Metallic ions in the upper atmosphere of Mars from the passage of comet C/2013 A1 (Siding Spring)

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Abstract We report the first in situ detection of metal ions in the upper atmosphere of Mars resulting from the ablation of dust particles from comet Siding Spring. This detection was carried out by the Neutral Gas and Ion Mass Spectrometer on board the Mars Atmosphere and Volatile Evolution Mission. Metal ions of Na, Mg, Al, K, Ti, Cr, Mn, Fe, Co, Ni, Cu, and Zn, and possibly of Si, and Ca, were identified in the ion spectra collected at altitudes of ~185 km. The measurements revealed that Na⁺ was the most abundant species, and that the remaining metals were depleted with respect to the CI (type 1 carbonaceous Chondrites) abundance of Na⁺. The temporal profile and abundance ratios of these metal ions suggest that the combined effects of dust composition, partial ablation, differential upward transport, and differences in the rates of formation and removal of these metal ions are responsible for the observed depletion.

1. Introduction

The close passage of comet Siding Spring (C/2013 A1, CSS) [McNaught et al., 2013] provided a unique opportunity to observe the close interaction between the dusty coma of a comet and a dense planetary atmosphere. Traveling on a highly inclined, hyperbolic orbit, the comet encountered Mars on 19 October 2014 at 18:29 UTC from a closest approach distance of ~134,000 km and with a relative velocity of 56 km s⁻¹ [Farnocchia et al., 2014]. During this event, the Mars Atmosphere and Volatile Evolution Mission (MAVEN) spacecraft [Jakosky, 2014] was ideally located and equipped to measure the response of the upper atmosphere of Mars to a strong and rapid cometary mass and energy disposition, and to assess the mechanisms by which the atmospheric system returns to equilibrium [Yelle et al., 2014]. It also offered the rare opportunity for direct characterization of cometary material being freshly delivered into the upper atmosphere and ionosphere of Mars. Of particular interest was the potential formation of metal ions from the ablation of cometary dust particles streaming through the atmosphere.

As on Earth and Jupiter, the presence of metal ions at high altitudes provides unequivocal signatures of the exogenic nature of the supplied material [Hughes, 1978; Grebowsky, 1981]. The formation of metal ions and their associated ionospheric layers by the ablation of meteoroids has been extensively observed in Earth’s upper atmosphere, and metal ions such as Mg⁺, Fe⁺, Na⁺, Al⁺, Ca⁺, and Ni⁺, have been detected in the ionosphere [Kopp, 1997; Grebowsky et al., 1998]. Several models mapped the chemical pathways by which these metal ions are recycled and ultimately removed [Molina-Cuberos et al., 2008, and references herein]. Similar mechanisms were predicted to form metal ions in the Martian ionosphere as the result of the sporadic meteor background [Pesnell and Grebowsky, 2000; Molina-Cuberos et al., 2003; Whalley and Plane, 2010]. Although the formation of transient ionospheric layers below 120 km at Mars was observed by Mars Express and Mars Global Surveyor [Pätzold et al., 2005; Withers et al., 2008], meteoric metal ions had not yet been directly detected. Radio occultation detects layers of electrons, and the corresponding positive ions in these low-lying layers are assumed to be metal ions. Predictions for the cometary dust flux into the Mars atmosphere were concerned primarily with impact hazard to spacecraft [Tricarico et al., 2014; Ye and Hui, 2014; Kelley et al., 2014; Farnocchia et al., 2014] and predicted an input dust flux comparable to the sporadic background with a maximum occurring on 19 October 2014 at 20:08 UTC. Withers [2014] pointed out that the high velocity of CSS dust implied the formation of a more robust ionosphere than formed in sporadic layers, which might be detectable by spacecraft. Such detection is important because it has the potential to validate models of ion transport, recycling, and removal mechanisms in the atmosphere of Mars, as well as allowing
the direct characterization of the refractory content of CSS. Compositional information of CSS will be valuable as it may provide the evidence that some of the Oort cloud comets come from the protoplanetary disks of other stars [Levison et al., 2010].

In this paper, we report the definitive detection of 12 metal ions of cometary origin and tentative detection of two additional ions in the upper atmosphere of Mars by the Neutral Gas and Ion Mass Spectrometer (NGIMS) in the hours that followed the close passage of CSS. The NGIMS instrument is a quadrupole mass spectrometer that utilizes a dual ion source designed to measure both surface reactive and inert atmospheric neutrals, and ion species in the mass range of 2–150 Da [Mahaffy et al., 2014]. Carried by the MAVEN spacecraft, this instrument aims to contribute to the mission goals of measuring the rate of atmospheric escape to space by characterizing the composition and the dynamics of neutrals and thermal ions in the upper atmosphere of Mars.

2. Detections of Metal Ions

Data of neutrals and ions were collected by NGIMS from 18 October to 22 October 2014 as part of the MAVEN Siding Spring observation campaign that took place immediately before and after the comet's encounter with the planet. During this campaign, NGIMS carried out 19 sets of special observations (also called activities). This special observation sequence devoted the majority of the instrument's observing time to a select set of atmospheric neutrals and ions but included 10 ion "survey" scans that were conducted at various altitudes. Each survey scan covered the full 2–90 Da mass range at a unit mass resolution. The NGIMS activities were conducted at regularly spaced intervals (orbits #108-128), interrupted only by a 10 h period at the peak of the cometary dust flux, when the instrument was temporarily turned off to minimize risk to the hardware. The NGIMS instrument operated when the spacecraft altitude was below 500 km. During the days that preceded and followed the encounter with CSS, the orbit of the MAVEN spacecraft remained nearly the same with respect to the Mars Solar Orbital Frame, with a periapsis located at 42° north latitude and 14:58 local mean solar time, a periapsis altitude of 185 km, and an orbital period of 4.6 h. However, due to the planet's rotation, the position of the spacecraft's tracks relative to the predicted region of cometary dust impacts was changing through the orbits (see Figure S1 in the supporting information for additional details).

The changes in the spectral signatures of ions in the upper atmosphere of Mars after the passage of CSS are depicted in two ion survey spectra collected at the same altitude (185–189 km). The first spectrum (Figure 1a) was captured on orbit #114, ~5 h prior to the peak dust flux of CSS, while the second (Figure 1b) was captured on orbit #119, ~19 h after the comet's peak dust flux. The difference between these two spectra (Figure 1c) reveals the emergence of several mass peaks that are characteristic of metal ions in the few hours that followed the CSS encounter. These ions were identified as Na⁺, Mg⁺, Al⁺, K⁺, Ti⁺, Cr⁺, Mn⁺, Fe⁺, Co⁺, Ni⁺, Cu⁺, and Zn⁺. The identity of most of these species (Mg⁺,
K+, Ti+, Cr+, Fe+, Ni+, Cu+, and Zn+) was established unambiguously by comparing measured isotope ratios to their relevant natural relative abundances established by the National Institute of Standard and Technology (NIST) (Figure 2).

Although it is possible to attribute the variability of the observed residuals at m/z = 27, 28, 29, and 40 to atmospheric ions (HCN+, CO+/N2+, and Ar+, to cite a few), their similar temporal evolution when compared to Mg+, Fe+, and Zn+ strongly support Al+, Si+, and Ca+ as being the originators of these spectral signatures. These NGIMS observations are also supported by concurrent detection of Mg+ and Fe+ reported by MAVEN’s Imaging Ultraviolet Spectrograph (IUVS) [Schneider et al., 2015], and of a transient dense ionospheric layer reported by the Mars Advanced Radar for Subsurface and Ionospheric Sounding (MARSIS) instrument on MEX and the Shallow Subsurface Radar (SHARAD) instrument on the Mars Reconnaissance Orbiter (MRO) [Gumett et al., 2015; Restano et al., 2015].

3. Temporal Evolution and Spatial Variation

All detected metal ions display comparable temporal evolution during the 48 h leading and following the passage of CSS (Figure 3 for Mg+, and figures in the supporting information for the rest of the species). None of these metal ions were found prior to the arrival of the comet. All were detected during the first NGIMS activity that followed the comet’s passage (initiated on orbit #117, 20 October 2014 05:02 UTC), and their highest recorded abundances occurred 19 h following the peak of dust flux (orbit #119). Although the measured abundances decayed exponentially afterward, metal ions were still detectable for 2.5 days after CSS passage. Previous models of metallic ions in the Martian atmosphere [Pesnell and Grebowsky, 2000; Molina-Cuberos et al., 2003; Whalley and Plane, 2010] have shown that sporadic meteors which enter with relatively low velocities (<15 km s⁻¹) ablate around 80 km. However, the CSS dust particles enter with velocities of 56 km s⁻¹. Assuming that these are low-density cometary dust particles with densities of less than 1000 kg m⁻³, the chemical ablation model CABMOD [Vondrak et al., 2008] predicts ablation around 115 km. A 1-D model for Mg⁺ [Whalley and Plane, 2010] shows that a combination of eddy and then ambipolar ion diffusion will transport ions from 115 km to 185 km fast enough to account for the abundance level measured by NGIMS.
Rapid redistribution by atmospheric winds would also explain the fact that the temporal evolution of the measured abundances seemed to be independent of the spacecraft location relative to the initial metal deposition region. At the predicted speeds, the winds can drag the outer envelope of the deposit hemisphere more than 90° in less than 10 h, which would tend to “dilute” and homogenize the metal distribution around the planet. The measured deviations on orbit #119 and #125 may reflect localized heterogeneities in the atmosphere as the redistribution was taking place.

The absolute abundances of the metal ions during their observed peak on 20 October 2014 14:26 UTC (orbit #119) are compiled in Table 1. Of all the “survey” scans, only data collected for Na⁺, Mg⁺, K⁺, and Fe⁺ on the first NGIMS activity following the comet’s passage exhibit signals above 200 km with levels high enough to permit the extraction of altitude profiles. Figure 4 shows the detector signal for each of the four species on that first NGIMS activity (orbit #117), revealing an interesting asymmetry between the inbound and the outbound leg. During the inbound leg, only K⁺ and Fe⁺ were detectable above 300 km. This is a surprising observation given that none of the Na⁺ or Mg⁺ could be detected above 216 km, and the first indication of the presence of these two species was obtained near periapsis. During the outbound leg, the abundances of all four species decayed asymptotically to ~1/15 of their peak value at periapsis and remained at that level until the last measurement was acquired at ~425 km. With the sparse sampling and limiting spatial information available, we cannot determine whether the apparent asymmetry is due to temporal variations in the metal ion densities or spatial variations along the track.

4. Ion Diffusion and Transport Effects

Figure 5 compares the relative NGIMS ion abundances to those in primitive CI carbonaceous chondrites [Lodders et al., 2009], both normalized to Na⁺. Note that all the elements are depleted with respect to the CI abundance of Na⁺. In view of the high signal-to-noise ratio for Fe⁺ and Mg⁺, and the fact that the chemical rates of formation and removal of these ions as well as those of Na⁺ have been measured [Plane, 2002; Whalley and Plane, 2010], we now focus on possible reasons for the depletions of Fe⁺ and Mg⁺ by factors of 44 and 24, respectively.

**Table 1. Abundance of Metal Ions as Derived From the NGIMS Measurements of 20 October 2014 14:26 UTC (Orbit #119) at 185 km Altitude**

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<tr>
<th>Ion Abundance (cm⁻³)</th>
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<tr>
<td>Na⁺ 201.2 ± 10.2 Cr⁺ 1.0 ± 0.1</td>
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<tr>
<td>Mg⁺ 153.6 ± 7.5 Mn⁺ 3.0 ± 0.2</td>
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<tr>
<td>Al⁺ 2.8 ± 0.4 Fe⁺ 69.0 ± 3.3</td>
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<tr>
<td>Si⁺ 318.2² Co⁺ 0.13 ± 0.03</td>
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<tr>
<td>K⁺ 3.4 ± 1.4 Ni⁺ 2.8 ± 0.2</td>
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<tr>
<td>Ca⁺ 2.0⁶ Cu⁺ 0.12 ± 0.03</td>
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<td>Ti⁷ 0.38 ± 0.07 Zn⁺ 0.62 ± 0.09</td>
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²Upper limit value.
5. Conclusions

The identification of metal ions in the ionosphere of Mars following the passage of CSS is a first-of-its-kind measurement conducted on another planet of the solar system. The characterization of the metal content in the CSS dust particles that ablated in the Martian atmosphere will require untangling the effects of the various mechanisms that interplayed to produce the signatures of metals observed by NGIMS. However, the large number of metal ion species detected by NGIMS along with published estimates that the dust fluxes from Siding Spring are comparable to the sporadic flux [Tricarico et al., 2014; Ye and Hui, 2014; Kelley et al., 2014; Farnocchia et al., 2014] suggests that the larger sporadic events are detectable at the MAVEN periapsis altitude. These measurements can place useful constraints on exogenic input to Mars.

References

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