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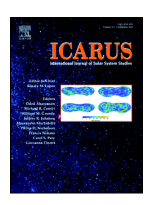
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Space Weathering on Inner Planetary Surface Analogues Induced by Swift Multicharged Heavy Ion Bombardment

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ABSTRACT

Silicates are ubiquitous in space. They dominate the surfaces of the inner rocky planets (Mercury in the case of the Solar System), the Moon, and asteroids, forming the major part of the non-volatile material. The physical and chemical properties of the rocky surfaces are determined not only by their initial composition but also by the processes occurring on them. Here we discuss one of these processes; irradiation by energetic cosmic particles that induces many effects among which structural changes and sputtering the latter contributing to the formation of exospheres. In the current work we report the 1 sull s of experiments conducted on anorthite, jadeite and nepheline silicates that have bein irradiated with energetic heavy ions with the aim to better understand the interaction of alactic cosmic rays, solar wind, and solar energetic particles with planetary and smill body surfaces. The sputtering effects induced by energetic (MeV/u) multicharged heavy ions (e.g., 105Rh and 140Ba) were analyzed by the PDMS-TOF-SIMS technique (plasi, a desorption mass spectrometry - time-of-flight secondary ion mass spectrometry'. Positive and negative secondary ionic species are identified: Na⁺, K⁺, Al⁺, Ca⁺, Si['] 1₂. Ejection of (SiO₂)_n and (AlSi)O_m cluster series are also observed. Less frequent, negative ion yields are one order of magnitude less than positive ones, or greater, which s the case for nepheline, with 0.671 ions impact⁻¹ for positive and 0.126 ions impact⁻¹ for negative ions. The results concerning ejection of ionic species show, for instance, that the Na^+/K^+ ratio is ~2.5, which is in very good agreement with that observed in the Hermean exosphere found to be ~ 2.3 .

Keywords: Moon, surface, Mercury, surface; Cosmic rays; Solar wind; Impact processes; Atmospheres, evolution

Silicate Irradiation, Energetic Heavy Ions, Space Weathering, Sputtering, Planetary surfaces.

1. INTRODUCTION

Among materials present in space, silicates are one of the most abundant. In the Solar System (SS), they are present in asteroids, meteorites and comets, where they comprise the major part of the non-volatile material (e.g. Hapke 2001, Strazzulla *et al.* 2005, Nuth & Johnson 2006). Their occurrence in other objects has been reported: in planets and their moons (e.g. Hapke 2001, Gudipati 2014, Cruikshank *et al.* 2015, Helled *et al.* 2020) and in trans Neptunian objects (Brucato *et al.* 2003). For instance, silicates are the dominant component of planet's crust (Henning 2010). In Mercury's crust silicates are even more abundant than iron. The large diversity in composition and surrection and surrection and processes occurring on them, like albedo or composition. Briefly, silicates play an important role in the cosmic life cycle of matter (Henning 2010).

Similarly to other materials in mace, solicates are modified by Solar Wind (SW), Solar Energetic Particles (SEP), and Ga'acta Cosmic Rays (GCR) irradiation (Strazzulla *et al.* 1995, Davoisne *et al.* 2008). In reinafter we use the terms cosmic rays (CR) to indicate energetic ions in space. Cosmic ray impacts induce physico-chemical modifications, particularly structural changes (like amorphisation) (Brucato *et al.* 2004, Szenes *et al.* 2010), and sputtering (Wiens *et al.* 1997, Baragiola 2004, Martinez *et al.* 2017). Sputtering is an erosion process that happens to any airless object exposed to the harsh environment of outer space (Brunetto 2009, Pieters & Noble 2016, Longobardo *et al.* 2020). It is of great importance because it causes major changes in optical properties of surfaces, like the Moon, where Space Weathering (SpWe) lowers the albedo, reddens the spectral slope, obscures absorption bands, and generates the characteristic magnetic electron spin resonance features of the lunar regolith (Cassidy & Hapke 1975, Hapke 2001).

SpWe effects due to sputtering have already been studied, in particular the erosion and surface alteration caused by the removal of atoms and/or molecules from a solid surface due to the interaction of projectile ions with target electrons and nuclei, as well as secondary cascades of collisions between target atoms (Sigmund 1981, Plainaki *et al.* 2009). In the Solar System, in several planetary environments, sputtering is one of the most important agents for the surface erosion; it occurs e.g. on Mercury (Cheng *et al.* 1987, Milillo *et al.* 2005, Martinez *et al.* 2017), the Moon (Hunten & Sprague 1997, Wurz *et al.* 2007), Europa (Eviatar *et al.* 1985) and Io (Wiens *et al.* 1997). Sputtering of the mineral components present at the surfaces of Mercury and the Moon appears to be among the most efficient source of sodium and potassium in their exospheres (e.g. Leblanc *et al.* 2007). Fillen *et al.* 2008). Na⁺ and K⁺ ions have been observed in Mercury's exosphere by Earth band and space observations e.g. by the MESSENGER spacecraft (Zurbuchen *et al.* 2008, Pral *et al.* 2010).

Effects induced by energetic ions 'ave been studied in laboratories since several decades with the aim to simulate and cosmic rays induced modifications. In particular to determine the importance of spracing by solar wind ions on the formation of a sodium exosphere around Mercury and the Moon, Dukes *et al.* (2011) irradiated Na bearing tectosilicates with 4 keV. He ions. Specifically space weathering has been analyzed and studied in laboratory; for instance, Marchi *et al.* (2005) performed ion irradiation experiments of silicates and compared laboratory spectra with those of asteroids and meteorites. They found similar mineralogy between most asteroids and meteorites, but different distributions of spectral slopes. All these experiments produced reddening and darkening of reflectance spectra, thus the variation observed in astronomical spectra were interpreted as space weathering induced by solar wind ion irradiation. Fu *et al.* (2012) performed ion irradiation of silicates samples with 50 keV He⁺, reflectance spectra at the VIS–NIR range showed also reddening and darkening of samples (for a review see Brunetto *et al.* 2015).

Several silicates were irradiated and analyzed as possible candidates for the surfaces of small objects and planets in the SS. Dukes *et al.* (2011) irradiated with 4 keV He⁺ projectiles albite, labradorite, and anorthoclase samples, as well as adsorbed Na layers deposited on albite and on olivine in order to quantify the Na sputtering at the surface and near surface. Nepheline, an aluminosilicate considered to be a good analogue for Mercury's surface (Shimoda *et al.* 2005, Wooden 2008), has also been irradiated with keV/u–MeV/u ion beams and measured sputtering yields and velocity spectra to better understand the evolution of silicate surfaces and the presence of Na and K vapor in the Herman exosphere (Martinez *et al.* 2017). As an additional process, to simulate the energy caused by micrometeorite bombardment, pulsed intense infrared lasers were used to irradiate olivine and pyroxene targets (Weber *et al.* 2020).

Ion sputtering is an important and active process in many SS planetary environments where a temporary plasma is generated and can precipitate on their surfaces. In this work we present experimental results concerning prorthite, jadeite and nepheline silicates irradiation with energetic heavy ions to contribute to a better understanding of the interaction of energetic cosmic particle irradiation of planetary and small body surfaces. The heavy ion projectiles are fission fragments (FF) enarge of yard 252Cf source. Experiments with FF (e.g., 105Rh and 140Ba) with atomic numbers close to that of heavy constituents of CR are indeed a good simulation of effects induced by the heavy ion fraction (a few percent) of cosmic rays (Farenzena et al. 2005, Martinez et al. 2014). Under these conditions, the FF-silicate interaction yields an electronic (inelastic) stopping power of about 5 keV/nm, i.e. much higher than the nuclear (elastic) stopping power one. The aim of the present work is to provide new insights about compositions of surfaces and exospheres of Mercury, Mars and the Moon, as well as desorption yields of ionic species (sputtering) due to the space weathering caused by cosmic rays. In addition, the results here presented may help to understand future observations of missions like Artemis to the Moon, BepiColombo to Mercury and Perseverance to Mars.

2. SPUTTERING - EXPERIMENTAL METHODS

The experiments were performed at the Van de Graaff Laboratory of the Pontifical Catholic University of Rio de Janeiro (PUC-Rio). The silicate samples were irradiated and analyzed in situ by using the technique PDMS-TOF-SIMS (plasma desorption mass spectrometry - time-of-flight secondary ion mass spectrometry). This technique uses as projectiles energetic (~0.5 MeV/u) multicharged heavy ions (e.g. 105Rh and 140Ba) emitted by a 252Cf source. The outline of the set-up is shown in Fig. 1 2ric fly, it consists of a time-offlight mass spectrometer with the sample holder located in mont of a ²⁵²Cf source. The FF impact analysis occurs by transmission mode (the FF passes through the sample). The experiments are performed under high vacuum (ρ) ssure ~ 1 × 10⁻⁶ mbar). Details of the technique can be found elsewhere (Ponciano et al. 2002, 2007). The projectile ions penetrate the target layers deposited on a substrate, a thin carbon foil. The sputtered ions are forward ejected with respect to the incident bean (forward sputtering process) and are extracted by means of an electrostatic field and directed onto a micro-channel plate detector. The induced electron avalanche at the position of the secondary ion impact generates a fast "STOP" signal. In addition, secondar, exercises produced at the target backside are accelerated and used to trigger "START" signais. Both the start and the stop signals are feed into a start-stop time-todigital converter (TDC) to generate TOF mass spectra of the secondary ions.

Nepheline, jadeite and anorthite targets were prepared *ex situ* at PUC-Rio. The different rocks were ground with a mortar and pestle and the grains were glued on carbon foils. With this technique, almost uniform thin films can be produced. The target thicknesses were typically ~ 1 µm determined by means of an optical microscope. In order to reduce surface contamination, the targets were heated *ex-situ*, (T = 60°C for 60 min) to allow evaporation of

water and other volatile contaminants and then reintroduced via a rapid airlock into the chamber. Finally the samples were irradiated at room temperature.

2.1 Heavy-ion impact irradiation

Cosmic rays are composed mainly of light ions (protons and α particles), but also of heavier ones (e.g. Fe and Xe). The main physicochemical phenomena that occur when an energetic (keV to GeV) ion impinges a solid are: compaction-amorphisation, electronic excitations (including ionization) in the bulk, chemical reaction—secondary electron emission and sputtering (secondary particle emission). Laboratory investigations about particle—solid interactions are numerous and have used charged energetic particles as projectiles (Brown *et al.* 1984; Baragiola *et al.* 2003; Strazzulla 2011; Raut & Faragiola 2013).

Experimental data from materials irradicted with MeV/u heavy ions is somehow scarce because of the technical difficulties in obtaining these ion beams. In this sense and for certain applications and studies (e.g. ion desorption yield measurements and mass spectrometry) the $^{252}_{98}Cf$ FF nuclei constitute an easily a ressible ion source. This particular radioactive nuclide is the most practical source due to its 2.64-yr half-life and affordable price (Macfarlane & Torgerson 1976).

The 252 Cf FF mass distribution has a characteristic camel-like shape with maxima at about 105 and 140 u (Nervik 1960; Kiesewetter *et al.* 1992; Schmidt *et al.* 2008). Thus, typically 105 Rh and 140 Ba, have energies of ~ 0.5 MeV/u and interact with the target atoms depositing energy mainly in the electronic regime. Because the Cf source is sealed for radiological safety reasons, the FF's kinetic energy is reduced by $\sim 25\%$ as they pass through the nickel capsule sealing. For the analysis and simulations here presented, the average kinetic energy per nucleon (E/m) of ~ 0.5 MeV/u was considered for the FF group. The electronic stopping powers for the three silicates irradiated with H, He, Fe and FF have been calculated

with the SRIM code¹ (see **Fig. 2**). Simulations show that Fe and FF projectile distributions are quite equivalent with a maximum stopping power (Bragg peak) at around $1x10^3$ eV (10^{-15} atoms⁻¹ cm²) or $7.6x10^2$ eV/Å. It is also important to note the stopping powers for H and He projectiles, at this energy, are around two orders of magnitude lower than for Fe or FF ones. **For heavy projectiles like Fe and FF, a small discontinuity feature is observed at 2.5 keV/nucleon, which is due to the SRIM calculation itself. This region is the limit of the TRIM calculation. For lower energies it is simply a straight line (in log-log) extrapolated to zero.**

3. SECONDARY ION FORMATION ON SUNFACES

Three silicate targets of astrophysical interest were prepared and irradiated with swift heavy multicharged ions. Anorthite is a naire ral silicate species of the tectosilicates subgroup, with formula CaAl₂Si₂O₈. It is the calcium end member of the plagioclase feldspar mineral series and may contain traces of Ti, Te, Na and K. Anorthite, on Earth, can be found in igneous rocks and is abundant on the Moon (Wenk *et al.* 1980, Smyth 1986, Wu *et al.* 2020). Also, feldspars, like anorthite, are the most plausible source of alkali metals in the exosphere of Mercury and may contribute to the calcium signal (Sprague *et al.* 2007, Wurs *et al.* 2010, Domingue *et al.* 2014).

Jadeite is a mineral species from the silicate family, a class of inosilicates, group of pyroxenes whose ideal chemical composition is NaAlSi₂O₆. It has a typical Mohs hardness 6.5 to 7.0, depending on the composition, with a specific gravity of about 3. This unusual high density was assumed be due to its occurrence in high-pressure sites; in fact jadeite is formed under high pressure (Harlow *et al.* 2015). In the early Solar System, such high-pressure polymorphs were formed by collisions and impact events on Mars and Moon surfaces (Ohtani

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¹ SRIM code version SRIM-2013.00 (http://www.srim.org).

et al. 2017), and on asteroids like Bennu² or the Chelyabinsk meteor, which microscopic examinations revealed rapid crystallization, meaning that the meteorite is the product of a collision of the original body (Ozawa et al. 2014).

Nepheline is a mineral species of the tectosilicate family, a group of feldspathoids: a silica-undersaturated aluminosilicate, containing sodium and potassium with the chemical formula (Na,K)AlSiO₄. Nepheline is well suited for simulating the surface of several objects in our Solar System. For instance, it is thought to be formed in mercorites parent bodies, by late thermal alteration event(s) of chondrules, as in the Allende CV2 meteorite (Lumpkin 1980, Shimoda *et al.* 2005, Ichimura *et al.* 2017). Also, among other silicate minerals discovered in Stardust Mission return samples, from comet 81P/Wild, tepheline was observed (Wooden *et al.* 2008). In planetary surfaces, nepheline also may be present. According to Emery *et al.* (1998), spectra analysis from Mercury's curtain indicates possible alkali-rich volcanic, like nepheline. They also show features common to igneous nepheline-bearing alkali syenite (Sprague 1998).

3.1. Sputtering of Positive Yons.

Positive ion mass spec ra from the silicate samples irradiated with energetic heavy ions (FF) are shown in **Fig. 3**. They are characterized by the presence of the three hydrogen peaks, $H^+ H_2^+$ and H_3^+ , being an unavoidable contaminant in every vacuum chamber. Another characteristic peak in all positive spectra is at 23 u, which is assigned to the Na⁺ ion.

In particular, nepheline spectrum is characterized by mass peaks at 39 u and 63 u which can be assigned to K^+ and $(Na_2O)H^+$. A small peak is observed for $A1^+$ at 27 u. Also a cluster series of molecules due to atom recombination upon ejection can be observed at 81 u, 97 u,

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² www.nasa.gov/feature/goddard/2002/osiris-rex-unlocks-more-secrets-from-asteroid-bennu

113 u, 129 u and 145 u mass peaks assigned to $Al_3O_n^+$ series, with n=0 to 4. For the peak at mass 103 u, the $AlSiO_3^+$ species is a plausible candidate.

On the other hand, anorthite and jadeite positive mass spectra are less rich in species. Besides the characteristic peaks, some others can be observed like the Al^+ , K^+ and Ca^+ for anorthite, and Al^+ , K^+ , $NaAl^+$ and $(Na_2O)H^+$ for jadeite. For silicates containing Na in its structure, the species $(Na_2O)H^+$, at 63 u, seems to be ubiquitous.

3.2. Sputtering of Negative Ions

Negative ion mass spectra are more abundant in species than positive ones. **Fig. 4** shows the mass spectra, from 50 to 185 m/z, for the three species. Interesting sputtered species can be observed in the three spectra, like the silicon dioxide (SiO₂) forming a cluster series, the $(SiO_2)_n$ with n=1 to 3, being important for an the three silicates. Another example of sputtered species is the aluminum silicon molecule linked to oxygen atoms, forming the cluster series $(AlSi)O_m$, with m=0 to 8, being more important for nepheline. The $Al(SiO_2)_2O_2$ molecule is also observed in the three spectra at 179 u.

In addition, each spectrum shows specific species for each silicate. For instance, the $(SiO_2)_2O^2$ molecule is owner ed when anorthite and jadeite targets are irradiated, while the anorthite spectrum shows the CaO_2^- specific species.

3.3. Sputtering Yields

Regarding the mechanism of ion induced desorption from planetary surfaces, the total desorption yield (Y_{tot}), which is defined as the number of molecules, neutral or ionized, ejected per impact, results from a sequence of events after the ion-surface interaction. Energetic heavy ions with velocities higher than the Bohr velocity (25 keV/u) transfer most of their energy to the target electrons. In insulators, the electronic excitations can lead to repulsive states, which cause, at the target surface, particle ejection. The PDMS-TOF-SIMS

technique is suited for the fragmentation pattern study during prompt induced desorption (i.e. sputtering), which occurs on time-scales of $\sim 10^{-11}$ s after the ion impact.

Table 1 shows the absolute desorption yields (number of desorbed ions per impact) for the most relevant positive and negative ion species. In addition, **Fig. 5** shows bar spectra of absolute desorption yields for positive ions, for each silicate, with mass-to-charge ratio up to 70 m/q. Since the majority of desorbed species have q = 1, the ratios m/q are in practice substituted by the ion masses, generating mass spectra. To estimate the absolute desorption yields for each discrete m/q, a flux of 60 ± 2 FF impacts s⁻¹ in the solid angle of the sample was used. Thus an analysis of a number of fragments of the cationic and anionic species, produced by ionization, dissociation and recombination processes, was performed.

The number of electronic excitations is proportional to $S_e = (dE/dx)_e$ (electronic stopping power). When the excitation d insity is high, cooperation between close excitation events along the nuclear track enhance the sputtering yields. Based on experiments, a power law $(Y \propto S_e^n)$ dependence of the sputtering yield (Y) on the energy deposition per unit path length (S_e) has been found. The range of reported powers (n) is wide, from 1 to 6, and they may be different for ejected for and neutrals. For instance Brown et al. (1984) and Shi et al. (1995) have shown the rundiatic dependence of the total (ionic and mainly neutral) sputtering yield (Y_{tot}) on the energy deposition per unit path length (S_e) , with MeV light ions (mainly using water ice-targets). Seperuelo-Duarte et al. (2010) extended the sputtering yield measurements on CO ice to heavy ion beams (50 and 537 MeV Ni), confirming the power law $Y_{tot} = a S_e^2$ for four order of magnitudes on S_e and six on Y. Regarding only ion yield dependence, de Barros et al. (2011), through H_2O ice irradiations, found a third power dependence of ion-sputtering yields on S_e . Concerning other materials, Toulemonde et al. (2020) have shown a quartic dependence of the total sputtering yield (Y_{tot}) of lithium fluoride

with swift heavy ions as projectiles, and Hijazi *et al.* (2014) a quadratic dependence for secondary ions (low Z projectiles), i.e. $Y_{ion} \propto S_e^2$.

Fig. 6 (upper panel) shows an estimate of the ion-sputtering yields for anorthite, jadeite and nepheline; considering the *quadratic power dependence*. The inset plot shows also the experimental values. The ion beam used for these simulations is the ¹⁴⁰Ba (0.46 MeV/u), which is considered as representative of the heavy FF mass group. Vertical arrows indicate the average energy per nucleon. Thus, **Fig. 6** shows the corresponding total ion sputtering yield occurring at E/m = 0.46 MeV/u, for ¹⁴⁰Ba, assuming the qualitatic power law to be fixed in this energy regime, related to the physical process of the electronic sputtering. The factor a in $Y_{ion} = a S_e^2$ is determined by using the experimental data and used to fine-tune the SRIM data of Fig. 2, to all energies. It varies for each target, but for silicate materials is approximately the same, being on the order of 5.0×10^{-9} (Å/eV)². The lower panel of Fig. 6 shows three simulations of the nepheline ion sputtering yield (Y_{ion}) dependence on the energy deposition per unit path length (S_e) to having the power law model $(Y_{ion} \alpha S_e^n)$. In fact, ion irradiation experiments with one silicate at several ion energies are being carried out. This will let us determine the right dependence of Y_{ion} on S_e , for silicates.

It should be noted that the same trend observed in **Fig. 2** is observed here as well, which means that heavy ions in the MeV/u range around the Bragg peak are several orders of magnitude more efficient for inducing desorption than protons and α particles, at the same velocity (E/m). This indicates that the very high desorption yields of MeV/u heavy ions must be taken into account for silicates chemical processing and space weathering.

4. ASTROPHYSICAL IMPLICATIONS

Space weathering of surfaces due to irradiation effects plays an important role in the

composition/modification of exospheres, mainly in objects where its magnetic field is weak or does not exist. Mercury and the Moon possess very thin atmospheres (exospheres) and magnetospheres weak enough to allow energetic charged particles to impinge directly into their surfaces depositing energy and physically altering materials present on them (Wurz *et al.* 2010, Paral *et al.* 2010, Milillo *et al.* 2010, Domingue *et al.* 2014). Thus space-weathering processes are tied to the exposure of a planetary surface to its space environment, generating and maintaining a surface-bounded exosphere. In this scenario, desorption of neutral and ionic species plays an important role. The current results concern the ionic contribution.

4.1. Sputtering in Planetary Surfaces

Desorbed species due to sputtering contribute of the exospheres, reflecting - in some extent - the surface composition. Particularly the presence of low energy ions in the exospheres near the surface of airless planetary bodies such as the Moon, Mercury, and asteroids should be largely indicative of the planetary surface elemental composition (Johnson and Baragiola 1991, Dukes and Baragiola 2015, Martinez 2017).

a) Effects on the Moon

The Moon has a reasonable gravity and no magnetic field; its surface is subjected to solar wind, solar energetic particles, and galactic cosmic ray irradiation, with a much higher flux than asteroids in the main belt. The lunar exosphere is a region of space above the surface reaching an altitude of ~100 km, with an atomic density as low as 10⁶ cm⁻³ (Dukes and Baragiola 2015). Via mass spectrometer measurements of the AMPTE, WIND and SELENE missions, charged species have been observed in lunar exosphere. They have locally identified ions such as Ar⁺, Na⁺, K⁺, Si⁺, Al⁺, C⁺, and O⁺ (Hilchenbach *et al.* 1993, Stern 1999, Yokota *et al.* 2009, 2014). More recently, measurements by the LADEE mission identified significant detections of lunar ions with mass per charge of 2, 4, 12, 20, 28, 39, and

40, moderate detections with 14 and 23, and weak detections with 24, 25, and 36 (Halekas *et al.* 2015). For these experiments, they used an outward pointing viewing geometry, ensuring that these ions originate from the exosphere.

Thus, determining lunar sputtering yields of ionic species is well justified. According to Dukes *et al.* (2015), sputtered ions may be more important in lunar exosphere in the sense that they continue to interact with the lunar surface by charging or neutralizing regions of the regolith and by self-sputtering (Poppe *et al.* 2013). Additionally, cation and anion yields are richer as they show molecular series. The Moon's exosphere hay depend on this too, in particular silicon oxides series, as shown later.

Considering that composition and stoichiometry of the targets are known and that, in first approximation, the sputtered species are proportional to the initial stoichiometry (Dukes et al. 2011), the sputtering yield of a species would represent a percentage of the initial (total) concentration, indicating the loss (depletion) of a particular element of the sample (Williams 1984, Dukes et al. 2011). In addition, the ion fraction of the sputtered species must be taken into account in this analysis. Neutrals are expected to be more abundant than ions. While sputtering measurements and model predictions for oxides and minerals have shown ionic fraction values close to 1, for several species (Benninghoven et al. 1972, Andersen et al. 1972). Similar SIMS measurement results were obtained by Elphic et al. (1991) by irradiating lunar soil analogs. Steinbrüchel et al. (1980, 1981) measured ion fractions of 0.4–0.8 from metal oxides, and more recently Dukes et al. (2011) have obtained results well in between by irradiating tectosilicates. A fraction of 0.5 will be considered here.

The PDMS-TOF-SIMS analyses of lunar soil analogs show an important surface depletion of Na⁺. For Jadeite (with 10% of Na by number), it is observed ~3.6% surface depletion ($Y_{ion} = 0.363$ Na ions impact⁻¹). And if we consider the, sputtered but not detected, Na⁰ species, the Na depletion increases to ~7.2%. For Nepheline (15% Na by number), a

~3.1% depletion is observed (0.251 ions impact⁻¹); and considering the neutral fraction, an erosion of ~6.2% is found; 14% less than for Jadeite. Na is not supposed to exist in Anorthite, but this mineral shows a very small emission yield of Na⁺ (0.0417 ions impact⁻¹) because it is a common impurity. Aluminum depletion is expected to be quite the same for the three silicates samples; Anorthite (with 15% of Al by number) shows a relative important surface depletion of ~0.51% (0.0340 ions impact⁻¹), ~1.02% taking into account neutrals. And Jadeite (10% Al by number) Al⁺ is depleted by $\sim 0.14\%$ (0.0138 ions impact⁻¹), 0.28% with neutrals; ~70% less depletion than the former. But interestingly, Nephence (12.5% Al by number) shows very low Al⁺ emission, of the order of 0.0037 ion mact⁻¹, implying a depletion of 0.05% (0.1% including neutrals); ~10 times lower than the one for Anorthite, for instance. This may indicate that planetary surfaces containing 'his kind of aluminosilicate will maintain exospheres with low quantities of Al. Concerning potassium, the ion depletion occurs for Nepheline (12% of K by number) samule only, with a relative high ~1.28% (0.1023 ions impact⁻¹) or ~2.56% considering neutrols. Anorthite and Jadeite samples show low emission yield of potassium ion since it is a common impurity. Similarly, Ca⁺ ion is only observed for Anorthite (7.7% of Ca by nu. ber) targets as expected, with a moderate ion depletion of $\sim 0.25\%$, or $\sim 0.5\%$ with neutron (0.0302 ions impact⁻¹).

b) Effects on the Me cury planet

Mercury has a relative strong gravity field. The gravitational acceleration at the surface is approximately 3.697 m s⁻², more than twice the value on the Moon (Wurz 2010, Paral 2010). Its magnetic field intensity is about 1% of the Earth's field intensity at the surface (Takahashi 2019). Thus, Mercury also presents an interesting environment to study and understand the interactions between the SW, SEP, and GCR, planetary surfaces and surface-bounded atmosphere as mediated by a not so strong internal magnetic field (Milillo *et al.* 2005). Although there is no direct information available about how space weathering affects

Mercury's regolith, like returned samples (Domingue *et al.* 2014), Zurbuchen *et al.* (2008) suggest clues on the composition of Mercury's ionized exosphere: the Fast Imaging Plasma Spectrometer (FIPS), the low energy portion of the Energetic Particle and Plasma Spectrometer (EPPS), detected Na⁺, O⁺, and K⁺ species, among other possible ions and molecular ions, consistent with expectations from observations of neutral species (Zurbuchen *et al.* 2008, Paral *et al.* 2010).

The PDMS-TOF-SIMS analyses of analogs for the Herr can soil shows relatively high sputtering yields of Na⁺ and K⁺ species, depending on the a valog. As stated by Dukes *et al.* (2015) and others, sodium and potassium are the predominant species sputtered from the Mercury surface (Potter *et al.* 1986, 1987, 1997, Gamborino 2019). For the first analog, Anorthite, those two species are natural contaminants. However, it could be a good source of ionized calcium Ca⁺, since it has already been detected in the Hermean exosphere by the MESSENGER mission, with abundances comparable to that of K⁺ (Zurbuchen *et al.* 2008). The Ca⁺ sputtering yield here measured is of the order of 0.0302 ions impact in, which is lower than necessary to explain the observation by Zurbuchen *et al.* (2008). In fact, Ca should be released less efficiently than in and K due to its bonds with oxygen atoms in the bulk/surface. On the other hand, the Jadem's sample shows an interesting Na⁺/Al⁺ ratio of ~26.3. The Na/Al ratio (neutral atom ratio) vas found to be between 1.3 and 1.7 (Wurz *et al.* 2010, Domingue *et al.* 2014). Comparing to the ionic ratio, those values are in agreement if we consider that Na is more likely to be ionized: the Al ionization energy (5.99 eV) is higher than that of Na (5.14 eV). On the contrary, Zurbuchen *et al.* (2008) did not report any observation of Al⁺.

Concerning the sodium and potassium emission, the spatial distribution of the Na/K exospheric column density ratio is temporally variable due to several factors like the interactions of solar wind and magnetosphere. In this sense, ratios already observed or calculated vary in a wide range with values ranging from ~26 to ~300 (Doressoundiram *et al.*)

2010, Leblanc *et al.* 2011, Domingue *et al.* 2014). Observations of Na⁺/K⁺ ratio reported by Zurbuchen *et al.* (2008) found a value of ~2.3, which is in very good agreement with the ~2.5 value that we measured for nepheline (see **Table 1**). This value is also in agreement with calculations made to estimate densities of species in the exosphere of Mercury; Wurz *et al.* (2010) found ionization Na⁺/K⁺ ratio of 2.4. This phenomenon may be due to the Na higher ionization energy.

4.2. Negative Ions in Exospheres

The formation processes of negative ions at surfaces relevant to astrophysical environments are not well studied yet. Moreover, collisional electron attachment processes would not be considered important, as pressures in most environments are far too low for this process to be significant (Millar *et al.* 2017). Here ever, particle-stimulated emission of anions from surfaces has been observed and studied in laboratory, at low temperatures and ultra high vacuum (e.g. de Castro *et al.* 1997, Andrade *et al.* 2008, Martinez *et al.* 2012, 2014, Oliveira *et al.* 2021).

Different mechanisms for the formation of negative ions have been proposed. Probably the most important is the excitron attachment, initiated by free electrons (Millar *et al.* 2017). Indeed, when heavy mult charged ions (e.g. cosmic rays, FFs) interact with a solid, emission of secondary electrons occur together with neutral and ionic species (Iza *et al.* 2005, Martinez *et al.* 2006); a plasma is temporary formed, favoring electron attachment (Macfarlane & Torgerson 1976, Oliveira *et al.* 2021). In other words, electron attachment cross sections are enhanced on the surface in part because of multiple scattering of electrons and adjacent species.

Laboratory results have demonstrated, for different materials, that cationic species desorption yields are much higher than anionic ones, at around two or three orders of

magnitude greater (Farenzena *et al.* 2006, Martinez *et al.* 2014). On the other hand, the here obtained absolute desorption yields Y_i (positive and negative ions, and the sum of all of them; see **Table 2**), show that, when silicate targets are irradiated by energetic heavy multicharged ions (cosmic rays analogs), high emission rates of SiO_2^- and O_1^- are observed, turning almost comparable the sputtering yields of cations and anions. Silicates are built of the same basic structural units, the SiO_4 tetrahedra. Metal cations in the silicate network leads to a partial destruction of the O bridges and the formation of nonbridging O (Henning 2010). This fact and the electron affinity of silicon dioxide (0.9 eV) favor the outstanding emission of SiO_2 and O anions (Williams 1965, Ballarotto *et al.* 2002). A coordingly, the three samples analyzed in this work confirm the high emission yields of SiO_2 and O mainly observed as aggregates or ion cluster series $(SiO_2)_n^-$ and $(AlSi)O_1^-$, as presented before. Molecular cluster emission from surfaces via sputtering, with a rephysical implications, has already been discussed before (Killen *et al.* 2005, $^{10}O_1^-$, Martinez *et al.* 2017). Recently, molecular aggregates of water (cosmic water na occlusters) have been considered to explain several processes occurring in the Universe (J hason 2021).

Mainly after the collected samples during the Apollo missions, Moon surface composition became very we'll known (Ringwood 1976, Dukes *et al.* 2015). Analyzes of those samples have also revealed that silicon dioxide is one major component, others being A1₂O₃, CaO and Na₂O. Moreover, Gu *et al.* (2018) reported the analysis of Apollo 15 soil grains by transmission electron microscopy, displaying silicon dioxides *nanoparticles*. Nevertheless, they state that those nanoparticles were not formed due to solar-wind generated vapor deposit or irradiated rim. In the current work, high emission yield of SiO₂ molecular clusters is observed when silicates are irradiated by heavy ions, which may indicate a molecular cluster redeposition. As mentioned before, the Moon's exosphere was analyzed by the LADEE spacecraft (ARTEMIS mission) and several lunar cations were identified (Halekas *et al.* 2015), proving an ionic exosphere. Nevertheless no anion was reported.

On the contrary, no returned samples or other direct information is available about Mercury's regolith and how it was affected by space weathering. In fact, Mariner 10 and MESSENGER missions and ground-based observations have given us information about surface and exosphere compositions. Ground-based observations of Mercury's surface indicate a heterogeneous surface composition with SiO₂ content ranging from 39 to 57 wt%. Also it is thought to be covered with highly space-weathered silicate material (Sprague *et al.* 2007). Fly-bys of the MESSENGER spacecraft provided insight into the spatial distribution of the heavy ion exosphere around the planet (Paral *et al.* 2010). More specifically, the Si⁺ abundance observed is too high, which means its source is either Mercury's surface or its exosphere (Zurbuchen *et al.* 2008). Likewise, no negative ions were reported until now. In other respect, sputtering of atomic or molecular cluster have not been observed directly in Mercury's exosphere, but it has been propose that a significant contribution to the Ca exosphere arises from sputtered CaO makerales (aggregates) (Killen *et al.* 2005). This is in agreement with the present results concerning Anorthite sample, from which CaO₂ ions were emitted (CaO + O' = CaO₂).

4.3. Ionic Sputtering Nate

The sputtering rate of Snic species by cosmic rays $R_{S,CR}$ of planetary surfaces analogues can be determined by adapting the analysis made by Mennella *et al.* (2003) and others (Godard *et al.* 2011, Dartois *et al.* 2017). Thus, to extend and apply the current results to astrophysical environments, like planetary surfaces, the total ionic sputtering yields ($Y_{i,total}$) must be summed over the cosmic ray ion abundances and energy distribution contributions:

$$R_{S,CR} = \sum_{Z} \int_{E_{min}}^{\infty} Y_{i,total}^{CR}(E,Z) \frac{dN}{dE}(E,Z) dE \quad (1)$$

where the second term of the integral is the differential flux of the cosmic ray element of atomic number Z, with a cutoff in energy E_{min} set at 100 eV/u. The differential cosmic ray flux

is given by the functional form given by the spectral shape data obtained from four instruments on NASA's Advanced Composition Explorer (ACE) (Mewaldt *et al.* 2001, 2007). Their data are empirically fitted by:

$$\frac{dN}{dE}(E,Z) = A_1 e^{-\eta \frac{E}{m}} + \frac{A_2}{\left(\frac{E}{m}\right)^2}$$
 (2)

where A_1 and A_2 are normalizing constants at a distance of 1 au (= 5.0×10^8 MeV⁻¹ cm⁻² s⁻¹ and 2.0×10^{-2} MeV u^2 cm⁻² s⁻¹ respectively), $\eta \sim 2.4 \times 10^3$ u MeV⁻¹, E/m is the cosmic ray kinetic energy in MeV per nucleon, and m is the ion's mas: (de Barros et al. 2011). Fig. 2 displays SRIM code predictions for the expected stopring rower (S_e) as a function of the atomic number Z and of the energy per nucleon E/n, and Fig. 6 (upper panel) the relation established in Section 3.3, linking the total ionic spattering yield $Y_{i,total}(E,Z)$ to $S_e(Z,E)$ through the relation $Y_{ion} = a S_e^2$. This allows determination of the total sputtering yield of ionic species when silicates are irradiated by cosmic ray particle analogues. By applying Eq. (1), the sputtering rate as a function of the cosmic particle (i.e., galactic CR and Solar particles) energy was determined as showed in Fig. 7, which is the product of cosmic particle flux times the total sputtering yield. At 1 au the differential flux of cosmic particles (galactic cosmic rays, solar wind, solar energetic particles) is given by the functional form given by the spectral shape data obtained from the instruments on NASA's Advanced Composition Explorer (ACE) (Mewaldt et al. 2001, 2007). The curves plotted in Fig. 7 are the result of the product of the cosmic particle flux times the total sputtering yield. The huge peak at E/m < 5 keV/u, reflects the fact that at these energy values the flux is very high (due to solar wind ions) but the sputtering yield is low. However we have to keep in mind that in the keV regime the nuclear stopping power is higher than the electronic stopping power and the relation $Y_{ion} = aS_e^2$ might no longer be valid. As pointed out by Martinez et al. (2019), the dependence of the sputtering yield on the total stopping power and, in particular, the contribution of the nuclear stopping power is not yet well known and

further experiments are needed to be fully understood. In Eq. (2) the parameters A_I and A_2 include the dependence of the flux on Z. The values of A_I and A_2 adopted in our calculations are taken from Table 9 in de Barros *et al.* (2011) where the A_I and A_2 values for various ions are reported taking into account their different abundance. In the present work, FF "simulate" Fe projectiles. To see the effects in space, we take into account the Fe abundance in space and correct the yield effects by its dependence on the higher stopping power of FF in relation to Fe.

Results for the three silicate samples are very similar that seem to merge in one curve, for each projectile (Fe and FF). At cosmic particle energies of 0.46 MeV/u, silicate ionic sputtering rate $R_{s,CR} = 6.3 \times 10^{-2}$ particles cm⁻² s⁻¹ is found. The corresponding sputtering timescales ($\tau = 1/R_{s,CR}$) associated with the destruction rates for E = 0.46 MeV/u are around $\tau \sim 15.9 \text{ s cm}^2$ ion⁻¹. Astrophysical implications for the evolution of planetary silicates subjected to cosmic rays, providing information on potential species to be released, show that silicates dust grains are relatively resilient against sputtering by cosmic rays.

The sputtering rate here estimated is valid for the Moon's surface (1 au). The evaluation of the sputtering rate at Mercury's surface (0.39 au) is more complex. In fact the energy spectrum and the flux of galactic cosmic rays is approximately the same everywhere in the Solar System. Also the energy spectrum of solar wind does not change but its flux varies as the inverse of the square of the distance from the Sun being then much larger at Mercury than at the Moon. In addition, Mercury's magnetic field partially inhibits the direct access of solar energetic ions to the surface. Then, sputtering rates and timescales on Mercury will be estimated when reliable data on cosmic particle flux on the surface of Mercury become available.

5. CONCLUSIONS

In order to determine ion-sputtering yields of astrophysical silicates bombarded by solar wind, solar energetic particles, and galactic cosmic rays, laboratory experiments with silicate analogs have been performed. Anorthite, Jadeite and Nepheline were exposed to ²⁵²Cf fission fragments and the absolute ion desorption yields have been measured and analyzed by the PDMS-TOF-SIMS technique. Current results show several ionic species emission due to this interaction, which is dependent on the irradiated silicate (see **Table 1**).

Desorbed species, neutral and ionic, contribute to exost bases, also reflecting - in some extent - surface composition. In this sense, the current results may represent sputtering in planetary surfaces. In fact, the observed ion species and the ion sputtering yield, Y_{ton} , refer to the magnitude of this depletion, which is sample dependent. Indeed, PDMS-TOF-C1M3 results from Anorthite sample, analogue of the Moon's crust, exhibit comparable and relative moderate depletions of Al⁺ and Ca⁺ (~0.51% and ~0.25% respectively). On the other hand, Jadeite and Nepheline reveal low and very low depletion rates of Al⁺, respectively. (~4 to ~10 times lower than the one for Anorthite). This indicates that planetary surfacts containing this kind of aluminosilicates will maintain exospheres with low quantities of Al⁺. Observation of Na⁺ in lunar exosphere indicates that others silicates must be present. Jadeite proves important surface depletion of Na⁺, with ~3.6% surface depletion ($Y_{ton} = 0.363$ ions impact⁻¹). Similarly, Nepheline shows a ~3.1% depletion of Na⁺ at the surface. The observation of the sodium cation is in agreement with initial measurements of the AMPTE, WIND and SELENE missions and, more recently, with measurements by the LADEE mission.

The PDMS-TOF-SIMS analyses of analogs for the Hermean soil evidence a Na⁺/Al⁺ ratio of ~26.3. This is unexpected because the Na/Al ratio (neutral atom ratio) was found to be between 1.3 and 1.7 (Wurz *et al.* 2010, Domingue *et al.* 2014). Comparing to the ionic ratio,

those values may be in agreement if we consider that Na is more likely to be ionized. Unexpectedly, Zurbuchen *et al.* (2008) did not report any observation of Al^+ . Concerning the sodium and potassium emission, observations of Na^+/K^+ ratio reported by Zurbuchen *et al.* (2008) found a value of ~2.3, which are in very good agreement with the ~2.5 current results measured with Nepheline. This value is also in agreement with calculations made to estimate densities of species in the exosphere of Mercury; Wurz *et al.* (2010) found an ionization Na^+/K^+ ratio of 2.4.

The three analyzed samples confirm the high anion emission yields of SiO₂ and O mainly observed as aggregates or ion cluster series (SiC₂)_n and (AlSi)O_m. This could be explained by the sputtered species electron affinity and by the temporary plasma formation when heavy multicharged ions (e.g. cosmic rays FFs) interact with a solid surface (emission of secondary electrons together with neutral and ionic species). This enhances electron attachment cross sections on the surface, is part because of multiple scattering of electrons and adjacent species. Gu *et al.* (2019), reported the presence of silicon dioxides *nanoparticles* in Apollo 15 soil grain samples The emission of (SiO₂)_n aggregates suggests a molecular cluster redeposition in the form of "nanoparticles". Nevertheless no anion was observed yet; neither on the lunar nor (n the Hermean surfaces.

The total ion yield $(Y_{t,ion})$ was also determined by means of the quadratic power dependence of ion-sputtering yields on S_e , i.e. $Y_{ion} = aS_e^2$ (**Fig. 6**). Sputtering rate dependence on the cosmic ray energy is analyzed by combining the total ion yield $Y_{t,ion}(E,Z)$ and the differential cosmic ray flux dN/dE(E,Z) given by the spectral shape data. Results for the three silicates irradiated with Fe and FF projectiles show comparable values and seem to merge in only one curve for each beam. Silicate sputtering rate of 6.3 x 10^{-2} particles cm⁻² s⁻¹ at cosmic ray energies of 0.46 MeV/u is found. The corresponding sputtering timescales is around $\tau \sim 15.9 \text{ s cm}^2$ ion⁻¹. These results indicate that the evolution of planetary silicates subjected to

cosmic rays, like silicates dust grains, are relatively resilient against sputtering by cosmic rays.

Ion irradiation experiments with one silicate at several ion energies are being carried out. This will let us determine or confirm the dependence of Y_{ion} on S_e , for silicates. In addition other silicates will be irradiated to confirm the anion emission values and its implication for astrophysical environments.

The presented results would be useful in understanding the results expected by the European space mission BepiColombo that will reach Mercery in December 2025. In fact among the scientific instrumentation on board the spacecraft there is SERENA 'Search for Exospheric Refilling and Emitted Natural Abundances'. It will investigate the complex particle environment of the Mercury'-s exosphere including thermal and directional neutral ionized species (see e.g. Orsini *et al.* 2016).

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Table 1: Absolute prompt desorption yields Y_{ion} of the most relevant ionic species desorbed from the silicate samples at room temperature after 252 Cf fission-fragment impacts.

SILICATE	Positive Ions			Negative Ions		
	m/q	Species	Ions impact ⁻¹	m/q	Species	Ions impact ⁻¹
Anorthite	1	H^{+}	0.2300	60	SiO ₂	0.03278
	23	Na^+	0.0417	120	$(SiO_2)_2$	0.00276
	27	Al^+	0.0340	180	$(SiO_2)_3$	0.00156
	39	K^{+}	0.0654	71	Al. iO	0.00291
	40	Ca^+	0.0302	87	$F1S1C_2$	0.00129
				103	A.SiO3	0.00236
				115	AlSiO ₄	0.00810
Jadeite	1	H^{+}	0.1338		SiO ₂	0.0337
	23	Na^+	0.363.	120	$(SiO_2)_2$	0.0034
	27	Al^+	0.0138	71	AlSiO	0.0054
	39	K ⁺	0.3374	87	AlSiO ₂	0.0010
	50	NaAl ⁺	0.0058	103	AlSiO ₃	0.0037
				119	AlSiO ₄	0.0207
Nepheline	1	F.	0.1026	60	SiO ₂	0.0460
	23	Na '	0.2511	120	$(SiO_2)_2$	0.0028
	27	Al^+	0.0037	180	$(SiO_2)_3$	0.0021
	2)	K^{+}	0.1023	Σ	(AlSi)O _n	0.0750
	46	$N{a_2}^+$	0.0072			
	Σ	$Al_3O_n^{^+}$	0.0777			

Table 2: Absolute prompt desorption yields Y_i of positive, negative and total species desorbed from the silicate samples at room temperature after ²⁵²Cf fission-fragment impacts.

SILICATE	$Y_{i,pos}$	$\mathbf{Y}_{\mathrm{i,neg}}$	$\mathbf{Y}_{i,total}$
Anorthite	0.56098	0.05401	0.61449
Jadeite	0.60177	0.06788	0.66965
Nepheline	0.67051	0.12585	0.79636

FIGURE CAPTIONS

- **Figure 1.** Schematic representation of the PDMS-TOF-MS experimental set-up. The ion beam impinges perpendicularly from behind on the silicate film. The sputtered ions are forward ejected.
- **Figure 2**. Electronic stopping power for each silicate: Anorthite, Jadeite, Nepheline; calculations were performed with the SRIM code. H, He, Fe and FF are considered as projectiles: Fe and FF projectile distributions are quite equivalent with a Bragg peak maximum at around $7.6x10^2$ eV/Å, for the three silicates.
- **Figure 3**. Mass spectrum of positive secondary ions desorted from Anorthite, Jadeite and Nepheline, irradiated by energetic multicharged heavy ons (FF at 0.46 MeV/u) at room temperature.
- **Figure 4**. Mass spectrum of negative secondary ions resorbed from Anorthite, Jadeite and Nepheline, irradiated by energetic multicharged neavy ions (FF at 0.46 MeV/u) at room temperature.
- **Figure 5**. Positive Ion Yield (ions/impact) bor spectra for the three analyzed silicates and for the principal atomic and small molecular species, with mass-to-charge ratio up to 70 m/q. They have been obtained from TOF mass spectra.
- **Figure 6.** Upper panel: Estimative of the ionic sputtering yields (Y_{ion}) after the impact of typical ²⁵²Cf fission fragments on Anorthite, Jadeite and Nepheline as a function of the projectile velocity, E/n; considering the quadratic power dependence $Y_{ion} \propto S_e^2$. The inset plot shows experimental values. Lower panel: Ionic sputtering yield (Y_{ion}) simulations for nepheline showing the energy deposition dependence per unit path length (S_e) , according to the power law model $(Y_{ion} \propto S_e^n)$, for three values of n.
- **Figure 7**. Sputtering rates as a function of cosmic ray energy for a typical ²⁵²Cf fission fragments and Fe projectiles. Results for each projectile are very similar.

FIGURES

Fig. 1

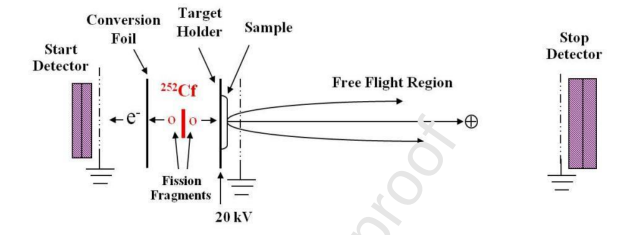


Fig. 2

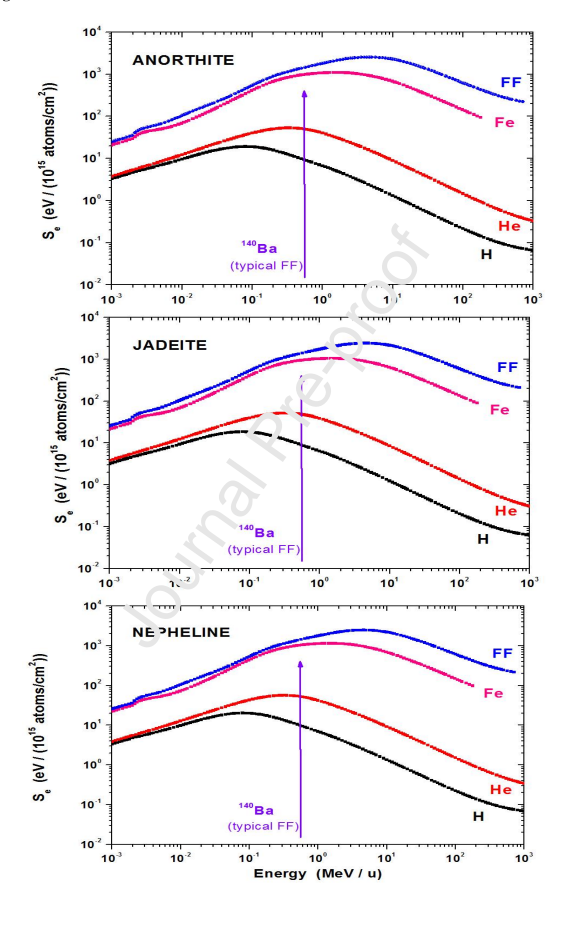


Fig. 3

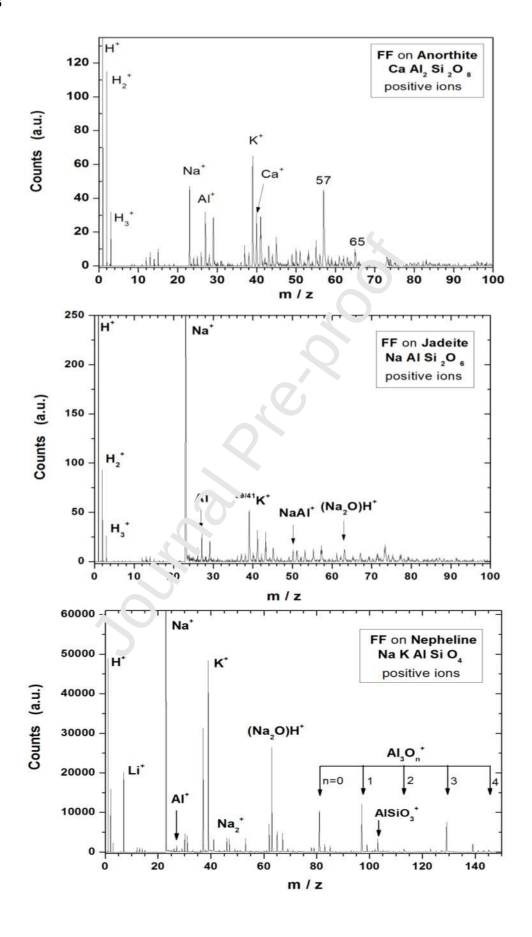


Fig. 4

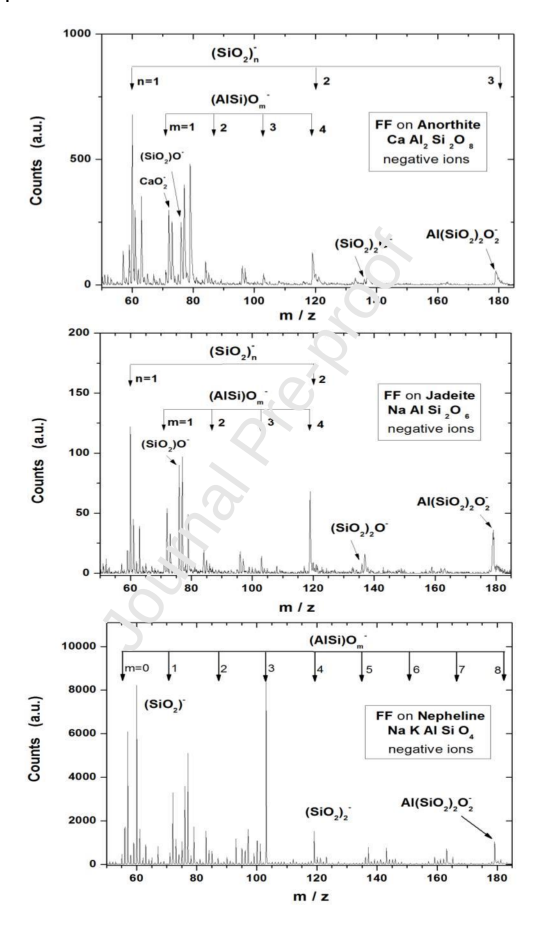


Fig. 5

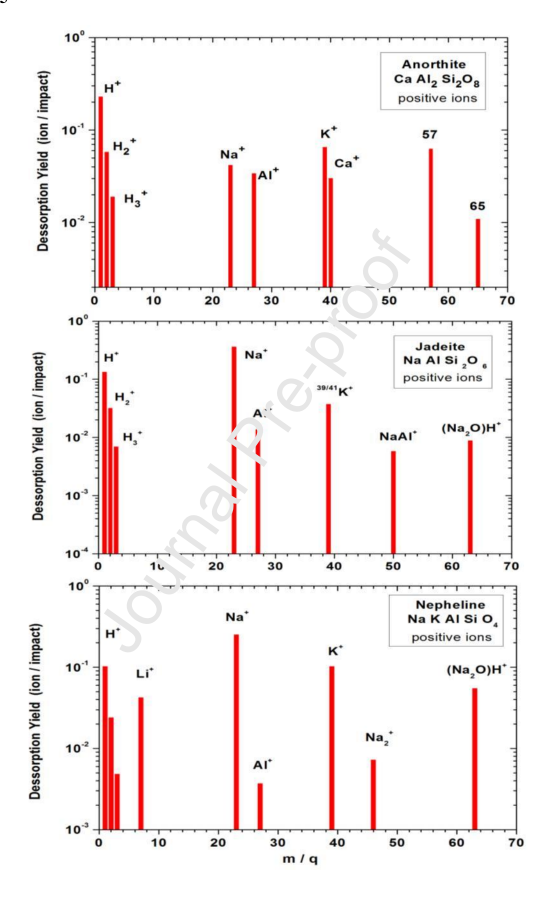


Fig. 6

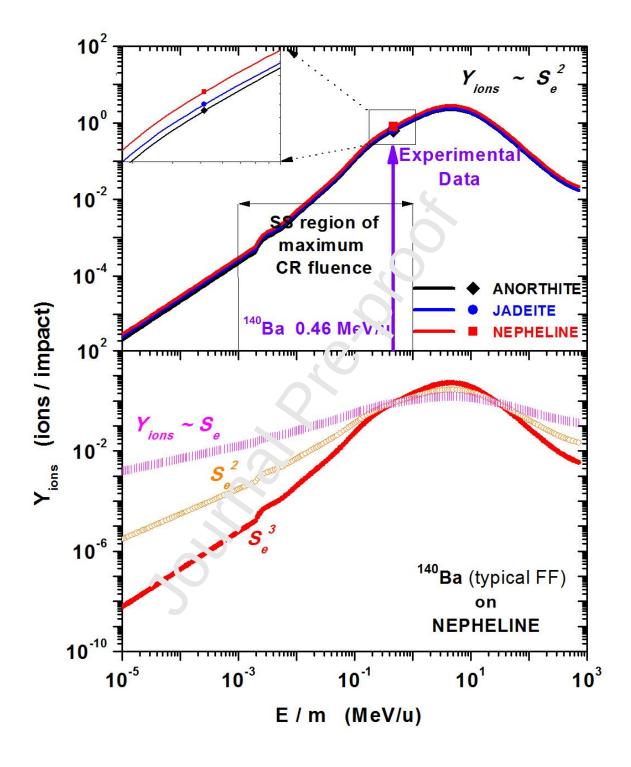
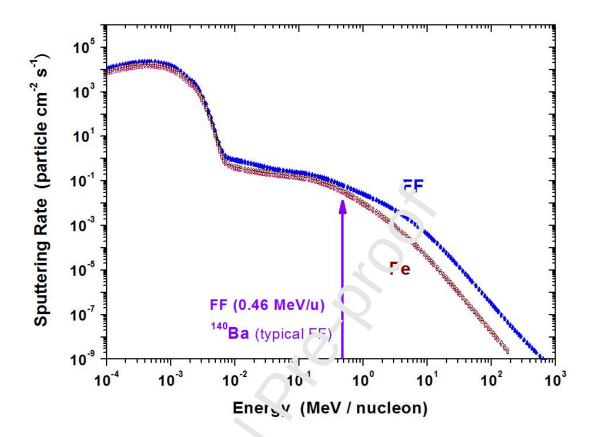


Fig. 7



Highlights

- At high energies, swift heavy ion irradiation deposits energy on silicate surfaces efficiently.
 Due to this interaction, several species are emitted from the surface.
- Low aluminum ion emission may indicate that planetary surfaces containing aluminosilicates
 (like nepheline) will maintain exospheres with low quantities of Al.
- A Na+/K+ ratio of ~2.5 was obtained for nepheline, which is in very good agreement with the one measured by the Messenger Mission, of around ~2.3. It is also in agreement with calculations made to estimate densities of species in the exospi are of Mercury, they found a Na+/K+ ratio of 2.4.
- The corresponding sputtering rates and timescales determined in the current work, indicate that the evolution of planetary silicates subjected to cosmic rays are relatively resilient against sputtering by cosmic particles