

Amorphization of S, Cl-salts induced by Martian Dust Activities

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Key points

- Amorphization of S and Cl salts was induced by electrostatic discharge (ESD) in a Mars chamber that simulates martian dust activities.
- Amorphization is commonly accompanied by dehydration of salts and the oxidation of Fe, Cl, and S species.
- Dust activities may have generated and deposited large quantities of S- and Cl-rich amorphous materials all over the martian surface.

Abstract

A high proportion of non-crystalline (X-ray-amorphous) components has been found in all samples analyzed by CheMin on the *Curiosity* rover at Gale crater on Mars, and such X-ray-amorphous components probably occur at all sites that have been investigated thus far by landers and rovers. The amorphous material at Gale crater is rich in volatiles (S, Cl, and H₂O), as indicated by other science payload elements (APXS, SAM). We demonstrate here that amorphization of S and Cl salts can be induced by energetic electrons and free radicals generated in a medium-strength electrostatic discharge (ESD) process during martian dust activities such as dust storms, dust devils, and grain saltation. Furthermore, we found that the amorphization is commonly accompanied by dehydration of the salts and oxidation of Cl, S, and Fe species. On the basis of experimentally observed rates of the above phase transformations and the mission-observed dust activities and wind speeds on Mars, we anticipate that similar phase transformations could occur on Mars within a time frame of years to hundreds of years. Considering the high frequency, long duration, and large areal coverage of Martian dust activities, our study suggests that the ESD induced by Martian dust activities may have contributed to some the S- and Cl-rich portion of X-ray amorphous materials observed in surface soils at Gale crater. Furthermore, dust activities in the Amazonian period may have generated and deposited a significant quantity of S- and Cl-rich amorphous materials all over Mars.

Plan language summary

Martian dust activities have been altering the look of the surface of Mars, by the blast of particles of various sizes. In addition, frictional electrification in these events would charge the particles, and the charged particles would be further spatially separated by wind at the same time, to form an electric field like those formed in the cloud layers on Earth. Electrostatic discharge (ESD) would occur when enough charges are accumulated. Unlike the lightning on Earth, another two types of ESD, Townsend dark discharge (TDD) and normal glow discharge (NGD) would more likely occur on Mars because of its thin atmosphere. The electrostatic discharge (ESD) would induce electrochemical reactions and change Martian surface materials. Our new simulated ESD experiments revealed three types of phase changes can occur in S and Cl salts. They are amorphization (damage of crystal structure), dehydration (loss of structural water), and oxidation of Cl, S, and Fe. Because of the high occurring frequency, the large area coverage, and long duration of dust events during the recent Amazonian period on Mars, our results imply that Martian dust activities may have generated and deposited a large quantity of S- and Cl-rich amorphous materials all over the surface of Mars.

1. Introduction

Among many great findings in martian mineralogy, the discovery of X-ray amorphous components in all samples analyzed by CheMin on the *Curiosity* rover at Gale crater (Bish et al., 2013, Blake et al., 2013, Vaniman et al., 2014) has been an eye-opening discovery. The proportion of X-ray amorphous components in different martian samples ranges from ~19-36 wt% in active and inactive dune materials, to ~20-56 wt% in all mudstones, and ~14-71 wt% in altered and less-altered Stimson formation samples (the lowest estimated percentages based on Table 1 and Table 10 of Morrison et al., 2018a, which were supported within the uncertainty range of measurements and analyses by Achilles et al., 2017; Morris, et al., 2016; Rampe et al., 2017, 2018; Yen et al., 2017), implying multiple geological processes for producing the X-ray amorphous components.

In the early history of Mars, several geological processes would form species with low crystallinity. These processes include volcanic activity, impacts, hydrothermal activity, and chemical (including acidic) weathering at low temperature that could free (or partially free) molecules or ionic groups in geological materials, but with insufficient time (or energy) for the newly formed phases to reach crystallographic equilibrium, i.e., an ordinary degree of crystallinity.

A phenomenon with equal significance was that the X-ray amorphous components in all Gale crater samples have a high concentration of volatile-element components (e.g., SO_3 and Cl), a conclusion from combined CheMin and APXS data analyses (Dehouck et al., 2014; Morris et al., 2016; Rampe et al., 2017, 2018; Yen et al., 2017; Achilles et al., 2017), based on a newly developed method to refine unit-cell parameters that has increased the accuracy in derived major-mineral chemistry (Morrison et al., 2018b). An overview published by Morrison et al. (2018a) revealed the highest concentration of $\text{SO}_3 + \text{Cl}$ in the amorphous component of a surface soil (Rocknest), among all 13 samples from Bradbury landing through Naukluft Plateau (from sol 69 to sol 1332) at Gale crater, to be 24.4 wt% for SO_3 and 3.5 wt% for Cl. In addition, the data from the SAM payload on the same set of collected samples support the existence of poorly crystalline magnesium and iron sulfates and the association of water with amorphous phases (Sutter et al. 2017, 2019). A relevant observation made by the *Spirit* and *Opportunity* rovers at Gusev crater and Meridiani Planum, was that a higher content of SO_3 and Cl were found in surface soils and unbrushed rock surfaces than in rock interiors (Yen et al. 2005; Gellert et al., 2006). The S and Cl enrichment nature and the similar molar S/C ratio in air-fall dust were further confirmed by APXS analyses of the *Curiosity* rover, and were implied to relate to global Martian dust (Berger et al., 2015, Schmidt et al., 2018). A key follow up question is: By what processes could some of the S- and Cl-bearing salts at the martian surface become (or form as) X-ray amorphous materials?

A sudden exposure (by impact or by rover trench, e.g., Byrne et al., 2009; Wang et al., 2006a) of subsurface hydrous sulfates to current atmospheric conditions at the martian surface, as simulated by vacuum desiccation of hydrous salts in laboratory experiments (Sklute et al., 2015; Vaniman et al., 2004; Wang et al., 2006b, Wang and Zhou 2014), could have formed amorphous sulfates directly from crystalline Mg, Fe^{2+} , and Fe^{3+} sulfates. Another process, a sudden release of subsurface brine(s) to the current martian surface with subsequent desiccation may also form non-crystalline salts. Geological processes on present-day Mars that might induce subsurface brine release could be Recurring Slope Lineae (RSL) (McEwen et al., 2014, Wang et al., 2019) and impacts. In a fast brine-dehydration laboratory simulation, amorphous ferric sulfates were first found to form at mid to high temperatures (293-323 K) (Sklute et al., 2015; Wang et al., 2012), and amorphous Mg, Fe^{2+} , Fe^{3+} (but not Ca, K, Na) sulfates formed at 77 K (Morris et al., 2015). Sklute et al. (2018) further revealed the formation of amorphous phases from pure FeCl_3 brine (but not from pure CaCl_2 , MgCl_2 , and NaCl brines), and from the brines of mixed salts, i.e., $\text{Fe}_2(\text{SO}_4)_3$ mixed with Na, Mg, Ca, Fe^{3+} chlorides and Na bicarbonate, at room temperature. Toner et al. (2014) observed that amorphous glasses of $\text{Mg}(\text{ClO}_4)_2$ and $\text{Ca}(\text{ClO}_4)_2$ formed near 153 K by cooling the relevant brines below their eutectic temperatures. Furthermore, both

amorphous Mg and Fe³⁺ sulfates can host structural H₂O to various degrees (up to three structural H₂O per Mg-sulfate molecule, and up to eleven structural H₂O per Fe³⁺-sulfate molecule), all of which are stable at low relative humidity (RH <11%) and in a wide temperature range (278-323 K) (Wang et al., 2009, 2012).

In addition, energetic particles from space, such as galactic cosmic rays and energetic UV photons, are capable of damaging the crystal structures of surface minerals on airless planetary bodies, while their effect on martian secondary minerals at the surface needs further investigation.

On the basis of the studies referenced above and our previous experimental investigations (Wu et al., 2018, Wang et al., 2020), we hypothesize that S- and Cl-rich X-ray amorphous materials at the surface of Mars may be very common, and one mechanism to produce them is by *multiphase redox plasma chemistry* (or simply, *electrochemistry*) induced by electrostatic discharge (ESD) that occurred during martian dust activities during the Amazonian period. In this manuscript, we report the results of 75 sets of ESD experiments on 22 Mars-relevant minerals under martian atmospheric conditions to explore and test our hypothesis.

We present some background about ESD in martian dust events in section 2, the studied samples and the experiments in section 3, and the results in section 4. We then report the finding of three phase transformation trends in section 5, and discuss the implications of our study in section 6.

2. Martian dust activities, ESD, free radicals, and electrochemistry

Frictional electrification of mineral particles and aerosols can occur in four types of martian surface processes: volcanic eruption, dust storm, dust devil, and grain saltation. Except for volcanic eruptions, the last three processes occur continuously on present-day Mars. For example, regional dust storms occur every martian year and a global dust storm occurs every 6-8 Earth years (Shirley, 2015; Wang and Richardson, 2015). Dust devils have been observed by all landed missions on Mars (Metzger and Carr, 1999; Ferri et al., 2000; Ellehoj et al., 2010; Greeley et al., 2006, 2010; Lemmon et al., 2017; Murphy et al., 2016), as well as remotely by orbital observations (Cantor et al., 2006; Choi and Dundas, 2011; Reiss and Lorentz, 2016; Verba et al., 2010; Whelley and Greeley, 2008). Grain saltation was first confirmed on Mars at Meridiani Planum and Gusev Crater (Sullivan et al., 2005, 2008), with laboratory simulations suggesting that they could be a ubiquitous occurrence on the martian surface (Sullivan and Kok, 2017).

A tendency for triboelectric charge, i.e., generation of negative charges on smaller grains, but positive charges on larger grains of similar composition, was revealed by experiments (Forward et al., 2009; Krauss et al., 2003). Separation of smaller grains from larger grains by convective martian dust events would generate a large-scale charge separation, i.e., *an active electric field (E-field)*. On Earth, E-fields of up to 166 kV/m was detected during grain saltation (Schmidt et al., 1998), and 60 kV/m during the passage of dust devils (Esposito et al., 2016; Farrell et al., 2004; Harrison et al., 2016; Jackson and Farrell, 2006).

When a local E-field accumulates beyond the *breakdown electric field threshold* (BEFT), electrostatic discharge (ESD) can occur. BEFT on Mars is estimated to be ~ 25-34 kV/m from measurements in Mars environmental chambers (Farrell et al., 2015; Yan et al., 2017), slightly higher than ~ 20-25 kV/m from a modeling study (Melnik and Parrot, 1998). This range is about 1% of the BEFT on Earth (~ 3000 kV/m), which matches with martian atmospheric pressure being < 1% that of Earth. Therefore, ESD occurs on Mars much more easily than on Earth. The low BEFT on Mars prevents the accumulation and separation of large amounts of charges, thus lightning would be unlikely to occur. Whereas the other two types of ESD (Gallo, 1975), *Townsend dark discharge (TDD)* and *normal glow discharge (NGD)* would more likely occur on Mars. In a set of experiments simulating grain saltation using silicates under Mars atmospheric composition and pressure, Bak et al. (2017) detected light emissions (red colored glow from

quartz sand and blue colored glow from basaltic sand) that are similar to normal glow discharge (NGD) in a Mars chamber simulated in our laboratory (Wang et al., 2020; Wu et al., 2018).

ESD generates a flux of energetic electrons with high speed, i.e., an electron avalanche. These electrons collide with gaseous molecules and atoms in the martian atmosphere, CO_2 , O_2 , N_2 , Ar, and H_2O , to cause electron impact ionization (EII) of CO_2 , electron/ CO_2 dissociation attachment (ED of CO_2), and electron/ H_2O dissociation attachment (ED of H_2O) (Jackson et al., 2010; Wu et al., 2018). The collision generates free radicals, including ions with positive and negative charges, neutral species at excited states, and additional electrons. The last one could cause further chain electron avalanches (Delory et al., 2006).

Above expectations were validated by our experimental observations (Wu et al., 2018) of CO^{2+} , CO^+ , O , H_{III} , H_{II} , OH , Ar, N_2 , N_2^+ by *in situ* plasma emission spectroscopy (also O_2 , NO , and O^+ because of the overlapping of plasma lines used for detection, Figure S8) and O_3 in the output gas by UV and mid-IR spectroscopy (Figure 5 of Wu et al., 2018), generated by normal glow discharge (ESD-NGD) under simulated Mars atmospheric composition and pressure. During a simulated saltation experiment on silicates upon contact of water (Bak et al., 2017), H_2O_2 and $\cdot\text{OH}$ were detected. Furthermore, Aerts et al. (2015) reported CO_2 splitting (to CO and O_2) by dielectric barrier discharge.

These energetic free radicals could react with the molecules in the martian atmosphere and in surface materials. As demonstrated by our previous work, these *multiphase redox plasma chemical reaction* (or simply *electrochemical reaction*) cause the oxidation of chlorine from chloride (Cl^{1-}) to chlorate/perchlorate (Cl^{5+} and Cl^{7+}) (Wu et al., 2018), and the release of Cl atoms at the first excited state (Cl_I) from common chlorides (Wang et al., 2020), instantaneously and apparently with high yields.

3. Samples and Experiments

3.1 Sample selection

In order to validate our hypothesis, i.e., to make our experiments relevant to the volatile portion (high wt% of $\text{SO}_3 + \text{Cl}$) of the X-ray amorphous component found on Mars (Dehouck et al., 2013; Morris et al., 2016; Morrison et al., 2018a; Rampe et al., 2017), we selected crystalline sulfates and chlorides as the starting phases for our ESD experiments. For the most part, hydrous salts were used, based on the association of H_2O with amorphous phases suggested by data analyses of SAM (Sutter et al., 2017, 2019). Salts with Mg, Fe^{2+} , Fe^{3+} , Ca, Al, Na, K were selected, based on the findings by recent missions of martian sedimentary minerals, i.e., they are enriched in Mg and Fe, less in Ca, and relatively depleted in Na and K (McLennan and Grotzinger 2008; McLennan, 2012; McLennan et al., 2019). A few relevant samples were added later, including two Na-sulfites, an Fe-sulfide (pyrite), and a Fe-hydroxide (akaganeite). These starting phases are listed in Table 1.

Each starting sample for an ESD experiment was ground and sieved. A grain size range of 63-88 μm was selected for all samples. Each powdered sample was placed into a fused- SiO_2 cell with an inner diameter of 22 mm and inner depth of 2 mm, i.e., about 760 mm^3 in volume. Depending on the density of different salts, a total mass of different salts in the range of 400-1100 mg was used. During each ESD experiment (Figure 1), the SiO_2 sample cell was placed in the lower electrode (Figure 1b), facing the energetic electrons from the upper electrode, and was entirely enveloped by the generated plasma (inset of Figure 1).

3.2 ESD experiments

A Planetary Environment and Analysis Chamber (PEACH) at Washington University in St. Louis (Sobron and Wang, 2012) was used for all ESD experiments designed for this study. A combination of needle and ball valves connect the PEACH with a CO_2 gas tank to regulate atmospheric pressure (and composition

when desired). The PEACH was first evacuated to 3×10^{-2} mbar to remove the air, and then filled with pure ultra-dry CO₂ for this study. The ESD experiments of this study were conducted with active evacuation and continuous CO₂ in-filling. The pressure was kept at 3 ± 0.1 mbar in each ESD experiment.

As described by Sobron and Wang (2012), the temperature of the sample cell can be controlled by a liquid nitrogen (LN₂) delivery system attached to the PEACH (Figure 1). The LN₂, stored in a dewar outside the PEACH, is heated by a resistor immersed in the LN₂ reservoir, and the evaporated N₂ gas at near-LN₂ temperature is directed via a feedthrough in the PEACH's wall into a toroid-shaped, double-walled copper block (referred to as *the cold plate* hereafter) that sits inside the PEACH. Another feedthrough allows for the evacuation of the nitrogen gas after its circulation through the cold plate. An electronic controller (OMEGA Engineering Inc. CN76000 autotune controller) monitors the temperature of the cold plate via a resistive thermal device and regulates the flow of cold N₂ gas that enters the cold plate. This setup allows the temperature of the cold plate to be kept relatively constant at a desired temperature between 21°C and -100°C with deviations of less than 0.5°C.

The normal glow discharge (NGD) in this set of ESD experiment was generated by two parallelly mounted electrodes in the PEACH (Fig. 1b). They are made of copper, with a diameter of 35 mm. The distance between the two electrodes was adjusted by a motorized precision translation stage (Thorlab PTI-Z8), 6 mm was used in all experiments of this study.

We used AC power (110 V, 50/60 Hz), a contact voltage regulator (No. 2090 VR), and a triggering neon power supply (CPI Advanced Inc., CPI-EZ12, max output 12 kV, 40 mA) that was directly connected to the ESD electrodes in the PEACH. Electric voltage from the upper electrode to the ground, and electric current through the pair of ESD electrodes were recorded every 30 minutes during an ESD experiment, using two multimeters (KEYSIGHT-U1251B). We found the values of electric current is ~ 22 mA, not affected by different starting salts, and is very similar to those reported in Wu et al. (2018) and Wang et al. (2020), i.e., with the same electron flux shown in Table S1.

The electron flux in our current experimental setting is $1.42 \times 10^{20} \text{ s}^{-1} \text{ m}^{-2}$, calculated from the electric current measurement (Table S1), indicating that we used an ESD with a strength midway between the two extreme cases, ESD-TDD and ESD-NGD (ranging from $9 \times 10^{16} \text{ s}^{-1} \text{ m}^{-2}$ (TDD) to $1.5 \times 10^{24} \text{ s}^{-1} \text{ m}^{-2}$ (NGD), detailed discussion in section 5.3 of Wu et al., 2018) to process all selected starting phases. Another important parameter is the kinetic energy of ESD generated electrons, for which we use the observed free radicals and relevant reactions to make estimation. For example, CO₂⁺ was the dominant species generated by the same ESD setup in CO₂ (Figure 4a, b, c, and Table 1 of Wu et al., 2018), and it is a product of electron impact ionization (EII) of CO₂ (Delory et al., 2006; Jackson et al., 2010). The occurrence of EII of CO₂ in our ESD experiment revealed that a considerable portion of ESD generated electrons has a kinetic energy >14 eV. Furthermore, a very strong H α line at 656.3 nm was observed by *in situ* plasma spectroscopy at extremely low P_{H2O} (Fig. 4b, d, e of Wu et al., 2018) or when the starting mineral is hydrous salt. This line is generated by a transition from H_{III} to H_{II} that indicates the presence of electrons with a kinetic energy > 17.19 eV to excite hydrogen to H_{III} (Delory et al., 2006, Itikawa and Mason 2005).

Specifically for this investigation, we used the cold plate to control the temperature of some starting salts during ESD so that is was lower than 30°C to avoid sample melting (melting point (MP) in Table 1, based on Lide 2001). The cold plate in the PEACH is electrically grounded, as well as the PEACH itself. On the other hand, the lower electrode must be isolated electrically from the cold plate; this was satisfied by using a Teflon holder between the lower electrode and the cold plate. The holder is thin enough (1.59 mm) to allow the temperature (T) of the lower electrode to be thermally controlled by the cold plate. A

thermocouple was inserted into a tunnel of 0.79 mm diameter in this Teflon holder to measure the T of the lower electrode while keeping its electric isolation from the cold plate.

During an ESD process in the PEACH, the equilibrated temperature T_{eq} of the lower-electrode sample cell was normally reached after > 30 minutes. For most hydrous S and Cl salts with melting-point temperatures above 130°C, we do not use LN₂ cooling. The resulting T_{eq} is in the range of 80-105°C (Figure 2). For a few selected salts (e.g., FeSO₄·7H₂O and some hydrous chlorides, Table 1), the ESD experiments were run using LN₂ to cool the cold plate and generate a T_{eq} in the range of 10-30°C, with the exact T_{eq} value depending on the type of salt (Figure 2). Those experiment products are marked as LT (low temperature) –ESD in figures.

3.3 Analysis methods of ESD reaction products

Almost all sulfates and chlorides show a color change after an ESD process of certain time duration (e.g., ≥ 7 hours, Table 1) under Mars conditions. For example, the photos in Figure 3 (a, b) are MgCl₂·6H₂O before and after 7h ESD, shown with different magnifications (Figure 3b, c, d). The color change only appears at the surface of ESD products, that is consistent with our previous finding of surface enrichment of ESD-generated species (NaClO₃ and NaClO₄, when using NaCl as the starting phase) based on Ion Chromatography (IC) analyses of the layers in the ESD-product (Figure 6b of Wu et al., 2018). Both color change and IC data suggest that the electrochemical reaction induced by ESD in a Mars chamber is an atmosphere-to-surface interaction, which also means the *as is* surface of an ESD product should be the best sampling site for the characterization of new species generated by ESD.

We used laser Raman spectroscopy (Raman), X-ray diffraction (XRD), Vis-Near-IR spectroscopy (VNIR), Mössbauer spectroscopy (MB), and Ion Chromatography (IC) to characterize the ESD products in this study. Among them, Raman and VNIR measurements were made on the *as is* surface. In particular, all micro-beam Raman analyses were made directly on the spots that show color changes, which were selected under the microscope of our Raman system. On the other hand, the field of view (FOV) of the VNIR probe matches well with the size of the SiO₂ cell that contains the ESD product, so a VNIR spectrum was taken from the whole *as is* surface of the ESD product. For XRD and Mössbauer analyses, the bulk powder sample from the full depth of the SiO₂ cell of an ESD product was re-ground and used. Bulk powder samples were also used for IC analyses. All these analyses were taken at room temperature.

Raman spectra of all samples were collected using a Renishaw inVia Raman system, with 532 nm excitation wavelength, spectral range of 50-4300 cm⁻¹ and spectral resolution better than 1 cm⁻¹. A 50× long-working-distance objective was used that generates an ~ 1 μm beam diameter at laser focus. Raman analysis of each ESD product was always taken on multiple spots in several areas at the *as is* surface of a sample, with a total of > 30 (at least) spectra per sample. A laser beam energy of 5 mW was normally used for the measurements of ESD products from Mg, Ca, and Na salts. A much lower laser energy (0.5-0.05 mW) was used on the ESD products from Fe-bearing salts. The Raman spectrometer is calibrated during each working day to keep the accuracy and precision of Raman peak positions within ± 0.5 cm⁻¹.

The VNIR spectra of the ESD products were acquired using Analytical Spectral Devices (ASD) FieldSpec4 spectroradiometer (Malvern Panalytical Company) with a contact probe. The obtained spectrum covers a wavelength range of 0.35 to 2.5 μm, with spectral resolution of 3 nm @ 700 nm and 10 nm @ 1400/2100 nm. A Spectralon target was used for absolute reflectance calibration before the sample measurements. The recording time of each spectrum was 1 second, and at least two spectra were taken from each sample for redundancy.

XRD measurements of ESD products were made using a Bruker D8 Advance diffractometer, with CuKα radiation ($\lambda = 1.54052 \text{ \AA}$) at 40 kV and 40 mA and a collecting angle of 3°. Each sample was ground

again to fine powder, and was put into a MTI zero background silicon holder. The XRD measurement routine was 0.02° step size, 1 second dwell time, and 15 rotation per minute, and 4° to 60° 2θ range. The Bruker XRD has a guaranteed calibration that is confirmed during installation and monitored by analysis of a NIST SRM 1976a Al₂O₃ standard. The alignment is also guaranteed and has been demonstrated to be within 0.03° 2θ of the absolute peak position of the NIST SRM.

Mössbauer spectral measurements were made on the ESD products from a few Fe-bearing phases. A plastic washer with an inner diameter of 1.2 cm was used to hold a mixture of sample plus sugar with Kapton® polyimide tape on either side. A source of ~80 mCi ⁵⁷Co in Rh on a SEE Co. (formerly WEB Research Co.) model WT302 spectrometer at Mount Holyoke College was used. Following standard practice, Compton scattering of 122 keV gammas caused by electrons inside the detector were measured with and without a 14.4-keV Al foil stop filter in the gamma beam. Absorption was corrected Compton scattering by dividing the uncorrected absorption by one minus the Compton fraction. This correction allows accurate determination of % absorption in the spectra without affecting the fits. The range of energy deposited in the detector by Compton events extends from 0 keV to 40 keV, and overlaps both the 14 keV and 2 keV energies deposited by the 14 keV gammas, thus requiring correction.

Mössbauer spectra were acquired in 1024 channels at 295K over 24 hours for each sample. Interpolation to a linear velocity scale was used to correct for non-linearity using the spectrum of a 31 mm Fe foil.. The WMOSS Auto-fold procedure fits a straight line to points at the published values of the Fe metal and the observed positions in channels (*x* values), then folds each spectrum around the channel value that minimizes the least squares sum difference between the first and second halves of each spectrum.

Ion chromatography was used to quantify the SO₄ production from ESD experiments using Na₂SO₃ and NaHSO₃ as starting salts. A 15-20 mg homogenized sample was dissolved in N₂-purged milliQ water for IC analysis. An A-Supp7-250 anion column (45°C, 3mM Na₂CO₃ eluent, 0.8mL/min, with suppression) on a Metrohm 881 Compact IC pro was used with a conductivity detector. We prepared five standards, from pure Na₂SO₄ from Sigma-Aldrich, at concentrations of 1.3 ppm, 10.6 ppm, 20.3 ppm, 51.4 ppm, and 100.6 ppm, and generated a calibration line with R² value of 0.9954. The detection limit for sulfate by IC analysis was 0.1 ppm. The concentration in solution (in ppm) is then converted to ppm in the solid (mg SO₄/kg sample). Due to the rapid oxidation of sulfite in solution to sulfate under ambient laboratory conditions, the sample solutions were analyzed immediately after preparation to minimize oxidation during sample handling. The stability of sulfite was tested by comparing the concentration in the freshly prepared solutions and the same solution after sitting in air for 1 hour. Slight oxidation occurs even in this short time; therefore, the error of these measurements is estimated as the standard deviation between these two replicate measurements and was found to be on average 0.6 ppm in solution. This equates to a conservative estimate of ~350 ppm error in the solid, although samples were prepared and analyzed within ~20 min.

4. Analysis results of the ESD products from S and Cl salts

4.1 ESD products from Mg, Fe²⁺, Fe³⁺, Ca, Na sulfates and sulfites

4.1.1. ESD products from MgSO₄·xH₂O (*x*=1, 4, 7)

As listed in Table 1, 0.25h, 1h, 2h and 7h ESD experiments were carried out on MgSO₄·xH₂O (*x*=1, 4, 7) powder. The ESD product from each experiment was analyzed using Raman on multiple spots of an *as is* surface and using XRD on the bulk sample (from the full depth of the SiO₂ cell).

Raman spectra in Figure 4a are from 36-spot analyses on the *as is* surface of a 0.25h-ESD product from epsomite MgSO₄·7H₂O. Data from the shortest ESD duration (0.25 hour) were purposely presented to show the intermediate species generated during the ESD process. In a Raman spectrum of sulfate, the

fundamental vibrational modes (ν_1 , ν_2 , ν_3 , and ν_4) of the $(\text{SO}_4)^{2-}$ unit are located in four spectral regions centered around ~ 1000 , 500 , 1150 , and 600 cm^{-1} , often with multiple peaks for each mode. Among the ν_1 Raman peaks near 1000 cm^{-1} in Figure 4a, the strongest sharp peak at 1000 cm^{-1} belongs to a crystalline starkeyite $\text{MgSO}_4 \cdot 4\text{H}_2\text{O}$ (4W), a weak sharp peak at 983 cm^{-1} belongs to a crystalline $\text{MgSO}_4 \cdot 6\text{--}7\text{H}_2\text{O}$ phase (6–7 W) (983.6 cm^{-1} for $\text{MgSO}_4 \cdot 6\text{H}_2\text{O}$ and 984.1 cm^{-1} for $\text{MgSO}_4 \cdot 7\text{H}_2\text{O}$, Table 3 of Wang et al., 2006), the broad peaks (from all sampling spots) centered $\sim 1030 \text{ cm}^{-1}$, $\sim 600 \text{ cm}^{-1}$ and $\sim 500 \text{ cm}^{-1}$ (marked as “Amor”) demonstrate the initiation of amorphization (based on Figure 20 of Wang et al., 2009). Figure 4b shows an overlay of the Raman spectra from a 37-spot analysis of the same 0.25h-ESD product in $3876 - 3072 \text{ cm}^{-1}$ spectral range. The occurrence of H_2O peaks from each sampling spots indicates retention of structural H_2O in ESD product, i.e., $\text{MgSO}_4 \cdot 7\text{H}_2\text{O}$ is partially dehydrated.

Raman spectra in Figure 4 reveal a fast dehydration from epsomite $\text{MgSO}_4 \cdot 7\text{H}_2\text{O}$ to crystalline starkeyite (4W) at least, and the initiation of amorphization. Experimental study of hydrous Mg-sulfates (Wang et al., 2009) revealed that amorphous $\text{MgSO}_4 \cdot x\text{H}_2\text{O}$ can hold up to three structural H_2O per MgSO_4 molecule, consistent with the observation of H_2O Raman peaks of different shapes from all sampled spots on the *as is* surface (Fig. 4b).

Amorphization in epsomite $\text{MgSO}_4 \cdot 7\text{H}_2\text{O}$ developed very fast with increased ESD time duration. After 1 hour of ESD processing in the PEACH, almost no crystalline Mg-sulfates can be detected in the fundamental vibrational spectra range (not shown, but similar to Figure 5a). On the other hand, a trace of the H_2O peak still remains (not shown) after 7h ESD process on epsomite. In other words, full amorphization was reached but the full dehydration was not reached at the *as is* surface.

Total amorphization at the *as is* surface was reached after 1.5 h of ESD process on crystalline starkeyite $\text{MgSO}_4 \cdot 4\text{H}_2\text{O}$. As seen in Figure 5a, amorphization is characterized by (1) shift of the ν_1 peak from 1000 cm^{-1} to $\sim 1030 \text{ cm}^{-1}$, (2) broadened peak width for every peak in the whole spectrum and in every spectrum from all sampled spots, (3) merged peaks of ν_2 , ν_4 , and the lattice modes (below 400 cm^{-1}) into three large spectral envelopes, and (4) severely reduced S/N ratio (because the Raman peak intensity of non-crystalline phase is 1–2 order of magnitude weaker than that of crystalline phase with similar composition, White 1975) which also appeared as a raised spectral background. In Figure 5b, the structural damage of a crystalline kieserite $\text{MgSO}_4 \cdot \text{H}_2\text{O}$ by a 1.5h ESD are presented by (1) broadened peak widths, (2) loss of minor peaks, and (3) reduction of S/N, but to a lesser degree when compared with Figure 5a from crystalline starkeyite of 1.5h ESD.

XRD measurements made on the bulk ESD products from $\text{MgSO}_4 \cdot x\text{H}_2\text{O}$ ($x=1, 4, 7$) are shown in Figure 6. Both XRD patterns on the top of Figure 6 have a raised “hump” from 10° to $40^\circ(2\theta)$, which can be fitted with three wide “bands”, centered at 14.2° , 21.9° , and 28.0° with widths of 5.4° , 11.4° , and 11.5° , respectively. A few sharp XRD lines remain on the top of the “hump”; some can be assigned as crystalline $\text{MgSO}_4 \cdot \text{H}_2\text{O}$ (1w) or $\text{MgSO}_4 \cdot 4\text{H}_2\text{O}$ (4w), with some lines unassigned since hydrous Mg-sulfates can have eight different hydration degrees. The difference in the degrees of amorphization seen in XRD patterns (Fig. 6) and Raman spectra (Fig. 5a) is caused by the sampling differences of the two analyses, with XRD sampled the bulk sample (from full depth of SiO_2 cell) and Raman sampled the *as is* surface only.

The third XRD pattern in Figure 6 is obtained from the 1.5h-ESD product from kieserite $\text{MgSO}_4 \cdot \text{H}_2\text{O}$ (8.5h-ESD product has the same pattern). When compared with the standard XRD pattern of kieserite, the 1.5h-ESD product has a broadened line width for every line in the whole XRD pattern. In addition, the multiple lines in the line groups at $25\text{--}30^\circ$, $35\text{--}40^\circ$, $\sim 44^\circ$, and $55\text{--}60^\circ 2\theta$ ranges are merged into envelopes with non-symmetric shape. XRD standard 00-001-0638 (kieserite, $\text{MgSO}_4 \cdot \text{H}_2\text{O}$, monoclinic) and 00-037-009 (caminite $\text{Mg}_2(\text{SO}_4)_2 \cdot 2\text{H}_2\text{O}$, tetragonal) in the PDF database have lines that may contribute the

merged line groups of the ESD product (Fig. 6). The observed XRD line widening and merging of line clusters suggest a heavily distorted crystal structure in the ESD product from kieserite, suggesting development of amorphization caused by the impact of energetic electrons and by the reaction with free radicals generated in the ESD process. This conclusion is consistent with Raman observations from Figure 5b.

Overall, the effects of ESD processes on hydrous Mg sulfates are dehydration and amorphization. The higher the hydration degree in an original salt, the higher the rate of amorphization by the ESD process, a phenomenon that we also observed for other salts in this study.

4.1.2. ESD products from $\text{FeSO}_4 \cdot x\text{H}_2\text{O}$ ($x=1, 4, 7$)

The melting point of melanterite $\text{FeSO}_4 \cdot 7\text{H}_2\text{O}$ is 60°C (Lide 2001, Table 1). So the ESD experiment of melanterite was made with LN_2 temperature control, $T_{\text{eq}} < 30^\circ\text{C}$ (Figure 2), and durations of 0.25h, 1h, 3h, and 7h. ESD products were analyzed by Raman and VNIR on the *as is* surface, and by XRD and Mössbauer on bulk samples. Very low laser power (0.5 mW) was used in Raman measurements to avoid overheating of these Fe-bearing phases by the laser beam, which was confirmed by visual inspection under the Raman microscope before and after each scan.

Raman spectra Figure 7 (a, b) shows the overlay of the first 60 Raman spectra from the 0.25h ESD and 7h ESD products. The obtained spectra from the 0.25h ESD product are compared in Figure 7a with the standard Raman spectra of $\text{FeSO}_4 \cdot x\text{H}_2\text{O}$ ($x=1, 4, 7$, Choi et al., 2007). The ν_1 mode of rozenite $\text{FeSO}_4 \cdot 4\text{H}_2\text{O}$ (4w) and szomolnokite $\text{FeSO}_4 \cdot \text{H}_2\text{O}$ (1w), 1018 cm^{-1} and 990 cm^{-1} , appeared after a 0.25 hour ESD process, while the Raman peaks of melanterite $\text{FeSO}_4 \cdot 7\text{H}_2\text{O}$ (7w) disappeared i.e., a total destruction of melanterite by 0.25h-ESD. The only unassigned Raman peak in Figure 7a is at 1035 cm^{-1} (marked by red arrow), which is an indication of early development of amorphization (Fig. 7 of Ling and Wang, 2010), similar to the case of $\text{MgSO}_4 \cdot 7\text{H}_2\text{O}$ after 0.25h-ESD (Fig. 4a). The peak position at 1035 cm^{-1} suggests an amorphous phase $\text{Fe}^{3+}_2(\text{SO}_4)_3 \cdot 5\text{H}_2\text{O}$ (based on Fig. 18 of Wang and Ling 2011). Because of the different T_{eq} achieved by ESD processes on epsomite and melanterite (Fig. 2), a direct comparison of the amorphization rate cannot be extracted for these Mg, Fe-sulfates.

Figure 7b shows the first 60 spectra from a 120-spots Raman scan on the *as is* surface of 7h ESD product from melanterite, which revealed the abundant amorphization in the product. The center of strongest peak moves to near 1075 cm^{-1} (Fig. 7b), shown as a large envelop that merged the multiple peaks of ν_1 and ν_3 modes of SO_4 unit together (the merge of ν_1 and ν_3 was less obvious in Mg-sulfates, Fig. 5, 6). Peaks of the ν_2 , ν_4 , and lattice modes are also merged into three large envelopes around 610 , 470 , and 240 cm^{-1} , respectively, with severely reduced S/N ratio. The ν_1 peak of crystalline szomolnokite $\text{FeSO}_4 \cdot \text{H}_2\text{O}$ (1w) at 1018 cm^{-1} remains in some spectra.

Raman spectra obtained from a scan on the 1.5h-ESD product from rozenite $\text{FeSO}_4 \cdot 4\text{H}_2\text{O}$ are shown in Figure 8a. The strong and wide envelope spanning 1300 to 950 cm^{-1} represents a merge of ν_1 and ν_3 modes of SO_4 unit, with the sharp ν_1 peak of $\text{FeSO}_4 \cdot \text{H}_2\text{O}$ remaining in some spectra. The other three wide envelopes (below 1000 cm^{-1}) are from the merged peaks of ν_2 , ν_4 , and lattice modes respectively, similar to 7h ESD products from melanterite (Fig. 7b). H_2O peaks centered at 3430 cm^{-1} were detected at almost all sampled spots on the *as is* surface (Fig. 8a), suggesting that full dehydration was not reached.

Very sharp Raman peaks remained in all spectra from the ESD products of szomolnokite $\text{FeSO}_4 \cdot \text{H}_2\text{O}$ (1w), even after 15.5h-ESD (Fig. 8b). When compared with the standard spectra of various ferrous and ferric sulfates, we found that the best match was with $\text{Fe}^{3+}\text{OHSO}_4$. The transformation from $\text{Fe}^{2+}\text{SO}_4 \cdot \text{H}_2\text{O}$ to $\text{Fe}^{3+}\text{OHSO}_4$ demonstrated the occurrence of oxidation ($\text{Fe}^{2+} \rightarrow \text{Fe}^{3+}$) and hydrolysis caused by the ESD

process. The major peak at 1094 cm^{-1} of $\text{Fe}^{3+}\text{OHSO}_4$ was also observed at some sampled spots (hidden in the spectral overlay of Fig. 8a) on the *as is* surface of 1.5h-ESD product from rozenite $\text{FeSO}_4\cdot 4\text{H}_2\text{O}$.

XRD measurements were made on the bulk samples of ESD products from $\text{FeSO}_4\cdot x\text{H}_2\text{O}$ ($x = 1, 4, 7$). Figure 9 shows the XRD pattern from the 7h ESD product from melanterite $\text{FeSO}_4\cdot 7\text{H}_2\text{O}$ and from szomolnokite $\text{FeSO}_4\cdot \text{H}_2\text{O}$. The top XRD pattern in Figure 9 overlays the characteristic XRD lines of $\text{FeSO}_4\cdot \text{H}_2\text{O}$ onto two large XRD “humps”, centered at 11.4° and 27.5° that are similar to the humps in Figure 6, but slightly narrower in widths, 5.4° and 9.0° . This XRD pattern reveals a partially amorphous bulk sample. The XRD pattern from the 1.5h-ESD product from $\text{FeSO}_4\cdot 4\text{H}_2\text{O}$ (not shown) has strong lines of crystalline rozenite and szomolnokite. The partial amorphization observed in Raman data (Fig. 8a) appears less obvious in XRD data, likely because a bulk sample from full depth of ESD product in the SiO_2 cell was used for XRD. The XRD pattern (2nd in Fig. 9) from the 1.5h-ESD product from $\text{FeSO}_4\cdot \text{H}_2\text{O}$ has the major lines of butlerite $\text{Fe}^{3+}\text{OHSO}_4\cdot 2\text{H}_2\text{O}$ and $\text{Fe}^{3+}\text{OHSO}_4$, in addition to the remaining lines of $\text{FeSO}_4\cdot \text{H}_2\text{O}$. The appearance of butlerite and dehydrated butlerite indicate the oxidation of Fe^{2+} to Fe^{3+} , and hydrolysis caused by ESD processes, consistent with Raman observations (Fig. 8b).

VNIR spectra (Fig. 10) were taken at the *as is* surfaces of 0.25h, 1h, 3h, and 7h ESD products from $\text{FeSO}_4\cdot 7\text{H}_2\text{O}$ ($T_{\text{eq}} < 30^\circ\text{C}$, Fig. 2). The first major spectral change with increasing ESD time is a decrease in absorption band depth near 1.0, 1.4, and 1.9 μm , suggesting gradual dehydration. Indeed, the narrow and strong VNIR doublet near 1.9 μm of the 0.25h-ESD product can be assigned to $\text{FeSO}_4\cdot 4\text{H}_2\text{O}$ (Fig. 3 of Wang et al., 2016), and the much wider and weaker doublets (near 1.9 μm) in the spectra of the 1h, 3h, 7h ESD products are similar to the doublet of $\text{FeSO}_4\cdot \text{H}_2\text{O}$ (Fig. 6 of Wang et al., 2016). The second major spectral change is a gradual decrease of spectral contrast, which is especially obvious in the spectrum of 7h ESD product. This phenomenon reflects the destruction from a crystalline structure, consistent with Raman and XRD observations. Notice a weak but characteristic VNIR spectral pattern of $\text{FeSO}_4\cdot \text{H}_2\text{O}$ (Fig. 10) was obtained from the 7h ESD product, while on the same *as is* surface, a multi-spot Raman scan shows almost total amorphization; only a few spots retained the 1018 cm^{-1} peak of szomolnokite (Fig. 7b). This phenomenon shows that the sensitivities of Raman and VNIR spectra for detecting structural damage are different. A previous study using XRD, Raman, Mid-IR, and VNIR on a set of saponite samples with different degrees of amorphization revealed that VNIR spectroscopy is less sensitive to the loss of crystallinity. From the saponite samples that show totally non-crystalline XRD patterns, sharp VNIR spectral peaks can still be seen at 1.4 μm , 1.9 μm and especially in the 2.2-2.4 μm spectral range (Fig. 1, 2 of Fu et al., 2017).

A Mössbauer spectrum was obtained from 59 mg of the powdered bulk 7h ESD product from $\text{FeSO}_4\cdot 7\text{H}_2\text{O}$. The measured spectrum was fit in a variety of ways using two different programs. The Mex_disdd solves the full hyperfine interaction Hamiltonian and minimizes the difference between modeled line shape independent peaks and the experimental spectrum using center shift, quadrupole splitting, and linewidth as variable parameters. Disd_3e makes velocity approximations rather than solving the full interaction Hamiltonian and uses a range of line-shape-independent quadrupole splitting distributions (QSD) to build the peaks. Because the peaks in these spectra overlap heavily and it is difficult to prioritize one model over the other, so both are shown and reported (Figure 11a, b). In both models, there are two Fe^{3+} and three Fe^{2+} distributions. Parameters are given in Table 2; errors on total % Fe^{3+} are $\pm 1\text{-}5\%$ absolute based on repeated fits to the same spectra, with a detection limit for Fe^{3+} of roughly 1%. Although it is known that differential recoil-free fraction (f) effects can affect the assumption that Mössbauer peak area directly represents the proportions of Fe in each site or valence state, f has not been determined for these materials. Thus, this paper assumes that recoil is the same for both Fe^{2+} and Fe^{3+} . Based on Mössbauer analysis, we obtained a $\text{Fe}^{3+}/\text{Fe}_{\text{total}}$ ratio of 44% in analyzed sample. This result indicates a strong oxidation has occurred during ESD process on melanterite $\text{FeSO}_4\cdot 7\text{H}_2\text{O}$. This conclusion is visually confirmed by the color change of the samples, from light green of

FeSO₄·7H₂O, to light yellow, to light brown, and finally to dark brown after 7h ESD, suggesting more oxidized iron (Fig. S1). The amorphization development in 7h ESD product from FeSO₄·7H₂O may be reflected in the details of Mössbauer parameters, e.g., CS = 1.21-1.29 mm/s and QS = 1.55-2.66 mm/s, which are different from the ranges in literature for FeSO₄·xH₂O (x=1, 4, 7), 1.16-1.31 mm/s for CS and 3.17-3.24 mm/s for QS (Cheetham et al., 1981; Dyar et al., 2013; Eissa et al., 1994a,b; Grant et al., 1966; Montano, 1981; Sakai et al., 1981; Sallam et al., 1994).

Overall, the effects of ESD processes on hydrous Fe²⁺ sulfates are dehydration, amorphization, and oxidation from Fe²⁺ to Fe³⁺.

4.1.3. ESD products from ferric sulfates

Given the importance of Fe³⁺-bearing phases for Mars surface mineralogy, we conducted ESD experiments in the PEACH on four Fe³⁺-bearing species, Na-jarosite NaFe₃(OH)₆(SO₄)₂, ferricopiapite Fe_{4.67}(SO₄)₆(OH)₂·20H₂O, akaganeite FeO(OH,Cl), and pyrite FeS₂ (Table 1).

4.1.3.1. ESD products from ferricopiapite

Figure 12b shows typical Raman spectra obtained from ESD products of ferricopiapite Fe_{4.67}(SO₄)₆(OH)₂·20H₂O, with the ESD durations of 0.25h, 1h, 2h, 3h, and 7h. When compared with Raman spectra of standard ferric sulfates (Fig. 12a, from Ling and Wang, 2010), we found the spectra from the 0.25h-ESD product can be assigned to ferricopiapite (#1 in Fig. 12b), rhomboclase FeH(SO₄)₂·4H₂O (#2 in Fig. 12b), and a phase whose most peaks are similar to those of ferricopiapite in positions and shapes, except its strongest ν₁ peak at 1018 cm⁻¹ (#3 in Fig. 12b) that can be regarded as a merged ferricopiapite doublet, with an upper-shifted peak center. Forming acidic rhomboclase from basic ferricopiapite is also a dehydration process with N_{H2O}/N_{SO4}=3.67 → 2. The shifting of ν₁ peak position to higher wavenumber is normally an indication of dehydration, observed in the Raman spectra of hydrous Mg, Fe²⁺, and Ca sulfates. The merge of the doublet suggests a damaged crystal structure, i.e., an early step towards the amorphization.

Typical Raman spectra from #4 to #8 in Figure 12b obtained from 1h, 2h, and 7h ESD products all have the strong and wide peak in ν₁ spectral range, but the central position of this peak gradually shifted towards higher wavenumber (indicated by a dotted arrow line in Fig. 12b), from 1018 cm⁻¹ to 1057cm⁻¹ (#3 to #8 in Fig. 12b). This peak shift is accompanied by the loss of details for all peaks below 800 cm⁻¹, which eventually become three wide envelopes centered around ~650 cm⁻¹, ~480 cm⁻¹, and ~250 cm⁻¹ (spectrum #4, #5, #6, #7, #8). These are typical Raman spectral pattern of amorphous ferric sulfates (#3 in Fig. 12a). Wang and Ling (2011) built a calibration curve (their Fig. 18) to quantify the number of structural H₂O per Fe₂(SO₄)₃ unit, from eleven to five, using the ν₁ peak position of amorphous ferric sulfates (from 1022 to 1035 cm⁻¹). Although there are no other experimental studies of amorphous ferric sulfates with structural water less than five, an educated guess is that the continuous ν₁ peak upper shift from 1035 cm⁻¹ to 1057 cm⁻¹ (Fig. 12b) is due to continuous dehydration from five to zero structural waters per Fe₂(SO₄)₃ unit, based on the assignment of spectra #9 and #10 in Figure 12b.

Spectra #9 and #10 (Figure 12b) were obtained from many Raman sampled spots on the *as is* surface of the 7h ESD product from ferricopiapite. Spectrum #10 is a perfect match with the standard Raman spectrum of crystalline anhydrous Fe₂(SO₄)₃ (#4 in Fig. 12a), and spectrum #9 is a mixture of standard spectra of two polymorphs, the anhydrous Fe₂(SO₄)₃ and mikasaite Fe₂(SO₄)₃ (#5 in Fig. 12a), both are crystalline materials (Ling and Wang 2010). The multi-spots Raman scans revealed that crystalline anhydrous Fe₂(SO₄)₃ appeared early in ESD products (e.g., 1h-ESD) and became abundant in the ESD product with longer time duration (e.g., 7h ESD).

Overall, the major effect of ESD process on ferricopiapite is dehydration, from $N_{H_2O}/N_{SO_4}=3.67$ to $N_{H_2O}/N_{SO_4}=0$. The ESD caused structural change is complicated, from crystalline to full amorphization, then back to crystalline again. During the early dehydration, the ferric sulfate also changed from basic ferricopiapite to acidic rhomboclase.

4.1.3.2. ESD products from Na-jarosite, akaganeite, and pyrite

It is quite surprising that no obvious mineral transformation occurred in Na-jarosite after long duration ESD processes (even after 64 hours). The color of post-ESD sample surface shows an obvious darkening (Fig. S2). The peaks of hematite appeared in the spectra from some spots in multi-spots Raman scans. However, 99% of Raman spectra obtained from the *as is* surfaces of eight ESD products from Na-jarosite maintain an almost perfect match with the Raman spectrum of the original Na-jarosite (Fig. S3). The Mössbauer spectrum of this post-64h ESD product revealed a set of typical CS and QS for Na-jarosite (Fig. S4).

Considering the importance of jarosite in martian surface mineralogy, we made two sets of 1:1 mixtures of Na-jarosite with $MgSO_4 \cdot 7H_2O$ and $MgCl_2 \cdot 6H_2O$, and did ESD on them, with a goal to study a potential catalysis effect. Again, no obvious mineral phase transformation was observed by Raman spectroscopy in the products from two additional sets of eight ESD experiments (Table 1, 2), except the spectrum of hematite was seen from some spots in the Raman scans on these ESD products. The Mössbauer spectra of the 7h ESD products from the two mixtures (not shown) support ~ 100% jarosite among the Fe-phases in bulk samples.

Similar to Na-jarosite, no obvious mineral transformation from akaganeite $Fe^{3+}O(OH,Cl)$ was found by Raman and Mössbauer analysis of the post-ESD products (Fig. S5). On the other hand, Raman scans on post-ESD products from a natural pyrite show minor changes in Raman peak positions and shapes (Fig. S6). However, the XRD pattern of the 14h ESD product from this natural pyrite does show an obvious change from standard pyrite, thus it cannot be compared with Raman spectra to suggest any significant phase transformation.

4.1.4. ESD products from Na_2SO_4

The Raman spectra obtained on the *as is* surface of the 7h ESD product from Na_2SO_4 powder have only minor changes from the standard spectrum (Fig. 13), such as the slight peak broadening of Raman ν_1 mode near 992 cm^{-1} (insert of Fig. 13), and slightly raised spectral background after 400 cm^{-1} . The broadening of the Raman peak indicates a damaged crystalline structure from original Na_2SO_4 . XRD measurements were not made on this sample.

4.1.5. ESD products from $CaSO_4 \cdot 2H_2O$

The Raman spectra (Fig. 14) obtained on the 7h ESD product of $CaSO_4 \cdot 2H_2O$ powder revealed the transformation from gypsum to γ - $CaSO_4$ phase (not anhydrite α - $CaSO_4$) mainly, that has a sharp peak at 1026 cm^{-1} shown in the insert of Figure 14. Ordinary γ - $CaSO_4$ would not be stable at ambient terrestrial laboratory conditions. The post-ESD product from $CaSO_4 \cdot 2H_2O$ was sealed in a glass vial, with the Raman measurements made through the glass wall. In addition to γ - $CaSO_4$, bassanite $CaSO_4 \cdot 0.5H_2O$ was identified in some sampled spots (peak at 1014 cm^{-1} in the inset of Fig. 14), that is also supported by the observations of structural H_2O peaks in Raman spectra (not shown) of some sampled spots. No XRD measurement was made on this sample. It is notable that a large amount of γ - $CaSO_4$ was found to be stable within the vein system of martian meteorite MIL03346 (Ling and Wang, 2015), as well as in soils of hyperarid regions on Earth, such as from the Atacama Desert (Wei et al., 2015) and the saline playa on the Tibet Plateau (Wang et al., 2018). The γ - $CaSO_4$ phase apparently can exist stably under ambient terrestrial laboratory conditions. We are currently investigating the structural and chemical reasons for the abnormal stability of the γ - $CaSO_4$.

4.1.6. *ESD products from Na_2SO_3 and NaHSO_3*

In order to evaluate the oxidation power of ESD processes on S-bearing species, we selected two sulfites, $\text{Na}_2\text{S}^{4+}\text{O}_3$ and $\text{NaHS}^{4+}\text{O}_3$, as the starting phases for a set of ESD experiments with 0.25h to 11h duration (Table 1).

Among the obtained Raman spectra from 252 spots-scan on the *as is* surface of 7h ESD product from Na_2SO_3 , about ~ 12% has an additional peaks at 997 cm^{-1} (insert of Fig. 15). On the other hand, all Raman spectra from a total of 188 spots-scan on the *as is* surface of the 7h ESD product from NaHSO_3 has the additional peak at 997 cm^{-1} (not shown). Besides, this peak position does not match with the ν_1 peak of Na_2SO_4 at 993 cm^{-1} (insert of Fig. 15). XRD measurements were made on both sulfites. The XRD pattern of the 7h ESD product of NaHSO_3 is different from that of original sample (#1 in Fig. 16). After the 7h ESD process, new XRD lines appeared, and multiple lines merged in four regions ($22\text{--}30^\circ$, $31\text{--}37^\circ$, $46\text{--}50^\circ$, and $57\text{--}61^\circ$ 2θ), suggesting structural damage of NaHSO_3 . In addition, XRD standards 00-037-1488 ($\text{Na}_2\text{S}^{4+}\text{O}_3$), 04-015-3684 ($\text{Na}_2\text{S}^{4+}_2\text{O}_5$), and 00-021-1371 ($\text{Na}_2\text{S}^{6+}_3\text{O}_{10}$) in PDF database (#2, #3, #4 in Fig. 16) have lines that might contribute the merged line groups that appear in the top XRD pattern of Figure 16. The potential contribution of $\text{Na}_2\text{S}^{6+}_3\text{O}_{10}$ (#4 in Fig. 16) to the merged line groups provides a hint for the oxidation of S^{4+} to S^{6+} .

Oxidation of sulfite ($\text{S}^{4+}\text{O}_3^{2-}$) to sulfate ($\text{S}^{6+}\text{O}_4^{2-}$) through the ESD process was finally established using the Ion Chromatography (IC) analyses of 1h, 3h, 7h ESD products from Na_2SO_3 (dissolved in N_2 -purged milliQ water and immediately analyzed by IC), shown in Figure 17. The time series shows a gradual increase as a function of ESD experimental duration, from 1 hour to 7 hours.

4.2. *Analyses of the ESD products from Mg, Fe, Ca, Al, Na, K- Chlorides*

We conducted ESD experiments on six chlorides (Table 1), two anhydrous (NaCl and KCl) and four hydrous (Mg- , Fe^{2+} -, Ca- , Al- chlorides). This selection was made on the basis of their potential existence in anhydrous or hydrous forms on Mars and their stable existence at ambient laboratory conditions making them experimentally feasible. Three hydrous chlorides (Mg- , Fe^{2+} -, Al-) among them have melting points T_{mp} below 100°C (Table 1); thus we ran the ESD experiments on these chlorides with LN2 temperature control with equilibrated temperature $T_{\text{eq}} < 30^\circ\text{C}$ (Fig. 2).

4.2.1. *ESD products from $\text{FeCl}_2\cdot 4\text{H}_2\text{O}$*

Figure 18 shows typical Raman spectra obtained from the *as is* surfaces of 0.25h, 1h, 3h, and 7h ESD products from $\text{FeCl}_2\cdot 4\text{H}_2\text{O}$. Spectral features from most spots on 0.25h-ESD product remained similar to those of crystalline chlorides, with an additional sharp Raman peak at 334 cm^{-1} (#2 in Fig. 18). The major H_2O peak at 3406 cm^{-1} and a peak-shoulder at 3445 cm^{-1} (not shown) suggest a change of hydration degree from $\text{FeCl}_2\cdot 4\text{H}_2\text{O}$ to probably $\text{FeCl}_2\cdot 2\text{H}_2\text{O}$. After 1h ESD, Raman peaks began to broaden (#3 in Fig. 18). In addition, an overall Raman spectral shape very similar to that of hematite $\text{Fe}^{3+}_2\text{O}_3$ occurs frequently after 3h ESD (#4 in Fig. 18). After 7h ESD, a few sampled spots have a peak near 395 cm^{-1} (#5 in Fig. 18) that is the major Raman peak of goethite, Fe^{3+}OOH , but most spots have extremely broad Raman “humps” near 1310 , 710 , and 300 cm^{-1} (#6 in Fig. 18). The H_2O peak in the $3300\text{--}3700\text{ cm}^{-1}$ range becomes wide and weak, to almost nonexistent (not shown). Overall, this set of spectra revealed a progressive loss of crystallinity, oxidation ($\text{Fe}^{2+} \rightarrow \text{Fe}^{3+}$), and dehydration.

The XRD pattern obtained from this 7h ESD product from $\text{FeCl}_2\cdot 4\text{H}_2\text{O}$ (Fig. 19) has three “humps” in 2θ ranges of $8\text{--}18^\circ$, $23\text{--}41^\circ$, and $45\text{--}60^\circ$, with broadened lines on top of the humps. These XRD lines match with $\text{FeCl}_2\cdot 2\text{H}_2\text{O}$ (pdf: 00-025-1040), FeCl_2 (pdf: 00-001-1106), and Fe^{3+}OCl (pdf: 00-039-0612) (#1, #2, #3 in Fig. 19), but have much wider linewidth and merged line groups. Therefore, XRD data supports the inferences of amorphization, oxidation, and dehydration that are based on Raman analysis.

The Mössbauer spectrum of bulk 7h ESD product from $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$ (Fig. 20) was modeled using the Mex_disdd program. These data are interpreted within the context of broader work on Mössbauer spectroscopy of minerals with similar structures. Only a handful of Mössbauer spectra could be found in the literature for iron dichloride tetrahydrate, and most of the papers do not provide explicit parameters (Ohshita et al., 2002) or focus on low-temperature (Shinohara, 1977) or magnetic field measurements (Johnson, 1966; Kandel et al., 1973; Spiering et al., 1978) that are not comparable to this study. Overall, the analyzed sample contains 88% of the total Fe as Fe^{3+} , which was 0% in the starting phase of $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$. The strong oxidation revealed by this Mössbauer result is consistent with Raman observation of hematite occurrence in 3h and 7h ESD products.

The Mössbauer result suggests that both Fe^{2+} and Fe^{3+} are present in two dissimilar sites. Most of the Fe is Fe^{3+} in sites with CS = 0.37-0.38 mm/s and Qs = 0.55 and 0.93 mm/s. A small amount of Fe^{2+} is also observed, with CS = 1.22 and 1.26 mm/s and Qs = 2.96 and 2.10 mm/s, respectively. Ôno et al. (1964) report parameters of CS = 1.26 mm/s and QS = 0.80 mm/s for FeCl_2 ; these seem unusual for this phase. Chandra and Hoy (1966) studied $\text{FeCl}_2 \cdot 2\text{H}_2\text{O}$ and gave parameters of CS = 1.03 mm/s and QS = 2.50 mm/s. Grant et al. (1966) gave parameters of CS = 1.22 mm/s and QS = 2.98 mm/s for $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$ – very similar to the parameters obtained in this study, though our modern analyses reveal two different doublets.

VNIR spectra (Fig. 21) were measured from the *as is* surfaces of 0.25h, 1h, 3h, and 7h ESD products from $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$ ($T_{\text{eq}} < 30^\circ\text{C}$, Fig. 2). As with the ESD products from $\text{FeSO}_4 \cdot 7\text{H}_2\text{O}$ (Fig. 10), there are two major trends of spectral changes following the increase of ESD duration. Absorption band depth near 1.0, 1.4, and 1.9 μm decreases, suggesting a gradual dehydration, and overall spectral contrast decreases, consistent with the destruction of a crystalline structure observed by Raman and XRD analysis. Furthermore, there is a reduction of spectral slope between 400 to 800 nm that is more obvious in Figure 21 than in Figure 10. This more obvious slope change could be a reflection of $\text{Fe}^{3+}/\text{Fe}_{\text{total}} = 88\%$ in 7h ESD from $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$, which is higher than $\text{Fe}^{3+}/\text{Fe}_{\text{total}} = 44\%$ in 7h ESD product from $\text{FeSO}_4 \cdot 7\text{H}_2\text{O}$, determined by Mössbauer analyses.

4.2.2. ESD products from Na, K, Mg, Al, Fe^{2+} -chlorides

The structural damage (i.e., development towards amorphization) of common chlorides by energetic electrons of the ESD process is strongly influenced by chemical bonding. XRD is the major tool to characterize these ESD products.

After running ESD experiments with the same time duration (Table 1), the product from a starting material of $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$ shows an obvious change in XRD pattern (Fig. 19), whereas the products from either NaCl or KCl as starting material (Fig. 22) still have perfect matches with the standard crystalline forms of NaCl (pdf: 00-005-0628) and KCl (pdf: 00-004-0587).

Similarly, the XRD pattern of the LT-7h ESD product from $\text{AlCl}_3 \cdot 6\text{H}_2\text{O}$ indicates a total amorphization (1st black curve in Fig. 23). On the other hand, the XRD pattern of the LT-7h ESD product from $\text{MgCl}_2 \cdot 6\text{H}_2\text{O}$ (2nd black curve in Fig. 23) has an uneven raised background, with lines that match with a mixture of $\text{MgCl}_2 \cdot \text{H}_2\text{O}$ (pdf: 00-061-0222), $\text{MgCl}_2 \cdot 2\text{H}_2\text{O}$ (pdf: 00-003-0765), and $\text{MgCl}_2 \cdot 4\text{H}_2\text{O}$ (pdf: 04-017-8711), suggesting dehydration from $\text{MgCl}_2 \cdot 6\text{H}_2\text{O}$. The raised background, the line broadening, and line merging indicate development of amorphization. Furthermore after a long duration (21 hours) ESD on $\text{MgCl}_2 \cdot 6\text{H}_2\text{O}$, the XRD lines of $\text{Mg}(\text{ClO}_4)_2 \cdot 6\text{H}_2\text{O}$ (pdf: 00-014-0022) appear (not shown), indicating oxidation from Cl^{1-} to Cl^{7+} as observed in NaCl by our previous study (Wu et al., 2018).

In contrast, the XRD pattern of the LT-7h ESD product from $\text{CaCl}_2 \cdot 2\text{H}_2\text{O}$ (3rd black curve in Fig. 23) almost perfectly matches standard $\text{CaCl}_2 \cdot 2\text{H}_2\text{O}$ (pdf: 04-010-1481). During the XRD measurement (total of 15 minutes) on the 14h ESD product from $\text{CaCl}_2 \cdot 2\text{H}_2\text{O}$, an obvious color change of the sample (from

white to semi-transparent) was observed (laboratory relative humidity > 55%) and an XRD pattern similar to standard $\text{CaCl}_2 \cdot 2\text{H}_2\text{O}$ was obtained. Based on these two XRD measurements, we estimate that the ESD process may have caused dehydration of $\text{CaCl}_2 \cdot 2\text{H}_2\text{O}$, while a positive identification of the ending phase is lacking at current stage.

Overall, these XRD data on 7h ESD products from Mg, Fe, Al, Ca, Na, K chlorides suggest an approximate grouping that reflects how easily the ESD process causes structural damage of these chlorides, with Al, Fe, and Mg chlorides more easily damaged than Ca, Na, and K chlorides. This grouping is consistent with the observed ease of Cl release induced by the ESD process from common chlorides as reported in Wang et al. (2020).

5. Discussion

Table 3 summarizes the major conclusions on phase identifications of the ESD products from each salt. The column of “mid-phase” after a short period of ESD (≤ 1.5 hour) shows the pathways of these ESD-induced phase transformations. The listed final phases were reached mostly after 7 hours of the ESD process, with a few exceptions. Table 3 shows that phase transformations have occurred to different degrees in most of the 22 minerals and their mixtures tested in this study (a total of 75 ESD experiments, Table 1), as induced by a medium-strength ESD process with limited time duration (normally 7 hours). There are three major trends in the phase transformations: dehydration, amorphization, and oxidation of Fe, S, and Cl, which will be discussed separately.

5.1. Dehydration

Dehydration of hydrous salts was determined on the basis of direct Raman and XRD identification in ESD products of phases with hydration degrees lower than the starting salts. For example, $\text{MgSO}_4 \cdot x\text{H}_2\text{O}$ ($x=1, 4$), $\text{FeSO}_4 \cdot x\text{H}_2\text{O}$ ($x=1, 4$), FeOHSO_4 , $\text{Fe}_2(\text{SO}_4)_3$, and $\gamma\text{-CaSO}_4$ were found in the ESD products of $\text{MgSO}_4 \cdot 7\text{H}_2\text{O}$, $\text{FeSO}_4 \cdot 7\text{H}_2\text{O}$, $\text{Fe}_{4.67}(\text{SO}_4)_6(\text{OH})_2 \cdot 20\text{H}_2\text{O}$, and $\text{CaSO}_4 \cdot 2\text{H}_2\text{O}$ (Fig. 4, 6, 7, 8, 9, 12, 14). $\text{FeCl}_2 \cdot 2\text{H}_2\text{O}$, FeCl_2 , and $\text{MgCl}_2 \cdot x\text{H}_2\text{O}$ ($x=1, 2, 4$) were found in the ESD products of $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$ and $\text{MgCl}_2 \cdot 6\text{H}_2\text{O}$ (Fig. 19, 23).

In addition, there are three specific ESD-induced dehydrations. The first is that the highly hydrated sulfates ($\text{MgSO}_4 \cdot 7\text{H}_2\text{O}$, $\text{FeSO}_4 \cdot 7\text{H}_2\text{O}$) have a higher dehydration rate than those with lower hydration degrees of the same type (i.e., $\text{MgSO}_4 \cdot x\text{H}_2\text{O}$, $\text{FeSO}_4 \cdot x\text{H}_2\text{O}$, $x=1, 4$). This was reflected by the shift of central positions of the ν_1 Raman peak in the products as a function of ESD duration. The second is the occurrence of hydrolysis induced by ESD, i.e., from $\text{FeSO}_4 \cdot \text{H}_2\text{O}$ to $\text{Fe}(\text{OH})\text{SO}_4$. The third is the difficulty in the dehydration of the salts (and minerals) without structural H_2O but only OH, such as jarosite $\text{NaFe}_3(\text{OH})_6(\text{SO}_4)_2$ and akaganeite $\text{FeO}(\text{OH}, \text{Cl})$, as well as the 1:1 mixtures of Na-jarosite with each of two hydrous salts ($\text{MgSO}_4 \cdot 7\text{H}_2\text{O}$ and $\text{MgCl}_2 \cdot 6\text{H}_2\text{O}$). The spectrum of hematite was seen from some spots in the Raman scans on these ESD products, but it did not affect the conclusion from Mössbauer spectra, i.e. ~ 100% jarosite among the Fe-phases in bulk samples. This result means that transformation to hematite from jarosite by medium-strength ESD must be at a very low rate compared with those of other Mg, Fe^{2+} , Fe^{3+} , Ca, and Na sulfates.

It is worth to mention that we did not put a control in PEACH (i.e., a starting salt of same type but to protect it from being affected by ESD generated plasma and free radicals) during each of 75 sets of experiments, mainly due to the experimental difficulties. Considering the vacuum desiccation experiments on hydrous salts (Sklute et al., 2015; Vaniman et al., 2004; Wang et al., 2006b, Wang and Zhou 2014), dehydrations caused by the simulated martian atmospheric conditions should have also occurred.

5.2. Amorphization

Amorphization of salts in this study is judged by Raman and XRD data analyses, with Raman analysis made on the *as is* surface of ESD products, and XRD analysis made on the bulk sample of ESD products. Three stages of amorphization can be distinguished based on Raman data analyses (Table 3), as Amor-I, Amor-II, and Amor-III. For the Amor-I group, such as those observed from Na_2SO_4 , $\text{CaSO}_4 \cdot 2\text{H}_2\text{O}$, and Na_2SO_3 after 7h ESD (Fig. 13, 14, 15), the obvious broadening of major Raman peaks indicates the initiation of structural distortion from a perfect crystalline phase, which is commonly accompanied by the raising of spectral background and reduction of signal to noise (S/N). Structural distortion would generate the observed changes in chemical bond lengths and bond angles in a crystal, causing a much wider Raman peak to be obtained from a vibrational mode. The envelope of many Raman peaks with slightly different central positions widens because of many slightly different chemical bonds in a distorted structure, and an overall reduced S/N. Raman spectra of Amor-III exhibit a total loss of spectral details, in many cases with merged peaks in ν_1 , ν_2 , ν_3 , and ν_4 modes, extensively broadened peak widths (> 10 times), and very low S/N, observed from $\text{MgSO}_4 \cdot y\text{H}_2\text{O}$ ($y=7, 4$), $\text{FeSO}_4 \cdot y\text{H}_2\text{O}$ ($y=7, 4$), $\text{Fe}_{4.67}(\text{SO}_4)_6(\text{OH})_2 \cdot 20\text{H}_2\text{O}$, $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$ after the ESD process (Fig. 5a, 7b, spectra #4 to #8 in Fig. 12b, spectra #3 to #6 in Fig. 18). The spectra of Amor-II have characteristics between those of Amor-I and Amor-III, such as the ESD product of $\text{MgSO}_4 \cdot \text{H}_2\text{O}$ shown in Fig. 5b.

Two stages of amorphization can be distinguished from the XRD data (Table 3), corresponding to Amor-III and the combined Amor-I and -II assigned by Raman results. Amor-III would appear as the “large hump” in the XRD patterns from the ESD products from $\text{MgSO}_4 \cdot y\text{H}_2\text{O}$ ($y=7, 4$) (Fig. 6), $\text{FeSO}_4 \cdot 7\text{H}_2\text{O}$ (Fig. 9), $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$ (Fig. 19), and $\text{AlCl}_3 \cdot 6\text{H}_2\text{O}$ (Fig. 23), consistent with Amor-III assigned by Raman analyses. Because XRD sampled the powder from the full depth of the SiO_2 cell including the bottom of cell that is less affected by energetic electrons, some XRD lines of crystalline phases may overlie the “large hump” (Fig. 6, 9, 19) while the Raman spectra from the *as is* surface show full amorphization (Fig. 5a, 18). The XRD pattern of the products assigned to Amor-I and Amor-II by Raman analysis would appear as line broadening and the merge of line groups, as seen in Figure 6 (1.5h-ESD from $\text{MgSO}_4 \cdot \text{H}_2\text{O}$), Figure 16 (7h ESD from NaHSO_3), and Figure 23 (LT- 7h ESD from $\text{MgCl}_2 \cdot 6\text{H}_2\text{O}$).

It is worth noting that the generation of Amor-III is normally accompanied by the rapid dehydration from sulfates originally having high degrees of hydration, such as from $\text{MgSO}_4 \cdot x\text{H}_2\text{O}$ ($x=4, 7$), $\text{FeSO}_4 \cdot x\text{H}_2\text{O}$ ($x=4, 7$), and $\text{Fe}_{4.67}(\text{SO}_4)_6(\text{OH})_2 \cdot 20\text{H}_2\text{O}$. Amor-I or Amor-II occurred in anhydrous sulfates (e.g., Na_2SO_4 , Na_2SO_3) or those with a low degree of hydration, e.g., $\text{MgSO}_4 \cdot \text{H}_2\text{O}$.

The rate of amorphization (i.e., the ease of damaging the crystalline structure) of common chlorides appears to be more related to the type of cations present, and reflected by a grouping of chlorides mentioned in section 4.2.2, i.e., higher rates in Fe, Mg, and Al chlorides than in Ca, Na, and K chlorides. The same grouping was found in experimentally observed rates of Cl-release induced by ESD from Mg, Fe, Al, Ca, Na, and K chlorides (Wang et al., 2020), which was correlated with the degree of M - Cl bond covalence that is usually quantified by the difference of electronegativity of M and Cl (Allred, 1961). These electronegativity differences range from 2.34 to 2.16 for KCl, NaCl, and CaCl_2 , and from 1.2 to 1.85 for FeCl_3 , AlCl_3 , FeCl_2 , and MgCl_2 (Wang et al., 2020). An apparent connection between amorphization and dehydration is observed in chlorides. Hydrous chlorides ($\text{AlCl}_3 \cdot 6\text{H}_2\text{O}$, $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$, $\text{MgCl}_2 \cdot 6\text{H}_2\text{O}$) appeared to amorphize more quickly (Fig. 19, 23) than anhydrous chlorides (NaCl, KCl, Fig. 22). Nevertheless, the ESD-induced Cl-release experiments were conducted strictly on anhydrous chlorides (KCl, NaCl, CaCl_2 , FeCl_3 , AlCl_3 , FeCl_2 , and MgCl_2), and the same grouping was found in two sets of experiments (Wang et al., 2020, and this study). We conclude that the rate of amorphization of the tested chlorides is fundamentally affected by the degree of M - Cl bond covalence.

5.3. Oxidation

Oxidation of Fe^{2+} to Fe^{3+} is evidenced by Mössbauer analyses of the ESD products from $\text{FeSO}_4 \cdot 7\text{H}_2\text{O}$ and $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$. The ratio of $\text{Fe}^{3+}/\text{Fe}_{\text{total}}$ in both cases changed from zero to 44%, and to 88%, respectively, after a medium-strength ESD process of only 7 hours. These Mössbauer results are confirmed by the slope change in the 400-800 nm range of VNIR spectra, particularly as seen in the ESD-product set from $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$ (Fig. 21). The hydrolysis from $\text{Fe}^{2+}\text{SO}_4 \cdot \text{H}_2\text{O}$ to $\text{Fe}^{3+}(\text{OH})\text{SO}_4$ revealed by Raman spectra (Fig. 8) and by XRD-based phase ID (Fig. 9), and the appearance of Fe^{3+}OCl in the ESD product of $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$ (Fig. 19), provide additional evidence of the oxidation of ferrous iron to ferric iron, induced by the ESD process.

During a normal glow discharge (ESD-NGD) in simulated Mars atmospheric composition and pressure, the free radicals CO^{2+} , CO^+ , O_b , H_{III} , H_{II} , OH , Ar_b , N_2 , N_2^+ (as well as O_2 , NO , and O^+ because of the overlapping of plasma lines used for detection, Figure S8) were detected instantaneously by *in situ* plasma emission spectroscopy. O_3 was also detected in the output gas by UV and mid-IR spectroscopy (Wu et al., 2018). These free radicals would induce the oxidation of Fe^{2+} to Fe^{3+} . Similarly, oxidation of Cl^- to Cl^{7+} was indicated in this study by the presence of $\text{Mg}(\text{ClO}_4)_2 \cdot 6\text{H}_2\text{O}$ in 21h ESD products from $\text{MgCl}_2 \cdot 6\text{H}_2\text{O}$ (section 4.2.2). In addition, Wu et al. (2018) observed the formation of NaClO_3 and NaClO_4 from NaCl , both induced by the ESD processes of medium strength (Table S1).

Finally, our hypothesis of S-oxidation induced by ESD process was validated by the analyses of ESD products from two sulfites, $\text{Na}_2\text{S}^{4+}\text{O}_3$ and $\text{NaHS}^{4+}\text{O}_3$. The XRD-based identification of $\text{Na}_2\text{S}^{6+}_3\text{O}_{10}$ in ESD products (Fig. 16) supports the oxidation hypothesis, as does the IC observation of a surge in SO_4 concentration with increasing ESD duration (Fig. 17).

Overall, the results of this study demonstrated that dehydration and amorphization (to different degrees depending on the properties of salts) were induced by the medium-strength ESD processes within a nominal time duration from 0.25 hours to 7 hours. In addition, oxidation of Cl, S, and Fe occurred in Cl-, S-, and Fe-bearing salts as a result of this type of ESD process.

6. Implications for martian dust activities

Martian dust activities have been physically altering the morphology of the martian surface. Some of them, especially the regional and global dust storms, have the potential to physically remove and re-deposit the top layer of regolith and even rocks if secondary minerals (salts and phyllosilicates) are the major components. When electrostatic discharge is induced by martian dust activity, it exerts two additional effects on surface minerals: physically impacting them with energetic electrons and chemically attacking them with free radicals and electrons. The potential effects in chemistry and biology of ESD induced by Mars dust activities have been investigated and reported by Wu et al. (2018), and Wang et al. (2020a, b and this study) and by another group in Denmark (Bak et al., 2016, 2017, 2018; Knakjensen et al., 2014; Thøgersen et al., 2019).

To date, there has been no actual measurement made of the electric properties of martian dust events. Thus, two important unknowns remain. First is the probability of the occurrence of an ESD during a dust event, which would be expressed as a percentage of time duration in a dust event. Second is the type of ESD, either *Townsend dark discharge (TDD)*, or *normal glow discharge (NGD)*, that would occur in a specific dust event. Our current investigations (Wu et al., (2018), and Wang et al., 2020a, b, c) revealed the types of phase transformations (with some rate information) that can be induced by a medium-strength NGD, to a group of important secondary martian minerals (S- and Cl-salts). However, the overall effect of these phase transformations on the “big picture” of martian surface mineralogy cannot be estimated, unless some assumptions on the above two important unknowns are made.

We will take a very conservative assumption for the first unknown, and consider only two extreme cases: grain saltation and global dust storm. Our terminology in using “dust” in following discussion lumps all particles (dust, sand, soil grains, etc.) that can be physically moved by these two processes

First, grain saltation (GS) can occur on Mars everywhere and all year around when wind speed is beyond a threshold. Atmospheric scientists use threshold friction speed (u_{*t}) to judge the generation of GS. An early experiment using a wind tunnel (Greely et al., 1980) found that the u_{*t} at martian condition (at 5 mbar, 95% CO₂, T = 150-240K) is about 10 times higher than on Earth, i.e., GS is more difficult to initiate on Mars than on Earth. For example, the u_{*t} for grain size of 100 μm would be 2.5-3.5 m/s, and the u_{*t} for grain size of 800 μm would be 4-5 m/s under Mars conditions (Figure 5 of Greely et al., 1980). When adding the effect of the lower gravitational field of Mars, however, a recent study (Sullivan and Kok, 2017) found that at a wind speed much lower than u_{*t} , “sporadically mobilized” grains on Mars can develop into “self-sustaining saltation.” This conclusion was validated by the observations made during the *Spirit*, *Opportunity*, and *Curiosity* rover missions (Sullivan et al., 2005, 2008; Sullivan and Kok, 2017).

Fortunately, a Rover Environmental Monitoring Station (REMS) was carried by *Curiosity* rover to Gale crater. A recent paper (Viudex-Mpreiras et al., 2019) published wind speed data at Gale crater collected by REMS from sol 9 to sol 1474. Their finding was that the wind speed probability density function at Gale crater matches quite well with the Weibull function, with a scale factor $c = 6.87$ m/s and a shape parameter $k = 1.73$ (Fig.1 of Viudex-Mpreiras et al., 2019; Table S2). Furthermore, they found these parameters fall into the ranges of two Viking landers (which landed on flat plain, $c = 2.55 - 7.9$ m/s, $k = 1.06 - 1.68$). On the basis of this REMS data set collected during 2.19 Mars years, the probability of wind speed > 3.5 m/s is 70% and of wind speed > 5 m/s is 53%, at Gale crater (Table S2). Therefore, even we take the most conservative consideration by using the high threshold friction speed (u_{*t}) derived by the Greely et al. (1980) experiments (> 5 m/s for grain size of 800 μm), the winds at Gale crater would induce grain saltation (GS) over half of a martian year (53%, Table 4).

Schmidt et al. (1998) measured the E-fields in saltating grains at a California field site and found a level as high as 166 kV/m ($\gg 25$ -34 kV/m BEFT measured in Mars environmental chambers, Farrell et al., 2015; Yan et al., 2017). Since there was no actual measurement of the electric properties of grain saltation made on Mars and counting the other uncertainties in ESD generation that beyond the scope of current study, we chose a very conservative number, 1%, to estimate the ESD occurrence probability induced by martian grain saltation (GS). Combined with the occurring probability of GS during a Mars year at specific sites (e.g., Gale crater), the resulting probability of a GS-induced ESD would be 5.3×10^{-3} , equal to 87 hours during a martian year (Table 4).

Other martian dust activities, e.g., dust devils and dust storms, would disturb the martian surface dust and sand to much larger degrees than that by grain saltation. For simplification, we use global dust storm (GDS) to make a rough numerical calculation for comparison purposes. In general, GDS occurs on average once every three martian years. Once it occurs, it can be roughly assumed to cover at least 80% of the martian surface and last 10% of a martian year, corresponding ~ 69 Earth days (Gierasch, 1974; Shirley, 2015; Wang and Richardson, 2015). Therefore, the probability of any location on Mars during a martian year encountering a GDS is 0.026, or 435 hours (Table 4).

Now the question is: what is the percentage of time during a GDS that ESD could occur? Based on decades of study of martian dust storms, we understand that GDS are driven by a set of hot cores. Within the cores, there is likely convective activity, which could generate large electric fields (E-field) that would eventually cause an ESD to occur. During a GDS, the dust in these cores then gets transported to high altitudes and covers the globe of Mars. Therefore in the regions away from the cores, the dust load increases, but it is not vigorously mixing dust. The amount of suspended small aerosols increases, but

there is probably not a lot of electrical activity away from the cores (Gierasch, 1974). Considering this model and many more remaining uncertainties in martian GDS than GS, we chose a percentage at two orders of magnitude lower, 0.01%, for the probability of ESD occurring during a global dust storm (Table 4). Thus, the probability of encountering ESD as induced by GDS at any location on Mars during a martian year is 2.6×10^{-6} that equals to 4.4×10^{-2} hour (Table 4).

The second unknown is the type of ESD (either *Townsend dark discharge (TDD)*, or *normal glow discharge (NGD)*) that would occur in a specific dust event. Based on gas discharge phenomena described in the literature (Fig. S7), the major difference between TDD and NGD is in electron flux. This is reflected by the electric current measured in a discharge event (Gallo, 1975), which was observed at μA level for TDD by Farrell et al., (2015), and at mA level for NGD by Wu et al., (2018), both under Mars atmospheric conditions. Based on a modeling study (Delory et al., 2006), the full range of estimated electron flux between martian TDD and NGD ranges over seven orders of magnitude, from $9 \times 10^{16} \text{ s}^{-1} \text{ m}^{-2}$ (TDD) and $1.5 \times 10^{24} \text{ s}^{-1} \text{ m}^{-2}$ (NGD) (detailed discussion in section 5.3 of Wu et al., 2018). An ESD-NGD was observed in our experimental setting, with an electron flux of $1.42 \times 10^{20} \text{ s}^{-1} \text{ m}^{-2}$ (Table S1) calculated on the basis of measured electric current across the two electrodes, a strength at mid-way between the two extreme cases.

On the other hand, the type of chemical reaction that could be induced by an ESD process depends on the kinetic energy (E_k) of electrons being generated, i.e., $E_k >$ an energy threshold for a chemical reaction to occur. As discussed in Wu et al. (2018) and discussed in section 3.2, the kinetic energy of electrons generated in our ESD experiments was estimated on the basis of plasma spectroscopic observations of CO_2^+ and H_α lines, with considerable portion of electrons having $E_k > 14 \text{ eV}$ and even $> 17.2 \text{ eV}$. Because the drift velocities of electrons in TDD and NGD are similar (Fig. 4a and equation (8) of Delory et al., 2006; Jackson et al., 2008, 2010), their electrons should have similar kinetic energy distributions. Therefore, the chemical reactions induced by NGD can also be induced by TDD. However, the same reaction induced by the TDD process will take much longer time than NGD to reach the same level of phase transformation because of its lower electron flux (Fig. S7). For example, the electron flux realized in our NGD experiments is about $\sim 10^4$ times of that of typical TDD, thus the mineral transformation produced by a 0.25h ESD-NGD in our experiments (Table 3) would only be seen after about 2500 hours of a typical TDD process.

We further assume that NGD would more likely be induced by dust storms, while TDD would more likely be induced by grain saltation, simply because the difference in the amounts of dust grains involved in these two extreme dust activities matches well with the differences in electron flux densities of NGD and TDD, which are at mA and μA levels, respectively, when measuring electric current under Mars conditions (Farrell et al., 2015; Wu et al., 2018).

Based on the above assumptions and considering only the grain saltation and the global dust storms (and the probabilities derived in the front part of this section), we can make very rough estimations on how long it would take on Mars to reach the levels of phase transformations induced by our medium-strength ESD-NGD process in PEACH. The last two rows of Table 4 show that the phase transformations produced by a 0.25h-ESD-NGD process in the PEACH (column #3 of Table 3) would likely be seen after > 29 martian years when considering only the grain saltation induced ESD-TDD (at typical TDD electron flux level, Table S1). The time would be > 5.7 martian years when considering only the global dust storms induced ESD-NGD (at the same electron flux level of our experiments, Table S1). The durations needed for the phase transformation levels reached by 7h ESD-NGD (column #4 of Table 3) would likely be seen on Mars after a few hundred martian years for both extreme cases (grain saltation and global dust storms).

Note that the choice of ESD-occurring probabilities in the above analyses (1% for grain saltation and 0.01% for global dust storms), although based on some knowledge, maintains certain arbitrary nature because no real measurements have yet been made on Mars. Our goal was to enable a rough estimation on the numbers of martian years (Table 4), after which the phase transformations observed in our experiments (dehydration, amorphization, and oxidations of Fe, Cl, and S) would be seen on Mars. The estimated time lengths would be 10 to 100 times longer if we chose to further reduce the probability by one to two order of magnitude for both extreme dust activities, which would result an estimation of 'thousands to tens' thousands martian years. Nevertheless, the cold and dry atmospheric conditions have prevailed during Amazonian period (~ 3 Ga), especially in the most recent tens' of million years, during which martian dust activities (dust storms, dust devil, and grain saltation) have been rampantly altering martian surface materials. Our experimental results suggest that ESD induced by martian dust activities may have contributed some of the S- and Cl-rich X-ray amorphous materials in surface soils at Gale crater.

Furthermore, compared with the high frequency of occurrence, large area, and long temporal coverage of martian dust activities in the current epoch on Mars, the other potential amorphization processes, i.e., the sudden exposure of subsurface hydrous salts or a sudden release of subsurface brines (section 1) that might be induced by impacts or by other events such as Recurring Slope Lineae (RSL) would result in mostly localized occurrences with a lower probability of occurrence. This comparison suggests that the ESD process induced by martian dust activities could be a very important process during the Amazonian period on Mars in causing the generation of S- and Cl-rich amorphous materials and the oxidation of Fe, S, and Cl. A direct implication of this conclusion is that we would anticipate significant amounts of S- and Cl-rich amorphous materials, with highly oxidized Cl, S, and Fe, over the entire surface of Mars.

7. Acknowledgements

All coauthors claim no conflict of interest in publishing this manuscript.

Additional data that support this manuscript can be found in Supporting Information document. The digital file corresponding the spectral data in figures of this manuscript is available (Wang et al., 2020b), no user ID and password are required to access these data.

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8. Authors contributions

AW designed the experimental investigations. YCY did the sample preparations and conducted the ESD experiments in the PEACH, with the help of HKQ. AW conducted the analyses using Raman, VNIR spectroscopy, and XRD diffractometry, with the help of EBS. DMD and JH performed Mössbauer and IC analyses of ESD products and data interpretations. WMF provided scientific support for the ESD process in martian dust activity, especially the implication study. BLJ and SMM examined the experimental results and the conclusions derived from them. All coauthors participated in the manuscript writing.

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10. Captions of tables and figures

Table 1. Salts used in the ESD process and analysis.

Table 2. Mössbauer parameters obtained from curve fittings.

Table 3. Compilation of analysis results of ESD products of the salts studied.

Table 4. Rough calculations on grain saltation and global-dust-storm-induced ESD probabilities based on mission observations and various assumptions.

Figure 1. Scheme of ESD experimental setup.

Figure 2. Examples of the equilibrated temperatures T_{eq} during ESD experiments on studied salts.

Figure 3. (a) starting $MgCl_2 \cdot 6H_2O$ in a SiO_2 cell; (b, c, d) LT-7h ESD product of $MgCl_2 \cdot 6H_2O$ (in a SiO_2 cell, and zoom-in)

Figure 4. Raman spectra of 0.25h ESD products from $MgSO_4 \cdot 7H_2O$ in (a) spectral range of fundamental vibrational modes; (b) spectral range of H_2O modes, which are compared with standard Raman spectra of $MgSO_4 \cdot 3H_2O$ (3w), $MgSO_4 \cdot 4H_2O$ (4w), $MgSO_4 \cdot 5H_2O$ (5w), $MgSO_4 \cdot 6H_2O$ (6w).

Figure 5. Raman spectra of 1.5h ESD products: (a) from $MgSO_4 \cdot 4H_2O$ compared with standard spectrum (4w); (b) from $MgSO_4 \cdot H_2O$ compared with standard spectrum (1w).

Figure 6. XRD results of ESD products from $MgSO_4 \cdot 7H_2O$, $MgSO_4 \cdot 4H_2O$, $MgSO_4 \cdot H_2O$, compared with standard XRD patterns of starkeyite ($MgSO_4 \cdot 4H_2O$), Kieserite ($MgSO_4 \cdot H_2O$), and caminite ($MgSO_4 \cdot xMg(OH)_2 \cdot (1-2x)H_2O$).

Figure 7. Raman spectra of ESD products from $FeSO_4 \cdot 7H_2O$ (using 1% laser power=0.5 mw, $T_{eq} < 30^\circ C$); (a) First 60 spectra from a 99-spots Raman analysis on *as is* surface of 0.25h ESD product, compared with standard Raman spectra; (b) first 60 spectra of a 120 Raman analysis on *as is* surface of 7h ESD product.

Figure 8. Raman spectra of ESD products: (a) 1.5h-ESD product from $FeSO_4 \cdot 4H_2O$; (b) products after 1.5h, 8.5h, 15.5h ESD process from $FeSO_4 \cdot H_2O$, compared with standard spectra.

Figure 9. XRD result of LT-7h ESD product (bulk sample) from melanterite $FeSO_4 \cdot 7H_2O$ and szomolnokite $FeSO_4 \cdot H_2O$, compared with standard.

Figure 10. VNIR spectra on *as is* surfaces of 0.25h, 1h, 3h, 7h ESD ($T_{eq} < 30^\circ C$) from melanterite $FeSO_4 \cdot 7H_2O$.

Figure 11. Mössbauer spectrum and curve fitting results (a, b) of a 7h ESD product from melanterite, $FeSO_4 \cdot 7H_2O$.

Figure 12. Raman spectra of ESD product from ferricopiapite, $Fe_{4.67}(SO_4)_6(OH)_2 \cdot 20H_2O$, compared with standards. (a) Standard spectra of ferricopiapite (ferri), rhomboclase, $FeH(SO_4) \cdot 4H_2O$ (rhomb), amorphous $FeS_2(SO_4)_3 \cdot 5H_2O$ (Am5w), anhydrous $Fe_2(SO_4)_3$ (anhy), and mikasaite, $Fe_2(SO_4)_3$ (mika) (Ling and Wang 2010); (b) Raman spectra of ESD products from ferricopiapite.

Figure 13. Raman spectra of 7h ESD product from Na_2SO_4 , compared with the spectrum of starting Na_2SO_4 .

Figure 14. Raman spectra of 7h ESD product from $\text{CaSO}_4 \cdot 2\text{H}_2\text{O}$

Figure 15. Raman spectra of 7h ESD products from NaSO_3 , compared with standard spectra.

Figure 16. XRD results of 7h ESD product from NaHSO_3 , compared with original salt, and standards.

Figure 17. Results from ion chromatography on 1h, 2h, 7h ESD products from Na_2SO_3 .

Figure 18. Typical Raman spectra of ESD-products from $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$, compared with a standard spectrum of starting $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$.

Figure 19. XRD pattern of 7h ESD product from $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$, compared with standards.

Figure 20. Mössbauer spectrum and curve fitting results of a 7h ESD product from $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$.

Figure 21. VNIR spectra from as is surfaces of 0.25h, 1h, 3h, 7h ESD ($T_{\text{eq}} < 30^\circ\text{C}$) from $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$.

Figure 22. XRD results of ESD-7h products from NaCl , KCl , which match with PDF:00-005-0628(NaCl) and PDF: 00-004-0587 (KCl).

Figure 23. XRD results obtained from LT-7h ESD on $\text{AlCl}_3 \cdot 6\text{H}_2\text{O}$, $\text{MgCl}_2 \cdot 6\text{H}_2\text{O}$ and $\text{CaCl}_2 \cdot 2\text{H}_2\text{O}$, compared with standards.