topographic feature in the 5 cm diameter circle that was abraded.

**Anomalous Behavior**

This entry highlights noteworthy deviations from the standard sampling activities or their expected results.
Additional Notes:

STOP List

To expedite the sample collection and acquisition process, the Mars 2020 Team developed a minimum set of observations to be performed in association with each sample. This Standardized Observation Protocol, or STOP list, was encoded into an optimized sample sol path for efficient and repeatable execution. STOP list observations form the main data set for each sample’s Initial Report. A standardized set of observations on each sample permits a templatized Initial Report format. Although the STOP list is likely to evolve as the team learns what observations provide highest science value, it is the intention that the overall format be retained.
Instrument and Mission References


Cite as:


Revision History:

This is Release 2.1 of the Mars 2020 Initial Reports series. Initial Reports 5-10 were added to an earlier release that included only Initial Reports 1-4. Changes made in release 2.1 include correction to sample designations to reflect mission sol number of coring activity. Details and images were added to the Explanatory Notes to better describe sample serial numbers.

Acknowledgement:

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INITIAL REPORT
M2020-120-1 WB1

Witness Tube Designation: M2020-120-1 WB1
Date of Initial Witness Exposure: Pre-launch
Date of Activation: Pre-launch
Date of Sealing: 21-June-2021
Mars Time of Witness Tube Sealing: 16:25:55 LMST, Sol 120, Ls 61.4
Latitude (N), Longitude (E), Elevation: 18.43906919, 77.44940206, -2568.073
Sample Serial Numbers: Tube SN061; Seal SN147; Ferrule SN090
ACA Temperature at Time of Sealing: 40°C
Estimated Rover-Ambient Pressure and Temperature at Time of Sealing: 758 Pa, 245
Estimated Amount of Martian Atmosphere Headspace Gas: $2.2 \times 10^{-6}$ mol
Anomalous Behavior: None

November 27, 2021
K. A. Farley, Y. Goreva, R. Moeller and the Mars 2020 Science Team
**Summary Description**

*Perseverance* carries 5 Witness Tube Assemblies (WTA’s) with which to document the evolving rover contamination environment over the course of the mission (Moeller et al., 2020). Four of these “witness tubes” are identical and are located in sheaths just like sample tubes within the ACA. The fifth witness tube, WB1, was handled differently. During final pre-launch activities, this witness tube was activated (the inner seal was punctured to begin accumulation) and placed in the Bit Carousel. This tube will therefore have accumulated contaminants for the entire duration of exposure from a few months before launch through cruise and EDL until it was sealed on the surface of Mars on Sol 120. This exposure is very unlike the sample tubes sheathed in the ACA. Sample tubes (and the four other witness tubes) are protected from particle accumulation by fluid mechanical particle barriers and are isolated from volatile organic molecules by a tortuous path of gettering surfaces (Moeller et al., 2020). Given its long exposure, it is likely that the inner surfaces of WB1 will be saturated with organic contaminants, i.e., they will be in adsorption equilibrium with their immediate surroundings in the rover (and or the entire spacecraft prior to landing). WB1 is therefore expected to have higher concentrations of contaminants, and potentially different contaminants, than the sample tubes.

**Operations**

1. This witness tube was initially exposed and activated in the ultraclean ACA cleanroom at JPL just prior to being installed in its sheath in the Bit Carousel, after which the Bit Carousel doors were immediately closed.

2. The Bit Carousel doors were first re-opened, on Mars, on Sol 24.

3. Limited Bit Carousel motion occurred on sols 88, 90, and 92.

4. On Sol 118, WB1 was rotated to the lower Bit Carousel door.

5. WB1 was processed and sealed on sol 120. Note that this tube was not subjected to simulated sampling, i.e., it was not placed in the corer and moved in a fashion similar to rock core acquisition. In this sense it differs from the expected use case of the remaining four witness tubes.
INITIAL REPORT

M2020-164-2 Roubion

(no core recovered, atmospheric sample)

Sample Designation: M2020-164-2 Roubion

Date of Coring: 5-Aug-2021

Mars Time of Sample Core Sealing: 19:11:35 LMST, Sol 164, Ls 81.1

Latitude (N), Longitude (E), Elevation: 18.42769340, 77.45165066, -2584.96 m

Campaign: Crater Floor

Region of Interest: Séítah Thumb

Lithology: Fine- to medium-grained mafic and likely igneous rock, possibly basalt or microgabbro (alternatively, basaltic sandstone). Primary minerals are plagioclase and pyroxene, also possibly apatite and FeTi oxides. Weathering and/or aqueous alteration is indicated by pits and crevices in abraded surface and abundant secondary minerals including iron oxide (possibly hydrated), sulfates, perchlorate, and possibly phosphate and halite, usually in distinct patches.

Estimated Volume Recovered: ~0 cm³ (some ~10 µm sized particles). No core recovery

Coring Bit Number: 5

Core Orientation: hade = 3.43°; azimuth = 208.74°; core roll = 289.35°

Sample Serial Numbers: Tube SN233; Seal SN025; Ferrule SN062

ACA Temperature at Time of Sealing: 40 C

Estimated Rover-Ambient Pressure and Temperature at Time of Sealing: 749 Pa, 221 K

Estimated Amount of Martian Atmosphere Headspace Gas: 4.9x10⁻⁶ mol

Abrasion Patch Name and Depth: Guillaumes, 8 mm

Anomalous Behavior: Core disintegrated; no core acquired

November 27, 2021

Summary Description

Collection of Roubion, the first sample of the Mars 2020 mission, was attempted in the Séítah Thumb region of the Jezero crater floor (Figure 1). Between Octavia E. Butler landing (OEB) and the sampling site, Perseverance traversed about 1 km southward over nearly continuous low-lying rocks typically forming meter-scale “paverstone” outcrops, with intervening regolith (Figure 2). In HiRISE orbital view, these rocks define a distinctive polygonal pattern, the lower lying of the several expressions of the Crater Floor Fractured Rough (Cf-fr) unit of Stack et al (2020).

Figure 1 | Regional context. (a) geologic map of Sun and Stack (2020) and units of Stack et al. 2020 and (b) HiRISE map with Perseverance traverse path leading to Roubion. Shown for reference are Octavia E. Buttl er (OEB) landing site, Séítah Thumb area, Polygon Valley, Mure, and Artuby ridge.

Prior to landing, the Cf-fr unit was variously interpreted to be igneous (lava or volcanoclastic) or sedimentary (fluviolaclastine or aeolian) in origin. Although the stratigraphic context of the fractured floor, and indeed the lithology itself, were unknown at the time of sampling, the Cf-fr unit was selected for sample acquisition because it is areally extensive and because it includes the most heavily cratered terrain to which the rover has access within Jezero crater. As such, a returned sample of this unit was thought to have high science value for understanding the geologic setting and timing of crater floor units, and possibly for calibration of the Mars crater chronology function.

As evidenced by the common whaleback morphology and surface polish and fluting, this expression of Cf-fr has been eroded to just above ground level by aeolian abrasion (Figure 2). Despite abundant outcrop, little or no visual evidence of sedimentary structure, clasts, or crystals were seen in natural exposures of these rocks. Supercam data on multiple outcrops along the traverse indicate an altered (hydrated, iron-oxide-bearing) mafic rock with crystal size large enough to create spot-to-spot variability in composition (i.e., > few hundred µm scale). These low-lying Cf-fr rocks are fairly homogenous in appearance and composition along the entire traverse from OEB to Roubion.
The *Roubion* coring attempt, and its companion *Guillaumes* abraded patch, were undertaken on a low relief rock at the tip of the *Séítah* Thumb region selected largely to meet first-time sampling requirements. *Roubion* is interpreted to be stratigraphically below several other expressions of Cf-fr, in particular the high-standing blocky, massive rocks to the east and north, and the layered rocks of *Mure* and *Artuby* ridge (Figures 2 and 3). *Roubion* is interpreted to be from a unit that overlies and therefore may be younger than the rocks of *Séítah* (Cf-f1 of Stack et al., 2020).

**Figure 2** | ZCAM images of distinct fractured crater floor surfaces. (a) on Sol 58 that exhibit outcrops of less altered “whalebacks” and areas of rounded boulders, representative of early workspaces in the north near OEB. (b) on Sol 157 that exhibit flatter surfaces and more-polygonal outcrops where *Roubion* was sampled in the south near *Séítah* Thumb area.

The dominant mineralogy of the *Guillaumes* abraded surface appears to be ~mm-size pyroxene and plagioclase crystals, consistent in composition and texture with a relatively fine-grained gabbro or holocrystalline basalt. An igneous origin is preferred for this rock, but it is not currently possible to rule out the alternative interpretation that it is a basaltic sandstone. Extensive brown iron-rich patches are indicative of aqueous alteration, especially around several deep pits and crevices penetrating below the abraded surface. White patches containing secondary minerals (sulfate, perchlorate, and possibly halite and phosphate) provide additional evidence of aqueous interaction. SHERLOC fluorescence
spectroscopy identified two types of aromatic organic molecules within Guillaumes, one distributed widely across the abrasion patch and a second localized in a few specific regions. Chemical characteristics of the abraded patch are consistent with observations of this lithology all along the traverse, providing confidence that the sampled rock is reasonably representative of this expression of the Cf-fr unit. However, the rock appears more altered (more reddish, and appearing to be more granulated and/or flaky) than most other examples of this lithology (e.g., compare examples in Figure 2).

Figure 3 | ZCAM images of crater floor stratigraphy near Roubion. (a) Mure Sol 168. (b) Artuby ridge Sol 116.

Coring produced a hole and cuttings pile of expected appearance, and the entire coring and sealing process completed nominally. However, the volume probe indicated no sample, a fact confirmed by a Cachecam image documenting an empty sample tube. Thus, no core was recovered. In the absence of core or core fragments on the ground, the most likely explanation for zero recovery is that the rock disaggregated during coring and wound up either in the cuttings pile or at the bottom of the hole. Further coring at this location was abandoned.

Although no rock was obtained, M2020-164-2 Roubion inadvertently provides a returnable sample of about 4.9 µmol of martian atmosphere.

**Stratigraphic and Geologic Context**

Coring of Roubion was attempted at a polygonal outcrop within a unit of the Jezero crater floor mapped in HiRISE orbital images as the “crater floor fractured rough” (Cf-fr) unit by the Mars 2020 Science Team (Stack et al., 2020). Roubion is positioned in a local topographic low, ~10 m downslope from blocky massive rocks to the ENE and resistant layered capping rocks of the Mure outcrop and Artuby ridge to the SSE and W, all of which were mapped as Cf-fr (Figure 1). Roubion is interpreted to be stratigraphically below these other expressions of Cf-fr, but the stratigraphic relationship between the higher-standing, blocky, massive rocks and the layered capping rocks is presently unknown.

Roubion is located ~80 m east of the inferred contact between Cf-fr and an extensive outcrop exposure called Séítah, included within a unit mapped by Stack et al. (2020) as the “crater floor fractured 1” (Cf-f-1). Although no exposed contact between these two units is observed in the immediate vicinity of Roubion, Roubion may be younger than the rocks of Séítah due to an apparent onlapping relationship.
observed in orbital images, and topographic data and subsurface radargrams from the RIMFAX instrument. The age and stratigraphic relationship of Roubion relative to the rocks of the Jezero delta are less certain. Goudge et al. (2015) interpreted the unit containing Roubion to be younger than the Jezero delta, and the youngest in the crater, while Sun and Stack (2020), Stack et al. (2020), and Holm-Alwmark et al. (2021) proposed that the rocks of the Jezero crater floor, including Roubion, pre-date the currently exposed Jezero delta deposit.

**Operations**

The desirability of sampling Cf-fr was apparent immediately after landing but abrading and sampling could not begin prior to completion of a series of hardware check-outs and first-time activities. The rover traverse and these preparatory activities were choreographed such that the first abrading and sampling effort would occur at the last obvious polygonal outcrops of Cf-fr along the planned southbound route, in the Séítah Thumb region (Figure 1).

Cf-fr in the OEB area occurs at an elevation of ~-2570 m. The elevation of this rock unit remained constant as Perseverance drove south, until Sol 131 (~650 linear m from OEB), when elevation started to decrease. By Sol 138, the elevation had decreased by 13 m to ~-2583 m in the Séítah Thumb area (1020 meters from OEB), the lowest elevation the rover is expected to visit on its entire mission.

Potentially sampleable polygonal outcrop was abundant in the Séítah Thumb area and ultimately the southernmost patch of outcrop was selected for sampling because it is adjacent to and can be directly related to a few-meter-high cliff of layered outcrop called Mure and its likely WNW continuation in Artuby ridge (Figure 3a). At this early stage of the mission, it was considered operationally prudent to target an outcrop with minimal surface roughness, thus limiting target options.

Perseverance arrived at the selected sampling target on Sol 156. Abrasion and sample collection were respectively conducted on Sols 160 and 164, as part of a single-core-sample STOP list sol path extending from Sol 157 to Sol 168. This was the first-time-activity for coring. In addition to the standardized STOP list observations, when it was recognized that no core was recovered, additional borehole imaging and inspection of the nearby surface was undertaken for anomaly resolution. On Sol 168 Perseverance executed its first drive towards and along Artuby ridge, leaving the Séítah Thumb region and the Roubion sampling site.

**Sample-Related Observations**

**Workspace Images**

Figure 4 shows the workspace in which both the Guillaumes abrasion patch and the Roubion coring attempt were undertaken. Figure 4a shows the southernmost cluster of Cf-fr polygons in the Séítah Thumb region in which these activities occurred. View is to the SSE. Note that the polygons appear to extend towards (and possibly beneath) the boulder-decorated hill named Mure. As evident in Figure 3, Mure is layered, distinct from Cf-fr pavestones as seen all along the traverse from OEB to Séítah Thumb. This layering also extends to the WNW in Artuby ridge (Figures 1, 3) indicating a notable transition in
surface appearance of rocks all mapped from orbit as Cf-fr. The transition appears to occur in or near the Séítah Thumb. Furthermore, in this area the polygons appear more granulated, flaky and flat, and less wind-polished, than the more typical whaleback morphology (e.g., as seen near OEB in Figure 2a).

**Figure 4** | **ZCAM workspace images at Roubion sampling site.** (a) Outcrop selected for abrasion and sample acquisition. (b) Close up showing candidate target locations.

![Figure 4a](image1.jpg) ![Figure 4b](image2.jpg)

**Figure 4b**, a close-up of the sampled polygonal outcrop, further emphasizes the rather degraded appearance of the rocks in this area. Especially notable are the polygon edges, in which individual rock fragments appear to be falling or flaking out in a fashion like grus in a weathered granite. This figure shows the four locations considered for abrasion and coring. Ultimately the location called Guillaumes was selected for abrasion and Roure was selected for coring (and the core was named Roubion). These target choices were largely dictated by engineering considerations (i.e., for minimum topographic relief to ensure safe instrument placement).

**Images Pre and Post Coring/Abrasion**

**Figure 5a** provides WATSON images at two different standoffs prior to abrasion and coring. The rock surface appears stucco-like, with likely loose granules lying on a rather bumpy rock surface. Finer and likely loose grains (sand?) lie in low spots. The wind abrasion that is interpreted to have sculpted many examples of Cf-fr along the traverse is not evident here, rather the rock appears to be disaggregating faster than it is being abraded smooth. In contrast to the abraded patch (Figure 5b, and see below), these natural surfaces show no evidence of crystals/grains nor of holes penetrating into the rock; if holes are present, they have been filled with indistinct materials and totally obscured from view. Similarly, in these images there is no compelling evidence for the presence of the purple surface layer or coating which was observed on many rocks of Cf-fr along the traverse (but see post-abrasion discussion).

**Figure 5c** shows at high resolution the textural features of the natural surface before coring. These features were thought adequate for reconstruction of original core azimuthal orientation provided they survived coring. Dedicated orientation marking with SCAM LIBS points was not undertaken as it would not have substantially improved the likelihood of successful orientation of this core; indeed the LIBS pits
Figure 5 | WATSON images. (a) pre-abrasion target surface. (b) post-abrasion (Guillaumes) target surface. (c) pre-coring (Roubion) target surface, here with its original target name, Roure.
would likely be very difficult to locate in this surface, and in any case would be lost if the surface features were obliterated upon coring.

Figure 6 shows WATSON images of Guillaumes after removal of abrasion cuttings using the gas dust removal tool (gDRT) at two different resolutions. Notable observations include:

a) mm-size light and dark grains or crystals comprise most of the abraded surface. White materials have at least two distinct morphologies and tones: some have angular shapes, including what may be elongated laths, others (especially those that are brightest white) have a more irregular outline. The latter suggests a secondary cavity-filling or replacement substance. Ignoring these patches as well as the brown material described in b, the light and dark grains appear roughly equal in grain size with no obvious spatial heterogeneity in relative abundance. In some cases, for example in parts of Figure 6b, the grains may be interlocking as if crystallized from a magma. Taken together these observations are reminiscent of the appearance of a fine-grained gabbro or holocrystalline basalt.

b) some parts of the abraded surface are smooth and dark brown in color with less-evident grain boundaries, suggesting an (Fe-rich?) stain or coating. This material is heterogeneously distributed and is preferentially found in topographic lows of the abraded surface.

c) several irregularly shaped ~5 mm diameter pits and crevices penetrate below the abrasion patch and seem to share the dark brown color and texture noted in b. These pits are at least a few mm deep. Note that the abrasion removed 8 mm of surficial material (from original topographic peak to abraded surface); if these pits extended to the pre-abrasion surface they would have been 1.3 cm deep, possibly more. However, as noted above, pits were not visible on the natural surface. These pits do not have the typical appearance/distribution of vesicles in basalt and seem more likely to indicate mass removal by
some form of weathering that may have occurred prior to the formation of the current rock surface seen in Figure 5c.

d) while the abrasion patch rim is generally ragged in appearance, in some places the distinctive purple coating can be seen near or on the original surface, presumably now revealed by the gDRT. It is reasonable to expect the surface of the intact Roubion core would have captured this material.

**Elemental Geochemistry - PIXL**

X-ray fluorescence mapping by PIXL was undertaken in a single region in the Guillaumes abrasion patch (Figure 7; observation footprint in Figure 6). A bulk sum analysis of the Sol 167 scanned abrasion surface is shown in **Table 1**. Note this analysis includes a mixture of what may be igneous mineral grains along with the white and brown patches suggestive of aqueous alteration. Guillaumes is clearly a silicate rock; compared to a typical basalt, it is low in SiO$_2$, Al$_2$O$_3$ and MgO, and rich in FeO, Na$_2$O, SO$_3$, and Cl.

The right-hand column of **Table 1** documents large point-to-point heterogeneity in elemental composition; the heterogeneity is much larger than analytical uncertainty, as is expected from a rock containing chemically distinct minerals of a size comparable to and/or larger than the 120 µm X-ray beam size. This point is further evident in elemental maps (Figure 10a-d); there are distinct ~500 µm size regions rich in FeO, or Al$_2$O$_3$, or CaO. Further work is necessary to fully characterize these compositions and map them to the WATSON images, but a tentative interpretation is that the calcic regions correspond to the dark minerals and are pyroxene (augite), the aluminous phase is light colored and tabular and corresponds to plagioclase, and the Fe-rich material is the brownish coating(?) and could be an iron oxide or iron silicate. FeTi oxides and sodium and chlorine rich patches (halite? Na perchlorate?) are also apparent.

**Figure 7** | PIXL X-ray element maps. (a) MgO-Al$_2$O$_3$-CaO. (b) SiO$_2$-FeO$_{total}$-TiO$_2$. (c) FeO$_{total}$-Al$_2$O$_3$-CaO. (d) MgO-Na$_2$O-Cl in Sol 167 analysis of Guillaumes. Refer to Figure 6 for scale and region investigated.

<table>
<thead>
<tr>
<th>Guillaumes bulk sum (00167)</th>
<th>wt%</th>
<th>wt% err</th>
<th>std</th>
<th>std%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Na$_2$O</td>
<td>5.55</td>
<td>0.28</td>
<td>3.10</td>
<td>55.8%</td>
</tr>
<tr>
<td>MgO</td>
<td>2.64</td>
<td>0.56</td>
<td>1.08</td>
<td>41.0%</td>
</tr>
<tr>
<td>Al$_2$O$_3$</td>
<td>7.56</td>
<td>0.38</td>
<td>3.02</td>
<td>40.0%</td>
</tr>
<tr>
<td>SiO$_2$</td>
<td>38.29</td>
<td>1.92</td>
<td>7.12</td>
<td>18.6%</td>
</tr>
<tr>
<td>P$_2$O$_5$</td>
<td>1.65</td>
<td>0.46</td>
<td>1.41</td>
<td>45.4%</td>
</tr>
<tr>
<td>SO$_3$</td>
<td>2.69</td>
<td>0.56</td>
<td>2.90</td>
<td>108.0%</td>
</tr>
<tr>
<td>Cl</td>
<td>3.40</td>
<td>0.54</td>
<td>4.14</td>
<td>121.8%</td>
</tr>
<tr>
<td>K$_2$O</td>
<td>0.75</td>
<td>0.26</td>
<td>0.47</td>
<td>62.4%</td>
</tr>
<tr>
<td>CaO</td>
<td>7.77</td>
<td>0.39</td>
<td>3.86</td>
<td>49.6%</td>
</tr>
<tr>
<td>TiO$_2$</td>
<td>1.47</td>
<td>0.43</td>
<td>2.27</td>
<td>154.8%</td>
</tr>
<tr>
<td>Cr$_2$O$_3$</td>
<td>0.03</td>
<td>0.06</td>
<td>0.07</td>
<td>246.9%</td>
</tr>
<tr>
<td>MnO</td>
<td>0.46</td>
<td>0.20</td>
<td>0.37</td>
<td>81.5%</td>
</tr>
<tr>
<td>FeO$_{total}$</td>
<td>18.72</td>
<td>0.94</td>
<td>10.94</td>
<td>58.5%</td>
</tr>
<tr>
<td>Ni</td>
<td>0.00</td>
<td>0.00</td>
<td>0.03</td>
<td>N/A</td>
</tr>
<tr>
<td>Zn</td>
<td>0.02</td>
<td>0.03</td>
<td>0.06</td>
<td>411.3%</td>
</tr>
<tr>
<td>Br</td>
<td>0.01</td>
<td>0.03</td>
<td>0.05</td>
<td>313.5%</td>
</tr>
<tr>
<td>Zr</td>
<td>0.02</td>
<td>0.03</td>
<td>0.03</td>
<td>122.7%</td>
</tr>
</tbody>
</table>

Table 1. PIXL Bulk Sum Composition
Figure 8 is a ternary diagram of elemental compositions (Al₂O₃-CaO+Na₂O+K₂O-FeO+MgO) for all the individual PIXL points and the bulk sum spectrum. Common mineral compositions are also plotted on the diagram. Most Guillaumes points lie in a triangle defined by minerals typical of mafic rocks (feldspar, pyroxenes, olivine). While there is little evidence of aluminum enrichment expected from open-system leaching and formation of aluminous clay minerals, there is a clear trend towards unidentified secondary phases which includes carbonates, halides, and sulfates. This plot supports the general conclusion that the analyzed rock is an aqueously altered but not intensely weathered basalt, gabbro, or basaltic sandstone.

Figure 8 | Guillaumes sol 167 PIXL data. Individual analysis points (grey) and bulk sum composition (blue) on a ternary diagram of molar abundances Al₂O₃–(CaO+Na₂O+K₂O)–(FeO+MgO). Common igneous minerals (olivine, pyroxene, feldspar, Fe-Ti-oxides) are typically found within the red triangle whereas common clay minerals fall above this triangle.

The PIXL point measurements and bulk sum spectra are plotted on a total alkali versus silica igneous classification diagram in Figure 9. While this diagram can be useful for naming specific igneous lithologies, the high abundance of alteration phases complicates such usage. For that reason, no igneous lithology is assigned here. Of additional note on this diagram is the prominent array of points trending towards high alkalis and very low SiO₂; this observation suggests the presence of alkali-rich salts such as NaCl or Na-perchlorate.
Figure 9 | Total alkalis vs silica plots for Guillaumes and Cf-fr. (a) PIXL of Guillaumes (b) SCAM LIBS of all analyzed Cf-fr rocks through sol 168 (light symbols) and for Guillaumes (dark). The individual data points likely represent a mixture of one or more primary and/or secondary minerals. Bulk sum composition of PIXL data and average value of SCAM data shown by blue circles. Endmember igneous mineral compositions are shown for reference. The underlying igneous classification scheme does not apply to individual data points but provides a frame of reference.
**Mineralogy and Organics-SHERLOC**

A SHERLOC UV Raman and florescence survey scan was obtained on Sol 162. An additional SHERLOC scan, not discussed here, was obtained on Sol 161. The analysis footprints are indicated in Figure 6.

As shown in Figure 10 and Table 2, SHERLOC Raman spectra indicate the likely presence of Ca-sulfate, Na-perchlorate, and possibly minor phosphate associated with the light-toned regions of Guillaumes (Figure 10d). An amorphous silicate phase widely distributed across the sample is also tentatively identified. These data are consistent with the multiple lines of evidence indicating aqueous alteration phases, especially concentrated in the bright white patches.

**Figure 10 | SHERLOC Spectral-textural correlations.** (a) SHERLOC ACI image. (b) Fluorescence and Raman maps of organic distribution. (c) Fluorescence spectra. (d) Raman mineral distribution and spectra

The average fluorescence spectrum indicates two peaks, one at 275 nm and the other at 340 nm (Figure 10c). Both are likely attributable to aromatic organic compounds in fairly low average abundance. The 275 nm peak is highly localized in a small number of spots (Figure 10b), but at the time of this writing it is not possible to link these hotspots to specific mineralogical or textural features of the rock. The 340 nm peak is more widely distributed across the analyzed region, suggesting it is a spatially dispersed organic constituent. Unlike fluorescence, there is no evidence from Raman spectroscopy for organic molecules in Guillaumes.
Table 2 SHERLOC Mineral Identifications in Guillaumes

<table>
<thead>
<tr>
<th>Sample</th>
<th>Certain</th>
<th>Almost certain (currently working to confirm)</th>
<th>Possible (not confirmed)</th>
<th>No detections found in the primary lithology</th>
</tr>
</thead>
<tbody>
<tr>
<td>Guillaumes</td>
<td>Ca-sulfate (minor phase, occurs in white patches secondary to primary lithology)</td>
<td>Na-perchlorate (main phase, occurs in white patches secondary to primary lithology)</td>
<td>Phosphate (minor phase)</td>
<td>Pyroxene, olivine</td>
</tr>
</tbody>
</table>

Elemental Geochemistry and Mineralogy - Supercam

SCAM LIBS and VISIR measurements of the pre- and post-abrasion surface, borehole, and borehole cuttings for the Roubion sample target are shown in Figure 11, a series of Harker oxide diagrams, supplemented with LIBS results from many Cf-fr rocks analyzed along the traverse from OEB to Roubion through sol 168. There are no striking differences among the Roubion borehole walls and cuttings and various Guillaumes abrasion patch LIBS datasets for MgO, Al₂O₃, and K₂O. Although considerable scatter exists, the borehole appears to show higher FeOT and lower Na₂O. This heterogeneity in composition within the cored and abraded rock is possible evidence for a greater concentration of salts near the rock surface. It is unclear whether FeOT may have been removed from the surface or enriched at depth. Despite these abrasion surface-borehole differences, the compositional tends from the abraded/cored rock are mostly in-family with those measured for the many Cf-fr rocks analyzed along the traverse. The main exception is the fact that the abrasion patch exhibits little evidence for SiO₂ concentrations higher than ~ 60 wt%, whereas such values are occasionally seen in other (unabraded) Cf-fr rocks. Such high SiO₂ concentrations may be restricted to natural rock surfaces. High MgO values like those of Entrevaux and Alguines (Figure 11, targets in the topographic low between Artuby ridge and Séítah) are absent in the abraded surface and are rare in Cf-fr rock analyses (the latter may reflect olivine regolith grains/sand on top of or trapped in Cf-fr paverstone rock crevices).

Many of the trends observed in these diagrams are reasonably attributed to mixing among igneous minerals, e.g. pyroxene and plagioclase. The 250 µm LIBS spots commonly interrogates more than one specific mineral (typical Guillaumes grain size is ~ 1 mm as described above).

To complement the PIXL data, the LIBS results from Cf-fr and from Guillaumes are also plotted on a total alkali vs. silica igneous classification diagram (Figure 9b). It is notable that within Guillaumes, PIXL picks up a strong trend towards high alkali and low silica (likely secondary salts) not seen by SCAM. Similarly SCAM LIBS of Cf-fr appears to yield more points with high SiO₂ concentrations (>50 wt %) than are observed by PIXL in Guillaumes. Both observations likely represent compositional heterogeneity among the different regions/rocks being analyzed.

SCAM VISIR spectra from the Guillaumes abraded patch have notable absorption features at 1.43 and 1.94 µm, attributable to hydration (Figure 12), a feature common to many of the Cf-fr rocks investigated.
along the OEB to Roubion traverse. Interestingly, these features are not evident or are less pronounced in the Roubion borehole walls and in the Roubion cuttings. This observation may indicate preferential hydration of the upper surface of the investigated rock. A 2.28 µm absorption feature observed in the abrasion patch may indicate Fe-bearing clay.

**Figure 11** | Harker oxide diagram. SCAM LIBS spot compositions of natural rock surfaces of fractured crater floor (Cf-fr), Guillaumes, and borehole + borehole cuttings of Roubion. Distribution reflects individual laser spots (~250 µm) that represent mixtures of one or more primary and secondary mineral. Theoretical pyroxene, feldspar, and olivine compositions shown for reference.
Figure 12 | Guillaumes SCAM VISIR. Abrasion surface shows absorptions at ~1.43 and 1.93 µm that indicate hydration. These features are weaker in the borehole wall and borehole cuttings of Roubion.
**Subsurface Structure**

The RIMFAX radargram for the sol 157-168 traverse path, including interpreted subsurface stratigraphy and densities from estimated permittivity values, are shown in Figure 13. Inferred densities are similar to basaltic material. The radargram shows surface-parallel reflectors and fine-scale layering below the paver-like morphology down to a burial depth of about a meter. Arcuate structures may be small, buried impact craters. Deeper reflectors (>3 m subsurface) appear to dip southward and may be associated with rocks of Séítah.

**Figure 13 | RIMFAX results from near Guillaumes/Roubion.** (a) Radargram for the sol 168 traverse including 3 m drive NE from the Guillaumes/Roubion site on sol 157 end of drive (EOD) followed by a 141° southward turn where the rover passes the target area at <2 m distance (“projected”). Base of the surface paver rock inferred to occur at ~1 m burial depth is denoted with a black line, and red lines denote curved reflector geometries that may be remnant impact craters. (b) Geological interpretation of reflector geometries in (a) illustrating subsurface stratigraphy with an ~1 m thick paver unit with decimeter scale surface-parallel layering overlaying south-dipping Séítah (?) reflectors. Location of permittivity measurements (ε’) from diffraction hyperbolae are denoted with red and yellow circles. (c) Densities derived from permittivity measurements annotated in (b) plotted vs depth given as two-way signal travel time below the antenna. Densities for basaltic rocks with zero to 20% porosity, feldspar, pyroxenes and olivine are shown for comparison. Paver unit densities are similar to basaltic rocks.

**Core orientation**

At the time of drilling and 7-cm pre-drilling WATSON imaging, the rover, corer, WATSON, and coring target (called “Roubion_162.PSC.0668”), had the following characteristics:

1. Coring target normal (RMECH frame): \( \mathbf{uvw}_{\text{RMECH}} = (-0.005, -0.024, -1.000) \)
2. Rover orientation quaternion at time of drilling (transferring from RMECH to SITE frame): \( \mathbf{bQll1} = (0.332886, 0.0174924, -0.00616092, 0.942785) \)
3. Gravity vector (RMECH frame): \( \mathbf{g}_{\text{RMECH}} = (-0.0356277, -0.00190381, 0.999363) \)
4. Coring Drill orientation quaternion (transferring from CD to RMECH frame): \( \mathbf{cdQb} = (0.665264, 0.233161, -0.661495, 0.255898) \)
5. Rover orientation quaternion at time of acquisition of WATSON image
SIF_0161_0681240940_632FDR_N0060000SRLC00702_0000LMJ01: (transferring from RMECH to SITE frame): \( \mathbf{bQll2} = (0.332934, -0.0148229, -0.00733084, 0.942805) \)

6. WATSON orientation quaternion at time of acquisition of WATSON image
SIF_0161_0681240940_632FDR_N0060000SRLC00702_0000LMJ01: (transferring from WATSON to RMECH frame): \( \mathbf{wQb} = (0.667723, 0.237418, -0.658963, 0.252084) \)

Items 1 and 2 give a coring drill pointing vector estimate of (Hade = 3.33°; Azimuth = 195.47°). Item 3 gives an alternate coring drill hade estimate of (Hade = 2.76°). Items 2 and 4 gives a coring drill pointing vector estimate of (Hade = 3.43°; Azimuth = 208.74°). We report this as the final coring drill orientation estimate since it was estimated for the drill after preloading. Items 5 and 6 give an estimate of the angle between the WATSON y-axis and the up-dip direction in SITE coordinates of (Fig 14) (core roll, \( \alpha = 289.35° \))

Fig 14. 7-cm standoff WATSON image of Roubion core target on sol 161. WATSON image SIF_0161_0681240940_632FDR_N0060000SRLC00702_0000LMJ01 Image scale is 31.1 μm pixel\(^{-1}\). Orientation compass gives WATSON frame (\( \mathbf{wx}, \mathbf{wy}, \mathbf{wz} \)). Core roll is clockwise angle of core y-axis, \( \mathbf{cy} \), from \( \mathbf{wy} \), given by \( \alpha = 289.35° \).
Preliminary Scientific Assessment

Synthetic sample description and preliminary interpretation

Sample type: basaltic lava flow or microgabbro (or possibly basaltic sandstone)

1. Relationship with surrounding rocks
   a. **Roubion** appears to be in family with the Cf-fr paverstone rocks investigated all along the traverse from OEB in terms of composition and texture. It is therefore considered representative of the low-lying morphology of Cf-fr.
   b. Within Cf-fr rocks, **Roubion** lies below both the high standing blocky and cratered expression to the E and N, and also the layered expression of Artuby ridge and Mure.
   c. **Roubion** is structurally higher than rocks of the Séítah region (Cf-f1) and is therefore younger, unless it is an intrusive.
   d. Although difficult to prove owing to partial burial, the sampled rock is thought to be in place or very nearly so.

2. Texture and fabric
   a. Grain size: approximately 0.5 to 1 mm (plagioclase, pyroxene, and alteration products).
   b. No obvious igneous flow banding or sedimentary structures were seen.
   c. Pits and crevices occur in the abraded patch but are not evident in the rock surface. They do not have the appearance of typical vesicles. These features may be voids created by some type of weathering.
   d. There is evidence for interlocking grain texture, suggesting crystallization from a magma.
   e. There is no compelling evidence of intergranular porosity or cements, common to many sedimentary rocks.

3. Mineralogy and chemistry
   a. Minerals present include pyroxene, feldspar, and FeTi-oxides, and secondary iron oxides or silicates, sulfates, Na-Cl salts (halite? Na-perchlorate), and possibly phosphate. The phosphate could alternatively be primary.
   b. The bulk composition is consistent with a basalt modified by aqueous alteration. The alteration is sufficiently intense that it is not possible to assign a specific igneous lithologic name without further analysis to isolate the least altered portions of the rock.
   c. Low concentrations of aromatic organics are present.

4. Alteration/secondary characteristics
   a. There is significant textural and mineralogical evidence in the abraded patch for interaction of the rock with aqueous fluids, alteration, and mineral hydration.
   b. Secondary mineralogy remains speculative, but likely includes Ca-sulfates, Na-perchlorates, Ca-phosphates, and halite.
   c. The absence of Al-rich phases argues for low water/rock ratios and little cation leaching.
Returned Sample Science Considerations

Outcrop morphology and texture, as well as the appearance, composition and mineralogy of the abraded patch suggest that *Roubion* represents a fine- to medium-grained mafic lithology, possibly a basaltic lava flow or a microgabbro. An alternative interpretation of this unit is volcanoclastic sandstone derived from basaltic rocks of the Jezero watershed. The striking absence of sedimentary textures in the well-exposed Cf-fr rocks from OEB to *Roubion* disfavors the latter interpretation and would constrain potential sedimentary depositional mechanisms if this were indeed a sedimentary rock. The crystallization age of volcanic rocks (or volcanic minerals in a sandstone) can be readily determined by radiometric methods undertaken on returned samples. If the timing of this rock’s deposition can be obtained (e.g., if *Roubion* is a lava flow), this age would quantitatively constrain the ages of other crater floor units: Séitah is older, and the layered rocks of Artuby ridge and Mure are younger. The latter presumes that *Roubion* is not an intrusive rock. The composition of igneous rocks and/or mafic minerals in this rock would provide a geochemical probe that would reveal the geophysical and geochemical nature of the planet’s interior at the time of eruption. Furthermore, if *Roubion* can be confidently related to the heavily cratered portions of Cf-fr (e.g., to the east of the traverse), and the surface exposure history of Cf-fr can be understood, then geochronology of a returned sample of this lithology may provide a test of (or calibration for) crater chronology models. However, we are not yet confident in tying *Roubion* to the heavily cratered part of Cf-fr.

Patches of water-soluble salts compellingly document aqueous alteration in this rock. There is no strong evidence for clay minerals in Guillaumes, suggesting that chemical weathering (i.e., open system behavior in which cation leaching yields clay minerals) is limited. There is no doubt this rock has experienced interaction with one or more aqueous fluids, possibly groundwater that may or may not be related to Jezero lake. Soluble salts (e.g., halite or perchlorate) almost by definition document the final aqueous interaction experienced by this rock. It is likely these water-rock interactions occurred in a habitable subsurface environment recorded in the alteration mineral assemblage. Analysis of the secondary minerals may be used to study the chemistry and conditions (e.g., pH, temperature, salinity, age) of aqueous fluids. Similar salts on Earth are known to contain organic molecules and even fossilized microscopic life.

For these reasons a returned sample of this unit has high science value for astrobiology, as a point in the geochemical/geophysical evolution of the martian interior and surface environment, and for understanding the geologic setting and timing of Jezero floor units.

The core sample was not recovered. The sealed tube includes martian atmosphere and a few tiny *Roubion* particles that are most likely fragments of the failed core. Thus, the sample tube could be used to study martian atmosphere. Alternatively (or perhaps in addition) the tube may provide a sample processing contamination blank.
References


Sample Designation: M2020-190-3 Montdenier

Date of Coring: 31-Aug-2021

Mars Time of Sample Core Sealing: 16:53:19 LMST, Sol 194, Ls 92.5

Latitude (N), Longitude (E), Elevation: 18.43074132, 77.44436502, -2574.553 m

Campaign: Crater Floor

Region of Interest: Artuby ridge

Lithology: Fine-grained mafic and likely igneous rock, possibly basalt (alternatively, basaltic sandstone). Primary minerals are plagioclase, pyroxene, and FeTi oxides. Aqueous alteration is indicated by abundant secondary minerals including iron oxide, Ca-sulfates (occasionally hydrated), phosphate, and minor carbonate.

Estimated Volume Recovered: 8.4 cm³

Coring Bit Number: 5

Core Orientation: hade = 9.42°; azimuth = 219.27°; core roll = 191.24°

Sample Serial Numbers: Tube SN266; Seal SN170; Ferrule SN099

ACA Temperature at Time of Sealing: 37 C

Estimated Rover-Ambient Pressure and Temperature at Time of Sealing: 702 Pa, 245 K

Estimated Amount of Martian Atmosphere Headspace Gas: 1.2x10⁻⁶ mol

Abrasion Patch Name and Depth: Bellegarde, 8 mm

Anomalous Behavior: Multiple sols between coring and sealing

November 30, 2021

Herd, J. Hurowitz, L. E. Mayhew, D. L. Shuster, K. M. Stack, M. Sylvestre, B. P. Weiss, P. A. Willis, R. Wiens,
M.-P. Zorzano, and the Mars 2020 Team
Summary Description

Collection of Montdenier and Montagnac, the pair of rock cores from the second sample target of the Mars 2020 mission, was conducted in the Citadel area, a relatively high-standing outcrop within the “crater floor fractured rough” orbital unit (Cf-fr of Stack et al., 2020; Figure 1). Citadel sits on the crest of Artuby, an ~1 km long NW-SE trending linear ridge exposing on its NE side a several meter high cliff of the crater floor stratigraphy. Artuby ridge borders the southern edge of an exposure of rocks called Séita (Cf-f-1 of Stack et al., 2020). After an unsuccessful first attempt to collect a core from low-lying outcrops at Roubion, Perseverance traversed about 450 m NW along the base of Artuby ridge, in a regolith-covered topographic low that defines the boundary between Séita and Cf-fr. Perseverance traversed to Citadel to access and sample crater floor rocks that were prominently exposed on the ridgeline (Figure 2) on the supposition that such rocks would be more resistant to disaggregation than Roubion.

Prior to landing, the Cf-fr unit was variously interpreted to be igneous (a lava flow or a volcanoclastic deposit) or sedimentary (fluviolacustrine or aeolian). The lithology of Cf-fr remains uncertain, though evidence from Roubion suggests that at least part of Cf-fr is igneous. Unlike outcrops of Cf-fr observed between the Octavia E. Butler (OEB) landing site and Roubion, rocks of Artuby ridge display fine and coarse semi-continuous layers and a distinctive knobby texture (Figure 3). Despite excellent exposure, there is no compelling evidence for sedimentary structures formed via traction transport, nor were clasts observed. Supercam data collected on these rocks reveal an altered (hydrated, iron-oxide-bearing) mafic composition with distinct crystals greater than the ~200 μm size of the LIBS shots.

The Montdenier and Montagnac cores, and their companion Bellegarde abrasion patch, were acquired on a small tabular boulder (Rochette, ~40 cm across) that is a member of a NW-SE-trending band of boulders or outcrop on the SW side of the Artuby ridge crest. These rocks are morphologically consistent with a degraded lava flow or layer (Figure 2b). Rochette moved slightly during coring, suggesting it is not currently anchored in-place. Although Rochette may have been displaced from its original location, the similarity and abundance of blocks near Rochette suggest displacement is likely small. Rochette was selected for coring primarily for its accessibility by the rover, limited relief, and perceived resistance to disaggregation during coring. Additionally, its position high on Artuby ridge places it stratigraphically above and potentially younger than other studied rocks including Roubion and Cf-f-1 rocks outcropping in Séita.

The Bellegarde abrasion patch is dominated by 0.2-0.5 mm pyroxene and plagioclase crystals, and is consistent in composition and texture with an aphanitic holocrystalline basalt (or microgabbro). The interlocking crystal texture and the lack of fine-grained matrix or intergranular cement suggest an igneous origin. Extensive iron-rich brown patches are indicative of aqueous alteration, especially evident around white patches that may reflect cavities filled by secondary minerals such as Ca-sulfate (occasionally hydrated) and phosphate. Carbonate and amorphous silicates were tentatively identified within Bellegarde by SHERLOC, as were three types of aromatic organic signals, one distributed widely across the abrasion patch and two localized in a few specific regions. Chemical characteristics of the abraded patch are generally consistent with other Cf-fr rocks observed along the traverse from OEB to Roubion, and along Artuby ridge from Roubion to Bellegarde, providing high confidence that the sampled rock is reasonably representative of the Cf-fr unit. The abraded rock appears less altered (has
no obvious pits or crevices, has less abundant secondary mineralization, is less reddish, and overall appears more intact) than Roubion, and therefore better represents the primary lithology.

**Figure 1 | Regional context.** (a) geologic map, Stack et al. 2020. (b) HIRISE map with Perseverance traverse leading to Bellegarde.
Given the failure to acquire core at Roubion, coring of Montdenier included a special step to confirm sample-in-the-tube with ZCAM images prior to tube sealing. This added four sols to the period between coring and sealing; in that period the sample was held in the drill, external to the rover. Coring of both Montdenier and Montagnac produced a hole and tailings pile of expected appearance, the ZCAM image of Montdenier revealed core in the tube, and the sealing process of both tubes completed nominally. The volume probe indicated full length cores (i.e., ~60 mm or ~8.5 cm³) were acquired in both cases.

**Figure 2 | ZCAM and NCAM images of Citadelle area.** (a) caprock of crater floor stratigraphy defining the crest of Artuby ridge (b) surface unit from which sample cores Montdenier and Montagnac were collected. (c) overview of (a) and (b) showing how the dip of the strata shallows from about 10° to near horizontal to the south away from Séítah.

**Stratigraphic and Geologic Context**

The Montdenier and Montagnac cores and Bellegarde abrasion patch were collected from Rochette, a ~40 cm wide tabular boulder that is part of an elongated exposure of similar rocks on the SW side of the Artuby ridge crest (Figure 2). Rochette and the boulders in its immediate vicinity are inferred to be part of a single degraded rock layer forming part of the caprock of Artuby ridge based on the similar alignment of boulder long axes, their similar thickness, and their nearly interlocking shapes. The strata
exposed by the Artuby scarp appear to be dipping ~10° to the southwest, roughly perpendicular to the strike of Artuby ridge. This dip appears to flatten out within tens of meters southwest of the ridge crest (Figure 2c). The dip and exposures are such that the rock layer containing Rochette may be a few tens of cm higher in the stratigraphy than rocks on the crest of Artuby ridge.

The contact between Cf-fr and Cf-f-1 locally coincides with Artuby ridge (Figure 1a). Cf-fr is distinguished from other units on the crater floor by its relatively high crater density, curvilinear margins, and lack of the strong olivine absorption characteristic of Cf-f-1 (Goudge et al., 2015). Rocks thus far examined along Artuby ridge, including Rochette, lack evidence for abundant olivine, suggesting the exposed rocks of the ridge are all part of the Cf-fr unit. Unlike the apparently structureless Cf-fr rocks from OEB to Roubion, the rocks forming Artuby ridge are clearly layered, with layers varying in thickness from a few cm to several tens of cm (Figure 3). The Cf-fr - Cf-f-1 contact is inferred to lie in a topographic low at the base of the ridge, where it is obscured by regolith.

Figure 3 | ZCAM images of layering of Cf-fr rocks exposed in near-vertical outcrop. (a) ~2 m cliff at Mure Sol 169. (b) ~2-3 m cliff at Artuby Sol 177.
Rochette is stratigraphically higher than rocks of Séítah and may therefore be younger. The relationship between the Artuby cap rocks and those observed at Mure, OEB, and north and east of OEB is presently unknown given the discontinuity of outcrop exposure and local disruption by impact craters, but these caprocks share morphological and compositional similarities suggesting they may be correlative. The age and stratigraphic relationship of Artuby caprocks relative to the rocks of the Jezero delta is also uncertain. Goudge et al. (2015) interpreted the unit containing Rochette to be younger than the Jezero delta, and the youngest in the crater, while Sun and Stack (2020), Stack et al. (2020), and Holm-Alwmark et al. (2021) proposed that the rocks of the Jezero crater floor, including Rochette and Roubion, pre-date the currently exposed Jezero delta deposit.

**Operations**

After failing to recover sample at Roubion (see Initial Report for M2020-164-2 Roubion), the mission team prioritized a sampling attempt on a rock more likely to resist disaggregation. Boulders of the upper layers of Artuby ridge were considered likely to have survived eons of exposure to wind abrasion, suggesting less friability than the low-lying degraded-appearing outcrop at Roubion. Accordingly, a traverse to the first location where such rocks were accessible along the rover’s strategically planned traverse, at a site named Citadelle, was undertaken.

On Sol 168 Perseverance executed its first drive NW along Artuby ridge, leaving the Roubion sampling site and gradually regaining about 10 meters of elevation along the base of the ridge. On Sol 178 Perseverance executed an autonomous drive that included a sharp turn up the NE-facing slope of Artuby ridge, concluding at Citadelle (Figure 1 and 2). After selecting Rochette as the preferred sampling target, abrasion at Bellegarde and collection of the first core Montdenier were respectively conducted on Sols 182 and 190. A paired-core, strategically developed sampling sol path was executed between Sols 180 and 196. Because visual proof of core acquisition was desired before ingesting the sample tube inside the rover, sealing of the Montdenier tube was delayed until Sol 194. In the intervening 4-sol period, the sample was exposed to the martian environment. This exposure is a notable deviation from expected practice in which a tube is sealed autonomously within hours of coring. The paired core Montagnac was collected and sealed on Sol 196 following the normal automated procedure and without visual confirmation prior to sealing.

On Sol 197, after successful collection of two cores, Perseverance executed its next drive NW along Artuby ridge, heading towards the planned entry point into Séítah.
Sample-Related Observations

Workspace Images

Figure 4 shows the workspace in which both the Bellegarde abrasion patch and the Montdenier and Montagnac coring were undertaken. Figure 4a shows the tabular boulder Rochette and its alignment with, and spacing from, other blocks near where these activities occurred. Rochette appears darker in tone than nearby blocks, likely because its surface is less dusty. The Rochette block is smooth, with sharp angular edges, abundant mm- to cm-scale pits, and parallel fluting suggesting wind abrasion. It stands ~20 cm higher than the surrounding surface, and on several sides overhangs the surrounding regolith. Rochette has a mottled grey and purple appearance. There is no evidence for layering or any other substantial heterogeneity across the boulder. Rochette appears to have broken off of, and is slightly displaced from, the triangular shaped rock behind it (Figure 4).

Figure 4b and 4c show Rochette after abrasion of the Bellegarde patch and after the acquisition of the two cores, respectively. All three targets (Bellegarde, Montdenier, and Montagnac) seem to have been acquired in compositionally similar material. Tailings slid off the rock during abrading and coring, and appear grey on the reddish brown regolith in front of Rochette. Behind Rochette, localized regolith subsidence and a steep crevice formed during abrasion. These features were further developed during coring.

Figure 4 | NCAM workspace images. (a) flat-lying outcrop containing resistant target rock Rochette. (b) Close-up of Rochette showing abraded surface Bellegarde. The act of abrading Rochette resulted in localized subsidence of regolith (arrow), and (c) Rochette with boreholes from acquisition of sample cores Montdenier (right hole) and Montagnac (left hole). Rochette is ~ 40 cm across.
**Pre and Post Coring/Abrasion Images**

WATSON images at two different standoffs prior to abrasion and coring are shown in Figures 5 and 6. At high resolution the rock surface appears very smooth and fairly free of dust except where it has collected along the edges of subtle sculpted surfaces and within rare pits. In contrast to the abraded patch (Figure 5b, and see below), these natural surfaces show no evidence of crystals or grains. A heterogeneous purple surface layer or coating is evident in these images (see post-abrasion discussion). A similar coating appears on many rocks along the traverse from OEB to Roubion.

**Figure 5 | WATSON images.** (a) pre-abrasion target surface. (b) post-abrasion target surface.
Figure 6 | WATSON images. (a) pre-coring target surface Montdenier. (b) pre-coring target surface Montagnac.
Figure 7 | WATSON images of abrasion surface. (a) ~25 cm standoff, showing PXL, SHERLOC, and SCAM instrument placement. Abrasion patch is 5 cm across. Inset (b) holocrystalline rock texture with regions of less altered intergrown plagioclase and pyroxene and secondary minerals potentially filling voids.

Figure 7 shows a WATSON image of the gDRT-cleared abraded patch Bellegarde. Notable observations include:

a) 0.2-0.5 mm-size dark and light grains or crystals comprise most of the abraded surface. White materials have at least two distinct morphologies and tones: some have angular shapes, others (especially those that are brightest white) have a more irregular outline. The latter suggests a secondary cavity-filling or replacement substance. Ignoring these patches as well as the brown material described in b, the dark and light grains appear roughly equal in grain size with no obvious spatial heterogeneity in relative abundance. In some cases, for example in parts of Figure 7b, the grains may be interlocking as if crystallized from a magma. Taken together these observations are consistent with the appearance of a holocrystalline basalt.

b) Some parts of the abraded surface are reddish brown in color with more diffuse boundaries, suggesting an (Fe-rich?) stain or coating. In some cases, this apparent staining is seen as haloes and in association with (or even partially covering) the irregular-shaped white phases that appear to fill cavities. This association can also be seen in CacheCam images showing the bottom of both the Montdenier and Montagnac cores (Figure 8). The Montagnac core has one especially large white patch in its CacheCam image, measuring more than 3 mm across and accounting for about 10% of the entire exposed surface of the bottom of the core.

c) The surface of Rochette and associated boulders appear to have a purple to tan coating (Figure 5b). This coating is particularly common on what appear to be wind abraded and likely older surfaces. The coating is less obvious, possibly thinner to absent, on fractured (younger) dark gray surfaces. These rough fractured surfaces also tend to exhibit more pitting.

d) Unlike the ragged appearance of the edge of the Guillaumes abrasion patch (Roubion sample), that of Bellegarde is sharp and angular. In some places the previously noted purple material can be seen near or on the original surface (Figure 9). It is reasonable to expect that the Montdenier and Montagnac cores contain some of this purple material.
Figure 8 | Cachecam images of the bottom of sample cores. (a) Montdenier on Sol 194. (b) Montagnac on Sol 196. Core diameter is ~1.3 cm. Note reddish haloes around secondary minerals potentially filling voids.

Figure 9 | WATSON images of abraded crater floor surfaces. (a) Large (up to ~7 mm) pits and ~1.0 mm crystals in Guillaumes were exposed on Sol 160 in the lower Cf-fr pavement surface. (b) the presence of a few, possibly filled pits and smaller ~0.2-0.5 mm crystals in Bellegarde were exposed on Sol 186 from the surface unit at the top of Artuby ridge. (c) close up of Guillaumes. (d) close up of Bellegarde. Degree of alteration appears less in Bellegarde.
**Elemental Geochemistry - PIXL**

X-ray fluorescence mapping by PIXL was performed on the *Bellegarde* abrasion patch (Figure 10; observation footprint in Figure 7). A bulk sum analysis of the scanned abrasion surface is shown in Table 1. Note this analysis includes a mixture of what are believed to be primary silicates (pyroxene and plagioclase) along with the bright white and reddish brown patches suggestive of aqueous alteration. *Bellegarde* is clearly a silicate rock; compared to a typical basalt, it is low in SiO$_2$, Al$_2$O$_3$ and MgO, and rich in FeO, Na$_2$O, SO$_3$, and Cl.

**Figure 10 | PIXL X-ray element maps.** (a) MgO-Al$_2$O$_3$-CaO. (b) SiO$_2$-FeO$_{\text{total}}$-TiO$_2$. (c) FeO$_{\text{total}}$-Al$_2$O$_3$-CaO. (d) MgO-Na$_2$O-Cl in *Bellegarde*. The bulk sum composition of this scan area is included as Table 1. See Figure 7 for scale and location.

Chemically distinct minerals of a size comparable to and/or larger than the 120 µm X-ray beam size can be seen in elemental maps (Figure 10a-d) where there are distinct regions up to ~500 µm in size rich in FeO, or Al$_2$O$_3$, or CaO. Further work is necessary to fully characterize these compositions and map them to the WATSON images, but a tentative interpretation is that the calcic regions correspond to the dark minerals and are pyroxene (augite), some iron and silicon bearing dark minerals may be an Fe-rich, Ca-poor pyroxene, the aluminous phase is tabular and light colored (bone white as opposed to the bright white secondary sulfate) and corresponds to plagioclase, and the Fe-rich material is the brownish coating(?) and could be an iron oxide or iron silicate. Apatite and FeTi oxides are also apparent in the PIXL data.

**Figure 11** is a ternary diagram of molar compositions Al$_2$O$_3$-(CaO+Na$_2$O+K$_2$O)-(FeO$_7$+MgO) for all of the individual PIXL points and the bulk sum spectrum. Idealized common mineral compositions are also plotted on the diagram. Most *Bellegarde* points lie in a triangle defined by typical mafic rock minerals (e.g., feldspar, pyroxene, olivine). While there is some evidence for unidentified secondary phases, there is little evidence of aluminum enrichment expected from open-system leaching (e.g., chemical

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**Table 1. PIXL Bulk Sum Composition**

<table>
<thead>
<tr>
<th>Oxide</th>
<th>wt%</th>
<th>wt% err</th>
<th>std</th>
<th>std%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Na$_2$O</td>
<td>4.67</td>
<td>0.34</td>
<td>1.59</td>
<td>34.0%</td>
</tr>
<tr>
<td>MgO</td>
<td>2.21</td>
<td>0.54</td>
<td>1.16</td>
<td>52.6%</td>
</tr>
<tr>
<td>Al$_2$O$_3$</td>
<td>6.97</td>
<td>0.35</td>
<td>3.62</td>
<td>52.0%</td>
</tr>
<tr>
<td>SiO$_2$</td>
<td>43.79</td>
<td>2.19</td>
<td>10.65</td>
<td>24.3%</td>
</tr>
<tr>
<td>FeO$_{\text{total}}$</td>
<td>2.76</td>
<td>0.56</td>
<td>2.46</td>
<td>89.1%</td>
</tr>
<tr>
<td>SO$_3$</td>
<td>3.21</td>
<td>0.56</td>
<td>5.84</td>
<td>181.9%</td>
</tr>
<tr>
<td>Cl</td>
<td>1.48</td>
<td>0.43</td>
<td>0.84</td>
<td>56.8%</td>
</tr>
<tr>
<td>K$_2$O</td>
<td>1.06</td>
<td>0.34</td>
<td>0.76</td>
<td>71.4%</td>
</tr>
<tr>
<td>CaO</td>
<td>7.62</td>
<td>0.38</td>
<td>4.34</td>
<td>57.0%</td>
</tr>
<tr>
<td>TiO$_2$</td>
<td>2.49</td>
<td>0.55</td>
<td>2.18</td>
<td>87.4%</td>
</tr>
<tr>
<td>Cr$_2$O$_3$</td>
<td>0.01</td>
<td>0.03</td>
<td>0.02</td>
<td>183.3%</td>
</tr>
<tr>
<td>MnO</td>
<td>0.44</td>
<td>0.21</td>
<td>0.21</td>
<td>47.9%</td>
</tr>
<tr>
<td>FeO$_7$</td>
<td>23.23</td>
<td>1.16</td>
<td>11.27</td>
<td>48.5%</td>
</tr>
<tr>
<td>Ni</td>
<td>0.00</td>
<td>0.00</td>
<td>0.01</td>
<td>N/A</td>
</tr>
<tr>
<td>Zn</td>
<td>0.01</td>
<td>0.03</td>
<td>0.04</td>
<td>260.4%</td>
</tr>
<tr>
<td>Br</td>
<td>0.00</td>
<td>0.01</td>
<td>0.03</td>
<td>742.6%</td>
</tr>
<tr>
<td>Zr</td>
<td>0.02</td>
<td>0.03</td>
<td>0.04</td>
<td>213.1%</td>
</tr>
</tbody>
</table>
weathering) and formation of aluminous clay minerals. This plot supports the general conclusion that the analyzed rock is weakly altered by interaction with water.

The PIXL point measurements and bulk sum spectra are also plotted on a total alkali versus silica igneous classification diagram (Figure 12). While this diagram can be useful for naming specific igneous lithologies, the high abundance of secondary and/or alteration phases in Rochette complicates such usage. For that reason, no specific igneous name is assigned here. Of additional note on this diagram is the prominent array of points trending towards very low SiO$_2$; this observation is consistent with the presence of sulfates or carbonates.

Figure 11 | Bellegarde PIXL data. Plotted as individual points (grey) and as bulk sum composition (blue circle) on a ternary diagram of molar abundances Al$_2$O$_3$–(CaO+Na$_2$O+K$_2$O)–(FeO$_{Total}$+MgO). Common igneous minerals (olivine, pyroxene, feldspar, Fe-Ti-oxides) are typically found within the dashed red triangle whereas common clay minerals fall above the upper red dashed line. Several additional secondary minerals (e.g., Fe-Mg-Ca-sulfates/carbonates, halite) are also plotted.
Figure 12 | Total alkalis versus silica plots. (a) PIXL data from scan of abraded surface Bellegarde. (b) SCAM LIBS laser spot data from abraded surface Bellegarde and from Cf-fr rocks obtained from sols 170 through 205. The individual data points represent small regions of the abraded surface (∼100 µm for PIXL; ∼250 µm for SCAM) and therefore represent a mixture of one or more primary and/or secondary minerals. Bulk sum composition of PIXL data and average value of SCAM data shown by blue circles. Endmember igneous mineral compositions shown as diamonds and lines for those with solid solution for reference. The underlying igneous classification scheme does not apply to the individual data points and simply provides a frame of reference for comparing samples.
Mineralogy and Organics-SHERLOC

A SHERLOC UV Raman and fluorescence survey scan was obtained on Sol 186. The analysis footprint is shown in Figure 7. As shown in Figure 13 and Table 2, SHERLOC Raman spectra indicate the likely presence of Ca-sulfate (occasionally hydrated) that appears to fill voids and/or replace primary mineral(s). SHERLOC also detected an unknown amorphous silicate, and minor carbonate and phosphate phases. These data are consistent with the multiple lines of evidence indicating that secondary phases are concentrated in the bright white patches (Figure 10d). Where PIXL observations overlap the SHERLOC area, they corroborate these secondary mineral identifications.

Figure 13 | SHERLOC fluorescence and Raman spectral results for Bellegarde. Spectra were collected with an HDR scan (7x7 mm, 100 points, 780 µm, and 500) and a survey scan (5x5 mm, 1296 points, 144 µm spacing, 15 pulses per point). (a) Colorized ACI image rectangles indicating scan areas: red=HDR, blue=survey, and white circles indicating ROIs highlighted in b and c. (b) Possible presence of organics. RGB map showing main regions of organic fluorescence (275, 305, and 340 nm). Note the 340 nm peak is distributed throughout, but the 275 and 305 nm peaks are found mainly in two ROIs (see panel a). No Raman features indicative of organics were detected. (c) Fluorescence spectra. The upper panel is the average fluorescence spectrum for the whole analysis area. The lower panel includes spectra from ROIs. (d) Mineral distributions. The left and middle panels show the results of the Raman survey and HDR scans, respectively. In each, the grid of red circles indicates the points analyzed, and the blue, white, and blue/white circles indicate locations of mineral detections, corresponding largely to lighter-toned features in ACI image. The legend in the survey scan panel applies to that of the HDR scan as well. The right panel includes selected Raman spectra from the two scans. The upper spectrum is the average spectrum of the HDR scan. The majority of features are caused by cosmic rays, but there is a possible detection of amorphous silicates based on a broad band centered near 1000 cm⁻¹. The spectra labeled No. 1, 2, and 3 correspond to the points in the map panels with the same numbers and are representative of spectra assigned to Ca-sulfate (1050 cm⁻¹), carbonate (1100 cm⁻¹), and (possibly) phosphate (~975 cm⁻¹), respectively. The spectrum labeled No. 1 also contains a possible peak for hydration at ~3550 cm⁻¹. Scale in panel a also applies to panel d.
The average fluorescence spectrum of the scanned area contains one major peak at 340 nm. Two other peaks at 275 nm and 305 nm are observed in the spectra from the selected region of interest (Figure 10c). All these peaks can likely be attributed to aromatic organic compounds. The relatively low intensity of fluorescence at all wavelengths and absence of Raman organics signature suggests that, if there are indeed organic materials present, they are in low abundance and generally dispersed with some localized hotspots.

**Table 2. SHERLOC Raman Spectra Mineral ID***

<table>
<thead>
<tr>
<th>Sample</th>
<th>Certain</th>
<th>Possible (not confirmed)</th>
<th>No detections found in the primary lithology</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bellegarde</td>
<td>Ca-sulfate, occasionally hydrated (occurs in white void fills or crystals, secondary to primary lithology)</td>
<td>Carbonate (very minor phase, only detected in a single spot)</td>
<td>Pyroxene, olivine</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Amorphous silicate (potentially major phase covering whole target)</td>
<td>We looked for these, but cannot find them</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Phosphate (minor phase)</td>
<td></td>
</tr>
</tbody>
</table>

*Elemental Geochemistry and Mineralogy - SuperCam*

SCAM LIBS and VISIR measurements of the pre- and post-abrasion surfaces, boreholes, and borehole tailings for the *Montdenier* and *Montagnac* sample targets are shown in Figure 14. These data are supplemented with LIBS results from many Cf-fr rocks acquired along the traverse, divided for clarity between those from OEB to *Roubion* (up until sol 168) and from *Roubion* to *Citadelle*. The *Montdenier* and *Montagnac* borehole walls and tailings and *Bellegarde* abrasion patch compositions are broadly similar. The compositional trends from the abraded/cored rock are consistent with those measured for the many Cf-fr rocks along the traverses. One notable distinction is that while SiO₂ values >55% are fairly common in Cf-fr targets, they are absent in the abraded and cored materials. The high SiO₂ values may thus be restricted to natural surfaces, possibly suggesting an association with a surficial coating. High MgO values comparable to those of *Entrevaux* and *Aiguines* (rocks at the base of *Artuby* ridge and likely exposures of *Séítah*) are absent in the abraded surface and are only present in regolith grains inadvertently analyzed in the Cf-fr targets. The ~250 µm LIBS spot most commonly interrogates more than one specific mineral. Given that the typical *Bellegarde* grain size is 0.2-0.5 mm, as described above, many of the trends observed in Figure 14 are reasonably attributed to a mixing among igneous minerals such as pyroxene and plagioclase.

To complement the PIXL data, the LIBS results from *Bellegarde* are also plotted on the total alkali vs. silica diagram (Figure 12b). PIXL detects more points rich in alkali elements and low in silica (tentatively interpreted as secondary salts) than are detected by SCAM. Similarly, no SCAM points exhibit SiO₂ values higher than ~55 wt % despite many such points detected by PIXL. These differences as well as distinct PIXL and SCAM bulk mean values (blue circles in Figure 12) could be related to different analysis regions on *Bellegarde*, but might also be related to the difference in spot size of the two measurement techniques.
SCAM VISIR spectra of the Bellegarde abraded patch have notable absorption features at 1.42 and 1.93 \( \mu \text{m} \), attributable to hydration (Figure 15), a feature common to many of the Cf-fr rocks investigated along the OEB to Roubion traverse. These features are not evident or are less pronounced in the Montdenier borehole walls and in the Montdenier and Montagnac tailings. This observation may indicate preferential hydration of the upper surface of the investigated rock or may arise from the different grain size of the tailings compared to the original rock. Absorption features in spectra at 2.28 \( \mu \text{m} \) could reflect carbonate and/or may indicate an Fe-bearing smectite clay.

Figure 14 | Harker oxide diagram. SCAM LIBS spot compositions of natural rock surfaces of fractured crater floor (Cf-fr), abraded surface Bellegarde sols 187-188, vertical borehole scan of Montdenier, two vertical borehole scans of Montagnac, and borehole tailings of Montdenier.
Figure 15 | Bellegarde SCAM VISIR. Abrasion surface shows evidence of hydration at ~1.42 and 1.93 um. These absorptions are much weaker in borehole tailings.

Subsurface Structure

The RIMFAX radargram acquired on sols 177-180, during the rover’s drive along Artuby ridge (177), ascent of Artuby ridge and arrival at Citadelle (178), and short bump to Rochette (180), can be seen in Figure 16. From left to right, the four panels in Figure 16 present results along a rover path that 1) begins NW, parallel to the NW-SE strike of Artuby ridge, then 2) turns left to cross the ridge, then 3) runs NW parallel to the ridge again, and 4) finishes with a nearly 180-degree turn. Figure 16 also includes an interpreted subsurface stratigraphy and rock densities based on permittivity. While driving along the base of Artuby ridge, RIMFAX observed a sequence of SW dipping reflectors that dominate the stratigraphy from approximately surface levels to >10 m burial depth. Starting on sol 178, the radargram shows a relatively thin horizontal capping layer, consistent with the Rochette outcrop at the surface. The estimated permittivity suggests that the horizontal, near surface layers are similar in density to basaltic...
material and those at depth exhibit greater density, consistent with an olivine-rich lithology (see Figure 16 caption for details).

**Figure 16 | Geological context for Citadelle area of Artubyridge including subsurface stratigraphy and permittivity.** (a) RIMFAX radargram for the sol 178 to 180 traverse. (b) Geological interpretation of reflector geometries in (a) with the Montdenier-Montagnac target indicated with a red star. A capping surface unit with decimeter-scale sub-horizontal layering appears to be present from ~80 m distance to the end of the sol 180 drive. A sequence with SW dipping reflectors dominates the stratigraphy from near surface levels to >10 m burial depth. Location of permittivity measurements ($\varepsilon'$) from diffraction hyperbolae are denoted with yellow circles. These and additional nearby permittivity measurements within sol 178 and 199 radargrams range between 7-8 and 10-14, translating to rock densities around 3.0-3.2 and 3.5-4.0 g/cm³, with the latter group suggesting olivine-rich lithologies (possibly Séítah).

![Figure 16](image)

**Core orientation**

At the time of drilling and 6-cm pre-drilling WATSON imaging, the rover, coring drill, WATSON, and coring target (called “Montdenier_188.PSC.0.0638”), had the following characteristics:

a) Coring target normal (RMECH frame): $uvw_{RMECH} = (-0.089, 0.104, -0.991)$

b) Rover orientation quaternion at time of drilling (transferring from RMECH to SITE frame): $bQll_1 = (0.696481, 0.0183586, -0.00168854, -0.717339)$

c) Gravity vector (RMECH frame): $g_{RMECH} = (-0.028215, -0.0211681, 0.999378)$

d) Coring Drill orientation quaternion (transferring from CORING DRILL to RMECH frame): $cdQb = (0.705714, 0.20918, -0.664503, 0.129009)$

e) Rover orientation quaternion at time of acquisition of WATSON image
   SIF_0188_0683643364_351FDR_N0070000SRLC00678_0000LMJ01: (transferring from RMECH to SITE frame): $bQll_2 = (0.696524, 0.0163777, -0.00351771, -0.717338)$

f) WATSON orientation quaternion at time of acquisition of WATSON image
   SIF_0188_0683643364_351FDR_N0070000SRLC00678_0000LMJ01: (transferring from WATSON to RMECH frame): $wQb = (0.706087, 0.213611, -0.663116, 0.126843)$
Items (a) and (b) gives a coring drill pointing vector estimate of (Hade = 9.98°; Azimuth = 218.87°). Item (c) gives an alternate coring drill hade estimate of (Hade = 8.24°). Items (b) and (d) gives a coring drill pointing vector estimate of (hade = 9.42°; azimuth = 219.27°). We report this as the final coring drill orientation estimate since it was estimated for the drill after preloading. Items (e) and (f) give an estimate of the angle between the WATSON y-axis and the up-dip direction in SITE coordinates of Figure 17: (core roll, α = 191.24°)

**Figure 17 | Core orientation.** 6-cm standoff WATSON image of Montdenier on sol 188. WATSON image SIF_0188_0683643364_351FDR_N0070000SRLC00678_0000LMJ01. Image scale is 27.8 μm pixel$^{-1}$. Orientation compass gives WATSON frame ($w_x$, $w_y$, $w_z$). Core roll is clockwise angle of core y-axis, $c_y$, from $w_y$, given by $\alpha = 191.24^\circ$. 5 mm scale bar.
Preliminary Scientific Assessment

Synthetic sample description and preliminary interpretation

Sample type: Basalt (or possibly basaltic sandstone)

1. Relationship with surrounding rocks
   a. Montdenier-Montagnac are part of the layered caprock of Artuby ridge, appearing to be in family, morphologically and compositionally, with caprocks investigated along the traverse from Roubion to Citadelle in terms of composition and texture. These cores are therefore considered representative of the Cf-fr unit.
   b. Within Cf-fr rocks, Montdenier-Montagnac probably lie below the high standing blocky and cratered expression to the east and north, and clearly above the granular rocks exposed at Roubion and along the NE-facing slope of Artuby ridge.
   c. Montdenier-Montagnac are stratigraphically higher than and therefore probably younger than rocks of the Séïtah region (Cf-f1).
   d. Although dislodged from the flat-lying outcrop at Citadelle, Rochette, the rock from which Montdenier-Montagnac were cored, is generally thought to be very close to its original outcrop location.

2. Texture and fabric
   a. Grain size: ~0.2-0.5 mm (plagioclase, pyroxene). Alteration products exhibit a greater size range appearing to fill void space (~primary minerals up to ~3 mm in size).
   b. The surface of Rochette is pitted. These pits could be the result of aeolian abrasion, volcanic vesicles enhanced by aeolian abrasion, or voids created by some type of chemical weathering or alteration. In the abraded surface, there are no obvious holes, although sulfate-filled voids are observed.
   c. There is evidence for interlocking grains, suggesting crystallization from a magma. But there is also evidence for more equigranular textures, and possible rounded grains.
   d. There is no compelling evidence of intergranular porosity or cements, common to many sedimentary rocks.

3. Mineralogy and chemistry
   a. Minerals present include pyroxene, feldspar, and FeTi-oxides, and secondary iron oxides or silicates, sulfates, and phosphate. The phosphate may be primary, secondary or both.
   b. The bulk composition is consistent with a basalt modified by aqueous alteration. Unlike Roubion, the alteration was not sufficiently intense to move it out of the basalt compositional field.
   c. Low concentrations of aromatic organics are present, both localized and distributed in the rock.

4. Alteration/secondary characteristics
   a. There is textural and mineralogical evidence in the abraded patch for interaction of the rock with aqueous fluids, resulting in chemical alteration and mineral hydration.
   b. Secondary mineralogy remains speculative, but likely includes Ca-sulfates and Ca-phosphates.

5. Core orientation
a. Sample cores were absolutely oriented to ~1° in martian geographic coordinates, although Rochette may have been differentially tilted with respect to the underlying bedrock.

**Returned Sample Science Considerations**

Outcrop morphology and texture, as well as the appearance, composition and mineralogy observed in the *Bellegarde* abrasion patch, suggest that *Rochette* is a fine-grained mafic igneous rock, possibly a member of a sequence of lava flows. An alternative interpretation is that it is a volcaniclastic sandstone derived from basaltic rocks of the Jezero watershed. The absence of sedimentary textures in the Cf-fr rocks exposed in cliffs at *Mure* and *Artuby* and the texture of *Rochette* disfavor the latter interpretation.

*Geochronology:* The timing of volcanic rock crystallization (and volcanic minerals preserved in a sandstone) will be quantifiable using laboratory-based, radio-isotopic geochronology of returned samples. Likewise, with knowledge of cosmogenic nuclide production rates in specific minerals on Mars, quantitative constraints on the sample’s surface exposure age may also be possible. The timing of *Rochette*’s deposition will constrain the emplacement ages of other crater floor units, e.g., *Séítah* is likely older, as is the stratigraphically lower *Roubion*. Furthermore, if *Rochette*’s lithology represents the heavily cratered surfaces of Cf-fr (e.g., to the east and north of the OEB), and the surface exposure and exhumation history of Cf-fr can be understood, then geochronology of returned samples may document the duration of crater accumulation and retention of the exposed Jezero floor. Such information could then be used to test, and potentially improve, the fundamental parameters assumed in the Mars crater chronology functions.

*Geochemistry:* The chemical composition and mineralogical properties of this igneous rock (or detrital igneous minerals, if sedimentary) will provide information on the geophysical and geochemical conditions of the rock’s source (i.e., at depth, within the martian crust or mantle) at the time of its primary eruption or intrusion. In addition, water-soluble salts, apparently filling holes, suggest aqueous alteration and/or addition of secondary phases at some point after deposition. Thus, this rock has likely experienced interaction with one or more aqueous fluids, possibly groundwater that may or may not be related to Jezero lake. The extent of alteration appears to be less than observed in *Roubion*. These water-rock interactions may have occurred in a habitable subsurface environment. Laboratory analyses of the secondary minerals will provide information on the chemistry and environmental conditions (e.g., pH, temperature, salinity, timing) during aqueous activity. However, at the time of writing, there is no strong chemical evidence for clay minerals in *Rochette*, indicating that any open system conditions (i.e., cation leaching yielding clay minerals) was limited. It is also unclear whether the observed phosphates are secondary or primary phases in the rock. Organic molecules within *Rochette* would provide information on the martian carbon cycle and organic preservation.

For these reasons, a returned sample of this unit has high science value for astrobiology, global evolution of Mars (interior and near surface environments), and for understanding the geologic setting and timing of Jezero floor units.
References


INITIAL REPORT
M2020-196-4 Montagnac

Sample Designation: M2020-196-4 Montagnac

Date of Coring: 7-Sept-2021

Mars Time of Sample Core Sealing: 20:35:07 LMST, Sol 196, Ls 95.6

Latitude (N), Longitude (E), Elevation: 18.43074132, 77.44436502, -2574.553 m

Campaign: Crater Floor

Region of Interest: Artuby Ridge

Lithology: Fine-grained mafic and likely igneous rock, possibly basalt (alternatively, basaltic sandstone). Primary minerals are plagioclase, pyroxene, and FeTi oxides. Aqueous alteration is indicated by abundant secondary minerals including iron oxide, Ca-sulfates (occasionally hydrated), phosphate, and minor carbonate.

Estimated Volume Recovered: 8.7 cm³

Coring Bit Number: 5

Core Orientation: hade = 17.0°; azimuth = 225.96°; core roll = 197.25°

Sample Serial Numbers: Tube SN267; Seal SN170, Ferrule SN074

ACA Temperature at Time of Sealing: 35 C

Estimated Rover-Ambient Pressure and Temperature at Time of Sealing: 707 Pa, 215 K

Estimated Amount of Martian Atmosphere Headspace Gas: 1.3x10⁻⁶ mol

Abrasion Patch Name and Depth: Bellegarde, 8 mm

Anomalous Behavior: None

December 3, 2021

This sample is paired with M2020-190-3 Montdenier. See section 3 (M2020-190-3 Montdenier) for details of this paired sampling. Only the core orientation for Montagnac is described below.

Core orientation

At the time of drilling and 7-cm pre-drilling WATSON imaging, the rover, coring drill, WATSON, and coring target (called “Montagnac_RP.PSC.0.0691”), had the following characteristics:

a) Coring target normal (RMECH frame): \( \mathbf{u}w_{RMECH} = (-0.205, 0.182, -0.962) \)
b) Rover orientation quaternion at time of drilling (transferring from RMECH to SITE frame): \( bQ_{ll1} = (0.696493, 0.0181555, -0.0019524, -0.717331) \)
c) Gravity vector (RMECH frame): \( \mathbf{g}_{RMECH} = (-0.028215, -0.0211681, 0.999378) \)
d) Coring Drill orientation quaternion (transferring from CORING DRILL to RMECH frame): \( \mathbf{c}dQ_{b} = (0.730214, 0.260758, -0.623407, 0.100777) \)
e) Rover orientation quaternion at time of acquisition of WATSON image SIF_0195_0684255577_390FDR_N0070000SRLC00701_0000LMJ01: (transferring from RMECH to SITE frame): \( bQ_{ll2} = (0.696528, 0.0164346, -0.00334345, -0.717334) \)
f) WATSON orientation quaternion at time of acquisition of WATSON image SIF_0195_0684255577_390FDR_N0070000SRLC00701_0000LMJ01: (transferring from WATSON to RMECH frame): \( wQ_{b} = (0.729438, 0.26773, -0.620981, 0.103072) \)

Items (a) and (b) gives a coring drill pointing vector estimate of (Hade = 17.98°, Azimuth = 225.73°). Item (c) gives an alternate coring drill hade estimate of (Hade = 16.43°). Items (b) and (d) gives a coring drill pointing vector estimate of (hade = 17.70°, azimuth = 225.96°). We report this as the final coring drill orientation estimate since it was estimated for the drill after preloading. Items (e) and (f) give an estimate of the angle between the WATSON y-axis and the up-dip direction in SITE coordinates of Figure 1: (core roll, \( \alpha = 197.25° \))
Figure 1 | Core orientation. 7-cm standoff WATSON image of Montagnac core target on sol 195. WATSON image SIF_0195_0684255577_390FDR_N0070000SRLC00701_0000LMJ01. Image scale is 32.0 μm pixel⁻¹. Orientation compass gives WATSON frame (wx, wy, wz). Core roll is clockwise angle of core y-axis, cy, from wy, given by α = 197.25°.
Sample Designation: M2020-262-5 Salette

Date of Coring: 15-November-2021

Mars Time of Sample Core Sealing: 20:22 LMST, Sol 262, Ls 127.5

Latitude (N), Longitude (E), Elevation: 18.43397°, 77.44301°, –2569.2 m

Campaign: Crater Floor

Region of Interest: Séítah South

Lithology: Medium- to coarse-grained poikilitic olivine cumulate rock. Primary minerals are olivine and pyroxene. Aqueous alteration phases include sulfates, carbonates and hydrated phases. Organic matter was detected.

Estimated Volume Recovered: 8.9 cm³

Coring Bit Number: 5

Core Orientation: hade = 17.78°; azimuth = 81.51°; core roll = 253.58°

Sample Serial Numbers: Tube SN246; Seal SN194; Ferrule SN107

ACA Temperature at Time of Sealing: 40°C

Estimated Rover-Ambient Pressure and Temperature at Time of Sealing: 643 Pa, 221 K

Estimated Amount of Martian Atmosphere Headspace Gas: 1.1 x10⁶ mol

Anomalous Behavior: None

Abrasion Patch Name and Depth: Dourbes, 8 mm
Summary Description

Salette and Coulettes, the pair of rock cores representing the third sample target of the Mars 2020 mission, were collected in the South Séítah region from Brac, a low-lying layered outcrop in the Caille locality at the easternmost end of the rover’s traverse into Séítah. Brac lies within a group of blocky outcrops near the crest of one of the NE-SW trending ridges that are common in Séítah, ~130 meters from the Máaz-Séítah contact at the base of Artuby ridge (Figures 1 and 2). Brac is likely to be in-place.

Séítah makes up the stratigraphically lowest rocks exposed on the floor of Jezero crater. In orbital mapping, Séítah corresponds to a light-toned olivine-bearing unit referred to as Crater floor-fractured-1 (Cf-f-1) by Stack et al., 2020. This unit may be correlated with a regional olivine-bearing unit exposed between Syrtis Major and Nili Fossae. The mineralogy, micro-texture, color and spectral characteristics of Brac and other outcrops observed throughout Séítah suggest that the Cf-f-1 unit is mostly or entirely an igneous olivine cumulate. The horizontal to gently dipping layers observed in Séítah may represent primary igneous layering as seen in layered igneous bodies, though no layer-bounding mineralogical or grain-size variations were observed.

The Dourbes abrasion patch, acquired on Brac, revealed abundant ~3–5 mm euhedral to subhedral olivine grains surrounded by clinopyroxene in a poikilitic texture with minor feldspars, magnetite, and phosphates. The rock is therefore interpreted to be an igneous olivine cumulate formed by the settling of olivine in a magma body such as a sill or thick lava flow. No evidence was obtained that could distinguish between intrusive and extrusive origins. Alteration phases include sulfates (most likely Ca- or Mg-dominated), which fill voids and are thus secondary to the igneous mineralogy. Carbonate, most likely Fe/Mg-carbonate, and amorphous/nanocrystalline material may also be present, and hydrated phases were detected. Organic matter is present in low abundance, dispersed throughout the rock matrix and concentrated in a few spots.

The returned sample science objectives of these cores include geochronology, paleomagnetism, geochemistry, and the past history of water and water–rock interactions in Jezero crater. This core pair is geochemically similar and likely genetically related to the other Séítah samples in the cache (Robine and Malay). Together with cores of the Máaz formation (Montdenier, Montagnac, Atsah, Hahonih), these samples constitute the entirely igneous Crater Floor Campaign sample suite.

Stratigraphic and Geologic Context

Salette and paired core Coulettes were collected from the Brac outcrop near the top of a local topographic high point in a region of the Jezero crater floor informally named Séítah, mapped in HiRISE images as the Cf-f-1 unit by the Mars 2020 Science Team (Stack et al., 2020; Figure 1). Prior to landing, the Cf-f-1 unit was interpreted to be either igneous or sedimentary. Several origin hypotheses have been suggested, including a pre-Isidis impact melt (Mustard et al., 2007), a pre-Isidis intrusive complex (Hoefen et al., 2003), basalt flows (Hamilton and Christensen, 2005; Tornabene et al., 2008), volcanic ash (Kremer et al. 2019), pyroclastic deposits (Mandon et al., 2020), and detrital sedimentary rocks (Rogers et al., 2018). Séítah has been correlated with a regional olivine-bearing unit exposed between Syrtis Major and Nili Fossae (Mandon et al., 2020).
Orbital mapping suggests that Séítah rocks are the stratigraphically lowest exposed rock unit on the crater floor and are overlain by both the Máaz formation and the delta. Rover observations confirm that Séítah lies below Máaz. Perseverance has not yet explored the relationship of the crater floor units to the delta. Consistent with its lower position in the stratigraphy, Séítah may be older than Máaz. However, because Séítah could be intrusive, the inverse is also possible.

**Operations**

Figure 2 shows the local context of Perseverance rover operations leading up to the sampling of Salette and Coulettes. Prior to approaching Brac, we conducted abrasion and proximity science of the nearby layered outcrop Bastide; however, we were unable to execute the full STOP list on Bastide and its Garde abrasion patch, thus the decision was made to seek another nearby rock for STOP list execution and sampling.

On Sol 248, we arrived at the Brac workspace and started the sampling sol path. On Sol 250, we selected four targets for abrasion (Lane, Traverses, Pont du Loup and Laupon) and acquired WATSON images of each. Pont de Loup was favored as the abrasion target because it appeared to cross multiple layers of the outcrop and because it may have offered a possibility to study later alteration phenomena in cross-cutting(?) veins. We attempted abrasion on Pont du Loup on Sol 251, but this operation faulted due to slip of the abrasion bit across the rock surface. Following recovery, we attempted abrasion on a new target, Dourbes, on sol 253. This target was chosen for its proximity to Pont du Loup and its potential to answer many of the same science questions as Pont du Loup. On Sol 255, we successfully abraded at Dourbes. PIXL and SHERLOC data were acquired on Dourbes on sol 257. We acquired the Salette core on sol 262.

After sampling Salette, we encountered a fault while picking up the abrading bit on sol 263; this precluded subsequent remote sensing and proximity science on the borehole. After performing a recovery on sol 267, we planned borehole proximity science on Salette, and pre-coring WATSON images on Coulettes, the second locality from which a core sample was to be obtained. The Coulettes core was acquired on sol 271.

We drove away from the Brac workspace on sol 277, heading in the direction of the Issole workspace.
Figure 1 | Regional context of Perseverance rover operations. (a) geologic map showing Jezero crater floor units defined by Stack et al. 2020. The light green shaded overlay indicates the location of the Jezero Crater Floor campaign. (b) Geological map showing the crater floor Cf-fr and Cf-f-1 units. Adapted from Stack et al. (2020).
Figure 2 | Local context of Perseverance rover operations during Salette and Coulettes sampling. HIRISE map showing the locations of sample collection together with other notable outcrops investigated during the sol path leading to Brac. Location of Brac is indicated with a blue circle.

Sample-Related Observations

Workspace Images

Figure 3a shows a Mastcam-Z mosaic of the Brac outcrop in its local context (view is to the SE). Figure 3b shows the Mastcam-Z image of the workspace after the Dourbes abrasion patch and before Salette and Coulettes were sampled. To the upper left of Brac in Figure 3b the outcrop transitions into apparently massive blocks and thickly layered outcrop. Brac and surrounding outcrop feature both smooth and sharper angular edges, surficial depressions, and some evidence for parallel fluting arising from wind erosion (cf. Figure 4–6). Although the natural surface of Brac is fairly free of dust, dust accumulations at the boundaries between layers obscure the layer–layer contacts (Figure 4). During workspace observations, the team proposed two scales of layering in Brac: i) layers several centimeters to tens of centimeters in thickness, delineated by dust-covered, recessively weathered contact; and ii) less distinct, thinner layers distinguished by ridges, parallel to the thicker layers, across the outcrop. No compelling evidence for layering was observed in the abrasion patch or the borehole.
**Figure 3 | Geology near the Brac outcrop.** a) Caille locality; red rectangle indicates the location of the Brac outcrop. Brac is ~70 cm in length and located below the red rectangle. b) Post-abrasion Mastcam-Z image of the Brac outcrop and workspace. Location of the Dourbes abrasion patch is indicated by the red arrow. The abrasion patch is 5 cm in diameter. File: zcam08261.0000a.caille_vignols_baleine_scam_rmi_visir_P28_L0R0_Z110.

RMI images of the outcrop (Figure 4) show large (0.1–0.4 cm), dark grey, angular minerals surrounded by a pale brown interstitial phase similar to that observed on relatively fresh surfaces at Bastide (e.g. the RMI target Cine). These observations are consistent with those undertaken on the Dourbes abrasion patch (see below). The interstitial phase may represent the dark brown phase between minerals observed in the Dourbes abrasion patch, dust trapped between grains, or a combination of the two.

Brac is surrounded by regolith and conceivably may have moved downhill slightly as evidenced by the regolith-filled crevice to the left of the outcrop and a lack of layer continuity with other adjacent outcrops. However, it is more likely that Brac is in-place because its layering has the same apparent orientation as the surrounding outcrops throughout this portion of South Séítah. Subsurface data confirm that Brac is not significantly out of place (if at all) and as such it is confidently interpreted to be bedrock. Regolith settled into the subvertical fracture to the immediate left of Dourbes after abrasion, but no motion of the outcrop was observed.
Figure 4 | SuperCam RMI mosaic of the Brac outcrop showing large gray grains surrounded by pale white-brown phases visible on less dusty surfaces. Natural color (a) and Gaussian stretch (b) images of the same region are shown. Regolith and loose grains have accumulated in depressions at the outcrop surface.

Pre and Post Coring/Abrasion Images

Figure 5 shows the natural surface of Brac prior to coring of Salette. Figure 6a shows the WATSON pre-coring image of the natural surface of Dourbes, Figure 6b shows a WATSON image of the gDRT-cleared abraded patch and Figures 6c–d show the corresponding WATSON ACI images.

Notable observations of high-resolution pre- and post-coring images include:

a) The natural weathered surface of Brac is generally smooth but has numerous small depressions that are filled with dust and other fine-grained, likely wind-blown, materials. There is some evidence of fluting caused by wind erosion. Few, if any, individual grains are visible in the natural surface images (Figures 5–6), but abundant grains were noted in RMIs of less-dusty surfaces (Figure 4).
b) 0.1–0.3 mm pale brown–white sub-angular phases dominate the Dourbes abrasion patch (Figure 6b). Other major constituents are 0.2–0.8 mm mid-brown crystals with sub-angular to angular morphologies, 0.1–0.4 mm grey–green crystals with angular and elongate morphologies, <0.1 mm white crystals and a single white crystal with a triangular cross-section. The pale brown–white, mid-brown, grey–green and white minerals are distributed throughout the abrasion patch with no obvious spatial heterogeneity in their relative abundances.

c) Most minerals within the abrasion patch are surrounded by mid-brown inter-crystal boundaries, suggesting a pervasive Fe-rich influence. Pale reddish rims also surround many olivine crystals. This apparent Fe staining is particularly strong in association with the pale brown–white phases that dominate much of the abrasion patch. This association is especially visible in the ACI images where its interstitial nature is obvious.

WATSON post-coring images were acquired on the Coulettes sampling site (Figure 7). Note the well-formed conical mound of tailings, and, on the underlying rock, the purple-hued coating frequently seen on crater floor rocks.

CacheCam images of the acquired samples in their sample tubes are shown in Figure 8. Although the Coulettes core end is slightly out of focus because the core was not full length, the image of the full length Salette core shows the core end to be very similar in color and texture to the Dourbes abrasion patch, suggesting little lithologic change to a depth of ~6 cm in Brac.
Figure 6 | **WATSON images of the Dourbes abrasion surface (sol 257).** (a) Pre-abrasion WATSON 8 cm standoff image showing the natural surface of Dourbes. (b) WATSON image of the entire abrasion target. (c-d) ACI images, grayscale and colorized, of a region of interest within the Dourbes abrasion patch. Note that the WATSON images are rotated with respect to the images obtained by the remote sensing mast.

Figure 7 | **Post-coring WATSON image of the Salette borehole.**
Figure 8 | CacheCam images of the *Salette* (left) and *Coulettes* (right) samples. Inset is a high-resolution montage of CacheCam frames of *Salette*.

*Elemental Geochemistry - PIXL*

Elemental geochemistry was determined using X-ray fluorescence (XRF) mapping by the PIXL instrument on the *Dourbes* abrasion patch. XRF rasters were acquired on sols 257 and 270 (*Figure 9*). This report includes only limited data from the sol 270 raster. Based on texture, bulk chemistry, and mineral chemistry, *Brac/Dourbes* is inferred to be a cumulate igneous rock formed by settling of olivine crystals and subsequent crystallization of interstitial basaltic magma.

The bulk sum analysis for the sol 257 raster, i.e. the average chemical composition, is given in *Table 2*. *Brac* is clearly a silicate rock, and high abundances of MgO, FeO, and Al₂O₃ suggest an igneous basaltic parentage. Compared to a typical basalt (terrestrial or martian), it is low in SiO₂, Al₂O₃ and MgO, and rich in FeO, Na₂O, SO₃ and Cl. Note that this bulk sum represents a mixture of what are believed to be anhydrous primary silicate minerals (olivine, pyroxene and plagioclase) and secondary (aqueous alteration) volatile-bearing minerals. In addition to the tabulated values, the bulk composition contains 550±200 (1s) ppm Ni; all other minor and trace elements are below detection limits.

On a total alkali versus silica (TAS) plot, *Figure 10*, PIXL analyses are distributed across the fields of olivine, microbasalt, and basalt. Some points also fall within the basaltic andesite field and a few points follow the trachybasalt–trachyte trend (a few points have SiO₂>55 wt%). Several also plot near the Fe-Ti oxide and apatite positions. The bulk sum of the PIXL analysis raster falls within the olivine–basalt field. *Figure 11* shows a ternary diagram of molar compositions Al₂O₃ – (CaO+Na₂O+K₂O) – (FeO⁺+MgO) for every individual PIXL point in the sol 257 raster and the bulk sum. Idealized common mineral compositions are also plotted on the diagram. The majority of points fall inside the triangle defined by...
typical mafic minerals (e.g., olivine, Ca-rich pyroxene, feldspar), and so are consistent with being mixtures of those minerals. PIXL data points along the olivine–clinopyroxene and olivine–feldspar lines represent binary mixtures of these minerals, arising because their analytical volumes included both minerals. Almost no points fall above the feldspar–olivine line, i.e. enriched in Al₂O₃ as would be expected for a weathered rock. Overall, this plot indicates that Dourbes is neither strongly aqueously altered nor intensely weathered.

Figure 9 | PIXL raster scans on Dourbes. Approximate locations of PIXL XRF scans on the Dourbes abrasion patch (field of view = ~5 cm across), annotated with the corresponding sol. Sol 0257 scan is 12.5 x 4 mm; sol 0270 scan is 7 x 5 mm. Olivine grains are pale buff, augite is greenish-black, and poorly diffracting Fe-Mg-poor silicate material is brown. Locations are overlain on WATSON image SIF_0257_0689757397_902FDR_N0080000SRLC00672_0000LMJ01, rotated 90° clockwise from its original orientation.
Table 1. Chemical compositions by PIXL XRF.

<table>
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<th>Bulk rasters 257&amp;270</th>
<th>Olivine salt-free</th>
<th>Augite salt-free</th>
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<tr>
<td>Wt %</td>
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</tr>
<tr>
<td>Mg#</td>
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<td>55</td>
<td>68</td>
</tr>
</tbody>
</table>

N indicates the number of analyses averaged. Err is uncertainty (1σ) from calibration and counting statistics. Mg# is molar Mg/(Mg+Fe), in percent.

PIXL multi-elemental maps show abundant euhedral to subhedral olivine grains (~Fo55) varying in size between 1 and 3.5 mm, together with anhedral augite (<8 mm). Olivine is the most abundant mineral, accounting for more than 60 vol% of the mafic material in the Dourbes abrasion patch (Figure 12). Olivine grains are partially enclosed by augite in a poikilitic texture commonly seen in igneous rocks on Earth, as well as in the chassignite Martian meteorites. Average chemical compositions of olivine and augite are given in Table 2. Back-reflection diffracted x-rays detected from augite regions indicate that they are monocrystalline and crystallographically oriented across non-contiguous areas separated by olivine in the abrasion patch. This feature is characteristic of pyroxenes in many olivine cumulate rocks. Other components identified in PIXL scans include mesostasis material among the olivine and augite, and secondary phases. The mesostasis consists of a mixture of Na-rich feldspar (<1 mm), Fe-Cr-Ti magnetite (≤0.25 mm), K-rich feldspar (<0.125 mm) and Ca-phosphates (<0.125 mm). Secondary phases include poorly diffracting (or very fine grained) silicate(s) with low Fe and Mg, Fe-Mg carbonate, Ca-sulfate, Mg-sulfate (likely MgSO₄·4H₂O), and Na chloride and/or perchlorate. These secondary materials, interpreted to be products of post-igneous aqueous alteration, occur in light-toned patches (primarily salt minerals) and in reddish brown patches (low Fe-Mg silicates) surrounding and invading anhydrous minerals, especially olivine.

PIXL Micro-Context Camera (MMC) images of the Dourbes abrasion patch (Figure 13) show more of the abrasion area than the XRF scans (Figure 13a) and are used to extrapolate phase proportions from those of the XRF scan areas to nearly the whole abrasion patch. The MMC image of NIR-G-B (Figure 13b) is analogous to WATSON visible images, and the G/NIR ratio image shows a clear distinction among the olivine, augite, and mesostasis phases (Figure 13c). These visible & qualitative relationships are
quantified in a graph of G versus NIR reflectances (Figure 13d), which allows extraction of the proportions of these phases in the whole MMC image area.

**Figure 10 | Total alkali versus silica (TAS) plot for the Dourbes abrasion, PIXL raster of sol 257.** Individual data points represent regions of the abraded surface (~100 µm) and therefore a mixture of one or more primary and/or secondary minerals. The bulk sum composition of PIXL data (Table 2) is shown by the blue circle. Reference end-member igneous mineral compositions are shown as diamonds and lines for those with solid solution series. The underlying igneous classification scheme does not apply to the individual data points and simply provides a frame of reference for comparing samples.
Figure 11 | Mineralogical interpretation of the Dourbes abrasion, PIXL raster of sol 257. Data are plotted by pixel and as bulk sum composition (blue circle, Table 1) on ternary diagrams of molar abundances of $\text{Al}_2\text{O}_3$–$(\text{CaO}+\text{Na}_2\text{O}+\text{K}_2\text{O})$–$(\text{FeO}_\text{T}+\text{MgO})$. Common pure mineral compositions are shown ($\text{opx} = \text{Ca}$-poor pyroxene, i.e. orthopyroxene; $\text{cpx} = \text{calcic pyroxene}$, i.e. augite). Most of the individual PIXL analysis points are consistent with being either pure minerals (olivine, cpx) or binary mixtures of olivine–cpx or olivine–feldspar, or ternary mixtures of olivine–cpx–feldspar. Aluminous alteration minerals, including common clay minerals, fall above the upper red dashed line. Several other secondary minerals (e.g., Fe-Mg-Ca-sulfates/carbonates, halite) are also plotted.
Figure 12 | PIXL X-ray multi-element maps for the Dourbes abrasion patch, PIXL raster sol 257; raster is 12.5 x 4 mm. (a) Red=MgO, Green=Al₂O₃, Blue=CaO. (b) Red=CaO, Green=Na₂O, Blue=K₂O. (c) Red=MgO, Green=Na₂O, Blue=Cl. (d) Red=FeO, Green=TiO₂, Blue=Cr₂O₃. Olivine is the most abundant material (red in panel a). Augite pyroxene is purple in panel a, and red in panel b. Feldspars are green in panel a. Salt is green in panel b, and white in panel c. Panel d shows Fe oxide minerals: Ti-rich oxides are yellow, Cr-rich oxides are purple.
Figure 13 | PIXL MCC (Micro-context camera) data for the Dourbes abrasion, sol 257. (a) MMC grayscale image showing a portion of the Dourbes abrasion. (b) MMC false color of abrasion area; Red = MMC Near-IR; Green = MMC Green; Blue = MMC Blue. Location of the PIXL XRF scan is shown. (c) MMC ratio image Near-IR/Green reflectances. Augite, A, is darkest; olivine, O, is mid-toned, and mesostasis and alteration materials have the lightest tone. Location of PIXL XRF scan shown. (d) MMC reflectance ratios.
**Mineralogy and Organics-SHERLOC**

SHERLOC Raman and florescence scans were performed on *Dourbes* on sols 257 (HDR and survey scans; **Figure 14**) and 269 (survey scan and three detail scans; **Figure 15**). The analytical footprints are shown in the corresponding figures.

In the sol 257 scans, Raman data show spectra with peaks assigned to Ca/Mg-sulfate (~1050 cm\(^{-1}\)), carbonate (~1090 cm\(^{-1}\)) and olivine (~840 cm\(^{-1}\)). Some spectra contain a possible hydration peak at ~3450 cm\(^{-1}\) and a broader peak at ~997 cm\(^{-1}\) assigned to pyroxene or sulphate. Fluorescence features at 290 and 330–340 nm are consistent with single and double ring aromatic organic molecules, respectively. The relatively low intensity of fluorescence at all wavelengths and absence of Raman organic signatures suggests that, if organics are present, they occur with low abundances and are generally dispersed with the exception of some localized hot spots.

In the sol 269 scans, Raman spectra show features assigned to Ca- and/or Mg-sulfate (1020 cm\(^{-1}\)) and Ca/Mg(Fe)-carbonate (1100 cm\(^{-1}\)). Some spectra also exhibit a peak for hydration at ~3450 cm\(^{-1}\). Detail scan 3 shows a possible detection of olivine based on a broad band centered near 840 cm\(^{-1}\). The fluorescence feature at 330–340 nm is consistent with double ring aromatic organic molecules. As in the sol 257 scan, the relatively low intensity of fluorescence at all wavelengths and absence of Raman organic signatures suggests that any organics present have low abundance and/or a dispersed distribution. SHERLOC-based mineral identifications are shown in **Table 2**.

**Table 2. SHERLOC mineral identifications in *Dourbes* abrasion patch.**

<table>
<thead>
<tr>
<th>Sample</th>
<th>Certain</th>
<th>Possible (not confirmed)</th>
<th>We looked for these, but cannot find them</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dourbes</td>
<td>Sulfate (most likely Ca or Mg), occasionally hydrated (occurs in white void fills or crystals, secondary to primary lithology)</td>
<td>Olivine (major phase, low Fe#) occurs sometimes associated with darker grains. Carbonate (major phase, most likely Ca/Mg/Fe) Amorphous/microcrystalline silicate Pyroxene</td>
<td></td>
</tr>
</tbody>
</table>

5-17
Figure 15 | SHERLOC fluorescence and Raman spectral results for Dourbes. Spectra were collected with an HDR scan (7x7 mm, 100 points, 780 µm spacing, 500 pulses per point) and a survey scan on sol 257 (5x5 mm, 1296 points, 144 µm spacing, 15 pulses per point, not shown). A) Colorized ACI image of Dourbes; blue rectangle indicates the HDR scan area (sol 257) and white circles indicate regions of interest (ROIs) correlating to the fluorescence data presented in B and C. B) HDR scan maps showing the possible presence of organics. Upper panel shows an RGB map indicating the main regions of organic fluorescence (290 and 330–340 nm). Both doublet peaks at 290 and 330 nm (ROI1) and the 330–340 nm peak (ROI 2) are localized primarily within two ROIs (see panel A). The lower panel is blank because no Raman features consistent with organic materials were detected. C) Fluorescence spectra. The upper panel shows the average fluorescence spectrum for the whole area of analysis and the lower panel includes spectra from selected fluorescence “regions of interest” (see A). D) Map showing the results of the Raman HDR scan. The grid of red unfilled circles indicates the points analyzed (cf. A), while blue, yellow, green, brown and purple filled circles indicate mineral detections. The right panel includes selected Raman spectra from the HDR scan. The upper spectrum is the average spectrum of the HDR scan. With the exception of cosmic ray artifacts, features within are rather weak, but do match some of the features in the spectra below. The spectra labeled Pts. 2, 1, and 18 correspond to the points in the map panels with the same numbers and are representative of spectra assigned to (Ca)/Mg-sulfate (~1050 cm\(^{-1}\)), carbonate (~1090 cm\(^{-1}\)) and olivine (~840 cm\(^{-1}\)), respectively. The spectrum labeled Pt. 2 also contains a possible hydration peak at ~3450 cm\(^{-1}\). The lowermost spectrum includes a broader peak at ~997 cm\(^{-1}\) assigned to potential pyroxene or sulfate (consistent with hydrated MgSO\(_4\)). The detection of pyroxene in this region by PIXL gives confidence to this interpretation, but given the broadness of the peak and absence of secondary peaks at lower wavenumber (not detectable with SHERLOC), sulfate is also a possibility. The scale in panel A also applies to the maps in panel D.
Figure 16 | SHERLOC fluorescence and Raman spectral results for Dourbes. Spectra were collected with a survey scan (5x5 mm, 1296 points, 144 µm spacing, 15 pulses per point, not shown) and three detail scans (1x1 mm, 100 points, 144 µm spacing, 500 pulses per point) on sol 269. A) Colorized ACI image of the abraded patch; blue rectangle indicates the survey scan area and red squares indicate detail scans 1 and 3. B) Detail scan 1 map showing the possible presence of organics. The upper panel shows an RGB map indicating the main regions of organic fluorescence (330–340 nm). The 330–340 nm peak (ROI 1) is localized mainly in “regions of interests” (ROI). The lower panel is blank because no Raman features consistent with organic materials were detected. C) Fluorescence spectra. The upper panel shows the average fluorescence spectrum for the whole analysis area and the lower panel includes spectra of selected fluorescence from ROI 1 (see A). D) Maps showing the results of the Raman detail scans 1 and 3. In each, the grid of red unfilled circles indicates the points analyzed (cf. A), and the blue, yellow, green, brown and purple filled circles indicate mineral detections. The right panel includes selected Raman spectra from the two scans. The spectra labeled Pts. 12 and 47 correspond to the points in the map panels with the same numbers and are representative of spectra assigned to Ca or Mg-sulfate (1050 cm⁻¹) and Ca/Mg-carbonate (1100 cm⁻¹). The spectrum labeled Pt. 12 also exhibits a peak for hydration at ~3450 cm⁻¹. The lower spectrum is the average spectrum of the points labeled with yellow circles in detail scan 3, which are interpreted to represent possible detections of olivine based on a broad band centered near 840 cm⁻¹. The scale in panel A also applies to the maps in panel D.
SCAM LIBS data from *Brac* and other Séítah targets, when plotted on the total alkalis versus silica (TAS) diagram (*Figure 17*), are scattered within the picrobasalt and basalt fields, and to a lesser extent within the olivine and basaltic andesite fields. The SCAM LIBS analyses targeted primarily natural surfaces, with only a small subset targeting the abraded patch. Olivine is the dominant detection and bulk sum average in the PIXL data obtained on the abraded patch, but the SCAM LIBS data differ, likely owing to natural variability in *Séítah*. Nonetheless, SCAM data are consistent with PIXL data in observing few points above SiO$_2$=55 wt%.

Harker oxide diagrams of SCAM measurements of *Brac* are plotted against a background of all *Séítah* points in *Figure 18*. These diagrams show that *Brac* is generally representative of *Séítah*; potentially with lower CaO, Al$_2$O$_3$ and Na$_2$O contents.

Note that the ~250 µm LIBS spot most commonly interrogates more than one mineral grain. Given that the typical Dourbes grain size is 0.2–0.5 mm, as described above, many of the trends observed in the diagrams shown in *Figures 18 and 19* can be reasonably attributed to mixing among mafic minerals, e.g., olivine and low-Ca pyroxene.

*Figure 17 | TAS plot of SCAM LIBS data.* Average value of all data is shown as a blue circle. All *Séítah* analyses, i.e., including those not from the *Brac* outcrop, are plotted as grey points.
Figure 18 | Harker oxide diagrams of SCAM data from Séítah. Data from the Brac outcrop are shown as colored points. All Séítah points, including those not from Brac, are shown as black points.

Figure 19 shows mean VISIR spectra of the natural surface, the Dourbes abraded patch, and the tailings of the Dourbes abrasion and Salette coring acquired between sols 247 and 278. The broad absorption at 0.8–1.8 µm denotes olivine in all observations. This signal is stronger in the abraded surface measurements relative to the natural surface. This is consistent with the stronger evidence for olivine in the PIXL abraded patch data relative to the SCAM LIBS natural surface data. The broad slope at 2.2–2.5 µm denotes the presence of hydrated minerals. Specifically, the absorption at 2.28–2.29 µm likely indicates Fe-bearing phyllosilicate, the absorption at 2.32 µm likely indicates Mg-phyllosilicate, and the broad absorption centered around 2.3 µm is consistent with Ca/Mg/Fe-carbonates and/or Fe/Mg-phyllosilicates.
Figure 19 | Mean VISIR spectra of the natural surface of Brac, the abraded patch Dourbes, the tailings of the Salette core and the tailings of the Dourbes abraded patch rasters. The broad slope between 2.2 and 2.5 µm is consistent with the presence of hydrated minerals. The broad absorption between 0.8 and 1.8 µm indicates olivine in all observations; the signal is stronger in the abraded surface relative to the natural surface. The absorption at 2.28–2.29 µm likely indicates Fe-bearing phyllosilicates, the absorption at 2.32 µm likely indicates Mg-phyllosilicates, and the broad absorption centered around 2.3 µm is consistent with Fe/Mg-phyllosilicates. Note that owing to temperature sensitivities in the IRS instrument, the calibration is uncertain for wavelengths longer than ~2.5 µm.
Subsurface Structure

Subsurface structure in the vicinity of the Brac outcrop was characterized by RIMFAX radargrams acquired on sols 237–239 during the rover’s drive across Séítah on a mostly easterly traverse heading toward Brac (Figure 20). From left to right, the three panels in Figure 20a present results along the rover path that 1) begins ENE past the Village region, 2) turns ESE toward Vignols, then turns NE toward Caille and the Brac outcrop. The interpreted subsurface stratigraphy and rock density is shown in Figure 20b. Throughout this traverse, the data indicate SW-dipping to sub-horizontal reflectors that dominate the stratigraphy at ~3–8 m below the surface, which is interpreted to represent an extension of the SW-dipping reflector unit extending NE from Artuby Ridge (see the Initial Report for samples Montdenier and Montagnac from the Rochette locality). Above this unit is a 2–3 m thick laminated sequence with dominantly sub-horizontal reflectors (the unit containing Brac). A particularly intense reflector may be an erosional unconformity between these two packages of reflectors. Permittivity measurements of the upper laminated sequence indicate rock densities of approximately 3.5 g/cm³, consistent with the olivine-rich lithology observed in the Brac outcrop.

Figure 20 | Geological context of the Brac outcrop from RIMFAX  
a) Radargram for the sol 237–239 traverse. 
b) Geological interpretation of reflector geometries in panel a with Brac at sol 240 EOD (annotated with red star). The stratigraphy is identified by SW-dipping to sub-horizontal reflectors in the lower unit, sub-horizontal reflectors in the upper unit (including Brac), and scattered reflection hyperbolae in between. The locations of permittivity measurements (\(\epsilon'\)) from reflection hyperbolae are denoted with yellow circles. Reflection hyperbolae 1–4 are from sol 237 and 1' and 2' are from sol 280. These and additional nearby permittivity measurements within the sol 237–280 radargrams cluster around \(\epsilon' = 10\). Reflection hyperbolae record the average permittivity within the overlying strata. Permittivity–density relationships are determined after Carrier III et al. (1991).
Core orientation

At the time of drilling and 6.6 cm pre-drilling WATSON imaging, the rover, coring drill, WATSON, and coring target (“Salette.PSC.0.0679”), had the following characteristics:

1. Rover orientation quaternion just after drilling but before unloading the stabilizers (transferring from RMECH to SITE frame): \( bQl1 = (0.8161970, -0.0570105, 0.0018349, 0.5749510) \);
2. Coring Drill orientation quaternion just after drilling but before unloading the stabilizers (transferring from CORING DRILL to RMECH frame): \( cdQb = (0.7611450, 0.1932350, -0.6043060, -0.1346610) \);
3. Rover orientation quaternion at time of acquisition of WATSON image \( SIF_0195_0684255577_390FDR_N0070000SRLC00701_0000LMJ01 \): (transferring from RMECH to SITE frame): \( bQl2 = (0.816367, -0.0553839, -0.000861533, 0.574871) \);
4. WATSON orientation quaternion at time of acquisition of WATSON image \( SIF_0195_0684255577_390FDR_N0070000SRLC00701_0000LMJ01 \): (transferring from WATSON to RMECH frame): \( wQb = (0.761644, 0.197503, -0.60257, 0.133416) \).

Items 1 and 2 give a coring drill pointing vector estimate of

\[
\text{hade} = 17.78^\circ \\
\text{azimuth} = 81.51^\circ
\]

Items 3 and 4 give an estimate of the angle between the WATSON y-axis and the up-dip direction in SITE coordinates (Figure 21) of

\[
\text{core roll, } \alpha = 253.58^\circ
\]

Figure 21 | 6.6-cm standoff WATSON image of Salette core target on sol 262. Image is 30.9 μm pixel⁻¹. Orientation compass gives WATSON frame \((w_x, w_y, w_z)\). Core roll is clockwise angle of core \(y\)-axis, \(c_y\), from \(w_x\), given by \(\alpha = 253.58^\circ\). Projection of Martian geographic north onto WATSON image plane is noted. WATSON image \( SIF_0195_0684255577_390FDR_N0070000SRLC00701_0000LMJ01 \).
Preliminary Scientific Assessment

Synthetic sample description and preliminary interpretation

Sample type: Olivine cumulate

1. Relationship with surrounding rocks
   a. *Salette* and *Coulettes* were obtained from a position high in the accessible stratigraphy of *Séítah*, near the location of overlying pitted rocks (e.g. the Hôtel rock).
   b. Based on its appearance, concordance with surrounding outcrops, and estimated dip consistent with that of the wider *Séítah* region, the *Brac* outcrop is interpreted to be an in-place portion of the *Séítah* bedrock.

2. Texture and fabric
   a. Grain size: <0.1–4 mm
   b. Rock fabric (natural surface): large olivine grains surrounded by interstitial pale materials exposed on relatively fresh surfaces, but minimal observations of grains on natural weathered surfaces.
   c. Rock fabric (abrasion patch): no obvious textural heterogeneity, main constituents include 0.1–0.3 mm pale brown–white sub-angular phases, 0.2–0.8 mm mid-brown sub-angular to angular crystals, 0.1–4 mm grey–green angular and elongate crystals, <0.1 mm white crystals.
   d. There is no compelling evidence of intergranular porosity or cements common to many sedimentary rocks.

3. Mineralogy and chemistry
   a. Minerals present: dominant olivine and low-Ca pyroxene (augite, potential ferrosilite). Mesostasis consists of a mixture of Na-rich feldspar (<1 mm), Fe-Cr-Ti magnetite (≤0.25 mm), K-rich feldspar (<0.125 mm) and Ca-phosphates (<0.125 mm). Secondary phases include poorly diffracting (or very fine grained) silicate(s) with low Fe and Mg, Fe-Mg carbonate, Ca-sulfate, Mg-sulfate (likely MgSO₄·4H₂O), and Na chloride and/or
perchlorate. Aromatic organic materials are present, but in low abundance and dispersed throughout the rock matrix except for a few hot spots.

b. The bulk composition is consistent with a mafic rock modified by a low degree of aqueous alteration and weathering. Point analyses by SCAM and PIXL plot mostly within the olivine, picrobasalt and basalt regions of the TAS diagram.

4. **Alteration/secondary characteristics**
   a. Interactions with aqueous fluids, alteration, and mineral hydration: some, e.g. carbonate and sulfate signatures.
   b. Secondary mineralogy: see point 3a above.
Returned Sample Science Considerations

Observations of olivine surrounded by clinopyroxene in a poikilitic texture within the Dourbes abrasion patch, along with remote observations of the mineralogy, texture, and spectral characteristics of Brac and other Séítah outcrops, suggest that the Salette (and Coulettes) core and much or all of Séítah are best interpreted as an igneous olivine cumulate. As such, the return of samples from Brac is expected to address a number of RSS themes of significance to the Mars 2020 mission:

**Geochronology:** The timing of volcanic rock crystallization can be quantified using laboratory-based, radiometric dating of returned samples. Likewise, assuming cosmogenic nuclide production rates in specific mineral on Mars can be established, quantitative constraints on the sample’s surface exposure age and/or erosion rate may also be obtained. Quantifying the timing of crystallization of Brac will also constrain the emplacement ages of other crater floor units, given their stratigraphic relationships (e.g. Séítah is likely older than Máaz from which the Montdenier and Montagnac samples were collected).

**Paleomagnetism:** The igneous lithologies of the Salette/Coulettes samples may enable absolute measurements of the paleointensity of the ancient Martian magnetic field at the time of crystallization. The orientations of the samples, combined with paleohorizontal indicators from outcrop foliation, may enable measurements of the absolute paleodirection of the ancient field. Paleomagnetic investigations of secondary ferromagnetic minerals if present could constrain the field at the time of aqueous alteration. Combined with measurements from other samples, these data could test the hypothesis that Martian atmospheric loss was driven by the cessation of an early dynamo, constrain the dynamo’s secular variation, and determine if the dynamo exhibited polarity reversals. Paleomagnetic analyses could also test whether Séítah rocks have experienced regional or global-scale tectonic tilting and displacements. Comparison of paleomagnetic directions with Máaz rocks (Montdenier/Montagnac and Atsah/Hahonih) could distinguish between the hypotheses that Máaz and Séítah form a single layered intrusion versus Máaz being an overlying lava flow deposited on Séítah.

**Geochemistry:** With a hypothesized origin as an olivine cumulate, understanding the petrogenesis of the Salette and Coulettes samples will place constraints on the nature of igneous processes on early Mars. These constraints can be applied to the origins of the extensive “Cf-f-1” unit that has been correlated with a regional olivine-bearing unit exposed between Syrtis Major and Nili Fossae. The diverse mineral assemblages of the obtained samples will be valuable for quantifying the conditions and timing of primary, high-temperature crystallization of igneous phases and later-stage alteration events. The bulk chemistry of the rock, as well as trace element and stable and radiogenic isotope geochemistry of the rock and its constituent igneous minerals, will enable comparisons with similar lithologies in the Martian meteorite suite, and insights into magmagenesis and mantle/crustal evolution more generally. The large grains observed in Brac may contain melt inclusions that are of sufficient size to track the petrogenesis of this material.

**Water–rock interactions and habitability:** Many hypotheses for the origins of olivine-carbonate rocks involve aqueous alteration of igneous rocks. The interaction of aqueous media with (ultra)mafic rocks produces secondary mineral phases that can be indicative of the temperature, redox state, and fluid chemistry. High resolution mineral identification and stable isotope measurements can place constraints on such aspects of the alteration history and environment and provide insights into the presence, distribution, and nature of water on Mars. Water/rock interactions could have occurred as result of
groundwater, possibly associated with Jezero lake, moving through these rocks and may have led to the establishment of microniches with water, energy, pH, nutrients and transition metals essential to life. Although likely a variable component in the history of these samples, the team has noted that low-temperature hydrothermal, serpentinizing and carbonation reactions may have sustained microbial life, particularly chemoautotrophs/chemolithotrophs that have been suggested as likely inhabitants of Martian biotopes. Although the fossil record of igneous rocks is highly variable and generally poor, the biosignature preservation potential of certain alteration phases associated with the Brac outcrop is higher, and it is possible that organic materials and/or biosignatures have been preserved within these phases.

References


INITIAL REPORT

M2020-271-6 Coulettes

Sample Designation: M2020-271-6 Coulettes

Date of Coring: 24-November-2021

Mars Time of Sample Core Sealing: 20:18 LMST, Sol 271, Ls = 132.0

Latitude (N), Longitude (E), Elevation: 18.43397°, 77.44301°, −2569.2 m

Campaign: Crater Floor

Region of Interest: Séítah South

Lithology: Medium- to coarse-grained poikilitic olivine cumulate rock. Primary minerals are olivine and pyroxene. Aqueous alteration phases include sulfates, carbonates and hydrated phases. Organic matter was detected.

Estimated Volume Recovered: 4.7 cm³

Coring Bit Number: 1

Core Orientation: hade = 14.50°; azimuth = 78.68°; core roll = 220.85°

Sample Serial Numbers: Tube SN284; Seal SN219; Ferrule SN189

ACA Temperature at Time of Sealing: 40°C

Estimated Rover-Ambient Pressure and Temperature at Time of Sealing: 636 Pa, 222 K

Estimated Amount of Martian Atmosphere Headspace Gas: 2.5 x10⁻⁶ mol

Anomalous Behavior: None

Abrasion Patch Name and Depth: Dourbes, 8 mm

January 27, 2022

Summary Description

This sample is paired with M2020-262-5 Salette. Only the core orientation for Coulettes is described below. See the Initial Report for M2020-262-5 Salette for sample details.

Core orientation

At the time of drilling and 6.6-cm pre-drilling WATSON imaging, the rover, Coring Drill, WATSON, and coring target (called “Coulettes_271.PSC.0.0656”), had the following characteristics:

1. Rover orientation quaternion just after drilling but before unloading the stabilizers (transferring from RMECH to SITE frame): \( \mathbf{bQl1} = (0.8161870000, -0.0570269000, 0.0016600700, 0.5749640000) \);
2. Coring Drill orientation quaternion just after drilling but before unloading the stabilizers (transferring from CORING DRILL to RMECH frame): \( \mathbf{cdQb} = (0.6692760, 0.3774960, -0.5735080, 0.2839970) \);
3. Rover orientation quaternion at time of acquisition of WATSON image SIF_0195_0684255577_390FDR_N0070000SRLC00701_0000LMJ01: (transferring from RMECH to SITE frame): \( \mathbf{bQl2} = (0.816343, -0.0553775, -0.000576532, 0.574906) \);
4. WATSON orientation quaternion at time of acquisition of WATSON image SIF_0195_0684255577_390FDR_N0070000SRLC00701_0000LMJ01: (transferring from WATSON to RMECH frame): \( \mathbf{wQb} = (0.669666, 0.383208, -0.570312, 0.281855) \).

Items 1 and 2 give a coring drill pointing vector estimate of

\[ \text{hade} = 14.50^\circ \]
\[ \text{azimuth} = 78.68^\circ \]

Items 3 and 4 give an estimate of the angle between the WATSON y-axis and the up-dip direction in SITE coordinates of (Figure 1):

\[ \text{core roll, } \alpha = 220.85^\circ \]
Figure 1 | 6.6-cm standoff WATSON image of Coulettes core target on sol 271. Image scale is 30.7 μm pixel\(^{-1}\). Orientation compass gives WATSON frame (\(\mathbf{w}_x, \mathbf{w}_y, \mathbf{w}_z\)). Core roll is clockwise angle of core y-axis, \(\mathbf{c}_y\), from \(\mathbf{w}_y\), given by \(\alpha = 220.85^\circ\). Projection of Martian geographic north onto WATSON image plane is noted. WATSON image SIF_0195_0684255577_390FDR_N0070000SRLC00701_0000LMJ01.
**INITIAL REPORT**

**M2020-295-7 Robine**

**Sample Designation:** M2020-295-7 Robine

**Date of Coring:** 18-December-2021

**Mars Time of Sample Core Sealing:** 17:22 LMST, Sol 298, Ls 143.4

**Latitude (N), Longitude (E), Elevation:** 18.43264°, 77.44133°, –2574.3 m

**Campaign:** Crater Floor

**Region of Interest:** Séítah South

**Lithology:** Medium- to coarse-grained poikilitic olivine cumulate rock. Primary minerals are olivine and pyroxene. Limited secondary aqueous alteration, including carbonate, hydrated sulfate, and other hydrated phases. Organic material is present, at least partially associated with sulfate phase(s).

**Estimated Volume Recovered:** 8.6 cm³

**Coring Bit Number:** 2

**Core Orientation:** hade = 5.36°; azimuth = 225.58°; core roll = 231.60°

**Sample Serial Numbers:** Tube SN206; Seal SN184; Ferrule SN064

**ACA Temperature at Time of Sealing:** 40°C

**Estimated Rover-Ambient Pressure and Temperature at Time of Sealing:** 608 Pa, 246 K

**Estimated Amount of Martian Atmosphere Headspace Gas:** 1.0 x10⁶ mol

**Anomalous Behavior:** Three sols between coring and sealing.

**Abrasion Patch Name and Depth:** Quartier, 8 mm

January 27, 2022

**Summary Description**

*Robine* and *Malay*, the pair of rock cores representing the fourth sample target of the Mars 2020 *Perseverance* rover mission, were collected in the *South Séítah* region on sols 295–298 and 337, respectively (Figures 1–2). Both samples were acquired from an outcrop named *Issole*, a ~5-20 cm high NE–SW-trending ridge of layered bedrock within the *Séítah* formation that parallels the larger ~2 m high *Artuby* ridge of the *Máaz* formation. *Issole* was collected in part because it was the *Séítah* outcrop closest to the *Máaz* contact along the South *Séítah* traverse.

*Séítah* is mapped as part of the olivine-bearing Crater Floor Fractured 1 unit defined from orbital observations (Cf-f-1 of Stack et al., 2020). The *Séítah* formation includes the stratigraphically lowest rocks exposed on the floor of Jezero crater, including a distinct light-toned olivine-bearing unit. This unit has been correlated with a regional olivine-bearing unit exposed between Syrtis Major and Nili Fossae.

The natural surface of *Issole* features large olivine grains and chemically-distinct purple patches. The *Quartier* abrasion patch contains abundant 1-4 mm size olivine crystals surrounded by augite, as well as feldspars, magnetite, and Ca-phosphates. *Issole* is interpreted to be an olivine cumulate formed either in a thick lava flow or in a sill. It is thus unclear whether *Issole* represents an extrusive or intrusive igneous rock. *Issole* shows evidence of limited aqueous alteration in the form of sulfates, carbonates, and chlorides or perchlorates. Organic compounds are present, especially in association with sulfates, but in low abundance.

*Issole* is interpreted to be in-place. Rover observations have shown conclusively that the *Séítah* formation, and in particular the reflector that outcrops at *Issole*, dip below the *Máaz* formation along *Artuby* ridge. Despite this relationship, the relative ages of *Máaz* and *Séítah* depend on whether the olivine cumulate is an intrusive or extrusive lithology.

*Issole* is located about 120 meters SW of *Brac*, a geochemically and mineralogically similar *Séítah* outcrop that yielded the *Salette* and *Coulettes* cores. Although we interpret both sets of samples to derive from a single olivine cumulate body, compared to the *Brac* cores, *Robine* and *Malay* appear to be less mafic and to have experienced a greater degree of aqueous alteration.

Sample processing and sealing of *Robine* occurred 3 sols after coring. *Malay* was acquired in a single sol.

The returned sample science objectives of this core pair are similar to those of *Salette* and *Coulettes*, and include geochronology, paleomagnetism, igneous geochemistry, and the past history of water and water–rock interactions in Jezero crater. Together with cores of the *Máaz* formation (*Montdenier, Montagnac, Atsah, Hahonih*), the four *Séítah* cores constitute the entirely igneous Crater Floor Campaign sample suite.
Figure 1 | Regional context of *Perseverance* rover operations during acquisition of *Robine* and *Malay*. (a) Geological map showing Jezero crater floor units. Light green shading indicates the location of the Jezero Crater Floor campaign. (b) Geological map showing the crater floor Cf-fr and Cf-f-1 units. Adapted from Stack et al. (2020).
Figure 2 | Local context of Perseverance rover operations during sampling of Robine and Malay. HIRISE map showing the locations of sample collection together with other notable outcrops observed or analyzed in the sols leading up to Issole. Rover position at Issole is highlighted by a blue circle. Issole is located along a ridge near the NW–SE-trending boundary with the Máaz formation (Cf-fr in Stack et al., 2020) along the ridge labelled Artuby West.

Stratigraphic and Geologic Context

The Robine core (and paired core Malay) were collected from the Issole outcrop, the second Séitah sampling locality. The Issole outcrop is located near the boundary between the Séitah (Cf-f-1) and Máaz (Cf-fr) formations in the south Séitah region (Figure 1; Stack et al. 2020). The formation boundary is covered by regolith but its location is known to within a few meters from outcrop lithology.

Orbital mapping suggests that Séitah rocks are the stratigraphically lowest exposed rock unit on the crater floor and are overlain by both the Máaz formation and the delta. Rover observations confirm that Séitah lies below Máaz. When this sample pair was collected, Perseverance had not yet explored the relationship of the crater floor units to the delta. Consistent with its lower position in the stratigraphy, Séitah may be older than Máaz. However, because Séitah could be intrusive, the inverse is also possible. For more details of Séitah context see Initial Report for M2020-262-5 Salette.
The Robine and Malay samples were cored from the Issole outcrop approximately 120 meters SW and about 5 meters lower than the Séitah formation cores Salette and Coulettes acquired from the Brac outcrop. Based on the dip of what is likely igneous layering, Issole may be stratigraphically higher than Brac by a few meters. However this is less apparent in subsurface observations (see Subsurface Structure section). Issole is approximately 200 meters northwest of, and at the same elevation as, Rochette, from which the Mâaz formation cores Montdenier and Montagnac were acquired.

**Operations**

Figure 2 shows the local context of Perseverance rover operations leading up to the sampling of Robine and Malay. To reach Issole after sampling Salette and Coulettes at the locality labeled Caille in Figure 2, the rover retraced its route through a narrow traversable corridor of South Séitah, returning to an outcrop (Issole) that had previously been imaged and analyzed remotely on sol 202 during the drive to Bastide and Brac.

On Sol 286, Perseverance arrived at the Issole workspace and started the sampling sol path. On Sol 287 WATSON imaging of abrasion and sampling candidates was acquired on the Quartier, Pauls, Robine, and Soberro targets. The Quartier abrasion patch was made on Sol 292. On Sol 293 post-abrasion WATSON and SHERLOC/PIXL scans were acquired on the Quartier abrasion patch, as well as pre-coring WATSON images on Robine and Pauls, our two coring candidates. On Sol 294, we completed the post-abrasion proximity science and performed other remote sensing activities to satisfy the requirements of the STOP list. Around this time, we also decided to split the sampling sol (nominally 1 sol) into 2 sols (similar to Montdenier), based on concerns from outcrop images and the abrasion patch that the Issole rock might be susceptible to fragmentation/disaggregation. By including imaging and a ground-in-the-loop assessment, we were able to verify that core was acquired before proceeding to seal the sample tube. Also, as a precaution, on this sol we acquired WATSON images of the Malay target located on a more resistant part of the rock, with the intention that if the first sampling attempt on Robine was not successful, we would try again on a potentially more resistant part of the rock at Malay. Successful core acquisition of Robine occurred on Sol 295, and sample processing and sealing on Sol 298.

On Sol 306, we attempted sampling of Pauls in a single sol, since we now had confidence that we could acquire an intact sample at this location. Although we successfully acquired the Pauls core, a fragment of the core slipped out of the sample tube during docking with the bit carousel, and prevented mating of drill to bit carousel. On Sol 322, we dumped the Pauls sample on to the ground to aid in the fault recovery process. Recovery activities ensued until sol 332.

On Sol 337, we successfully sampled Malay, our second Issole core, using the same (now-empty) sample tube used for the Pauls sampling attempt.

On sol 340 we drove away from the Issole workspace.

**Sample-Related Observations**

**Workspace Images**
Figure 3a shows a Mastcam-Z mosaic of the Issole outcrop in its local context and Figure 3b shows the Mastcam-Z image of the workspace in which the Quartier abrasion patch and the Robine and Malay cores were acquired. All abrasion and sampling was conducted on the same block and on the same surface (Figure 4). Issole features shallowly dipping layers, generally 1–5 cm, but up to several tens of centimeters, in thickness. These layers have a rough texture with sharp, angular edges, apparent recessive weathering beneath ledges, and surficial depressions. Although Issole appears layered at the outcrop scale, no layering was observed in the abrasion patch or borehole observations.

The continuity of the several tens of meters long ridge on which Issole is located and the similarity between the dip of Issole and that of nearby outcrops suggest that Issole is in-situ bedrock. WATSON images show that the natural surface of Issole is well-exposed, although dust and wind-blown sediment have accumulated in surficial depressions and beneath overhangs (Figures 5–7). Wind polishing and fluting are present on the higher relief parts of Issole, whereas the lower relief portions of the outcrop have an abundance of small pits partly filled with sand (Figure 3b). RMI images of Issole (Figure 4) show dark gray grains, mostly around 0.1–0.5 cm, and abundant smooth brown coatings. Where not dust covered, mineral faces are visible. The surface appearance of Issole is similar to that observed on Bastide and on Brac.

Figure 3 | Local and outcrop images of Issole. a) Issole locality; location of the sampled block within the outcrop is indicated. b) Post-abrasion and coring Mastcam-Z image of the Issole outcrop and work space. The abrasion patch is 5 cm in diameter and the Robine borehole is 2.7 cm in diameter.
This block where Quartier/Robine/Pauls occurs is ~70 cm across.
Figure 4 | RMIs of the Issole outcrop. Gaussian stretch RMIs of two different targets from Issole illustrate a) the presence of coarse-grained textures with a dusty surface (Castiglione target) and b) smooth brownish coatings (Gacia target).
Figure 5 | WATSON pre-coring images of the Robine target at ~7 and 35 cm standoff distances.

Figure 6 | WATSON pre-coring images of the Malay target at ~6 and 25 cm standoff distances.

Pre- and Post-Coring/Abrasion Images

Pre- and post-abrasion WATSON and ACI images of the Quartier abrasion patch are shown in Figures 7. Figure 7A–B shows WATSON pre-abrasion images of the natural surface of Quartier, Figure 7C shows a WATSON image of the gDRT-cleared abraded patch and Figure 7D shows a greyscale ACI image superimposed onto the abraded patch. The ACI image corresponds to the locations of SHERLOC scans shown in subsequent figures. Notable observations from RMI, WATSON and ACI images include:
Figure 7 | WATSON images of the Quartier abrasion surface (sol 257). (a-b) Pre-abrasion WATSON ~7 and 35 cm standoff images showing the natural surface of Quartier. (c) WATSON image of the entire abrasion target. (d) WATSON image of the abrasion patch overlain with greyscale ACI image. Note that the WATSON images are rotated with respect to the images obtained by the remote sensing mast.

(a) The natural weathered surface has numerous large and small protuberances, grains and pebbles (between 0.1 and 1.5 cm), as well as small depressions that are filled with dust and other fine-grained, wind-blown materials. Few, if any, individual grains are visible in the WATSON pre-coring natural surface images, but note that abundant grains are visible in the RMIs of relatively dust-free surfaces.

(b) 0.2–3 mm pale brown–white sub-angular phases comprise most of the abraded surface. Other major constituents are 0.2–0.8 mm mid-brown crystals with sub-angular to angular morphologies, 1–4 mm
grey–green crystals with angular and elongate, <0.1 mm white crystals. The pale brown–white, mid-brown, gray–green and white minerals are distributed irregularly throughout the abraded patch with no obvious spatial heterogeneity in their relative abundances.

c) Some minerals within the abraded surface are surrounded by mid-brown and grey inter-crystal boundaries, potentially suggesting Fe-rich alteration phenomena. Pale reddish rims also surround or are spatially associated with many grey–green crystals.

WATSON and Mastcam-Z post-coring images of the Robine and Malay sampling sites are shown in Figures 8-9. The location of the unsuccessful Pauls sampling attempt, and the dumped core fragments, are also visible in Figure 9. Mastcam-Z and CacheCam images of the samples within the coring bit and secured within their sample tubes are shown in Figures 10-11. The Mastcam-Z images of the three Issole cores (Robine, Pauls and Malay) show clear visual similarities, confirming that Malay was an appropriate substitute for the disaggregated Pauls core.

Figure 8 | Post-coring WATSON image of the Robine borehole.
Figure 9 | Mastcam-Z images showing the Issole outcrop before (left) and after (right) drilling of the Malay core; images taken on Sol 335 and 337, respectively. Within the field of view, the Quartier abrasion patch (5 cm diameter) is visible in the lower middle of the image, the Robine borehole is to the right, the Pauls borehole is in the middle, and the Malay borehole is to the left in the right-hand image (boreholes = 2.7 cm diameter). In the lower right quadrant of the image, some fragments of the discarded Pauls core are visible between the abrasion patch and boreholes.

Figure 10 | Mastcam-Z images of Robine (left), Pauls (middle), and Malay (right) cores inside the drill bit, taken on Sols 295, 306 and 337, respectively.

Figure 11 | CacheCam images of the Robine and Malay cores. Rightmost image shows a more in-focus image of Malay.
**Elemental Geochemistry - PIXL**

Elemental geochemistry of the *Issole* rock was obtained by PIXL X-ray fluorescence (XRF) mapping on the *Quartier* abrasion patch. Two XRF rasters were acquired, on sols 292 and 300 (Figure 12). The sol 292 raster focused on a patch of white alteration material (informally called the ‘polar bear’), while the sol 300 raster emphasized silicate (igneous) mineralogy. *Issole* is inferred to be a cumulate igneous rock formed by accumulated olivine crystals with crystallized basaltic magma among the olivine grains; later alteration by one or more aqueous solutions deposited salt minerals and produced hydrous silicates and carbonates (see similar observations in the *Salette* and *Coulettes* Initial Reports).

**Figure 12 | PIXL raster scans on Quartier.** Approximate locations of PIXL XRF scans on the *Quartier* abrasion patch (~5 cm across), annotated as to scan sol. On this image, salt minerals are off-white, olivine grains are pale buff, augite is greenish-black, and poorly diffracting Fe-Mg-poor silicate material is brown. Locations overlain on WATSON image SIF_0293_0692957360_269ECM_N009000SRLC00479_0000LMJ01, rotated 90° clockwise from original orientation.
The average chemical composition for the sol 300 raster on Quartier is given in Table 1. Quartier has a silicate composition. Compared to a typical basalt (terrestrial or Martian), the Quartier sol 300 scan is low in SiO$_2$, Al$_2$O$_3$ and MgO, and rich in FeO and Na$_2$O. Note that this bulk sum represents a mixture of igneous silicate minerals (olivine, pyroxene and plagioclase) and secondary salts, carbonates, and hydrous silicates. Besides the tabulated values, no minor or trace elements were detected.

Table 1. Chemical compositions obtained using PIXL XRF.

<table>
<thead>
<tr>
<th></th>
<th>Bulk Raster 0300</th>
<th>Olivine</th>
<th>Augite</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wt %</td>
<td>N=3333</td>
<td>Err 1σ</td>
<td>N=93</td>
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<tr>
<td>Na$_2$O</td>
<td>1.87</td>
<td>0.50</td>
<td>0.53</td>
</tr>
<tr>
<td>MgO</td>
<td>12.8</td>
<td>0.6</td>
<td>17.86</td>
</tr>
<tr>
<td>Al$_2$O$_3$</td>
<td>2.36</td>
<td>0.55</td>
<td>0.92</td>
</tr>
<tr>
<td>SiO$_2$</td>
<td>40.3</td>
<td>2.0</td>
<td>35.14</td>
</tr>
<tr>
<td>P$_2$O$_5$</td>
<td>0.28</td>
<td>0.23</td>
<td>0.08</td>
</tr>
<tr>
<td>SO$_3$</td>
<td>1.66</td>
<td>0.47</td>
<td>0.68</td>
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<tr>
<td>Cl</td>
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<td>0.3</td>
<td>0.38</td>
</tr>
<tr>
<td>K$_2$O</td>
<td>0.20</td>
<td>0.19</td>
<td>0.01</td>
</tr>
<tr>
<td>CaO</td>
<td>2.94</td>
<td>0.56</td>
<td>0.61</td>
</tr>
<tr>
<td>TiO$_2$</td>
<td>0.99</td>
<td>0.32</td>
<td>0.01</td>
</tr>
<tr>
<td>Cr$_2$O$_3$</td>
<td>0.29</td>
<td>0.23</td>
<td>0.00</td>
</tr>
<tr>
<td>MnO</td>
<td>0.58</td>
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<td>0.62</td>
</tr>
<tr>
<td>FeO-T</td>
<td>25.7</td>
<td>1.3</td>
<td>41.20</td>
</tr>
<tr>
<td>Sum %</td>
<td>94.1</td>
<td>49.04</td>
<td>97.2</td>
</tr>
<tr>
<td>Mg#</td>
<td>47</td>
<td>44</td>
<td>60</td>
</tr>
</tbody>
</table>

N are number of analyses averaged. Err is uncertainty (1σ) from calibration and counting statistics. Mg# is molar Mg/(Mg+Fe), in percent.

On total alkali versus silica (TAS) plots (Figure 13), the PIXL spot analyses for Quartier are distributed widely. For the sol 292 scan, which included abundant salt minerals, points spread toward the origin. These points represent mixtures of igneous silicates with non-silicate igneous minerals (Fe-Ti-Cr oxides, Ca-phosphates) and abundant salt minerals, including Fe-Mg-Ca carbonates, sulfates, and chlorides (or oxychlorine compounds). Points on that plot with low silica and moderate alkali contents likely represent alkali-bearing sulfates, chlorides, and/or oxychlorine compounds. A few spot analyses show high SiO$_2$ (>65%) and relatively low alkali content, suggesting that the alteration assemblage includes a silica-rich phase. Other points with SiO$_2$ > 40% likely represent diverse igneous phases (described below) and mixtures thereof. On the TAS diagram for the sol 300 raster the bulk composition lies within the olivine field, consistent with the interpretation of Quartier as an olivine cumulate igneous rock. Most analysis points are consistent with mixtures of olivine, augite pyroxene and alkali-rich feldspar. Several
points extend toward the origin and represent mixtures of olivine and other mafic minerals with non-silicate minerals (Fe-Ti-Cr oxides, Ca-phosphates) and/or salt minerals.

Figure 13 | Total alkalis vs silica plots for the Quartier abrasion; data from PIXL. Individual data points represent regions of the abraded surface (~125 µm) and most are mixtures of multiple minerals. Bulk sum compositions shown by blue circles. Reference end-member igneous mineral compositions are shown as diamonds and as lines for those with solid solution series. The underlying igneous classification scheme does not apply to the individual data points and simply provides a frame of reference for comparing samples. Upper panel) Scan of sol 292 (see also Figures 15 and 17), including the ‘polar bear.’ Points with >35% SiO₂ likely represent anhydrous silicate minerals (olivine, augite, intermediate feldspar) or mixtures thereof, as well as hydrous silicate alteration materials. The points at SiO₂ <35% represent oxide and salt minerals, mostly sulfates and carbonates. Lower panel) Scan of sol 300 (see also Figures 16 and 17), focused on igneous silicate materials. Most individual points are consistent with olivine and/or augite compositions with some mixture of plagioclase and/or salts. Individual points at SiO₂ <35% represent oxide and salt minerals, mostly sulfates and carbonates.
Figure 14 shows ternary diagrams of molar compositions $\text{Al}_2\text{O}_3 - (\text{CaO+Na}_2\text{O+K}_2\text{O}) - (\text{FeOT+MgO})$ for PIXL spot analyses and scan averages. Idealized common mineral compositions are also plotted on the diagram. In this projection, the sol 292 (Figure 14a) and sol 300 (Figure 14b) plots appear very similar, with the vast majority of points near the (FeOT+MgO) apex. For the sol 292 scan, these points are rich in Fe-Mg bearing sulfates and carbonates. For the sol 300 scan, these points represent areas rich in olivine. Effectively all spot analyses for both scans fall in the triangle feldspar–augite–(FeOT+MgO), suggesting that Issole/Quartier experienced little loss of Ca+alkalis, i.e., no points are enriched in Al as would be expected through open-system weathering processes. Overall, this plot indicates that Quartier is neither heavily weathered nor altered, although it does contain a significant proportion of salt minerals and carbonate (Figure 15).

PIXL multi-elemental maps are shown in Figures 15-16. Figure 15, of the sol 292 raster, includes the sulfate-rich ‘polar bear’ area (see also Figure 12). The core of the ‘polar bear’ appears to be Na-bearing Mg-sulfate (orange in Figure 15b), it is surrounded by Mg-rich sulfate (Figure 15a). Fe-rich sulfate may be present, and Fe-Mg carbonate is inferred to be widespread outside the ‘polar bear’ because PIXL analytical totals are low. Chlorine is not commonly associated with Na, but rather with Mg (Figure 15a–b). Sodium is most commonly associated with Al in alkali feldspar (Figure 15a–b). Little Ca-bearing sulfate is apparent. These secondary materials, interpreted to be products of post-igneous aqueous alteration, occur in light-toned patches (primarily salt minerals) and in reddish brown patches (low Fe-Mg silicates) surrounding and invading anhydrous minerals, especially olivine.

Figure 16 shows multi-element maps of the sol 300 PIXL scan, which targeted an area with a small amount of salt minerals. Olivine of intermediate Fe/Mg ratio (~Fo44) is the most abundant mineral, and occurs in euhedral to elongate subhedral to euhedral shapes (to ~5x2 mm). The olivines contain multi-mineral inclusions, possibly crystallized melt inclusions, along their midlines and elsewhere. Olivine grains are partially enclosed by anhedral augite grains in a poikilitic texture similar to that observed in the Brac rock, and as seen in some igneous rocks on Earth and the chassignite Martian meteorites. Augite grains are up to ~10 mm in size. Back-reflection diffracted X-rays from augite regions indicate that they are monocrystalline and crystallographically oriented across non-contiguous areas separated by olivine in the abrasion patch. Average chemical compositions of olivine and augite are given in Table 1. Other components identified in the sol 300 raster include mesostasis material among the olivine and augite and secondary phases. The mesostasis consists of alkali feldspar (rich in Na and K, <1 mm), Cr-rich and Ti-rich Fe oxides (≤0.25 mm), and Ca-phosphates (<0.125 mm). Secondary phases include poorly diffracting (or very fine grained) silicate(s) with low Fe and Mg, and salt minerals as described above.
Figure 14 | Mineralogical interpretation for the Quartier abrasion; data from PIXL. Data plotted by point and as bulk sum composition (blue circle, Table 2) on ternary diagrams of molar abundances of Al₂O₃–(CaO+Na₂O+K₂O)–(FeO+MgO). Common pure mineral compositions are shown (cpx = Ca-poor pyroxene, i.e orthopyroxene; cpx = calcic pyroxene, i.e. augite). Most PIXL analysis points are consistent with pure minerals (olivine, cpx) or binary mixtures of olivine–cpx or olivine–feldspar, or ternary mixtures of olivine–cpx–feldspar. Aluminous alteration minerals, including common clay minerals, fall above the upper red dashed line. Several salt minerals (e.g., Fe-Mg-Ca-sulfates/carbonates, halite) are also plotted. a) Sol 292. In this projection, the Mg-bearing sulfate minerals of the ‘polar bear’ plot near the ‘FeOT+MgO’ corner of the triangle b) Sol 300. Most of the points are consistent (in this projection) with mixtures of olivine and cpx.
Figure 15 | PIXL X-ray multi-element maps for the Quartier abrasion patch, sol 292. Raster area is 7x5 mm. a) Red=\text{MgO}, Green=\text{SO}_3, Blue=\text{Cl}. b) Red=\text{SO}_3, Green=\text{Na}_2\text{O}, Blue=\text{Cl}. c) Red=\text{MgO}, Green=\text{Al}_2\text{O}_3, Blue=\text{CaO}. d) Red=\text{FeO}_x, Green=\text{TiO}_2, Blue=\text{Cr}_2\text{O}_3. The ‘polar bear’ comprises mostly sulfate minerals (panels a, b) with little Ca (panel c); its rind is rich in Mg (orange in panel a), and its core contains Mg (panel c) and Na (orange, panel b), but little Ca (panel c). Chlorine-rich spots are scattered (panels a, b); most are not associated with Na (panel b). Most of the rock outside the ‘polar bear’ is olivine (red, panel c) with interstitial augite (purple, panel c) and feldspar (green, panel c). This area contains few oxide minerals: a grain of Fe-Ti-Cr oxide to the upper left, and a Cr-rich grain to the lower right.
Figure 16 | PIXL X-ray multi-element maps for the Quartier abrasion patch, sol 300. Raster area is 12.5x4 mm. a) Red=MgO, Green=Al₂O₃, Blue=CaO. b) Red=CaO, Green=Na₂O, Blue=K₂O. c) Red=CaO, Green= P₂O₅, Blue=Cl. d) Red=FeO₇, Green=TiO₂, Blue=Cr₂O₃. Olivine is the most abundant material (red in panel a), with interstitial augite (purple in panel a) and plagioclase (green in panel a). Only one feldspar is present; panel b shows only one blue-green color correlated with the Al in panel a. A few salt grains (green in panel b) are present, as is a little Mg-Fe sulfate (blue in panel c), and a grain of phosphate (yellow in panel c). Panel d shows Fe oxide minerals: Ti-rich oxides are teal-green, and Cr-rich oxides are purple.
PIXL Micro-Context Camera (MMC) images of the Quartier abrasion patch (Figure 17) show more of the abrasion area than the XRF scans and may be used to extrapolate phase proportions from those of the XRF scan areas to nearly the whole abrasion patch. The MMC image of NIR-G-B (Figure 17b) is analogous to WATSON visible images, and the G/NIR ratio image shows a clear distinction among the olivine, augite, and mesostasis phases (Figure 17c). These visible and qualitative relationships are quantified in a graph of G versus NIR reflectance (Figure 17d), which allows extraction of the proportions of these phases over the whole MMC image area.

**Figure 17 | PIXL MCC (Micro-context camera) data for the Quartiers abrasion.** a) MMC grayscale image, a portion of the Quartiers abrasion. b) MMC false colour of abrasion area; Red = MMC Near-IR; Green = MMC Green; Blue = MMC Blue. Location of PIXL XRF scans shown. c) MMC ratio image Near-IR/Green reflectances. Augite is darkest; olivine is next, and mesostasis and alteration materials are lightest tone. Location of PIXL XRF scan is shown. d) MMC reflectance ratios.
**Mineralogy and Organics-SHERLOC**

SHERLOC Raman and florescence scans were obtained on sols 293 (survey and HDR scans; Figure 18) and 304 (detail scan; Figure 19). In the sol 293 and 304 scans, several mineral identifications were made (Table 2). Spectral features were assigned to carbonate (~1090 cm\(^{-1}\)), olivine (~840 cm\(^{-1}\)) and Mg/Ca/Fe-sulfate (~1050 cm\(^{-1}\)), the latter corresponding to the white material observed in the ACI image (Figure 18A). Possible hydration peaks between 3100 and 3600 cm\(^{-1}\) were also observed in some spectra. Fluorescence features at 303 nm, 325 nm, and 330–340 nm are consistent with single- and double-ring aromatic organic molecules. The Raman peak at ~1664 cm\(^{-1}\) is consistent with the presence of aromatic organic carbon and is exclusively found in association with fluorescence features at 303 and 330 nm. This signal is present in several spots associated with hydrated sulfate, but is not uniformly found throughout the sulfate detected in the scan regions. The only significant difference between the spectra obtained on sol 293 and 304 is in the hydration region between 3100 and 3600 cm\(^{-1}\). Differences in spectral structure might be caused by a change in hydration state during the eleven sols between analyses, during which the abrasion patch was exposed to the martian atmosphere potentially causing mineral dehydration.

**Table 2. SHERLOC mineral identifications in Quartier.**

<table>
<thead>
<tr>
<th>Sample</th>
<th>Certain</th>
<th>Possible (not confirmed)</th>
<th>We looked for these, but cannot find them</th>
</tr>
</thead>
<tbody>
<tr>
<td>Quartier</td>
<td>Sulfate (most likely Mg, Ca and/or Fe), virtually always hydrated (occurs in white void fills or crystals, secondary to primary lithology)</td>
<td>Olivine (low Fo#) occurs associated with one darker grain.</td>
<td>Pyroxene</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Carbonate (major phase, could be Mg/Ca/Fe)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Amorphous/microcrystalline silicate</td>
<td></td>
</tr>
</tbody>
</table>
Figure 17 | SHERLOC fluorescence and Raman spectral results for the Quartier abrasion surface. Spectra were collected with a survey scan (5x5 mm, 1296 points, 144 µm spacing, 15 pulses per point, not shown) and a HDR scan (7x7 mm, 100 points, 780 µm spacing, 500 pulses per point) on sol 293. A) Colorized ACI image of the abraded patch in which the blue rectangle indicates the HDR scan area from sol 293, and the white circles indicate regions of interest (ROIs) that correlate with fluorescence data shown in B and C. B) HDR scan maps show the possible presence of organics. Upper panel is an RGB map showing the main regions of organic fluorescence (303–330–340 nm). Both doublet peaks at 303 and 325 nm (ROI1) and the 330–340 nm peak (ROI 2) are localized to the ROIs in panel A. Lower panel shows Raman features at 1664 cm⁻¹ that indicate the presence of organic materials (see Raman data in D). C) Fluorescence spectra. Upper panel shows the average fluorescence spectrum for the whole analysis area; lower panel shows spectra from selected fluorescence ROIs (see panel A). D) Map showing the results of the Raman HDR scan. The colored circles correspond to mineral and organic detections. The right panel includes selected Raman spectra from the HDR scan, offset on the y-axis for clarity. The uppermost spectrum corresponds to Pt. 34 in the Raman map and is representative of spectra assigned to Mg/Ca/Fe-sulfate (~1050 cm⁻¹), which corresponds to the white material in the ACI image. The second spectrum is also from Pt. 34, but the vertical scale has been exaggerated by 2000x to show the presence of peaks at 1664 and ~1400 cm⁻¹ that are assigned to organic carbon. These peaks were also observed in several points in the sol 304 SHERLOC map (see Figure 18). The third spectrum is an average of all spectra marked with light blue circles on the map in D and is assigned to carbonate (~1090 cm⁻¹). The bottom spectrum is an average of those from Pts. 65 and 66 and is assigned to olivine (~840 cm⁻¹). The spectrum from Pt. 34 (and all spectra marked with purple dots on the map) also contains possible hydration peaks between 3100 and 3600 cm⁻¹. The scale in panel A also applies to D.
**Figure 18 | SHERLOC Raman spectral results for Quartier abrasion surface.** Spectra shown were collected with the HDR scan described in Figure 17 (7x7 mm, 100 points, 780 µm spacing, 500 pulses per point) on sol 293, and a detail scan (1x1 mm, 100 points, 144 µm spacing, 500 pulses per point) on sol 304. A) Colorized ACI image of the abraded patch where the blue box indicates the HDR scan area from sol 293 and the red box indicates the detail scan from sol 304. B) Sol 304 ACI image of the scanned region indicating the locations of point 34 from the 293 detail scan (blue circle) and point 45 from the sol 304 detail scan #1 (black circle). Note the overlap of analysis spots. C) Raman spectra from point 34 (sol 293 HDR scan; blue) and point 45 (sol 304 detail scan #1; black). Note the high degree of similarity between these two spectra, including the peaks at 1664 and ~1400 cm⁻¹, confirming the identification of organic carbon. The only significant difference between the two spectra is in the hydration region between 3100 and 3600 cm⁻¹. This difference might be caused by a change in hydration state during the eleven sols between analyses.

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**Elemental Geochemistry and Mineralogy - SuperCam**

SCAM LIBS data are shown plotted on a total alkalis vs silica diagram in Figure 19. Points are scattered within the picrobasalt and basalt fields, and to a lesser extent within the olivine and basaltic andesite fields. The SCAM LIBS analyses targeted primarily natural surfaces in Séítah, with only a small subset targeting the abraded patch. Olivine is the dominant detection and bulk sum average in the PIXL data obtained on the abraded patch. SCAM LIBS data differ but are consistent with PIXL data in that both contain few points with SiO₂ > 50 wt%.

Harker oxide diagrams of SCAM measurements of Issole are plotted against a background of all Séítah points in Figure 20. These diagrams show that Issole is generally representative of the wider Séítah region with potentially lower contents of CaO and MgO. Additionally, relatively few points in the borehole and abrasion tailings have SiO₂ values below 40 wt%.

7-23
Figure 19 | TAS plot of SCAM LIBS data on Séítah rocks. Average value of all analyses is shown as a blue circle. Individual Séítah analyses, i.e., including those not from the Issole outcrop, are plotted as grey points. The LIBS-derived composition of the natural surfaces indicates the presence of grains on a pyroxene to olivine gradient, hydration and distinct coatings enriched in Al$_2$O$_3$, TiO$_2$ and Na$_2$O+K$_2$O compared to the surrounding rock.

Note that the ~250 µm SCAM LIBS spots most commonly interrogate more than one specific mineral. Given that the typical Quartier grain size is larger than this size, as described above, many of the trends observed in the diagrams shown in Figures 19 and 20 can be reasonably attributed to mixing among several mafic minerals, e.g. olivine and pyroxene.

Figure 21 shows the mean VISIR spectrum of the natural surface of Issole, the Quartier abrasion patch, and the tailings of the Quartier abrasion and Robine tailings acquired between sols 294 and 303. The broad slope between 2.2 and 2.5 µm in all VISIR data is consistent with the presence of hydrated minerals. The broad absorption between 0.8 and 1.8 µm indicates olivine, consistent with PIXL and SHERLOC data on the abraded surface described above. The olivine signal is stronger in the abraded surface than on the natural surface. The absorption near 2.3 µm is consistent with carbonates and/or Fe/Mg-phyllosilicates. The broader single absorption (as in Quartier sol 297) is more consistent with carbonate, whereas the narrower 2.28–2.29 and 2.32 µm absorptions (as in Quartier sol 303) are more consistent with Fe-phyllosilicate and Mg-phyllosilicates, respectively.
Figure 20 | Harker oxide diagrams of SCAM data from Séítah. Data relating to the Issole outcrop are shown in color. All Séítah points, i.e., including those not from the Issole outcrop, are shown in black.
Figure 21 | Mean SCAM VISIR spectra of the unabraded surface of Issole (targets Sagnes, Calern, Aurent, Castiglione and Gacia), the abraded patch Quartier and its tailings and tailings from the Robine borehole. Note that owing to temperature sensitivities in the IRS instrument, the calibration is uncertain for wavelengths longer than ~2.5 µm.

Subsurface Structure

Subsurface structure in the vicinity of the Issole outcrop is shown in Figure 22. Three radargrams were acquired during sols 201–202 while traversing from the Artuby ridge eastward into Séität (Figure 22b), on sols 285 and beyond (Figure 22c), and sols 280–281 (Figure 22d). Issole is an outcrop expression of a series of SW dipping high-amplitude reflectors shown in the sol 201 radargram and labeled Issole member (Mb). These dips clearly project the Issole Mb beneath the Máaz formation layers on Artuby ridge. A magnification of reflector geometries in the subsurface extension of the Issole outcrop (Figure 23) suggests an intimate relationship between SW-dipping and antithetic NE-dipping reflectors as well as crosscutting horizontal surfaces. Although these reflector geometries are consistent with a clastic
depositional origin for *Issole* (e.g., water-lain sediment, pyroclastics, or impactoclastics), compelling observations elsewhere in this document indicate *Seitha* has a cumulate igneous origin.

Further east toward the *Brac* outcrop (*Salette* and *Coulettes* sampling site) the reflectors become less apparent. One possible interpretation of the radargram with the *Bastide* Mb (*Brac* outcrop) capping the *Issole* Mb above an unconformity is shown in the middle and lower panels. In contrast to this interpretation, projection of dip angles from both outcrop and the radargram near *Issole* would indicate that *Issole* lies a few meters above the *Bastide* Mb. If our currently preferred explanation that *Brac* and *Issole* are layers of a single layered igneous body is correct, this distinction is not of great significance.

**Figure 22 | Geological context and RIMFAX-derived subsurface stratigraphy from *Issole* to *Brac*.** a) Location of *Issole* and *Brac* together with selected RIMFAX traverses used. b) Radargram for sols 201 to 202 (yellow traverse in panel a) showing dipping high-amplitude reflectors. The black box denotes the subsurface extension of *Issole* shown in detail in **Figure 23**. c) Radargram along sol 285 and onwards traversing the *Issole* outcrop (blue traverse in panel a) showing the boundary between the *Bastide* and *Issole* members. d) Radargram along sols 280 and 281 (green traverse in panel a) showing the *Brac* outcrop and a possible erosional unconformity at the top of the *Issole* member and base of the *Bastide* member extending SE into the region shown in panel c.
**Figure 23** | Subsurface extension of *Issole* enlarged. a) Subsurface extension of *Issole* outcrop in sol 201 radargram (location given in Figure 22). b) Line tracing of reflector geometries in panel a showing intimate relationship between SW-dipping and antithetic NE-dipping reflectors, as well as crosscutting horizontal surfaces.

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**Core orientation**

At the time of drilling and 5.9 cm pre-drilling WATSON imaging, the rover, coring drill, WATSON, and coring target (“Robine_coring.PSC.0.0341”), had the following characteristics:

1. Rover orientation quaternion just after drilling but before unloading the stabilizers (transferring from RMECH to SITE frame):  
   \[
   bQ_{ll1} = (0.4283390, 0.0740838, 0.0263520, -0.9001910);
   \]

2. Coring Drill orientation quaternion just after drilling but before unloading the stabilizers (transferring from CORING DRILL to RMECH frame):  
   \[
   cdQ_{b} = (0.6646180, 0.1631090, -0.7068370, 0.1790540);
   \]

3. Rover orientation quaternion at time of acquisition of WATSON image SIF_0293_0692957683_398FDR_N0090000SRLC00643_0000LMJ02: (transferring from RMECH to SITE frame):  
   \[
   bQ_{ll2} = (0.428458, 0.0711518, 0.0249948, -0.900409);
   \]

4. WATSON orientation quaternion at time of acquisition of WATSON image SIF_0293_0692957683_398FDR_N0090000SRLC00643_0000LMJ02: (transferring from WATSON to RMECH frame):  
   \[
   wQ_{b} = (0.666062, 0.167862, -0.704872, 0.177028).
   \]

Items 1 and 2 give a coring drill pointing vector estimate of

- hade = 5.36°
- azimuth = 225.58°

Items 3 and 4 give an estimate of the angle between the WATSON y-axis and the up-dip direction in SITE coordinates (Figure 24) of

- core roll, \( \alpha = 231.60° \)
**Preliminary Scientific Assessment**

**Synthetic sample description and preliminary interpretation**

Sample type: *Olivine cumulate*

1. **Relationship with surrounding rocks**
   a. *Robine* and *Malay* were obtained from the *Issole* outcrop, which has a stratigraphic position close to, but below, the boundary/contact with the *Máaz* formation.
   b. Based on its occurrence as part of a NE–SW-trending ridge, concordance with surrounding outcrops, and estimated dip consistent with that of the wider *Séítah* region, *Issole* is confidently interpreted to be an in-place portion of the *Séítah* bedrock.

2. **Texture and fabric**
   a. Grain size: <0.1–4 mm
   b. Rock fabric (natural surface): large olivine crystals exposed on relatively fresh surfaces, but few definitive observations of grains on natural weathered surfaces.
c. Rock fabric (abrasion patch): some textural heterogeneity, main constituents include <0.1–3 mm pale brown–white sub-angular phases, 0.2–0.8 mm mid-brown sub-angular to angular crystals, 1–4 mm grey–green angular and elongate crystals, <0.1 mm white crystals. Secondary minerals (including a large region of sulfates) are particularly concentrated within the central region of the abrasion patch.

3. **Mineralogy and chemistry**
   a. Minerals present: olivine, pyroxene (augite), Ca/Mg-sulfate, carbonate, Na-rich and K-rich feldspars, Fe-Cr-Ti magnetite, Ca-phosphates; also, aromatic organic materials sometimes associated with secondary mineral phases. Mesostasis consists of Na-rich alkali feldspar, Cr-rich and Ti-rich Fe oxides, and Ca-phosphates. Secondary phases include poorly diffracting (or very fine-grained) silicate(s) with low Fe and Mg, carbonate, and salt minerals (sulfates, perchlorates).
   b. The bulk composition is consistent with a mafic/ultramafic rock modified by limited aqueous alteration. Point analyses by SCAM and PIXL plot mostly within the olivine, picrobasalt and basalt regions of the TAS diagram.

4. **Alteration/secondary characteristics**
   a. Interactions with aqueous fluids, alteration, and mineral hydration: evident, but limited.
   b. Secondary mineralogy: see point 3a above.

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**Returned Sample Science Considerations**

Observations of olivine surrounded by clinopyroxene in a poikilitic texture within the Dourbes and Quartier abrasion patches, along with remote observations of the mineralogy, texture, and spectral characteristics of Brac, Issole and other Séítah outcrops, suggest that the Robine and Malay samples (as well as the Salette and Coulettes samples), and much or all of Séítah, are best interpreted as an igneous olivine cumulate. As such, the return of these samples is expected to address a number of RSS themes of significance to the Mars 2020 mission:

*Geochronology:* The timing of volcanic rock crystallization can be quantified using laboratory-based, radiometric dating of returned samples. Likewise, assuming cosmogenic nuclide production rates in specific mineral on Mars can be established, quantitative constraints on the sample’s surface exposure age and/or erosion rate may also be obtained. Quantifying the timing of crystallization of Brac and Issole will also constrain the emplacement ages of other crater floor units, given their stratigraphic relationships (e.g. Séítah is likely older than Cf-fr from which the Montdenier and Montagnac samples were collected).

*Paleomagnetism:* The igneous lithologies of the Salette/Couletes/Robine/Malay samples may enable absolute measurements of the paleointensity of the ancient Martian magnetic field at the time of crystallization. The orientations of the samples, combined with paleohorizontal indicators from outcrop foliation, may enable measurements of the absolute paleodirection of the ancient field. Paleomagnetic investigations of secondary ferromagnetic minerals if present could constrain the field at the time of aqueous alteration. Combined with measurements from other samples, these data could test the hypothesis that Martian atmospheric loss was driven by the cessation of an early dynamo, constrain the dynamo’s secular variation, and determine if the dynamo exhibited polarity reversals. Paleomagnetic
analyses could also test whether Séitah rocks have experienced regional or global-scale tectonic tilting and displacements. Comparison of paleomagnetic directions with Máaz rocks (Montdenier/Montagnac and Atsah/Hahonih) could distinguish between the hypotheses that Máaz and Séitah form a single layered intrusion versus Máaz being an overlying lava flow deposited on Séitah.

Geochemistry: With a hypothesized origin as an olivine cumulate, understanding the petrogenesis of the Salette and Coulettes samples will place constraints on the nature of igneous processes on early Mars. These constraints can be applied to the origins of the extensive “Cf-f-1” unit that has been correlated with a regional olivine-bearing unit exposed between Syrtis Major and Nili Fossae. The diverse mineral assemblages of the obtained samples will be valuable for quantifying the conditions and timing of primary, high-temperature crystallization of igneous phases and later-stage alteration events. The bulk chemistry of the rock, as well as trace element and stable and radiogenic isotope geochemistry of the rock and its constituent igneous minerals, will enable comparisons with similar lithologies in the Martian meteorite suite, and insights into magmatogenesis and mantle/crustal evolution more generally. The large grains observed in Brac may contain melt inclusions that are of sufficient size to track the petrogenesis of this material.

Water–rock interactions and habitability: Many hypotheses for the origins of olivine-carbonate rocks involve aqueous alteration of igneous rocks. The interaction of aqueous media with (ultra)mafic rocks produces secondary mineral phases that can be indicative of the temperature, redox state, and fluid chemistry. High resolution mineral identification and stable isotope measurements can place constraints on such aspects of the alteration history and environment and provide insights into the presence, distribution, and nature of water on Mars. Water/rock interactions could have occurred as result of groundwater, possibly associated with Jezero lake, moving through these rocks and may have led to the establishment of microniches with water, energy, pH, nutrients and transition metals essential to life. Although likely a variable component in the history of these samples, the team has noted that low-temperature hydrothermal, serpentinizing and carbonation reactions may have sustained microbial life, particularly chemoautotrophs/chemolithotrophs that have been suggested as likely inhabitants of Martian biotopes. Although the fossil record of igneous rocks is highly variable and generally poor, the biosignature preservation potential of certain alteration phases associated with the Brac outcrop is higher, and it is possible that organic materials and/or biosignatures have been preserved within these phases.

The two pairs of cores from the Séitah formation (Brac: Salette and Coulettes, and Issole: Robine and Malay) are interpreted to be samples of a single olivine cumulate body. However, there are notable differences between them. Compared to Dourbes (the abrasion patch for the Brac samples), Quartier (Issole samples) has more Fe and Ca, and less Mg. This appears to arise from a higher ratio of augite to olivine in Quartier. In addition, Quartier has a lower abundance of Raman detected igneous phases and a higher abundance of alteration products (especially hydrated sulfates and carbonate) compared with Brac. One possible interpretation of these differences is that relative to Brac, Issole is a less-mafic layer of the Séitah igneous body that has experienced a greater degree of aqueous alteration.
References


Sample Designation: M2020-337-8 Malay

Date of Coring: 30-January-2022

Mars Time of Sample Core Sealing: 19:25 LMST, Sol 337, Ls 166.6

Latitude (N), Longitude (E), Elevation: 18.43264°, 77.44133, –2574.3 m

Campaign: Crater Floor

Region of Interest: Séítah South

Lithology: Medium- to coarse-grained poikilitic olivine cumulate rock. Primary minerals are olivine and pyroxene. Limited secondary aqueous alteration, including carbonate, hydrated sulfate, and other hydrated phases. Organic material is present, at least partially associated with sulfate phase(s).

Estimated Volume Recovered: 4.3 cm³

Coring Bit Number: 2

Core Orientation: hade = 2.72°; azimuth = 77.64°; core roll = 274.92°

Sample Serial Numbers: Tube SN261; Seal SN053; Ferrule SN062

ACA Temperature at Time of Sealing: 40°C

Estimated Rover-Ambient Pressure and Temperature at Time of Sealing: 642 Pa, 223 K

Estimated Amount of Martian Atmosphere Headspace Gas: $2.7 \times 10^{-6}$ mol

Anomalous Behavior: None

Abrasion Patch Name and Depth: Quartier, 8 mm

January 27, 2022

Summary Description

This sample is paired with M2020-298-7 Robine. Only the core orientation for Malay is described below. See the Initial Report for 2020-298-7 Robine for sample details.

Core orientation

At the time of drilling and 6 cm pre-drilling WATSON imaging, the rover, coring drill, WATSON, and coring target (“Malay_337_snc_coring”), had the following characteristics:

1. Rover orientation quaternion just after drilling but before unloading the stabilizers (transferring from RMECH to SITE frame): \( \mathbf{bQll1} = (0.9452490, 0.0016222, 0.0085824, 0.3262330) \);
2. Coring Drill orientation quaternion just after drilling but before unloading the stabilizers (transferring from CORING DRILL to RMECH frame): \( \mathbf{cdQb} = (0.6986380, 0.1504020, -0.6775800, 0.1736920) \);
3. Rover orientation quaternion at time of acquisition of WATSON image SIF_0337_0696859111_167FDR_N0090276SRLC00636_0000LMJ01: (transferring from RMECH to SITE frame): \( \mathbf{bQll2} = (0.945265, 0.00189065, 0.00584646, 0.326245) \);
4. WATSON orientation quaternion at time of acquisition of WATSON image SIF_0337_0696859111_167FDR_N0090276SRLC00636_0000LMJ01: (transferring from WATSON to RMECH frame): \( \mathbf{wQb} = (0.700578, 0.153594, -0.675732, 0.17025) \).

Items 1 and 2 give a coring drill pointing vector estimate of

- \( \text{hade} = 2.72^\circ \)
- \( \text{azimuth} 77.64^\circ \)

Items 3 and 4 give an estimate of the angle between the WATSON y-axis and the up-dip direction in SITE coordinates (Figure 1) of

- \( \text{core roll, } \alpha = 274.92^\circ \)
Figure 1. 6-cm standoff WATSON image of Malay core target on sol 337. Image scale is 28.7 μm pixel⁻¹. Orientation compass gives WATSON frame (wx, wy, wz). Core roll is clockwise angle of core y-axis, cy, from wy, given by α = 274.92°. Projection of Martian geographic north onto WATSON image plane is noted. WATSON image SIF_0337_0696859111_167FDR_N0090276SRLC00636_0000LMJ01.
INITIAL REPORT

M2020-371-9 Hahonih

Sample Designation: M2020-371-9 Hahonih

Date of Coring: 6-Mar-2022

Mars Time of Sample Core Sealing: 19:21:57 LMST, Sol 371, Ls 185.2

Latitude (N), Longitude (E), Elevation: 18.44386406, 77.45242176, -2568.357 m

Campaign: Crater Floor

Region of Interest: Ch’al member of Máaz formation, east of the Octavia E. Butler landing site

Lithology: Likely basaltic andesite. Few mm-size white feldspar laths and pyroxene grains are present, as possibly are olivine and quartz. Limited evidence of secondary alteration includes interstitial iron oxides, carbonate, possible phyllosilicates and chlorite, and hydration. Low concentrations of organic matter were detected.

Estimated Volume Recovered: 9.24 cm³

Coring Bit Number: 3

Core Orientation: hade = 21.41°; azimuth = 73.18°; core roll = 126.60°

Sample Serial Numbers: Tube SN262, Seal SN172, Ferrule SN129

ACA Temperature at Time of Sealing: 40°C

Estimated Rover-Ambient Pressure and Temperature at Time of Sealing: 672 Pa, 227 K

Estimated Amount of Martian Atmosphere Headspace Gas: 0.98x10⁻⁶ mol

Anomalous Behavior: None

Abrasion Patch Name and Depth: Alfalfa, 9 mm

May 8, 2022

Summary Description

The Hahonih and Atsah sample cores were collected about 100 m east of the Perseverance landing site, as the seventh and eighth rock samples of the mission and the last pair acquired during the Crater Floor Campaign. The Hahonih and Atsah samples were cored from the top of a boulder named Sid, associated with an abrasion patch named Alfalfa on the side of the Sid rock. The Hahonih and Atsah samples were selected to represent the Ch’al member of the Máaz formation, the stratigraphically highest Máaz member, expressed as massive, blocky rocks that appear to cover much of the Jezero crater floor as seen from orbit.

Figure 1 shows the local context of Perseverance rover operations leading up to the sampling of Hahonih and Atsah. Perseverance arrived in front of Sid on sol 363, abraded the Alfalfa target on Sol 367, and collected samples Hahonih (Sol 371) and Atsah (Sol 374/377; three-sol delay in sample processing due to arm fault between coring and sealing). Perseverance also conducted STOP list and opportunistic science observations on Sid and other nearby targets (Fig. 1).

Images of the Alfalfa abrasion patch and associated chemistry and mineralogy data reveal interlocking white feldspar laths, pyroxene, possibly olivine, and possibly quartz. Limited evidence of secondary alteration includes interstitial iron oxides, carbonate, possible phyllosilicates and chlorite, and hydration. Low concentrations of organic matter are present in the abrasion patch. Drill parameters during abrasion show that Sid is the hardest rock so far abraded by Perseverance. The Hahonih and Atsah cores are believed to be the least altered samples acquired from the Jezero crater floor.

Since the Hahonih and Atsah samples are the most compellingly in-place cores acquired from the crater-retaining Máaz formation, radiometric dates of these samples could be compared to crater-density based age models. Radiometric dates could also place constraints on the ages of the apparently overlying delta and the apparently underlying Séitah formation. The Hahonih and Atsah cores are oriented, so the magnetic field present at the time of crystallization could potentially be characterized. As the Hahonih and Atsah samples are the least altered samples acquired from the Máaz formation, they could be especially useful in characterizing martian igneous petrology and the geochemical evolution of the mantle or crust. Cosmogenic nuclide measurements could constrain Máaz formation erosion rates, as well as inform expectations of the impact of galactic cosmic radiation on the preservation and modification of organic molecules in samples from Mars.
**Figure 1** | Local context for the *Hahonih* and *Atsah* sample collection campaign and associated observations. The green star in the overhead view denotes the location of the rover during sampling, slightly east of the *Sid* rock. *Sid* (approx. 40 cm across) is shown in the lower left panel.
Figure 2 | Regional context of Perseverance operations. (a) Geologic map of western Jezero Crater (Stack et al., 2020) over CTX basemap; (b) Zoom of the rectangle in (a) shows the Perseverance Crater Floor campaign area with Máaz fm sample locations; dark green is mapped as Cf-F-1 (now Séítah fm) and tan is mapped as Cf-fr unit (now Máaz fm). Hahonih and Atsah were chosen to represent the Ch’al member, the extensively-cratered, upper unit that extends east of the landing site. (c) In situ geologic map (Crumpler and others, 2022) showing locations where polygonally-jointed Nataani type and massive Ch’al type rocks occur near the Perseverance landing site. The labeled Sid rock is the source of the Hahonih and Atsah sample cores.
Stratigraphic and Geologic Context

_Hahonih_ and its paired core _Atsah_ were collected from the _Sid_ outcrop, a representative of the _Ch’al_ member (mb), the uppermost unit of the _Máaz_ formation (fm). This formation was mapped as crater floor fractured rough (Cf-fr) in Stack et al. (2020) (Fig. 2). The _Máaz_ fm has also been referred to as the volcanic floor (Schon et al., 2012; Goudge et al., 2015), the dark-toned floor, and the mafic floor. Orbital detection of olivine and pyroxene in the unit, and the unit’s lobate margins and embayment relationships with the basin perimeter and other exposed floor units, have been offered as evidence of a volcanic origin (Goudge et al., 2015; Horgan et al., 2020a; Schon et al., 2012; Shahrzad et al., 2019). Upon landing, the prevalence of loose rocks and boulders with mafic mineralogy and a “pitted” appearance interpreted by some members of the science team as vesicles, provided potential evidence that lava flows were present in the vicinity of the landing site. However, alternative hypotheses, including a volcaniclastic and/or sedimentary origin, were also posited based on the proximity to the Jezero delta, history of water filling the crater, thermal inertia of the crater floor unit, and the apparent absence of volcanic landforms that could have sourced lava flows (Stack et al., 2020).

The _Ch’al_ member of the _Máaz_ formation is likely the stratigraphically highest member of the formation, lying above the _Nataani_, _Rochette_, _Artuby_, and _Roubion_ members. _Ch’al_ member outcrops like _Sid_ consist of massive rocks that form mounds and ridges east of the Octavia E. Butler landing site and south of the rover’s traverse to _Séítah_. The relatively large size of _Sid_ and other similar rocks in the area, and their direct contact with the underlying, polygonally jointed, flat rocks of the _Nataani_ member, suggest that _Sid_ is derived directly from the receding _Ch’al_ member and is unlikely to have been moved or rotated significantly after deposition. In many places, dark-toned, ventifacted “nubs” project upward from exposed _Nataani_ member outcrops and may be examples of _Ch’al_ rocks weathered in place (Fig. 3).

Earlier in the mission, we attempted to collect a sample of one of the low-lying, polygonally jointed members of the _Máaz_ fm ( _Roubion_ member), but that attempt failed when the rock disaggregated such that no core was retained in the sample tube (see _Roubion_ IR). Subsequent collection of the _Montdenier_ and _Montagnac_ cores from the _Rochette_ member of _Máaz_ was successful. Rover observations indicate that the _Séítah_ formation lies below the _Máaz_ formation, suggesting that _Séítah_ is older. Thus the four _Séítah_ formation cores ( _Salette_, _Coulottes_, _Robine_, and _Malay_ ) are likely older than all of the _Máaz_ cores. However, if _Séítah_ is a younger unit that intruded the _Máaz_ formation, the relative age relationship of the samples would be inverted. The crater floor units - _Máaz_ and _Séítah_ - are interpreted to be stratigraphically below the delta sedimentary rocks, but this interpretation will need to be tested as Perseverance explores the crater floor - delta contact.

Although the lithology and stratigraphy of the crater floor were unknown before landing, the Perseverance team had intended to sample the areally-extensive Cf-fr unit because of the high science value of radiometric ages for constraining the timing of many crater floor units, and for correlation with crater density to provide a comparison with crater-based chronologies (e.g., Beaty et al., 2019; Cohen et al., 2021; Werner, 2019).
**Operations**

Because the *Montagnac* and *Montdenier* samples were cored from a rock (*Rochette*) that may be slightly displaced from its original orientation, the Perseverance team had a strong desire to collect a sample of an in-place rock with unambiguous identification as the uppermost member of the *Máaz* fm prior to leaving the crater floor and climbing up the delta. Several locations were considered for sampling as the rover traversed north on the last leg of the Crater Floor Campaign, including examples of the “nubby” outcrops and topographic highs covered with *Ch’al* - type rocks that were inaccessible to the rover. The team ultimately chose to target an easily accessible *Ch’al*-type block close to the rover’s previous traverse. The team prioritized candidates based on their morphologic and topographic similarities to the *Ch’al* member, and evidence that the rock was in place.

**Figure 3 | Sid outcrop images.** a) Mastcam-Z merged sol 363-369 enhanced color mosaic of Sid and surroundings showing the darker, “bluer” nature of the blocky *Ch’al* type rocks overlying the paler, tan, polygonally-jointed rocks of the *Nataani* member. b) Sol 367 Front Hazcam image of the workspace; the *Sid* block (with the *Alfalfa* abrasion patch) is ~40 cm across. *Sid* rests on a low mound that appears to be composed of finely-layered material (in shadow) that may represent the *Nataani* member. Images: zcam08394/08386/08401 L0, FLF_0367_0699524043_237RAS _N0110108FHAZ02008_0A00LLJ01.IMG.
As the rover passed southeast of the Octavia E. Butler landing site, the team identified several blocks of Ch’al-type material in direct contact with underlying Nataani rocks, including Sid (Fig. 3). The relatively large size of the Sid rock and others in the area, and their direct contact with the Nataani member, suggest that Sid is derived directly from the receding Ch’al member and likely was not significantly moved or rotated after deposition. Table 1 provides details of the execution of STOP list observations made on Sid. Hahonih was cored on Sol 371, Atsah on Sol 374.

Table 1 | STOP list observations for the Hahonih and Atsah samples.

<table>
<thead>
<tr>
<th>Instrument</th>
<th>Rationale</th>
<th>Target</th>
<th>Uplink Sol</th>
</tr>
</thead>
<tbody>
<tr>
<td>ZCAM</td>
<td>Stereo imaging of the workspace, including abrasion and coring targets</td>
<td>Sid</td>
<td>362</td>
</tr>
<tr>
<td>WATSON</td>
<td>Pre-and post-abrasion imaging for rock texture documentation</td>
<td>Alfalfa pre-abrasion, Alfalfa post-abrasion</td>
<td>366, 370</td>
</tr>
<tr>
<td>SHERLOC</td>
<td>Organic/mineralogical map on abraded patch. SHERLOC scan co-registered with PIXL</td>
<td>Alfalfa</td>
<td>370</td>
</tr>
<tr>
<td>PIXL</td>
<td>Overnight elemental chemistry map on abraded patch. PIXL footprint co-registered with WATSON and SHERLOC - for lithology, chemistry, and mineralogy</td>
<td>Alfalfa</td>
<td>369</td>
</tr>
<tr>
<td>ZCAM</td>
<td>Multispectral observation of abraded patch for mineralogy and to support SCAM observations</td>
<td>Alfalfa</td>
<td>368</td>
</tr>
<tr>
<td>SCAM</td>
<td>SCAM RMI / LIBS / VISIR / RAMAN 3x3 raster on abraded patch for lithology documentation</td>
<td>Alfalfa</td>
<td>370, 375</td>
</tr>
<tr>
<td>WATSON</td>
<td>Coring target documentation (pre-drill) at each core site for paired samples</td>
<td>Hahonih</td>
<td>371, Atsah</td>
</tr>
<tr>
<td>WATSON</td>
<td>Nighttime image for borehole documentation, only one borehole imaging for paired samples</td>
<td>Hahonih</td>
<td>373</td>
</tr>
<tr>
<td>SCAM</td>
<td>10x1 SCAM RMI / LIBS for chemistry and mineralogy. Only required on one borehole for paired samples</td>
<td>Hahonih</td>
<td>373</td>
</tr>
<tr>
<td>ZCAM</td>
<td>Multispectral image of borehole and tailings for mineralogy. Only required on one borehole for paired samples</td>
<td>Hahonih</td>
<td>374</td>
</tr>
</tbody>
</table>

Sample-Related Observations

Workspace Images

Sid sits among an accumulation of similar Ch’al-type rocks, along with abundant sand and regolith. The surrounding area is characterized by flat-lying, polygonally-jointed bedrock, interpreted to be the Nataani member (Fig. 2c). In the bedrock below Sid, fine (~5 mm), parallel layers are present, interpreted to represent laminations or exfoliation in the Nataani member. The fine laminations below
and around Sid do not appear disturbed by the presence of the boulder, and there is no nearby ridge that Sid could have fallen from. This strongly suggests that Sid represents an in-place remnant of the Ch’ał member.

The natural surface of Sid (Fig. 4, 5) is wind polished and generally dust-free, though dust and wind-blown sediment have accumulated in depressions and in the regions beneath overhangs. Sid has patchy coatings which are co-located with modern accumulation of regolith/dust in local topographic lows. Few, if any, individual grains are visible in the WATSON pre-coring natural surface images, but individual grains and pits are observed in the RMIs of relatively fresh/broken surfaces on Sid and other nearby targets (Fig. 5). Where individual grains can be identified, an interstitial reddish-white material is present. This phase may represent an interstitial phase and/or surficial dust caught between grains. The few grains/crystals that could be measured ranged in size from 0.2 to 1.5 mm. Distinctive features at the surface of Sid were adequate for reconstruction of original core azimuthal orientation, so no additional core orientation marking was performed.

Figure 4 | Mastcam-Z image of Sid. Note its glossy, fluted exterior texture and the massive character. The yellow rectangle shows the location of the inset WATSON 7 cm standoff image of the Alfaifa natural surface texture (lower left) showing ventifact ridges and clinging dust.
**Figure 5 | Surface textures of Sid.** Stretched Mastcam-Z image of the Sid block and nearby targets with SCAM RMI images of natural surfaces showing grains and other textures.
Pre and Post Coring/Abrasion Images

The team evaluated four potential locations for proximity science (abrasion and/or coring). The Spidergrass and Bullgrass locations on the top of the Sid block (Fig. 1) were preferred for coring, in order to sample any internal structure correlated to the gravity vector. The Alfalfa target (Fig. 1) was selected for a 9-mm depth abrasion. Drill operation parameters during abrasion suggest that Sid was the hardest rock encountered by the mission to date. The abrasion yielded a complete, flat-bottomed disk with a crisp edge. The internal texture of the rock was well exposed, showing a crystalline texture with lath-like grains, dark grains, and a fair amount of reddish-brown material (Fig. 6). The sample cores Hahonih (Navajo for “perseverance”) and Atsah (Navajo for “eagle”) were collected from the top of the Sid block.

ZCAM multispectral observations of the Sid natural surface, Alfalfa abrasion patch, and Hahonih tailings are shown in Fig. 7. The uncoated wind-polished surface shows a very flat spectrum with a shallow absorption band centered <900 nm. Purple coatings show a similar band shifted to longer wavelengths (~900 nm). The Sid natural surface, average abraded patch, and tailings from Alfalfa all show a broad band centered near 860 nm. Hematite may be contributing to the position of the band center (some type of iron oxide or oxyhydroxide is supported by the band at ~535 nm), but the broadness is likely due to orthopyroxene. The tailings show a slight flattening or downturn at 1000 nm that could be related to hydration or scattering by fine particles.

The ZCAM multispectral data of Sid are consistent with mafic mineralogy (Fig. 8). These observations also show significant internal oxidation in Sid, which causes the overall red appearance of the core and tailings. Within the abraded patch, the strong red slopes at short wavelengths that provide the overall red color are associated with hematite, even though little crystalline hematite is detected at Mastcam-Z scales. The abraded patch exhibits bright grains with no clear iron absorptions (likely feldspar) and dark/red grains. The Hahonih tailings are redder than the wind-polished rock surface, likely due to liberation of fine-grained hematite from the interior. Similar effects have been observed by the Curiosity rover at Vera Rubin ridge in Gale crater (Horgan et al., 2020b). This is the first sample site for which we have seen this type of reddening in Jezero. For comparison: Brac was very magenta (olivine-dominated), Issole was very blue (CPX-dominated), Roubion was very green (OPX-dominated), and Rochette was very magenta (CPX-dominated).
Figure 6 | The Alfalfa abrasion (Left) Location of the Alfalfa abrasion patch in WATSON images (notice the sharp edges and complete abrasion for the 9 mm deep patch). (Bottom) nighttime WATSON image of the Alfalfa abrasion patch showing the interlocking, crystalline nature of the rock with large (several mm) white feldspar laths, pyroxene and other mafic minerals in shades of gray, and pervasive, interstitial rusty-brown iron oxides.
Figure 7 | Mastcam-Z multispectral image of Alfalfa. Images on left are approximate natural color with regions of interest for spectra at right indicated by colored boxes. (top) Average spectra of the natural rock surface (cyan/pink), abrasion tailings (yellow/red), and abraded surface (green), all exhibiting broad and weak absorptions centered near 860-900 nm due to orthopyroxene, hematite, or some combination, with the presence of bands near 525 nm supporting some ferric iron. (bottom) Detailed analysis of the abraded patch showing diversity in individual phenocrysts. Bright white regions (purple/green) show limited absorption due to iron consistent with feldspar, while red regions (magenta, orange, yellow) show broad bands consistent with orthopyroxene, either with Fe3+ substitutions or mixed with hematite. Dark black regions show limited spectral contrast, possibly due to opaque oxides or fine-grained mafic minerals. Sol 368 Zcam03328 taken at Z110 focal length.
**Figure 8 | ZCAM multispectral mineralogy.** (a) Sid natural surface and Alfalfa abrasion patch, and (b) Hahonih tailings. Red = R0R/R1 (630/800 nm) Green=BD910 (band depth) Blue=R1/R5 (800/978 nm), color over blue Bayer filter grayscale image. Mineral interpretation is magenta= olivine or CPX, blue/green= OPX or CPX, green/black= ferric dust/purple coatings. The Hahonih tailings have weaker mafic signatures than the Sid rock surface both due to a shallower 910 nm band and a much redder slope.

**Elemental Geochemistry - PIXL**

Elemental geochemistry for Sid from X-ray fluorescence (XRF) mapping by the PIXL instrument on the Alfalfa abrasion patch performed on sol 369 is shown in Fig. 9. Alfalfa is inferred to be a basaltic igneous rock, likely a ferroan trachyandesite or basaltic trachyandesite (Fig. 10), with large phenocrysts of X-ray-diffracting sodic plagioclase.

The average chemical composition for the sol 369 observations is given in Table 2. These data indicate a silicate rock that is quite rich in ferrous iron compared with terrestrial rocks of comparable SiO₂ contents; its Mg# (molar Mg/(Mg+Fe)) is only 9%. We know of no comparable martian compositions, except for other rocks of the Máaz formation (Guillaumes, Bellegarde). In addition to the tabulated values, the Alfalfa bulk composition contains 650±290 (1σ) ppm Sr, which is consistent with its significant proportion of plagioclase feldspar.

On a total alkali versus silica (TAS) plot (Fig. 10), PIXL analyses for Alfalfa are distributed across the fields of basalt, trachybasalt, trachyandesite, and andesite; this range reflects mixing of the abundant plagioclase with mafic materials (augite, K- and Fe-rich mesostasis materials, and their alteration products). The average composition for Alfalfa is near the boundary between trachybasalt and trachyandesite classifications. The scattered points at low SiO₂ contents represent igneous Fe-Ti oxide and Ca-phosphate minerals, the Fe-rich mesostasis material, and a small proportion of salt minerals.
Figure 9 | PIXL placement on Alfalfa, sol 369. PIXL XRF scan, measuring 7x7 mm, on the abrasion patch (~5 cm across). On this image, feldspar is white to greenish, augite pyroxene is dark green, and orthopyroxene and altered material are brown. WATSON image SIF_0367_0699544277_375FDR N0110108SRLC08029_0000LMJ01.

Figure 11 shows a ternary diagram of molar compositions Al₂O₃ – (CaO+Na₂O+K₂O) – (FeOT+MgO) for the average composition and for each individual PIXL spot analysis in the sol 369 observation. Idealized common mineral compositions are also plotted on the diagram. The vast majority of points fall in the triangle defined by typical mafic minerals, and the majority of those project (more or less) near binary mixtures of plagioclase + augite, or plagioclase + an iron and magnesium rich phase. These binary mixtures suggest grains are small enough to permit many PIXL analytical spots to span two grains, but large enough that relatively few spots span three grains. Only a few PIXL analytical points plot above the plagioclase--[FeOT-MgO] line, i.e., are enriched in Al₂O₃ relative to typical igneous minerals. Those points are close to a chlorite mineral composition, but could be interpreted as mixtures of the FeOT-MgO-rich material and a small proportion of aluminous clay minerals. Overall, this plot indicates that Alfalfa is not intensely weathered.

PIXL multi-element maps of the Alfalfa scan area (Fig. 12) show euhedral rectangular crystals of plagioclase and subhedral (partially irregular) augite pyroxene grains in a poorly crystalline (or fine-grained) Fe-rich and K-Al-rich groundmass. Figure 12a emphasizes the plagioclase crystals (light green), and Figure 12b shows that they are chemically zoned. The plagioclase grains show strong diffraction peaks, indicating that they are crystalline. They are commonly surrounded by rinds of
Figure 10 | Total alkali vs. silica diagram for PIXL analyses of the Alfalfa abrasion patch (gray circles) and SuperCam LIBS observations of Alfalfa, the Sid natural surface, and other nearby similar rocks (black squares). The relatively large feldspar crystals were of particular interest to the team, so their inclusion in the PIXL scan may bias the bulk composition toward the alkali-rich rock classes. The underlying igneous classification scheme does not apply to the individual data points and simply provides a frame of reference for comparing samples. The weighted mean of the observations (colored symbols) are consistent with Sid being a basaltic andesite to trachyandesite.

Material poor in Al and K. Among the plagioclase crystals are intergrown augite crystals and K-Al-rich (non-diffracting) mesostasis (purple and dark green respectively in Fig. 12a). Minor igneous minerals include Ca-phosphate (Fig. 12c) and Fe-Ti oxide (Fig. 12d); no Cr-rich oxides were observed (Fig. 12d). The Fe-rich material (red in Figs. 12a & d) does not show X-ray diffractions detectable by PIXL, has analytical total near 100%, and contains Mn but minimal proportions of other elements (alkalis, Ca, S, P, Cr, and Ti). The scan area on Alfalfa contains only a small proportion of salt minerals, recognizable as Ca- or Mg-sulfates. Although sulfates in Alfalfa are sparse relative to previous samples, chlorine is present at relatively high concentrations in most spots other than the plagioclase crystals. One-quarter of the spot analyzes contain Cl concentrations consistent with moderate levels of Cl-bearing salts, e.g., ~10 wt% of an analytical volume could be a perchlorate salt.

PIXL’s Micro-Context Camera (MMC) images of the Alfalfa abrasion patch (Fig. 13) show more of the abrasion area than the XRF scans (Fig. 13a), and are used to extrapolate phase proportions from those of the XRF scan areas to nearly the whole abrasion patch. The MMC image of NIR-G-B (Fig. 13b) is analogous to WATSON visible images, and the G/NIR ratio image shows a clear distinction among the phases plagioclase, augite, and mesostasis (Fig. 13c). These visible & qualitative relationships are quantified in a graph of G vs NIR reflectances (Fig. 13d), which allows extraction of the proportions of these phases in the whole MMC image area.
Table 2. Bulk sum (average) chemical compositions by PIXL XRF.

<table>
<thead>
<tr>
<th></th>
<th>Bulk Sum 369</th>
<th>Plagioclase</th>
<th>Augite Salt Free</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wt %</td>
<td>N=3249</td>
<td>Err 1σ</td>
<td>N=304</td>
</tr>
<tr>
<td>Na₂O</td>
<td>4.5</td>
<td>1.0</td>
<td>7.6</td>
</tr>
<tr>
<td>MgO</td>
<td>0.73</td>
<td>0.31</td>
<td>0.31</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>11.6</td>
<td>0.6</td>
<td>22.1</td>
</tr>
<tr>
<td>SiO₂</td>
<td>57.1</td>
<td>2.8</td>
<td>58.4</td>
</tr>
<tr>
<td>P₂O₅</td>
<td>0.84</td>
<td>0.24</td>
<td>0.25</td>
</tr>
<tr>
<td>SO₃</td>
<td>1.00</td>
<td>0.23</td>
<td>0.27</td>
</tr>
<tr>
<td>Cl</td>
<td>2.08</td>
<td>0.40</td>
<td>0.38</td>
</tr>
<tr>
<td>K₂O</td>
<td>1.90</td>
<td>0.40</td>
<td>0.73</td>
</tr>
<tr>
<td>CaO</td>
<td>4.31</td>
<td>0.42</td>
<td>6.70</td>
</tr>
<tr>
<td>TiO₂</td>
<td>0.59</td>
<td>0.18</td>
<td>0.01</td>
</tr>
<tr>
<td>Cr₂O₃</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>MnO</td>
<td>0.28</td>
<td>0.15</td>
<td>0.02</td>
</tr>
<tr>
<td>FeO-T</td>
<td>13.24</td>
<td>0.78</td>
<td>1.73</td>
</tr>
<tr>
<td>Sum %</td>
<td>98.2</td>
<td>98.6</td>
<td>100.6</td>
</tr>
<tr>
<td>Mg#/An</td>
<td>9.</td>
<td>33.</td>
<td>9.</td>
</tr>
</tbody>
</table>

N are number of analyses averaged. Err is uncertainty (1σ) from calibration and counting statistics. Mg# is molar Mg/(Mg+Fe), in percent. An for plagioclase is molar Ca/(Ca+Na), in percent.
Figure 11 | Mineralogical interpretation of the Alfalfa abrasion, PIXL observations on sol 369. Data plotted by pixel and as bulk sum composition (blue circle, Table 1) on ternary diagram of molar abundances of Al2O3–(CaO+Na2O+K2O)–(FeOT+MgO). Common pure mineral composition are shown (opx = Ca-poor pyroxene, i.e orthopyroxene; cpx = calcic pyroxene, i.e. augite). Most of the individual PIXL analysis points are consistent with pure minerals (olivine, cpx, Fe-oxide) or binary mixtures of feldspar--cpx, or feldspar--[FeOT+MgO], or ternary mixtures of cpx--feldspar--[FeOT+MgO]. Aluminous alteration minerals, like common clay minerals, fall above the upper red dashed line. Several other secondary minerals (e.g., Fe-Mg-Ca-sulfates/carbonates, halite) are also plotted.
Figure 12 | PIXL X-ray multi-element maps for the *Alfalfa* abrasion patch on sol 369. (a) Red=FeO, Green=Al₂O₃, Blue=CaO. (b) Red=CaO, Green=Na₂O, Blue=K₂O. (c) Red=CaO, Green=P₂O₅, Blue=SO₃. (d) Red=FeOT, Green=TiO₂, Blue=Cr₂O₃. The rock consists of Na-rich plagioclase crystals (green in frames a & b), augite (purple, frame a), alkali-rich groundmass (blue, frame b), and material of orthopyroxene composition (dark red, frame a; black, frame b). Ca-rich material (red in frames b & c), possibly carbonate, is scattered through the groundmass. Apatite and Ca-sulfate are present but uncommon (frame c) Frame d shows oxide minerals; the brightest red areas are iron-rich with minimal Ti or Cr, and show minimal diffractions, yellow-green is likely ilmenite; the area contains no Cr-rich oxide minerals.
Figure 13 | PIXL MCC (Micro-context camera) data for the Alfalfa abrasion patch, sol 369. (a) MMC grayscale image, a portion of the Alfalfa abrasion. (b) MMC false color of abrasion area; Red = MMC Near-IR; Green = MMC Green; Blue = MMC Blue. Location of PIXL XRF scans shown. (c) MMC ratio image Near-IR/Green reflectances. Augite is darkest and mesostasis and alteration materials are lightest tone. Location of PIXL XRF scan shown. (d) MMC reflectance ratios.

Mineralogy and Organics-SHERLOC

SHERLOC observed the Alfalfa abrasion patch on sol 370. Spectra were collected with a survey scan (5x5 mm, 1296 points, 144 µm spacing, 15 pulses per point, not shown) and a HDR scan (7x7 mm, 100 points, 780 µm spacing, 500 pulses per point). Results are shown in Fig. 14 and Table 3.

The average Raman spectrum of all 100 points in the scan area shows a broad band assigned to disordered/microcrystalline silicates (1000 cm$^{-1}$). Peaks in spectra at selected locations are consistent with the presence of carbonate (1090 cm$^{-1}$) and poly-crystalline carbonate, feldspar (515 cm$^{-1}$), olivine (low Fo#), and quartz (475 cm$^{-1}$), though weakness of the possible feldspar and quartz peaks could be due to the fact that they are below the nominal cut-off of the SHERLOC edge filter so only very intense signals would be detectable. The observed fluorescence features (275-280 nm, 330-340 nm) are similar to what has been observed in Guillaumes and other targets in the Máaz fm and are consistent with single and double ring aromatic organic molecules, respectively.
**Figure 14 | Sol 370 SHERLOC fluorescence and Raman spectral results on Alfalfa.** A) Colorized ACI image of the abraded patch with a blue rectangle indicating the HDR scan area, and three white circles indicating regions of interest (ROIs) that correlate to the fluorescence data shown in B and C. B) Upper panel is an RGB map showing main regions of organic fluorescence (275-280 and 330-340 nm). The blank lower panel indicates no detection of organic materials by Raman spectroscopy (see Raman data in part D). C) Fluorescence spectra for the whole analysis area (upper) and from selected ROIs (lower) (see A and B). D) Map of the Raman HDR scan (left) and selected spectra (right). In the map, the grid of small red points indicates analysis points, and the large colored circles indicate the locations of mineral detections. In the panel of spectra, the uppermost spectrum is the average spectrum of all 100 points in the scan area. The second spectrum is an average of all the points labeled with blue and light blue on the spectral map. The third spectrum is an average of all the spectra within the two dark green polygons on the map, which enclose two apparent feldspar laths. The lowermost spectrum is from Point 5 (orange circle on map). Each spectrum has been baseline subtracted and offset along the y-axis for clarity. Note in each the presence of a laser artifact near 650 cm$^{-1}$. The scale in panel A also applies to the map in panel D.
Table 3. SHERLOC Raman spectra mineral ID.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Certain</th>
<th>Possible (not confirmed)</th>
<th>We looked for these, but cannot find them</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alfa1a</td>
<td>Disordered and/or microcrystalline silicates (widespread throughout)</td>
<td>Olivine (low Fe/o) occurs in two spots, but not obviously associated with crystal habit.</td>
<td>Pyroxene</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Carbonate (could be Mg/Ca/Fe)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Feldspar, difficult to confirm because of weak signal, but largely associated with crystal habit</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Quartz, only found in one point spectrum, weak signal</td>
<td></td>
</tr>
</tbody>
</table>

**Elemental Geochemistry and Mineralogy – SuperCam**

The SuperCam LIBS observations for *Sid* and nearby targets (Fig. 15) show that *Ch’al* member rocks are heterogeneous, consistent with being coarse-grained on the scale of RMI images (Fig. 5) and LIBS point spacing (generally a few mm). The overall elemental trends are consistent with the presence of feldspar and pyroxene, with some Ca-pyroxene and the possible presence of olivine and/or Fe-oxide, with additional unrecognized phases. All points show evidence of hydration, some points have evidence for Fe-OH band and possible Al-phyllosilicates (2.2mm band) indicating some degree of aqueous alteration. Generally, the composition of *Sid* and nearby targets is in family with previously encountered *Ch’al*-like rocks (Wiens et al., 2022).

SuperCam VISIR spectra on *Sid* (natural surface, abraded surface and tailings from coring activity; Fig. 12) also reveal an overall mafic mineralogy, where the broad band between 0.7-1.8 microns is due to mafic minerals like olivine, pyroxene, or glass. The SCAM VISIR technique is highly sensitive to hydrated minerals, so the signatures of hydration are present in the spectra even when these minerals are a minor proportion of the rock. Fig. 16 shows the absorption bands that indicate the presence of hydrated minerals in *Sid*. The main signal of hydration is present at 1.94 um, whereas the absorptions at 2.29 µm coupled with absorptions centered near 1.4, 1.9 and 2.39 µm likely indicate Fe-bearing phyllosilicate minerals. The weak absorption at 2.32 µm in the tailings likely indicates a Mg-OH phase and the 2.46 µm can possibly be explained by the presence of akaganeite. The VISIR spectra are greatly affected by the grain size, contributing to the differences in the tailings and rock spectra.
Figure 15 | SuperCam LIBS major element oxide analyses of Sid and all Máaz fm natural surface analyses from sols 15–201 and sols 343-379. Solid solution fields for olivine, pyroxene, and feldspar compositions are included.
**Figure 16** | Mean reflectance spectra of the SuperCam VISIR observations on **Sid** (natural surface, abraded surface and tailings). Saturated and shadowed points were excluded from averaging. Lighter dots near 2 µm indicate the spectral range possibly affected by CO$_2$ absorption features that could remain after the atmospheric correction. Note that owing to temperature sensitivities in the IRS instrument, the calibration is still uncertain for wavelength longer than ~2.5 µm.

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**Subsurface Structure**

The subsurface structure in the vicinity of the **Sid** outcrop is revealed by RIMFAX radargrams acquired on sols 360–362, inbound to the outcrop, and sol 379, outbound from the outcrop (Fig. 17a and b). **Sid** is a **Ch’al**-type rock that outcrops as blocky material on top of topographic highs overlying recessive polygonal rocks, such as **Nataani** and **Roubion**, that appear to form a coherent layer in the subsurface as seen in RIMFAX radargrams (Fig. 17 c and d). In the radargrams, the uppermost ~2 m represents the **Máaz** fm, where the deposits are dominated by horizontal layering and polygons visible at the surface. Polygon-bounding fractures are not visible in the radargrams, suggesting that these fractures may be very shallow. Bowl-shaped reflector geometries in the **Máaz** fm are interpreted to represent remnant (eroded) impact structures (Fig. 17c and d). The underlying **Séítah** fm strata occur between ~2 and 7 m burial depth. The boundary between the base of the **Máaz** fm and the top of the **Séítah** fm is interpreted as an erosional unconformity based in part on the presence of eroded impact structures. To avoid safety hazards, the rover traverse stayed clear of high standing **Ch’al** member outcrops, so this unit was not characterized by radar.
Figure 17 | Geological context and RIMFAX-derived subsurface stratigraphy of the Sid area. (a) Location of the Sid area, southeast of the Octavia E. Butler (OEB) landing site. (b) Detailed map of the area around Sid showing rover drive sections (green traverses) used to create radargram sections; (c) Radargram of the inbound traverse toward Sid on sols 360–362 with annotations. (d) Radargram of the outbound traverse away from Sid on sol 379 with annotations. Each radargram covers similar stratigraphy and record a mixture of horizontal, dipping, and bowl-shaped reflector geometries.

Core orientation

The Hahonih core was collected on Sol 371 using nominal operations for drilling and caching. The core broke off cleanly at the base of the borehole with a slight “mushroom” stem (Fig. 18).

Features of the natural surface of the coring targets on Sid were used to measure core orientation (Fig. 19). The core roll (α) is defined as the angle between the WATSON y-axis and the up-dip direction in SITE coordinates. At the time of drilling and 5.6-cm pre-drilling WATSON imaging, the rover, coring drill, WATSON target, and coring target (called “Hahonih.PSC.0.0729”), had the following characteristics:

a) Rover orientation quaternion just after drilling but before unloading the stabilizers (transferring from RMECH to SITE frame): bQll1 = (0.1500300, 0.0144798, -0.0209876, -0.9883530)
b) Coring Drill orientation quaternion just after drilling but before unloading the stabilizers (transferring from CORING DRILL to RMECH frame): $\mathbf{cdQ}_b = (0.5911830, 0.1944370, -0.7818390, 0.0377382)$

c) Rover orientation quaternion at time of acquisition of WATSON image

\[ \mathbf{bQ}_l \mathbf{l}_2 = (0.1500950, 0.0115230, -0.0213943, -0.9883730) \]

d) WATSON orientation quaternion at time of acquisition of WATSON image

\[ \mathbf{wQ}_b = (0.5908020, 0.1961150, -0.7817180, 0.0375390) \]

Figure 18 | Hahonih borehole and core images. (left) SHERLOC WATSON image of the Hahonih borehole and (right) cropped CacheCam image of the bottom of the Hahonih core. The sample core is 13 mm in diameter.

Items 1 and 2 give a coring drill pointing vector estimate of

\[
\begin{align*}
\text{hade} &= 21.41^\circ \\
\text{azimuth} &= 73.18^\circ
\end{align*}
\]

Items 3 and 4 give an estimate of the angle between the WATSON y-axis and the up-dip direction in SITE coordinates of (Fig. 19):

\[
\text{core roll, } \alpha = 126.60^\circ
\]
Figure 19 | Hahonih core orientation. 5.6-cm standoff WATSON image of Hahonih core target on sol 370. Image scale is 27.3 μm pixel⁻¹. Orientation compass gives WATSON frame (wx, wy, wz). Core roll is clockwise angle of core y-axis, cy, from wy, given by $\alpha = 126.60^\circ$. Projection of Martian geographic north onto WATSON image plane is noted. WATSON image SIF_0370_0699795188_894RAS_N0110108SRLC00663_0000LMJ01.

**Preliminary Scientific Assessment**

**Synthetic sample description and preliminary interpretation**

Sample type: Basaltic andesite

1. Stratigraphic and Geologic Context

   a. Stratigraphic position: Hahonih and Atsah represent the Ch’al member of the Máaz formation. The Ch’al member appears to be the stratigraphically highest member of the Máaz fm, overlying the recessive rocks of the Nataani member.

   b. Source rock: Hahonih and Atsah are derived from the Sid rock. Sid is interpreted to be an in-place block of the Ch’al member bedrock.
2. Texture and fabric

a. Rock fabric (natural surface): Massive, ventifacted, individual grains or crystals rarely visible
b. Rock fabric (abrasion patch): Holocrystalline, no compelling evidence of intergranular porosity or cement
c. Grain Size: Large euhedral feldspar crystals a few mm long; subhedral crystalline ortho- and clinopyroxene ≤1 mm.

3. Petrology, Mineralogy and Elemental composition

b. Bulk composition: Consistent with a basaltic andesite.
c. Organic molecules: Present, low abundance, probably aromatic ring compounds.
d. Alteration and hydration: Minimal. Evidence of hydration and some hematite coloration.

4. Core orientation

e. Sample cores were absolutely oriented to ~1° in martian geographic coordinates. Cores were taken from the top of the Sid rock.

Returned Sample Science Considerations

The collection of the Hahonih and Atsah samples from the Ch’al member completes the suite of samples from the Máaz fm and the Crater Floor Campaign. The Hahonih and Atsah samples are currently believed to be the best candidates for the least altered, and possibly most compellingly in-place, samples of the Máaz fm.

**Geochronology:** The Hahonih and Atsah samples appear to have limited alteration relative to other Máaz fm samples, which may make them the most robust Crater Floor samples for radiometric dating. The Ch’al member’s stratigraphic position and the timing of its emplacement will help constrain the age of the other units in Jezero crater, such as the (apparently overlying) delta and (apparently underlying) Séítah fm. The Máaz formation and especially the Ch’al mb can be related to relatively high crater densities observed on the Jezero crater floor (Quantin-Natif and others, 2022), therefore ages determined by radiometric and possibly cosmogenic methods of the Hahonih and Atsah samples may provide a comparison point with crater chronology age estimates. Published crater retention model ages for the Máaz fm range from 1.8 Ga to older than 3 Ga (Goudge et al., 2012; Holm-Alwmark et al., 2021; Marchi, 2021; Shahrzad et al., 2019) with the spread possibly due to erosional exposure of the unit.

In addition, cosmogenic nuclide measurements of these two samples will constrain their integrated duration of exposure to galactic cosmic rays while residing near the martian surface. Cosmic-ray exposure dating may help reveal the erosion rates at the (modern) Martian surface, as the Ch’al member is apparently eroding away from the underlying Nataani member surface. Such information
would also inform the expected preservation and modification of organic molecules present in these rocks.

**Paleomagnetism:** The igneous lithologies of the Sid cores may enable absolute measurements of the paleointensity of the ancient martian magnetic field at the time of crystallization. The orientations of the cores, combined with paleohorizontal indicators from outcrop foliation, may enable measurements of the absolute paleodirection of the ancient field. Paleomagnetic investigations of secondary ferromagnetic minerals could constrain the field at the time of (limited) aqueous alteration. Combined with measurements from other samples, these data could test the hypothesis that Martian atmospheric loss was driven by the cessation of an early dynamo, constrain the dynamo’s secular variation, and determine if the dynamo exhibited polarity reversals. Comparison of paleomagnetic directions with Séitah rocks (Salette/Coulletes and Malay/Robine) could distinguish between the hypotheses that Máaz and Séitah form a single layered intrusion versus Máaz being an overlying lava flow.

**Geochemistry:** The geochemical and isotopic composition of the Sid cores can be used to characterize the evolution of the martian mantle/crust, and melting processes. The Hahonih and Atsah samples appear to be slightly altered, in contrast to the more intense aqueous alteration in Máaz and Séitah fm cores. In tandem with other Máaz fm cores, laboratory analysis of the alteration minerals, formation temperatures, and ages will provide a history of aqueous alteration conditions in Jezero crater post-crystallization. A particularly compelling comparison will be between the Ch’āl member samples and the Rochette member samples, which appear to have a similar primary mineralogy, but different extents of alteration. Given that Hahonih and Atsah represent the least altered igneous rocks from the crater floor and contain only a weak organic signal, laboratory analyses may help establish the background level of reduced carbon in igneous crater floor rocks (including the nature and distribution of such carbon).

**Comparison with other Máaz samples:** Abraded proximity science was conducted on four rocks in the Máaz fm: Roubion, Rochette, Rimplas, and Sid. Roubion is an example of the flat-lying, polygonally jointed material of the Máaz fm. The Guillaumes abrasion patch revealed that Roubion was a fine- to medium-grained mafic rock, composed of plagioclase and pyroxene with minor apatite and FeTi oxides. Roubion was pervasively altered, as evidenced by pits and crevices in the abraded surface and abundant secondary minerals including iron oxide (possibly hydrated), sulfates, perchlorate, and possibly phosphate and halite. Both Rochette (abraded patch Bellegarde) and Rimplas were located along the feature named Artuby Ridge, a resistant ledge within the Máaz fm that can be traced from near the Roubion site along the southern edge of the Séitah fm contact. Proximity science showed that the Rochette and Rimplas rocks were similar to each other and broadly mineralogically similar to Roubion, being fine-to-medium grained mafic rocks dominated by plagioclase and pyroxene. The Rochette and Rimplas rocks also contained abundant secondary phases likely resulting from aqueous alteration, including iron oxide, Ca-sulfates (occasionally hydrated), phosphate, and minor carbonate. However, Rochette and Rimplas appear to be substantially less altered than Roubion, from which it was not possible to collect a sample. The Alfalfa abrasion patch on the Sid rock, a representative of the Ch’āl member of the Máaz fm, appears to be relatively coarser-grained, more alkalic, and less altered than other Máaz fm samples. It also contains large feldspar laths that are lacking in the other Máaz fm samples. The Sid samples appear significantly less altered than all of the other Máaz fm samples including the Rochette samples Montdenier and Montagnac.
The Hahonih and Atsah samples are currently believed to be the best candidates for the least altered, and most compellingly in-place, sample of the Máaz fm. The Montdenier and Montagnac samples seem to have a similar primary mineralogy, and contain more abundant secondary phases that will better reveal the post-emplacement history of water interaction with the Máaz fm.

**Summary:** the return of the Ch’al member samples will directly address several major RSS objectives and markedly improve our understanding of the earliest history of igneous processes on Mars. Returned samples of this unit have particularly high science value for determining the igneous petrogenesis of possible lava flows, the global evolution of Mars (interior and near surface environments), and for understanding the geological setting and relative and absolute chronology of Jezero floor units.

**References**


Sample Designation: M2020-374-10 Atsah

Date of Coring: 9-Mar-22

Mars Time of Sample Core Sealing: 21:12:02 LMST, Sol 377, Ls 187

Latitude (N), Longitude (E), Elevation: 18.44386406, 77.45242176, -2568.357m

Campaign: Crater Floor

Region of Interest: Ch’al member of Máaz formation, east of the Octavia E. Butler landing site

Lithology: Likely basaltic andesite. Few mm-size white feldspar laths and pyroxene grains are present, as possibly are olivine and quartz. Limited evidence of secondary alteration includes interstitial iron oxides, carbonate, possible phyllosilicates and chlorite, and hydration. Low concentrations of organic matter were detected.

Estimated Volume Recovered: 8.46 cm³

Coring Bit Number: 4

Core Orientation: hade = 18.02°; azimuth = 69.87°; core roll = 110.96°

Sample Serial Numbers: Tube SN202, Seal SN168, Ferrule SN074

ACA Temperature at Time of Sealing: 40°C

Estimated Rover-Ambient Pressure and Temperature at Time of Sealing: 669 Pa, 215 K

Estimated Amount of Martian Atmosphere Headspace Gas: 1.3x10⁻⁶ mol

Anomalous Behavior: Three sols elapsed between sample coring and sealing.

Abrasion Patch Name and Depth: Alfalfa, 9 mm

May 8, 2022

Summary Description

This sample is paired with M2020-190-9 Hahonih. Only the core orientation for Atsah is described below. See the Initial Report for M2020-190-9 Hahonih for sample details.

Core orientation

At the time of drilling and 5.7-cm pre-drilling WATSON imaging, the rover, Coring Drill, WATSON, and coring target (called “Atsah.PSC.0.0687“), had the following characteristics:

a) Rover orientation quaternion just after drilling but before unloading the stabilizers (transferring from RMECH to SITE frame): \( \mathbf{bQll1} = (0.150025, 0.0148175, -0.020817, -0.988352) \)
b) Coring Drill orientation quaternion just after drilling but before unloading the stabilizers (transferring from CORING DRILL to RMECH frame): \( \mathbf{cdQb} = (0.58264, 0.235836, -0.76571, 0.136382) \)
c) Rover orientation quaternion at time of acquisition of WATSON image SIF_0373_070065505_402RAS_N01110108SRLC00657_0000LMJ01: (transferring from RMECH to SITE frame): \( \mathbf{bQll2} = (0.1501010, 0.0114020, -0.0207122, -0.9883880) \)
d) WATSON orientation quaternion at time of acquisition of WATSON image SIF_0373_070065505_402RAS_N01110108SRLC00657_0000LMJ01: (transferring from WATSON to RMECH frame): \( \mathbf{wQb} = (0.5801300 , 0.23807000, -0.7670220 , 0.1358290) \)

Items 1 and 2 give a coring drill pointing vector estimate of

\[ \text{hade} = 18.02^\circ \]
\[ \text{azimuth} = 69.87^\circ \]

Items 3 and 4 give an estimate of the angle between the WATSON y-axis and the up-dip direction in SITE coordinates of (Fig. 1):

\[ \text{core roll, } \alpha = 110.96^\circ \]
**Figure 1 | Atsah core orientation.** 5.7-cm standoff WATSON image of Atsah core target on sol 373. Image scale is 27.6 μm pixel$^{-1}$. Orientation compass gives WATSON frame ($\mathbf{w}_x$, $\mathbf{w}_y$, $\mathbf{w}_z$). Core roll is clockwise angle of core y-axis, $\mathbf{c}_y$, from $\mathbf{w}_y$, given by $\alpha = 110.96^\circ$. Projection of Martian geographic north onto WATSON image plane is noted. WATSON image SIF_0373_0700655505_402RAS_N01101085RLC00657_0000LMJ01.