Detection of Carbonates in Martian Weathering Profiles

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Abstract Noachian surfaces on Mars exhibit vertical assemblages of weathering horizons termed as weathering profiles; this indicates that surface water caused alteration of the rocks that required a different, warmer climate than today. Evidence of this early Martian climate with CO2 vapor as the main component causing greenhouse warming has been challenged by the lack of carbonate in these profiles. Here we report the analysis of Compact Reconnaissance Imaging Spectrometer for Mars L-detector data leading to the detections of carbonates using a spectral signature exclusively attributed to them. The carbonates are collocated with hydroxylated minerals in weathering profiles over the Martian surface. The origin of CO2 for the formation of carbonates could be the atmosphere. The widespread distribution of weathering profiles with carbonates on the surface of the planet suggest global interactions between fluids containing carbonate bicarbonate ions with the surface of Mars in the presence of atmospheric water until around 3.7 billion years ago.

Plain Language Summary The oldest surface of Mars witnessed an impact of dense and humid atmosphere leading to the formation of hydrated minerals. The analysis of remote sensing data allows identifying carbonates mixed with the hydrated minerals. These associations of carbonates and hydrated minerals are widespread on the surface of the planet. This indicates a planetary-scale process of formation involving the presence of fluids containing inorganic carbon. This finding allows a better understanding of the environment of Mars during the period suspected to be the most likely to have host habitable environment on the planet.

1. Introduction

Several morphological features (valley networks, Howard et al., 2005; open-basin lakes, Goudge et al., 2012; fluvial pebble conglomerates, Williams et al., 2013) indicate that liquid water once shaped the Martian surface. Moreover, widespread weathering leading to the formation of clay minerals documents that surface water percolated through Martian basaltic rocks (Carter et al., 2013). Among clay minerals formations, Carter et al. (2015) described weathering profiles. Weathering profiles are vertical assemblages of weathering horizons of a few centimeters to near a hundred meters depending on the local climatic conditions. Weathering profiles are open systems affected by percolation of meteoric water. The nature of the weathering profiles is a complex response to climatic and geologic controls and to long-term changes in surface environmental conditions (Pinti, 2011; Velde & Meunier, 2008). Some weathering profiles on Mars were preserved because they are protected by a datable capping unit made of unaltered volcanic rocks. This allows us to suggest that they formed no later than 3.65 billion years ago (Carter et al., 2015; Loizeau et al., 2012). The most complete weathering profiles are characterized by a sequence of horizons topped with Al-rich clay minerals, overlying Al,Fe-rich clay minerals, and Fe,Mg-rich clay minerals at the bottom (i.e., Carter et al., 2015). Such profiles are better explained by chemical weathering by an acidic fluid (Carter et al., 2015; Gaudin et al., 2011; Loizeau et al., 2012; Zolotov & Mironenko, 2016). On Earth, such sequences are indicative of humid climates with above-freezing temperatures and sustained precipitation over geological timescales (Allen, 1997; Navarre-Stitchler & Brantley, 2007). By analogy, meteoric waters (aqueous solution in equilibrium with atmosphere) on Mars must have percolated through the parent rock over millions of years to explain tens of meters thick horizons of clays as suggested by Earth studies (Righi & Meunier, 1995). However, Bishop et al. (2018) discuss the possibility of forming these profiles with a...
succession of sporadic short-term climate optima producing warm and wet conditions. This would be coherent with impact-induced rainfall (Zolotov & Mironenko, 2016). If Mars experienced a humid climate for a long time, a dense atmosphere mostly composed of CO₂ and lower amounts (few percent) of other greenhouse gases is required (Wordsworth et al., 2017). Under such climatic conditions, both geochemical models (Schaefer, 1990; Zolotov & Mironenko, 2016) and laboratory experiments (Gaudin et al., 2018; Viennet et al., 2017) have shown the coprecipitation of hydrated minerals and small amounts of carbonates. Therefore, the apparent absence of carbonates in these weathering profiles remains enigmatic and debates on inhibited formation or subsequent dissolution of the carbonates are yet inconclusive (Edwards & Ehlmann, 2015). Current carbonate detections on Mars are mostly restricted to subsurface hydrothermal settings implying little to no exchange with the atmosphere (i.e., Wray et al., 2016). Recently, several additional carbonate detections have been reported using Compact Reconnaissance Imaging Spectrometer for Mars (CRISM) near-infrared reflectance data in regions analyzed in our study (Amador et al., 2018), but the methodology of detection precludes a detailed context of the carbonate formation. Furthermore, a limited amount of carbonates, possibly formed by surface weathering or more likely by evaporative precipitation at low temperature, is documented (i.e., Boynton et al., 2009; Morris et al., 2010; Ruff et al., 2014). However, evidence for long-term atmosphere-surface interaction on global scale leading to surface weathering associated with formation of carbonates was lacking.

Here we report the presence of carbonates associated with clay minerals in weathering profiles. This suggests that the alteration fluid, containing inorganic carbon, interacted with the surface of Mars on a planetary scale. In section 2, the method for mineral detection is presented with an emphasis on the carbonates case. We present tests of the method used for the detection of carbonates mixed with hydrated minerals. Section 3 gives an overview of the results obtained from the analysis of the weathering profiles. Then, we focus on three well-preserved sites to describe the weathering profiles in more details. Finally, we discuss the formation process of the weathering profiles and its coherence with the current knowledge on the evolution of Mars.

2. Method

2.1. Data Selection

For the morphologic characterization of the sites studied, we used high-resolution image data collected by the Context Camera (CTX) and High Resolution Imaging Science Experiment (HiRISE) instruments both on board Mars Reconnaissance Orbiter. Altimetry data are derived from Mars Orbital Laser Altimeter (MOLA, Mars Global Surveyor) data to investigate the stratigraphy of the weathering profiles.

To select the portion of CRISM (Murchie et al., 2007) images to analyze, we relied on the well-documented study of Carter et al. (2015) to choose in the described area from all CRISM images. However, since the identification of carbonates is more difficult with low signal-to-noise ratio (SNR) data, we restrained our search on full-resolution targeted (FRT) data acquired before 2010 (2010 excluded). Nili Fossae has been excluded because in this region carbonates detected were probably formed in a hydrothermal context (Brown et al., 2010; Ehlmann et al., 2008; Viviano et al., 2013). Nevertheless, such area has been used to constrain our new criterion for carbonates detection. The number of CRISM images analyzed is 144. Table S1 in the supporting information summarizes the data used and the interpretation associated.

2.2. CRISM Processing

Using near-infrared spectral data (L-detector) obtained by CRISM on board the Mars Reconnaissance Orbiter, we successfully searched for carbonates within weathering profiles distributed over the Martian surface (Figure 1). The data were collected in the targeted mode in the spectral range of 1.00 to 3.92 μm, with a spectral resolution of ≈6 nm and a full spatial resolution of ≈18 m/pixel.

All data cubes were downloaded via the MarsSI application (Quantin-Nataf et al., 2018) and processed with CRISM Analysis Toolkit version 7.3.1 (Murchie et al., 2007). This process includes a photometric correction and a “volcano scan” correction for atmospheric gas absorptions (Langevin et al., 2005; McGuire et al., 2009). A spatial and spectral noise removal method is then applied to the data (Parente et al., 2008). The presence of dust on the surface weakens the mineralogical signal. Hence, a median ratio was used to highlight absorptions. Each pixel of a column is divided by the median spectrum calculated from the whole column following...
the procedure described in Bultel et al. (2015). All spectra were then analyzed by a minimal spatial median on 5 × 5 pixels for each region of interest. The spectral criteria, presented in Table 1, are adapted from Viviano-Beck et al. (2014) to map the signals of interest.

2.3. Mineral Identification

The hydrated minerals detected in this study exhibit absorption features in the 1.00 to 4.00 μm spectral domain but are typically identified on Mars in the 1.00 to 2.60 μm spectral domain (Clark et al., 1990). The absorptions near 1.40 and 1.90 μm are common to all hydrated minerals presented here, and they correspond to H2O stretching. The absorption near 2.16–2.21 μm indicates an Al-OH vibration (or Si-OH, depending on the shape of the absorption), while the absorption near 2.27–2.35 μm indicates a (Fe,Mg)-OH vibration. They usually have weaker overtones near 2.38 μm (linked to the Fe,Mg ratio) and near

Table 1

<table>
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<th>Liaison</th>
<th>Comment</th>
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<tr>
<td>BDa2200</td>
<td>$\frac{R(2.20)\times 2}{R(2.12) + R(2.23)}$</td>
<td>—</td>
<td>Al-OH</td>
<td>Highlights an absorption near 2.20 μm</td>
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<tr>
<td>BDa2300</td>
<td>$\frac{R(2.29) + R(2.36)}{R(2.23) + R(2.42)}$</td>
<td>—</td>
<td>(Fe, Mg)-OH</td>
<td>Highlights an absorption near 2.30–2.34 μm</td>
</tr>
<tr>
<td>Carb3900</td>
<td>$\frac{\text{Med}[R(3.81) : R(3.86)]}{\text{Med}[R(3.54) : R(3.75)]}$</td>
<td>$\text{NaN}$ if $\frac{\text{Med}[R(3.60) : R(3.70)]}{\text{Med}[R(3.60) : R(3.70)]}$</td>
<td>C-O$_3$</td>
<td>Highlights a spectrum with a bulge centered near 3.60–3.70 μm and a lowered reflectance near 3.40–3.50 and 3.80–3.85 μm.</td>
</tr>
</tbody>
</table>

Note. Med[R(X, XX); R(Y, YY)] is the median of reflectance on all the wavelengths between the two wavelengths indicated. R(X, XX) corresponds to the reflectance at the wavelength X, XX. “NaN” corresponds to “not a number.” $\alpha$ is a factor that can be adjusted in case of low absorption near 3.50 μm or large absorption near 3.00 μm hiding the carbonate feature and that is generally lower than 0.1.
2.50 μm. These different absorptions and their shapes aid in distinguishing the different phyllosilicates from Al- to Fe,Mg-rich phyllosilicates (Clark et al., 1990; Hunt, 1977). A single absorption near 2.40–2.50 μm relates to H2O stretching and AlCa-OH bending (Cloutis et al., 2002) or O-H, H-O-H, and S-O vibration (Cloutis et al., 2006) and indicates either zeolite or sulfate. While carbonates were previously identified on Mars by a paired absorption at 2.30 and 2.50 μm, we searched for a specific absorption pattern in the 3.40 to 3.80 μm wavelength range where absorptions band are the deepest (Figures 2a and 2b). This pattern can be described as reflectance inflections in the 3.40 to 3.50 and 3.80 to 3.90 μm ranges combined with higher reflectance near 3.60–3.70 μm. Moreover, the first identification of Martian carbonate has been possible thanks to this feature (Ehlmann et al., 2008). Second, such a spectral signature is uniquely explained by CO3^2− overtone vibrations, as documented in numerous laboratory measurements (Gaffey, 1987; Hexter, 1958; Hunt & Salisbury, 1971). Finally, the presence of absorptions near 3.40–3.50 and 3.80–3.90 μm causing a broad peak near 3.60–3.70 μm is systematic in all carbonate spectra for all the particle sizes considered, while it is not the case for the signature at 2.30 and 2.50 μm (Figure 2). In the following part we explain why the 3.40 to 3.92 μm range is preferred to detect carbonates.

In general, all carbonate spectra have a distinctive signature between 3.40 and 3.80 μm corresponding to a bulge-like feature with a maximum at ~3.65 to 3.70 μm. This feature is defined as follows in the literature (Cloutis et al., 2002; Hunt & Salisbury, 1971): a broad spectral peak near 3.60–3.70 μm, which is due to the deep 3.40 to 3.50 and 3.80 to 3.90 μm carbonate bands on either side. The median values of the...
reflectance near 3.40 and 3.80 μm are similar. This feature can only be attributed to carbonates (X-CO$_3$$^{2-}$; Gaffey, 1987; Hexter, 1958; Hunt & Salisbury, 1971; Hunt, 1977). Among the minerals most commonly found on Mars, which include olivine, pyroxenes, iron oxides, smectites, carbonates, and zeolites, the carbonates are the only one to show a clear distinctive feature in the 3.40 to 3.80 μm range. The other minerals listed above are showing a constant increase of reflectance from 3.00 to 4.00 μm (Figure 2a). Figure 2b shows two spectra of pure Fe-carbonate (FeCO$_3$ and siderite) of two grain size ranges (0–45 and 125–500 μm). Figure 2c presents different spectra of phyllosilicate and carbonate mixtures with different grain size ranges and carbonate contents, as well as a pure serpentine spectrum, for comparison. While the absorptions near 2.30–2.50 μm due to metal-carbonate ion compound (X-CO$_3$$^{2-}$) decrease with decreasing grain size range, the 3.40 to 3.80 μm absorptions due to C-O$_2$$^{2-}$ vibrations increase with decreasing grain size range (Figure 2b). This evolution has already been seen by Roush (2010), who showed that a decrease in grain size (from 500 to <45 μm) causes a decrease of the absorptions near 2.30 and 2.50 μm, and by Salisbury et al. (1987) and Salisbury and Walter (1989), who showed the amplification of the absorptions near 3.40 and 3.80 μm. The decrease of the absorptions near 2.30 and 2.50 μm with the grain size was also described in Hopkinson et al. (2017) for a grain size range from <3 to 121 μm.

The absorptions obtained on a mixture of phyllosilicates and carbonates are similar to the absorptions of a pure phyllosilicate in the 1.00 to 2.60 μm range (Figures 2b and 2c). The carbonate features, especially in the 1.00 to 2.60 μm range, are hidden by the absorption of phyllosilicates (Blaney & McCord, 1989; Bonello et al., 2004; Salisbury et al. 1987; Salisbury & Walter, 1989; Sutter et al., 2007, Wagner & Schade, 1996). Therefore, the identification and the characterization of the metal component in carbonate is not possible (see section 2.6). Sutter et al. (2007) report analysis of samples by infrared spectroscopy in the 1.00 to 4.00 μm range. The authors suggest that because of a complex mixing between carbonates, nitrates, sulfates, and phyllosilicates, the carbonate can uniquely be identified based on the 3.90 μm band in the spectral range studied. Indeed, the identification of carbonate mixed with various components can be uniquely identified using the 3.40 to 3.80 μm feature (Figure 2c). This also suggests that the grain size range plays a major role for carbonate absorptions band depth at 2.30–2.50 μm (respectively at 3.40–3.80 μm), which decreases (respectively increasing) with the decrease of the grain size range. Using such an absorption range (3.40–4.00 μm), Wagner and Schade (1996) suggest that more than 15 wt% of siderite mixed with palagonite (with a grain size <25 μm) is needed to allow making the bands due to carbonates visible in their study. Therefore, in the present study, the identification of carbonate is based on the shape of the spectra between 3.40 and 3.80 μm. Based on these specific signatures, we developed a spectral parameter to identify and map carbonates. We defined the spectral criterion Carb3900 to detect carbonate, which focused on the signal between 3.40 and 3.80 μm with a similar value for the medians of the reflectance intensity near 3.40 and 3.80 μm and a higher value for the median of the reflectance intensity near 3.60–3.70 μm. In section 2.4, we detail the validation of this spectral identification by comparison with previous carbonate detections in Nili Fossae (Ehlmann et al., 2008). Moreover, we have verified the presence of this CO$_3$$^{2-}$ signature by using overlapping CRISM spectral images in order to discard any possible artifacts at such positions (see section 2.5).

2.4. Carbonate Detection Validation at Nili Fossae

To further test whether the Carb3900 criterion is able to detect carbonates on previous detections (Ehlmann et al., 2008), we analyzed the CRISM data FRT0000B072 (Figure 3). The mineralogical maps compare the detection for the carbonate criteria described by Viviano-Beck et al. (2014), referred to as MIN2295_2480 (Figure 3a) and MIN2345_2537 (Figure 3a); BDCARB (Pelkey et al., 2007; Figure 3c), used in most of the publications with detections of carbonates on Mars with CRISM (Harner & Gilmore, 2015; Viviano et al., 2013; Wray et al., 2016); and Carb3900 (Figure 3b), newly used in this study. An RGB image is also present (Figure 3d) to help with the comparison. While the MIN2295_2480, MIN2345_2537, and BDCARB criteria are focusing on the absorptions near 2.30–2.50 μm, the Carb3900 criterion is focusing on the presence of a spectral feature near 3.40–3.80 μm. The criterion BDCARB highlights the carbonates; however, it contains false positives (see Viviano-Beck et al., 2014). Therefore, the MIN2295_2480 and MIN2345_2537 criteria are used here for the comparison. While the areas highlighted by the Carb3900 criterion are smaller than the one of the MIN2295_2480 or MIN2345_2537 criteria, there is no new area highlighted via the Carb3900 criterion. This is an illustration of how our criterion is more restrictive. The results displayed in
the maps confirm that the 3.40 to 3.80 μm feature highlighted by the Carb3900 criterion is linked to carbonate detections made in previous studies with BDCARB (Ehlmann et al., 2008). This example shows the ability of our criterion to map the presence of carbonates in places on Mars with earlier confident carbonate detections.

To understand why the Carb3900 criterion is more restrictive than are the MIN2295_2480 and MIN2345_2537 criteria, Figure 4 shows two spectra acquired in the Nili Fossae region from the image of Figure 3. The light green spectrum, identified as carbonates, is highlighted only by the MIN2295_2480 and MIN2345_2537 criteria. The light green spectrum, while having absorptions at 2.30, 2.51, and 3.80 μm and being detected with the MIN2295_2480 and MIN2345_2537 criteria, cannot be interpreted as carbonate with the Carb3900 criterion because of the missing clear 3.40 to 3.80 μm carbonate absorption shape. Indeed, a clear absorption near 3.40 μm is missing, and the reflectance at ~3.40–3.50 μm is higher than the reflectance at ~3.80–3.85 μm. The lack of feature at ~3.40 μm could be explained here by the presence of water, coatings, or mixture with other minerals as explained in Ehlmann et al. (2008). The light blue spectrum is interpreted as carbonates probably mixed with phyllosilicates (absorptions near 1.90 and 2.38 μm) with the Carb3900 and MIN2295_2480 and MIN2345_2537 criteria. Finally, the

Figure 3. (a) Color composite on CRISM cube FRT0000B072 with the following: in red, MIN2295_2480 and in green, MIN2345_2537 (Viviano-Beck et al., 2014) highlighting carbonates. (b) Criterion Carb3900 on CRISM cube FRT0000B072 highlighting carbonates. (c) Criterion BDCARB on CRISM cube FRT0000B072. (d) RGB composite on CRISM cube FRT0000B072 with the following: in red, 2.4303 μm; in green, 1.5066 μm; in and blue, 1.1127 μm.
Carb3900 criterion appears more restrictive as it needs a clear carbonate absorption shape in the 3.40 to 3.80 μm spectral range. This also explains why the mapping with Carb3900 reveals a smaller spatial extent of carbonates (Figure 3).

2.5. Detection Validation at Studied Areas

To verify the consistency of the acquired signal (especially after 3 μm), data for which two CRISM data cubes overlapped during two different orbits were compared. This approach evaluates the possible instrumental artifacts and any potential degrading process of instrument. The consistency of the signal over the entire spectral range of the CRISM L (long) channel (1.00 to 3.92 μm) is confirmed by the superimposition of the spectra obtained (see Figures 5 and 6, and Table S2 illustrate the regions of Mawrth Vallis, North Hellas, and Eridania Basin). Since the signal is consistent on the entire CRISM L spectral range, it demonstrates that the signal is related to characteristics of the surface itself (i.e., absorption due to interaction of the incident beam and mineral structures). Specifically, a co-occurrence of the 3.40 to 3.80 μm feature with the detection of weathering products is observed over Mawrth Vallis (Figure 5) while being systematically absent on nonaltered surfaces. Hence, this feature is an alteration product and is interpreted as carbonate.

Using two other examples within the sites of Eridania Basin and North Hellas, we compare different spectra obtained on the same coordinates with different CRISM acquisition characteristics (Figure 6). On the data from North Hellas (Figure 6a), the acquisitions are separated by 2 months, which explain the similarities of the two spectra. The same absorptions near 1.40, 1.90, 2.30, and possibly near 3.80 μm are visible. Note that the absence of a clear feature near 3.40 prevents the interpretation of carbonate signature. For Eridania Basin (Figure 6b), compared with for North Hellas, the larger difference in date of acquisition (1 year) led to a larger dissimilarity in the signal. However, as for the previous case, the same absorptions are visible. Those examples point out the consistency on the signal on the entire range of the CRISM L channel (1.00 to 3.92 μm) as a function of time.

2.6. The Challenge of the Carbonate Composition Characterization

Theoretically, the cation bonded to the CO$_3^{2-}$ ion can be identified based on the position of the bands near 3.40 and 3.90 μm (i.e., De Angelis et al., 2019). These features have recently been used to identify carbonate composition detected on Ceres (Carozzo et al., 2018; De Sanctis et al., 2015, 2016; Palomba et al., 2017). The distinction between different carbonate compositions can be done by determining the centers of absorptions near 3.40–3.50 and 3.90–4.00 μm. However, in the CRISM data, the right shoulder of the absorption near 3.90 μm is not visible, and only the position near 3.40–3.50 μm could be used. A center of absorption near 3.42 μm would be interpreted as a Mg-rich carbonate, near 3.45–3.50 μm as a Ca- or Fe-rich carbonate, and near 3.50 as a Na-rich carbonate. The presence of the other center of absorption near 3.90–4.00 μm is helping to clearly make the distinction. Distinguishing between Ca/Fe and Na-carbonates requires information on the position of the center of absorption near 3.90–4.00 μm, which is not completely covered here. Only Mg-rich carbonates could then be clearly identified. However, in case of a mixture with hydrated/hydroxylated minerals, the presence of a strong absorption near 3.00 μm (whose shape depends on the nature of hydrated/hydroxylated mineral present in the mixture) disturbs the position of the center of the absorption near 3.40–3.50 μm and hinders any characterization of the carbonate composition. Therefore, our study focuses on the detection of a spectral feature characteristic of carbonates and not on the characterization on the carbonate composition.
3. Results

3.1. Composition and Occurrence of the Weathering Profiles

We describe in this paragraph the common spectral characteristics of the apparent stratigraphy (see spectra in Figure 7). All the horizons of the weathering profiles have absorptions near 1.40 and 1.90 μm, both corresponding to the presence of hydrated minerals. They differ only in the 2.10 to 2.60 μm range and in the 3.40 to 3.80 μm range. The following description focused on these two ranges to identify minerals present in the weathering profiles.

The weathering profiles are characterized by a top horizon of Al-rich clay minerals having diagnostic absorption near 2.16–2.21 μm, indicating an Al-OH vibration liaison. The horizon beneath has the same absorption near 2.20 μm and an additional feature in the 3.40 to 3.80 μm spectral range diagnostic of a C-O$_2^-$ vibration liaison. This horizon can be interpreted as a mixture of Al-rich clay minerals and carbonates. It overlies a horizon having a pair of absorptions near 2.20 to 2.30 μm diagnostic of a combination of Al-OH and Fe-OH vibrations liaisons. It is frequently associated with an additional feature in the 3.40 to 3.80 μm spectral range. This horizon is interpreted as composed of Al,Fe-rich clay minerals mixed with carbonates when having a diagnostic feature near 3.40–3.80 μm. The horizon beneath has absorption near 2.27–2.32 μm diagnostic of (Fe,Mg)-OH vibration liaison and a feature near 3.40–3.80 μm. It is interpreted...
as a mixture of Fe,Mg-rich clay minerals with carbonates. Finally, the bottom horizon presents an absorption near 2.27–2.32 μm, which is interpreted as a horizon composed of Fe,Mg-rich clay minerals. In the Eridania Basin region, an additional bottom horizon can be present. It only presents a spectral shoulder near 2.50 μm in the 2.10 to 2.60 μm spectral range (only shown with Eridania Basin here but also described in Mawrth Vallis by Bishop et al., 2013). This can be interpreted as a sulfate or a zeolite signature. The top part of this horizon is associated with a feature near 3.40–3.80 μm, indicating the presence of carbonates.

Our investigation confirms the sequence of mineral assemblages described by previous authors (Carter et al., 2015). We additionally identified carbonates by the signature in the 3.40 to 3.80 μm spectral range in association with parts of the stratigraphic sequence. Indeed, the weathering profiles exhibit different horizons from top to bottom composed of Al-rich clay minerals, a mixture of Al-rich clay minerals and carbonates, Al,Fe-rich clay minerals mixed with carbonates, Fe,Mg-rich clay minerals mixed with carbonates, and finally, Fe, Mg-rich clay minerals.

The weathering profiles studied here are classified into three categories depending on their degree of preservation evaluated from a spectroscopic point of view. When exhibiting all the horizons, they are classified as a clear and complete stratigraphy (53 cubes). Because of erosion, some profiles exhibit only the bottom part of...
the stratigraphy (i.e., the top corresponding to Al-rich clay minerals without carbonate, a mixture of Al-rich clay minerals and carbonates, and Al,Fe-rich clay minerals mixed with carbonates is eroded). They are classified as having a partial stratigraphy (31 cubes). The third categories (60 cubes) concern the weathering profiles that do not exhibit a clear and complete or partial stratigraphy from a spectroscopic point of view (blue dots in Figure 8). The quality of these CRISM data allows us only to confirm the presence of Fe,Mg-rich clay minerals, although the morphology of the profile could indicate the presence of the top of the stratigraphy (Carter et al., 2015). In these cases, the signal was too noisy to allow a confident interpretation of the evolution of the profile mineralogy or any evidence of carbonate. Therefore, they were considered as not exhibiting any stratigraphy. However, the lack of confident detections of the top part of the stratigraphy or of the carbonate feature could be due to the low SNR.

Table 2 summarizes the results sorted by region, and a more detailed table sorted by CRISM cubes is presented in the supporting information (Table S1). In the supporting information, we describe in more details the different aspects of the profiles for each region and their degree of preservation. As shown in Figure 8, the first two categories are widespread on the surface of the planet and can be sorted into the following regions: North Hellas, North Argyre, Eridania Basin, Mawrth Vallis, Oxia Planum, Shalbatana

Figure 7. (a) CRISM data from FRT0000A2B8 for the North Hellas region. (b) CRISM data from FRT000092F7 for Eridania Basin. (c) CRISM data from FRT00003BFB for Mawrth Vallis. The reflectance between 1.00 and 2.60 μm is shifted compared with the 2.80 to 3.90 μm range for visual clarity. From top to bottom: black spectra correspond to unaltered material; red spectra, Al-rich clay minerals; purple spectra, Al-rich clay minerals and carbonates; blue spectra, Al,Fe-rich clay minerals and carbonates; cyan spectra, Fe,Mg-rich clay minerals and carbonates; green spectra, Fe,Mg-rich clay minerals; and brown spectra, sulfates and carbonates or zeolites and carbonates. Note that the color code used is the same as in Figures 8 and 9. The dashed lines indicate the position and interpretation of the absorptions (X corresponds to Fe or Mg) and are located at 1.39, 1.90, 2.19, 2.29, 2.50, 3.49, and 3.82 μm. The gray areas cover spectral ranges of strong atmospheric absorption near 2.60–2.80 μm and an instrumental artifact at ~3.2 μm. The site coordinates of spectra are given in Table S3.
Valley, Libya Montes, and South Coprates. Except in the region of Coprates Chasma, all the regions studied exhibit at least one example of clear and complete stratigraphy. The lack of CRISM data coverage and the low SNR of the data present in the Coprates Chasma region allow us to detect a partial stratigraphy. It appears that regions being both (i) the most covered by CRISM data and (ii) the most preserved are Mawrth Vallis, Eridania Basin, and North Hellas. In the following section, we focus on the three best preserved regions to describe in more detail the local variation in the mineralogy.

3.2. The Best Preserved Weathering Profiles

We report here the results on portions of three of the regions investigated. They represent the best preserved sites: North Hellas (i.e., Carter et al., 2015), Eridania Basin (i.e., Carter et al., 2015), and Mawrth Vallis (i.e., Loizeau et al., 2012). The spectra are summarized in Figure 7. In this section, the presence of carbonates is implied by the presence of the feature in the 3.40 to 3.80 μm range as explained in section 2. This feature consists of a high reflectance near 3.65 to 3.70 μm compared with the reflectance near 3.40 and 3.80 μm. The characterization of the clay minerals is based on the methodology described in section 2.3. We describe some individual variations in the mineralogy, as follows:

1. The North Hellas region (Figures 7 and 9): The carbonate feature (in the 3.40 to 3.80 μm range) is visible locally in the horizons composed of Al-rich, Al,Fe-rich, and Fe,Mg-rich clay minerals. For the unaltered rock below, the spectrum is featureless in the 2.80 to 3.92 μm range, implying the absence of carbonates.

<table>
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</tr>
<tr>
<td>Oxia Planum</td>
<td>1</td>
<td>3</td>
<td>3</td>
<td>7</td>
</tr>
<tr>
<td>Shalbatana Valley</td>
<td>3</td>
<td>0</td>
<td>0</td>
<td>3</td>
</tr>
<tr>
<td>Libya Montes</td>
<td>2</td>
<td>1</td>
<td>3</td>
<td>6</td>
</tr>
<tr>
<td>South Coprates</td>
<td>0</td>
<td>6</td>
<td>7</td>
<td>13</td>
</tr>
</tbody>
</table>
2. The Eridania Basin region (Figures 7 and 9): The sequence possesses a top horizon composed of Al-rich clay minerals. An Al,Fe-rich clay mineral horizon is occasionally present beneath, and the carbonate feature is associated in parts of this horizon. Below, an Fe,Mg-rich clay mineral horizon, exhibiting often the carbonates signature, is present. The bottom horizon, which is mixed with carbonates at its top, is composed of sulfates or zeolites. The unaltered surface with featureless spectrum is also visible (no carbonate features).

3. The Mawrth Vallis region (Figures 7 and 10): The top horizon composed of Al-rich clay minerals is better preserved and both laterally and vertically more extended compared with the other two sites. It overlies a horizon of Al-rich clay minerals mixed with carbonates. Below, the horizon composed of Al,Fe-rich clay minerals is mixed systematically with carbonates, while the horizon composed of Fe,Mg-rich clay minerals below is mixed with carbonates at its top only. This stratigraphic sequence is in many places still protected by a capping unit, likely responsible for the good preservation. This complete and most representative sequence seen at Mawrth Vallis contains carbonates in mixture in its middle part, leaving the top and bottom of the sequence without carbonates.

Of note, on all sites a nonaltered surface or horizon is present and shown for reference (Figure 7). Unlike the altered material, they neither exhibit the carbonate feature at 3.40–3.80 μm nor hydrated minerals in the 1.90 to 2.50 μm range.

Because the sites of North Hellas and Eridania Basin are not covered by a capping unit, they were less preserved. However, in some parts, it is still possible to visualize the sequence from Al-rich clay minerals to Fe,
Mg-rich clay minerals and carbonates in the middle part of the sequence as seen in Mawrth Vallis. Figures 9a and 9b illustrate a representative context of the North Hellas region. In this case, the weathering profile is present on the rim of a crater highly eroded. The highest part of the rim represents the top part of the weathering profile (highlighted in red with the color criteria used here). Figures 9c and 9d show the context of the detections in Eridania Basin. This area has several topographic depressions exhibiting on their walls the weathering profiles as described before. The altered material is present in these depressions and appears brighter than the surrounding surface. The top part of the depressions represents the top part of the weathering profiles (Al-rich clay minerals, in red with the color criteria used here). The bottom part of the weathering profile is composed of carbonates and zeolite or sulfate. It appears blue at its top only because the carbonate presence is limited in the top part of this horizon. The bottom part of this horizon is not highlighted in Figures 9c and 9d. The distinction between the hydrated sulfates and the zeolites spectra is not possible because they are often similar (Carter et al., 2015).

Morphological investigation of the sequences provides additional information for their formation as shown by the best preserved site (Mawrth Vallis; Figure 10). There is no unconformity visible between the altered horizons at the HiRISE submeter scale. Across the horizons, similar polygonal-shaped fractures are present, and their exposed surfaces appear brighter than do those of the unaltered unit. All altered horizons resemble each other in texture and display a gradual spectral change of the mineralogy across the sequence. The three globally distributed sites, described here in detail, exhibit a similar sequence from Al-rich to Fe,Mg-rich clay mineral zones without any unconformity between these horizons. The bottom of the Fe,Mg-rich clay mineral horizon represents the weathering front separating the entire weathering profile from the basement unit (Figure 11). This description is sketched with the stratigraphy presented in Figure 11 where local variations in mineralogy from sites to sites are not represented.

4. Discussion
4.1. Surface Weathering in a CO₂-Rich Environment
To better characterize the environment of weathering, we use here the mineral associations as a proxy and discuss the conditions of the system drawn from a terrestrial analog study (Gaudin et al., 2011), geochemical

Figure 10. (a) CTX mosaic on Mawrth Vallis with overlapping CRISM image FRT00003BFB. The white rectangle identifies the zoom in (b) and (c). (b) HiRISE image PSP0020742025 on Mawrth Vallis. (c) Same as (b) but overlaid by a color composite of CRISM image FRT00003BFB. (d) Schematic color code of the mineral mapping (red = Al-rich clay minerals; green = (Fe,Mg)-rich clay minerals; blue = carbonates) with the associated spectral parameters.
modeling (Zolotov & Mironenko, 2016), and laboratory experiments (Gaudin et al., 2018; Viennet et al., 2017).

Carter et al. (2015) suggest that weathering with humid climate is the best explanation for the sequence from Al- to Fe,Mg-rich clay mineral. Indeed, the absence of high-temperature minerals in weathering profiles suggests that hydrothermalism is unlikely to be responsible for the sequences. Moreover, the presence of the stratigraphy at elevated altitudes is inconsistent with lacustrine/basin or any detrital deposits. As shown in the analog study of Gaudin et al. (2011), the sequence systematically consisting of Al-rich clay minerals on top of Fe,Mg-rich clay minerals and the lack of unconformities between the horizons show that weathering of a single basaltic unit from top to bottom without periods of erosion/deposition is the most likely process. Moreover, this stratigraphy indicates an evolution of the pH from acidic to neutral/alkaline conditions from the top to the bottom (Gaudin et al., 2011 and references therein; Zolotov & Mironenko, 2016). Furthermore, carbonate precipitation only in part of the profiles can be explained by the pH increasing from the top to bottom of the weathering profiles as suggested by geochemical models (Zolotov & Mironenko, 2016).

On Earth, pedogenic carbonates are common in arid to subhumid environments (Landi et al., 2003). Interestingly, carbonates often form from calcareous parent rock but are not restricted to this environment (Doner & Lynn, 1989). For the formation of carbonates, cations (mostly calcium) are provided by dissolution of minerals and meteoric water percolation (Doner & Lynn, 1989). On noncalcareous soils, carbon is supplied by vegetation, microbes, and atmosphere (Cerling & Quade, 1993). Precipitation and dissolution of carbonates in soils is related to the pCO₂ and water-rock ratio of the system (Doner & Lynn, 1989). The latter is controlled by the climate (temperature and water availability) and the capacity of the water to interact with the protolith (e.g., porosity and slope; Doner & Lynn, 1989). In arid climate, the carbonates are mostly present in the top horizon of the profiles, while in humid climate, the carbonates are dissolved in the top part and occur deeper in the profiles. In general, the more the leaching is important, the deeper the carbonates will be. In case of very strong leaching, carbonates can be completely removed from the profiles. Several parameters between current Earth and early Mars are extremely different. First, these weathering profiles are developed in volcanic rocks, not calcareous soils. Second, no vegetation is present on Mars, and the question of the presence of microbes remains hypothetic. These are two reasons to have less carbon in the system. Third, the tens of meters of some weathering profiles is not consistent with arid to subhumid environment. However, the high variation of thickness from one profile to another could reflect a variation of the mean precipitation. Finally, it is unlikely that the atmosphere to consider is the same. Hence, no comparison can be made with current soil formation on Earth. Elements of answer could come from weathering profiles formed on early Earth. However, paleoweathering profiles on Earth often suffered from later alteration (diagenesis, for example; i.e., Nesbitt & Young, 1989). For these reasons, comparing paleoweathering profiles on Earth and on Mars is not straightforward. Therefore, the sole presence of carbonates in weathering profiles on Earth and Mars cannot be directly compared. We suggest here to conduct the discussion by using the mineralogical associations.

As explained in Zolotov & Mironenko (2016), the initial pH and the composition of alteration solution can be assessed from minerals associated with phyllosilicates (e.g., sulfates, amorphous silica, and carbonates). Concerning the sulfates, their presence cannot be confirmed as a bottom layer here because of the similitude of the sulfate and zeolite spectra (see section 2.3 and Carter et al., 2015). Indeed, as explained in Carter et al. (2015), both zeolite and sulfate could have formed during the weathering profiles formation, either by precipitation or by accumulation. The bottom layer (of zeolite/sulfate mixed with carbonate) is not systematically present in all the weathering profiles studied here. However, sulfates, such as bassanite (Ca-sulfate; Wray et al., 2010) or jarosite (Fe-sulfate; Farrand et al., 2009, Pajola et al., 2016), have been detected locally in some of the weathering profiles (e.g., in Mawrth Vallis and Eridania Basin). None of these detections

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**Figure 11.** Sketch of the stratigraphic column (not at the scale) of the weathering profile. The dotted line indicates the weathering front between the Fe,Mg-rich clay minerals horizon and the basement. The black arrow symbolizes the progression of the weathering front.
suggest that sulfates are present globally and systematically in one or multiple horizons of the weathering profiles. Amorphous silica is not clearly identified in this study but is suspected to be present at least in the Al-rich clay mineral horizon following previous studies (i.e., Bishop et al., 2013). Regarding the carbonates, we report here their presence in the middle part of the stratigraphy. In their models, Zolotov & Mironenko (2016) show that carbonates can form under a wide range of pCO₂ and that the pCO₂ can only be determined due to the identification of the carbonate compositions and their amount. However, Zolotov and Mironenko (2016) show that more carbonates form at higher pCO₂ and that the detectable amount of carbonates in the Martian weathering profiles suggests an elevated pCO₂. Zolotov and Mironenko (2016) suggest that minor carbonates precipitate at depth from solution formed through neutralization of initially very low pH in a CO₂-H₂SO₄-HCl-rich fluid. The coformation of clay minerals and carbonates is possible when the solution becomes neutral. Ca-sulfates are predicted to form in the middle part of the profile, and Mg-sulfate solutions are predicted to form in deeper part of the profile when the pH becomes neutral. Because none of the horizons in the weathering profiles are systematically composed of sulfates, the presence of sulfate ions might not be systematic in the formation of the weathering profiles. However the widespread presence of carbonates in the middle part of the profiles can suggest the presence of carbonate/bicarbonate ions in the alteration solution. Therefore, we suggest that the formation of the weathering profiles involved at least the presence of carboonic acid in the starting solution and possibly local presence of H₂SO₄ where sulfates are detected.

The experimental study of such system is challenged by the time needed to produce a significant amount of altered material. Nevertheless, a few studies have been performed (Gaudin et al., 2018, and references therein). Gaudin et al. (2018) reported experimental results of olivine weathering under a high-pCO₂ atmosphere (1 bar of pCO₂ and between 15 and 40 °C). The experiment in a high-pCO₂ environment leads to the formation of amorphous silica, Al,Fe,Mg-rich clay minerals, and carbonates. Although at a high temperature, Viennet et al. (2017) experiments on basalt alteration in high-pCO₂ environment demonstrate that depending on the pCO₂, and its subsequent pH, the alteration of a basaltic rock can lead to different associations of diverse Al,Fe,Mg-rich clay minerals, carbonates, and zeolites. Finally, experimental results indicate that alteration in a high-pCO₂ environment leads to the coprecipitation of clay minerals and carbonates. It also suggests that resulting mineralogical associations in such system is dependent on the pH induced (at least partly) by the pCO₂. The previous idea why the carbonate was absent leads to suggest sulfuric acid as an acidic source (with low pCO₂) for the formation of the weathering profiles (Zolotov & Mironenko, 2016). However, the presently reported widespread presence of carbonates in the stratigraphy indicates that the alteration solution contained inorganic carbon. Hence, the weathering sequence is coherent with a chemical weathering involving inorganic carbon. Finally, taken together, experimental and modeling results tend to indicate an evolution from the top to bottom of the profiles of the dissolved inorganic carbon species and of the resulting pH. Note that the initial (meteoritic water) pH and the solution composition could also have been controlled by other strong acids such as HCl and H₂SO₄ at least locally. While this is coherent with an atmosphere wetter and denser in CO₂ than today, the undetermined species and amount of carbonates do not allow estimating the pCO₂. Moreover, it does not exclude the presence of strong acids (HCl and H₂SO₄) in meteoric waters.

4.2. A Planetary-Scale Process

In the following paragraph, we discuss the potentially planetary-scale character of the weathering profiles formed by fluids rich in carbonate/bicarbonate ions. Carter et al. (2015) have described that the process leading to the formation of the weathering profiles (i.e., composed of Al-rich clay minerals overlying Fe, Mg-rich clay minerals) was a planetary-scale phenomenon. However, our study describes the presence of the weathering profiles containing carbonates in specific regions that are less widespread than the profiles described in Carter et al. (2015; Figure 7). Several facts can explain the distribution of the weathering profiles and an underestimation of the number of clear and complete stratigraphies: (i) First, the limitation of the coverage of high-resolution CRISM data acquired before 2010 reduces our capacity to fully investigate all the weathering profiles described in Carter et al. (2015). (ii) An important part of the CRISM data cubes investigated in this study had a too low SNR to allow visualizing clearly a spectral gradual change from the top to bottom of the stratigraphy (Table 2). (iii) The presence of dust globally on the planet can hinder the investigation of the weathering sequence. (iv) The resurfacing by volcanic activity on the planet can have a buried part of the
weathering profiles hiding them from remote sensing investigation, and we rely on erosion windows to detect and investigate the sequences. (v) The top layers of Al-rich clay minerals, Al-rich clay minerals with carbonates, and Al,Fe-rich clays minerals with carbonates are generally thinner than the rest of the profiles. Hence, such horizons are more easily impacted by erosion, which explains the high proportion of partial stratigraphy present in some regions (Table 2). While, for example, Mawrth Vallis profiles were protected by a capping unit, this was not the case for other regions such as North Hellas where impact craters and erosion reduce the capacity to preserve a clear and complete stratigraphy. Finally, the overall widespread distribution of the profiles suggests a planetary-scale weathering with a fluid rich in dissolved inorganic carbon species and possibly locally together with other strong acids (HCl and H₂SO₄) that affected the surface of Mars at least at the midlatitudes.

4.3. Implications for the Geological Evolution of Mars

Although we present here indications of a planetary-scale weathering related to a fluid rich in carbonate/bicarbonate ions, its coherence with regard to our present knowledge of the geological evolution of the planet should be addressed. The presence of a nonhydrated capping unit dated from around 3.7 Gyrs (Loizeau et al., 2012) allows dating the weathering process as a Noachian environment. It coincides well with morphological studies suggesting the Noachian as the most efficient period of water-related resurfacing processes (Quantin-Nataf et al., 2019; Werner et al., 2011). The presence of carbonates associated with planetary-scale weathering during its wetter period coincides well with climate models suggesting a denser CO₂-rich atmosphere as a part of explanation for adequate conditions to have liquid water at the surface of Mars during the Noachian (Wordsworth et al., 2017). However, the sole detection of carbonates in the weathering profiles does not imply a pCO₂ as high as what is suggested in Wordsworth et al. (2017). More in-depth investigation is needed to determine precisely the composition and amount of the carbonates to suggest any pCO₂ values. Alternatively, the (late) presence of liquid water could be punctual and related to volcano or impact-generated climate optima (i.e., Wordsworth et al., 2013). We suggest that the observation of carbonates in these weathering profiles is a new important step to reconcile climatic models with the mineralogical and morphological interpretations. However, the debate of episodic periods of warm enough environment to sustain liquid water versus a permanent warm period over the entire Noachian cannot be solved here and neither is the reason for the end of the warm and wet conditions.

4.4. Implications for Habitable Conditions on Mars

Of note, we show that the weathering profiles constitute environments with diverse conditions (composition, pH, and pCO₂). Such chemical heterogeneity alerts interest for astrobiology as it encompasses both a diversity of potentially habitable environments and chemical gradients that are essential for microbial life metabolism (Cleaves, 2013). Moreover, an analog study on carbonates in the Mojave Desert by Bishop et al. (2011) suggests that carbonate rocks can provide protection of organic matter from solar radiation. However, weathering profiles are subject to loss of nutrients by leaching in the weathered part of the soil, and this could reduce their habitability potential (Duddy, 1980; Nesbitt & Markovics, 1980). On the other hand, weathering profiles are known to have hosted life early on Earth and can be considered as of interest for astrobiology (ca. 3 billion years ago; Lenton & Daines, 2017).

5. Conclusion

Using a new spectral criterion, we mapped the presence of carbonates mixed with various clay minerals in weathering profiles. We used the particular variation of the mineralogical assemblages (i.e., from top to bottom: Al-rich clay minerals, a mixture of Al-rich clay minerals and carbonates, Al,Fe-rich clay minerals mixed with carbonates, Fe,Mg-rich clay minerals mixed with carbonates, and finally, Fe,Mg-rich clay minerals) as a proxy to apprehend the conditions of the water-rock interactions. Our observations indicate that the profiles formed by a weathering containing inorganic carbon (and possibly strong acids) with a pH solution gradually increasing with depth within the sequence toward neutral/alkaline values on a planetary scale. Based on comparison with geochemical models, we suggest an elevated pCO₂ to lead to the mineralogical sequence observed.

Future in situ investigation of the best preserved weathering profiles at Mawrth Vallis or on the margins of Eridania Basin should provide unprecedented details on the atmosphere composition and more specifically
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ancient atmospheric CO2 pressure, as well as the composition and evolution of the aqueous solution required to form these mineralogical associations.

References


