

UV Irradiation and Near Infrared Characterization of Laboratory Mars Soil Analog Samples: the case of Phthalic Acid, Adenosine 5'-Monophosphate, L-Glutamic Acid and L-Phenylalanine Adsorbed onto the Clay Mineral Montmorillonite in the presence of Magnesium Perchlorate

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Abstract

The search for molecular biosignatures at the surface of Mars is complicated by an intense irradiation in the mid- and near- ultraviolet (UV) spectral range for several reasons: (i) many astrobiologically relevant molecules are electronically excited by efficient absorption of UV radiation and rapidly undergo photochemical reactions; (ii) even though the penetration depth of UV radiation is limited, aeolian erosion continually exposes fresh material to radiation; and (iii) UV irradiation generates strong oxidants such as perchlorates that can penetrate deep into soils and cause subsurface oxidative degradation of organics.

As a consequence, it is crucial to investigate the effects of UV radiation on organic molecules embedded in mineral matrices mimicking the martian soil, in order to validate hypotheses about the nature of the organic compounds detected so far at the surface of Mars by the *Curiosity* rover, as well as organics that will be possibly found by the next rover missions *Mars 2020* and *ExoMars 2020*. In addition, studying the alteration of possible molecular biosignatures in the martian environment will help to redefine the molecular targets for life detection missions and devise suitable detection methods.

Here we report the results of mid-UV irradiation experiments of Mars soil analog samples obtained adsorbing relevant organic molecules on a clay mineral that is quite common on Mars, i.e. montmorillonite, doped with 1 wt% of magnesium perchlorate. Specifically, we chose to investigate

the photostability of a plausible precursor of the chlorohydrocarbons detected on Mars by the *Curiosity* rover, namely phthalic acid, along with the biomarkers of extant life L-phenylalanine and L-glutamic acid, which are proteomic amino acids, and adenosine 5'-monophosphate, which is a nucleic acid component.

We monitored the degradation of these molecules adsorbed on montmorillonite through *in situ* spectroscopic analysis, investigating the reflectance properties of the samples in the Near InfraRed (NIR) spectral region.

Such spectroscopic characterization of molecular alteration products provides support for two upcoming robotic missions to Mars that will employ NIR spectroscopy to look for molecular biosignatures, through the instruments SuperCam on board *Mars 2020*, ISEM, Ma_Miss and MicrOmega on board *ExoMars 2020*.

1 Introduction

Searching for molecular biosignatures is one of the primary goals of international Mars exploration programs (Des Marais et al., 2008; Walter et al., 2016).

Two upcoming robotic missions, NASA *Mars 2020* (Williford et al., 2018) and ESA *ExoMars 2020* (Vago et al., 2017), will employ space flight instruments based on Near InfraRed (NIR) vibrational spectroscopy to inspect the presence of organic matter at the respective landing sites. Specifically, *Mars 2020* will be equipped with a remote-sensing instrument located on the “head” of the rover’s mast, SuperCam, which is a combined remote-sensing instrument implementing several techniques (Wiens et al., 2017): Laser Induced Breakdown Spectroscopy (LIBS) (1064 nm laser), to measure elemental composition of targets up to 7 m distance from the rover; Raman (532 nm laser), Time-Resolved Fluorescence (TRF), visible and Near InfraRed (NIR) reflectance spectroscopy (in the spectral ranges 400-900 nm and 1.3-2.6 μm , respectively), to investigate mineralogy and presence of possible organics.

ExoMars 2020 will have 3 NIR instruments, namely ISEM (Infrared Spectrometer for ExoMars), Ma_MISS (Mars Multispectral Imager for Subsurface Studies), and MicrOmega (Vago et al., 2017). ISEM is a spectrometer mounted on the “head” of the rover’s mast, co-aligned with the panoramic camera PanCam, which will measure IR spectra of solar light reflected by surficial targets in the spectral range from 1.15 to 3.3 μm , to determine mineralogical composition at hundreds of meters distance, with a spatial resolution of 3-10 cm (Korablev et al., 2017). Together with PanCam, ISEM will be tasked to select remotely the target site to be analysed more closely with other payload instruments. Ma_MISS is a miniaturized spectrometer integrated with the rover’s drill, capable of reaching 2 m depth, which will acquire reflectance spectra in the spectral range from 0.5 to 2.3 μm , with a spatial resolution of 120 μm , and study stratigraphy and mineralogical composition as a function of depth (De Sanctis et al., 2017). The capability to reach the subsurface will facilitate investigation of environments with higher preservation potential, and Ma_MISS *in situ* measurements will provide information on subsurface materials in their natural setting before crushing and subsequent analysis with MicrOmega. MicrOmega is a visible and NIR imaging microspectrometer, capable of acquiring monochromatic images at high resolution (20 $\mu\text{m} \times 20 \mu\text{m}$ per pixel) on a 5 mm \times 5 mm field of view, with 20 cm^{-1} spectral sampling from 0.95 to 3.65 μm , along with few images at discrete wavelengths from 0.5 to 0.9 μm (Bibring et al., 2017). This instrument will characterize mineralogical composition and presence of organics at a scale of mineral grains, identifying non-destructively regions of interest for further examination with other payload instruments.

Given the importance of spectroscopic analysis to select the most interesting samples to further study with other payload instruments, a correct interpretation of vibrational spectroscopic data is essential for the success of these missions. However, the great variety of environmental factors and physico-chemical interactions that occur between organic molecules and minerals make vibrational spectra quite complex (Fornaro et al., 2013b). Vibrational spectroscopy is particularly sensitive to molecule-mineral interactions because different chemical functional groups give rise to characteristic vibrational bands that undergo significant changes as consequence of such interactions. Indeed, shifts of vibrational frequencies of molecular functional groups and mineral surface sites involved in the interaction can be observed to higher values in case the chemical bonds of these functional groups are strengthened, or to lower values in case they are weakened by the interaction. Band intensity variations can occur depending on molecular spatial arrangements as imposed by surface selection rules. New bands appear in the spectra due to formation of chemical bonds as a consequence of chemisorption. As an example, in the case of important biomarkers such as the components of nucleic acids, remarkable changes can be observed in the following cases: (i) the phosphate group forms covalent bonds with surface sites that lower symmetry and a splitting of its symmetric stretching mode is observed (Fornaro et al., 2018a); (ii) the carbonyl or amino groups of the nucleobases are involved in hydrogen bonds that weaken their chemical bonds and their stretching vibrational modes undergo red-shifts up to hundreds of cm^{-1} (Fornaro et al., 2016; Fornaro et al., 2015a; Fornaro et al., 2015b); and (iii) nucleobases are adsorbed parallel to the surface, cancelling the dynamic dipole moment perpendicular to the surface for vibrational modes in the plane of the aromatic ring, and no bands in the spectrum will be observed for these modes (Fornaro et al., 2013b). The nature of physico-chemical interactions established between molecules and minerals are affected by multiple factors: molecular properties; mineral characteristics; molecule-mineral concentration ratio; and environmental conditions of temperature, pressure, ionic strength, pH, presence of reactive species etc. (Fornaro et al., 2013a; Fornaro et al., 2018b). For instance, at very high molecular concentrations, nucleotides and related molecules tend to form clusters on the mineral surface through strong intermolecular interactions of stacking or hydrogen bonds among nucleobases, with spectroscopic characteristics similar to those of the molecular solid. Therefore, in these cases, it is not possible to observe the spectroscopic features due to molecule-mineral interactions. Moreover, pH influences the molecular protonation state and surface charge and, hence, the possible involvement of electrostatic interactions in the adsorption process. Furthermore, some ions can inhibit bonding to surface sites by interacting with specific molecular functional groups, or favor molecular adsorption by acting as bridges with the mineral surface sites and forming surface complexes with the molecules.

Organic molecules and minerals on Mars are also affected by chemical weathering due to irradiation and oxidative degradation (Fornaro et al., 2018c and references therein; Pavlov et al., 2002). Indeed, the present atmosphere of the planet enables mid- and near-UV radiation, galactic cosmic rays (GCR) and solar energetic protons (SEP) to reach the surface. The RAD instrument on board the *Curiosity* rover has measured a flux of ionizing radiations from GCR of $233 \pm 12 \mu\text{Gray}$ per day, which has a negligible effect on survivability of possible microbial lifeforms on Mars, with sporadic events of SEP that last a few days and increase the dose of about $50 \mu\text{Gray}$ (Hassler et al., 2014). Moreover, even though GCR can penetrate several meters in the Martian soil, measurable effects on organic matter apparently require hundreds of millions of years (Gerakines et al., 2012). On the other hand, organic molecules effectively absorb UV radiation, causing electronic excitation and photochemical reactions at very short timescales of a few hours (Fornaro et al., 2013b; Fornaro et al., 2018a; ten Kate, 2010). In addition, even though UV penetrates only a few μm or mm into the soil, aeolian erosion constantly exposes fresh material to radiation. As a consequence, it is crucial to investigate the effects of UV radiation on organic molecules embedded in mineral matrices mimicking the martian soil, in order to validate hypotheses about the nature of the organic compounds detected so far at the surface of Mars

by the *Curiosity* rover (Eigenbrode et al., 2018; Freissinet et al., 2015; Glavin et al., 2013; Steele et al., 2018, 2012), as well as organics that will be possibly found by the next rover missions *Mars 2020* collecting surficial samples or *ExoMars 2020* collecting subsurface samples. The *ExoMars 2020* landing site, Oxia Planum, underwent an extensive aeolian erosion that exposed at the surface material of less than 100 million years (Vago et al., 2017). As estimated by previous studies (Fornaro et al., 2018c), we would expect that GCR and SEP had a limited effect on possible organics in this site, while UV degradation occurred at much shorter timescales but only in the near surface. However, UV also causes secondary effects such as photo-induced formation of oxidants in the soil (Lasne et al., 2016), which might cause oxidative degradation up to a depth in the martian soil that is difficult to predict. Specifically, perchlorates have been detected both by orbital observations and *in situ* analyses by the Phoenix lander and the *Curiosity* rover, at a concentration between 0.05 and 1.05 wt% (Clark and Kounaves, 2016). Perchlorates have also been found in the martian meteorite EETA79001 (with a concentration of 0.6 ± 0.1 ppm by mass) (Kounaves et al., 2014), confirming their ubiquity on Mars. It is also true that *ExoMars* will analyse samples that have been in the subsurface for many millions of years at constant temperature of about -60°C , at which perchlorates are more stable and oxidation kinetics is very slow. Nevertheless, it is not possible to exclude that the interactions of perchlorates with GCR (capable of penetrating many meters) cause decomposition of perchlorate also at low temperature, giving rise to oxychlorine species, radicals, molecular oxygen and hydrogen peroxide, which in turn may degrade organics. Hydrogen peroxide is characterized by low thermal stability and can produce hydroxyl and peroxide radicals in the presence of ferrous minerals through the Fenton process (Foustoukos and Stern, 2012). Such chemical weathering phenomena might cause profound alterations of the original material, such as breakage of chemical bonds and formation of new molecular species, significantly influencing relative spectroscopic features. Such transformations must be taken into account to redefine molecular targets for life detection.

For these reasons, assignment of spectroscopic features cannot be based solely on databases for pure minerals and pure molecules measured under terrestrial ambient conditions; rather, it is indispensable to perform laboratory studies with analogs of the martian soil, prepared simulating realistically the martian environment, to interpret accurately mission data. In addition, NIR data and relative vibrational mode assignments for organics, both pure and adsorbed on mineral matrices, are scarce in the literature (Clark et al., 2009). Indeed, the only fundamental vibrational modes in this spectral region for organics are O-H, N-H and C-H stretching, and they occur around 3000 cm^{-1} where water gives strong and broad absorption, potentially obscuring molecular features. Beyond these fundamental bands, organics present overtones and combination bands in the NIR, but these modes are forbidden by primary vibrational selection rules, so they are usually very weak and often covered by much more intense mineral bands. As consequence, most literature studies for organics focus on the mid-IR spectral range, where very intense fundamental bands and the fingerprint region for organic molecules are present. Therefore, there is a great need of complementing vibrational studies for organics in the NIR, in order to be able to detect and identify organics on Mars from data acquired in this spectral range.

In this paper, we present laboratory analog studies concerning: (i) preparation of analog samples, obtained adsorbing relevant organic molecules – i.e. a plausible precursor of the chlorohydrocarbons detected on Mars by *Curiosity*, namely phthalic acid, as well as the biomarkers of extant life L-phenylalanine and L-glutamic acid, which are proteomic amino acids, and adenosine 5'-monophosphate, which is a nucleic acid component - at various concentrations on a common martian clay mineral, montmorillonite, doped with 1 wt% of magnesium perchlorate; (ii) mid-UV irradiation processing, to investigate possible transformations of organics and define more accurately targets to search in surficial samples; (iii) Near InfraRed characterization *in situ*, aided by computational spectroscopy at Density Functional Theory level to facilitate assignment of vibrational modes.

These investigations represent the starting point of a systematic study with analog samples composed of several combinations of relevant minerals, salts, oxidants, and different classes of organic compounds at various concentrations, prepared with methods mimicking natural processes that possibly occurred through time at the landing sites selected for *ExoMars 2020* and *Mars 2020*, i.e. Oxia Planum and Jezero Crater, respectively. These studies will allow the assembly of custom databases of vibrational features to be used as a reference dataset to deconvolve complex features of actual martian samples and interpret *in situ* NIR measurements to assist *ExoMars 2020* and *Mars 2020* in identifying organics at the respective landing sites.

2 Materials and Methods

2.1 Preparation and Characterization of Mars Soil Analog Samples

As mineral substrate, we used a Ca-rich montmorillonite ("Cheto"), from County of Apache, State of Arizona, USA, called "Cheto" SAZ-1, purchased from the Clay Mineral Society. Mineral phases were characterized at the Earth and Planets Laboratory of the Carnegie Institution for Science (Washington DC, USA) by making use of: (i) X-Ray Fluorescence (XRF) using a SPECTRO XEPOS (AMETEK) instrument to determine chemical composition; (ii) powder X-Ray Diffraction (XRD) using a Bruker D2 Phaser instrument to confirm mineralogy; and (iii) Isotope Ratio Mass Spectrometry (IRMS) using a Thermo Scientific Delta V Plus mass spectrometer interfaced with a CE Instruments NC 2500 series Elemental Analyser (EA) to estimate the organic content originally present in the montmorillonite sample. Then, we devised a cleaning procedure to extract the organic contamination from this mineral through four extraction steps with different solvent systems: 1) Methanol + sonication for ½ h; 2) 40% Acetone in water + sonication for ½ h; 3) 0.1 M sodium pyrophosphate ($\text{Na}_4\text{P}_2\text{O}_7 \cdot 10\text{H}_2\text{O}$) + sonication for ½ h; 4) Water + rotation for ½ h. We used a mineral concentration of 150 g L^{-1} in each step, and we alternated each extraction step with centrifugation and supernatant separation. Eventually we oven-dried the pellet at 50°C .

Given the ubiquity of perchlorates in the martian soil, we also prepared montmorillonite samples containing 1 wt% of magnesium perchlorate. Specifically, we impregnated the previously treated montmorillonite with 1 wt% of $\text{Mg}(\text{ClO}_4)_2 \cdot 6\text{H}_2\text{O}$ (Fisher Scientific) dissolved in milli-Q water. We agitated the suspension through magnetic stirring to homogenize and then we dried in an oven at 40°C .

To verify possible changes in mineral properties due to the presence of 1 wt% of magnesium perchlorate, we characterized the montmorillonite samples before and after impregnation with magnesium perchlorate through several techniques: XRD, IR, Raman, Transmission Electron Microscopy (TEM), Brunauer-Emmett-Teller (BET) surface area analysis, and solid state Nuclear Magnetic Resonance (NMR). IR measurements were performed using a single beam double pendulum interferometer Vertex 70v (Bruker), equipped with a Praying MantisTM Diffuse Reflection Accessory (Harrick DRIFT), in order to measure diffuse reflectance spectra, available at INAF-Astrophysical Observatory of Arcetri (Florence, Italy). Raman spectra and images were collected using a Witec α -Scanning Near-Field Optical Microscope that has been customized to incorporate confocal Raman spectroscopic imaging, available at the Earth and Planets Laboratory of the Carnegie Institution for Science (Washington DC, USA). The excitation source is a frequency-doubled solid-state YAG laser (532 nm) operating between 0.3 and 1 mW output power (dependent on objective), as measured at the sample using a laser power meter. Objective lenses used included a x100 LWD and a x20 LWD with

a 50 μm optical fiber acting as the confocal pin hole. Spectra were collected on a Peltier-cooled Andor EMCCD chip, after passing through a f/4 300mm focal length imaging spectrometer typically using a 600 lines/mm grating. The lateral resolution of the instrument is as small as 360 nm in air when using the x100 LWD objective, with a focal plane depth of ~ 800 nm. A cosmic ray reduction routine was used to reduce the effects of stray radiation on Raman images, as was image thresholding to reject isolated bright pixels. Fluorescence effects were inhibited by the use of specific peak fitting and flattening. TEM measurements of the samples were performed using a JEOL100CX microscope, available at Sorbonne Université (Paris, France). Specifically, samples in the form of bulk powders were suspended in ethanol and then deposited on 400-mesh copper grids covered with an ultrathin carbon membrane of 2-3 nm thickness. N_2 adsorption-desorption isotherms at -196°C were obtained on a Micromeritics ASAP 2020 instrument, available at Sorbonne Université (Paris, France). Prior to measurement, the samples (40 mg) were degassed under vacuum for 3 hours at 250°C . The Brunauer–Emmett–Teller (BET) surface areas were calculated from the BET equation for a relative pressure (P/P_0) range between 0.05 and 0.25. ^1H and ^{27}Al and ^{29}Si Magic angle spinning (MAS) nuclear magnetic resonance (NMR) spectra were obtained on a Bruker Advance III spectrometer equipped with a 2.5 mm and 4 mm H-X MAS probe, operating at frequency of 500.07 MHz (^1H), 99.36 MHz (^{29}Si) and 130.30 MHz (^{27}Al), available at Sorbonne Université (Paris, France). Chemical shifts were calibrated using the carboxyl signal of adamantane (38.52 ppm) for ^1H , TMS (0 ppm) for ^{29}Si and $\text{Al}(\text{NO}_3)_3$ (0 ppm) for ^{27}Al as external standard. The ^1H experiment using 90° pulse was recorded with a spinning rate of 14 kHz, with a number of scans of 128 and 1 s of recycle delay. Over an acquisition time of 40 ms, the number of scans to obtain the spectra depends on the S/N obtained for each sample. The ^{27}Al experiment using 30° pulse was recorded in the same condition of spinning rate (14 kHz), with 2048 scans, 263 kHz spectral width, and 500 ms of recycle delay. The ^{29}Si experiment using 90° pulse with a ^1H decoupling spin was recorded in the same condition of spinning rate. The number of scans was 8192 and the recycling time D1 was 10 s. The Spectra were processed with a zero-filling factor of 2 and without an exponential decay.

We used treated montmorillonite samples as substrates for performing adsorption experiments of adenosine 5'-monophosphate sodium salt (AMP), L-glutamic acid monosodium salt monohydrate 98% (GLU), L-phenylalanine $\geq 98\%$ (PHE), and phthalic acid $\geq 99.5\%$ (PHA) (all purchased from Sigma-Aldrich), both as individual molecules and as a mixture (MIX), at three different concentrations, i.e. 5 wt%, 1 wt% and 0.1 wt%. Chemical structures for these molecules are shown in Figure 1.

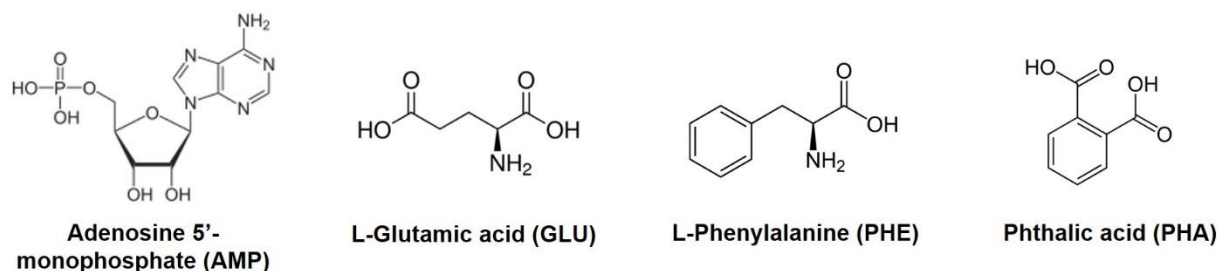


Figure 1. Chemical structures for adenosine 5'-monophosphate (AMP), L-glutamic acid (GLU), L-phenylalanine (PHE), and phthalic acid (PHA).

In order to simulate the formation of molecule-mineral complexes in diluted aqueous environments presumably present on early Mars, we performed equilibrium adsorption experiments in the aqueous phase (Fornaro et al., 2018a). Specifically, we dissolved specific amounts of the above-mentioned molecules, both individually and as a mixture, in a volume of milli-Q water much higher than the mineral porous volume, within molecular solubility, and then we mixed the mineral powder with such aqueous solutions using a mineral concentration of about $120 \text{ g}\cdot\text{L}^{-1}$. We vortexed such suspensions to favour homogeneous interaction between molecules and mineral, and then we put the suspensions on a test tube rotator for a time sufficient to reach the equilibrium state ($\approx 24 \text{ h}$). After equilibrium adsorption, we dried in an oven at 40°C to simulate a desiccation process of liquid water bodies on early Mars. With this method, all molecules in the initial aqueous solution adsorb on the solid substrate. Specifically, we chose to prepare montmorillonite samples with three different molecular concentrations, i.e. 5 wt%, 1 wt% and 0.1 wt%. This preparation method was advantageous because it is time-effective and it enables us to control the amount of molecules adsorbed on the solid substrate and the experimental conditions. In addition, these procedures maximize molecular interactions with mineral surfaces as well as molecular diffusion inside the mineral matrix; therefore, molecules establish physico-chemical interactions with mineral surfaces as in natural processes of adsorption.

2.2 UV Irradiation Experiments and Near InfraRed Characterization

We subjected the Mars soil analog samples at 5 wt% molecular concentration on montmorillonite containing 1 wt% of magnesium perchlorate to mid-UV irradiation processing. We chose the samples with the highest molecular concentration in order to observe the most obvious variations in the spectroscopic features due to UV irradiation, and with magnesium perchlorate to simulate realistically martian surficial samples. We used an experimental setup assembled at INAF-Astrophysical Observatory of Arcetri (Florence, Italy) that allows us to monitor the photochemistry *in situ* by Diffuse Reflectance Infrared Fourier Transform Spectroscopy (DRIFTS) analysis. Specifically, the UV light emitted by a Newport Xenon enhanced UV 300 W lamp with purified Xenon at 5-20 bar (wavelength range 185–2000 nm) is focused directly on the sample through an optical fiber of 800 μm spot size. With this setup, the irradiated spot of the sample presents an area of 7.07 mm^2 and the UV flux focused on the sample is $2.75 \times 10^{17} \text{ photons s}^{-1} \text{ cm}^{-2}$ in the 200-400 nm spectral range, as measured through a single monochromator Spectro 320 scanning spectrometer (Instrument System). The optical fiber is inserted into the sample chamber of a single beam double pendulum interferometer VERTEX 70v (Bruker), equipped with a Praying MantisTM Diffuse Reflection Accessory (Harrick DRIFT), in order to measure diffuse reflectance spectra of the sample *in situ* at regular intervals during UV irradiation (Figure 2).

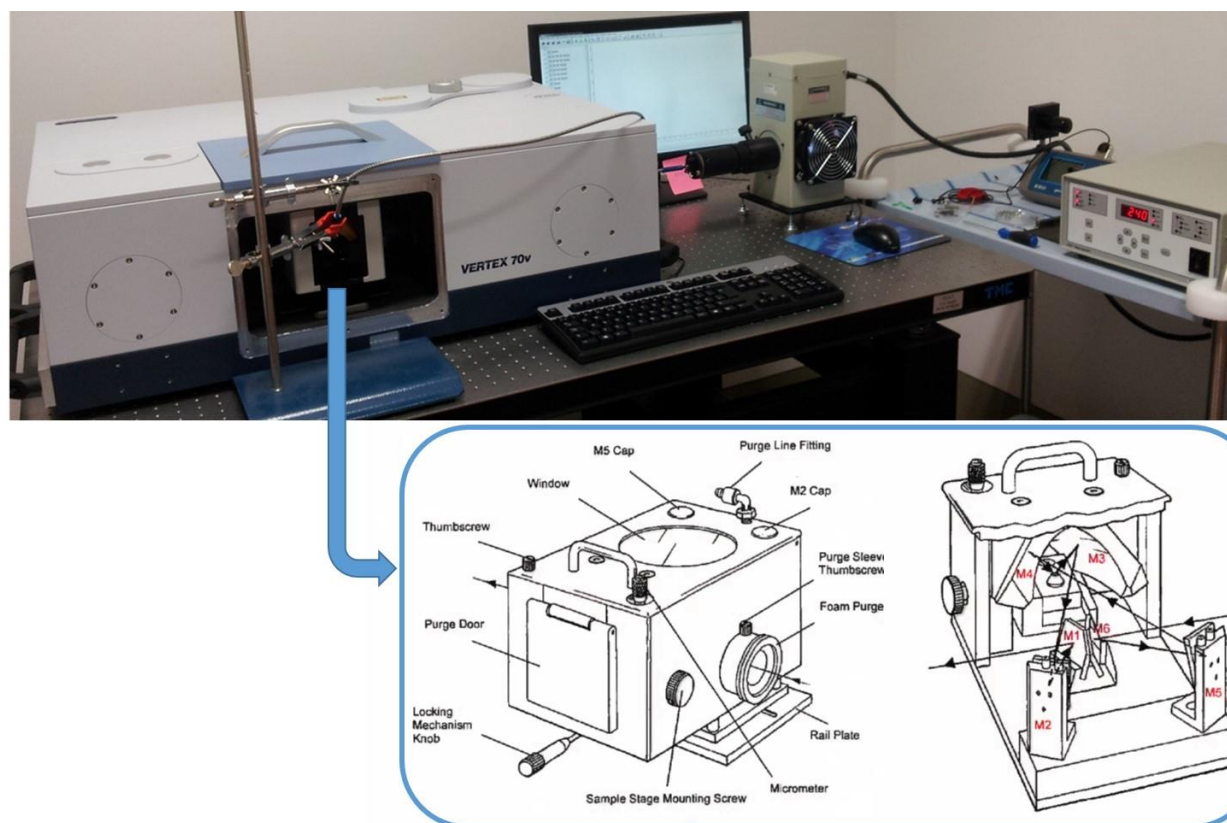


Figure 2. Experimental setup used for *in situ* UV irradiation experiments, with scheme of the Praying Mantis™ Diffuse Reflection Accessory.

For these experiments, we acquired reflectance spectra pre-irradiation and during UV irradiation in the spectral range $400\text{--}8000\text{ cm}^{-1}$, using a resolution of 4 cm^{-1} , 100 scan, Globar source, KBr beamsplitter, and a DTGS detector, but we focused our analysis only on the Near InfraRed (NIR) spectral region above 2000 cm^{-1} , which is relevant to space flight instruments such as SuperCam (spectral range $1.3\text{--}2.6\text{ }\mu\text{m} + 0.4\text{--}0.85\text{ }\mu\text{m}$) on board *Mars 2020*, Ma_MISS (spectral range $0.5\text{--}2.3\text{ }\mu\text{m}$), ISEM (spectral range $1.1\text{--}3.3\text{ }\mu\text{m}$) and MicrOmega (spectral range $0.95\text{--}3.65\text{ }\mu\text{m} + 0.5\text{--}0.9\text{ }\mu\text{m}$) on board *ExoMars 2020*. Furthermore, we acquired reflectance spectra also in the $8000\text{--}20000\text{ cm}^{-1}$ spectral region, using a Tungsten source, CaF₂ beamsplitter and InGaAs detector.

We irradiated for a total time of 70 minutes under a flux of nitrogen at room temperature. Such short irradiation experiments at room temperature simulate the most intense martian irradiation that occurs in the equatorial region around noon.

We evaluated the percentages of degradation after 70 min of UV irradiation as:

$$\% \text{ degr} = \frac{[(A_{pre} - A_{post}) \times 100]}{A_{pre}}$$

where A_{pre} is the area of the molecular peak pre-irradiation and A_{post} is the area of the same molecular peak post-irradiation.

2.3. Anharmonic Computational Spectroscopy Calculations

To assign vibrational features in the NIR, it is necessary to carry out calculations at the anharmonic level that predict frequencies and intensities of overtones and combination bands. The anharmonic vibrational frequencies and IR intensities were obtained via a generalized second-order vibrational perturbation theory approach (GVPT2) (Barone, 2005; Barone et al., 2014; Bloino, 2015; Bloino et al., 2015), following the procedures recommended in Refs. (Bloino et al., 2016). According to the experimental conditions, the geometry optimizations started from conformers present in the solid state. All computations have been performed by means of density functional theory employing the B3LYP functional (Becke, 1993) with the double-zeta basis sets SNSD (Barone et al., 2014), and including Grimme's dispersion correction D3 (in conjunction with Becke-Johnson damping) (Ehrlich et al., 2011; Grimme et al., 2011, 2010). This approach has been well tested for structural and spectroscopy properties of biological molecules of astrochemical interest (Barone et al., 2015; Biczysko et al., 2018; Fornaro et al., 2014; Fornaro et al., 2015a; Fornaro et al., 2015b).

In GVPT2 computations, the default criteria for anharmonic resonances have been exploited, and variational corrections have been applied to both energies and IR intensities. Moreover, considering that the large amplitude-motion (LAM) vibrations, which cannot be well-described by perturbative approaches, may contaminate the overall VPT2 treatment and higher frequency vibrations, LAM-free VPT2 scheme have been used in this work (where all LAMs anharmonic constants are excluded from the VPT2) (Biczysko et al., 2018). For all molecules, vibrations with harmonic frequencies below 200 cm^{-1} have been considered as LAMs.

All calculations were performed with Gaussian 16 suite of computer codes (Frisch et al., 2016). VMS-Draw (Licari et al., 2015) was used to visualize the normal modes and analyze in detail the outcome of vibrational computations and IR spectra.

3 Results

3.1 Mineral Characterization

XRF measurements showed the following chemical composition (%): $\text{SiO}_2 = 69.1$, $\text{Al}_2\text{O}_3 = 18.1$, $\text{TiO}_2 = 0.22$, $\text{Fe}_2\text{O}_3 = 1.75$, $\text{MnO} = 0.090$, $\text{MgO} = 6.49$, $\text{CaO} = 3.71$, $\text{Na}_2\text{O} = 0.19$, $\text{K}_2\text{O} = 0.26$, $\text{P}_2\text{O}_5 = 0.01$, $\text{SO}_3 = 0.059$, $\text{K}_2\text{O} = 0.26$.

We did not purify such montmorillonite sample from other mineral phases, so 1-2 wt% of quartz was also present, as confirmed by the presence of a peak at 26.6° in the XRD pattern shown in Supplementary Figure 1 of Supplementary Material.

The carbon content originally present in the montmorillonite sample, measured through IRMS-EA, was 0.11 wt%. After the cleaning treatment, the carbon content was below the IRMS detection limit for Montmorillonite SAz-1.

XRD measurements showed that the treatment with sodium pyrophosphate caused cation exchange of doubly-hydrated Ca^{2+} with singularly-hydrated Na^+ , reducing montmorillonite interlayer distance of the basal plane (001) from 14.63 Å to 12.54 Å (Supplementary Figure 2). Based on a cation exchange capacity (CEC) of 120 meq/100g for montmorillonite and the amount of $\text{Na}_4\text{P}_2\text{O}_7 \cdot 10\text{H}_2\text{O}$ used for extraction, we estimated the presence in the interlayers of about 60% Na^+ and 40% Ca^{2+} . We also

observed appearance of new peaks at 5.04, 5.45 and 8.56 Å in the XRD pattern of treated montmorillonite, likely due to salts precipitated as consequence of the cleaning treatment.

As shown in Supplementary Figure 3, XRD patterns of montmorillonite (MONT) and montmorillonite doped with 1 wt% magnesium perchlorate (MONTTP) present almost the same interlayer distance, intensity and band width. We only observed slight expansion of the interlayer distance of the (001) planes from 12.54 Å for MONT to 12.63 Å for MONTTP ($\Delta d(001) = 0.09\text{\AA}$), that cannot be explained with intercalation of Mg^{2+} , but it is likely due to hydration. We also observed disappearance of the peaks at 5.04, 5.45 and 8.56 Å, likely due to dissolution by perchlorate of the salts previously precipitated.

We did not observe any modifications of the mineral by IR or Raman characterization, as shown in Supplementary Figures 4 and 5.

TEM images of MONT and MONTTP showed stacked layers with an interlayer spacing in agreement with XRD measurements (Supplementary Figure 6). Differences between samples before and after attack with perchlorate is not obvious since only 1 wt% of the latter was used. We suggest that 1 wt% of perchlorate does not strongly affect the montmorillonite layers. Statistics of layer spacings were done on several regions. The micrograph shown in Supplementary Figure 6 (on the right) exhibits some irregular layers compared to the ones before addition of perchlorate. A discontinuity in the layers can be underlined. These results should be carefully analysed since 1 wt% perchlorate cannot affect bulk properties of the sample.

However, BET analysis indicated that magnesium perchlorate affected montmorillonite surface area, which increased from 43 m²/g to 82 m²/g. As can be seen on the adsorption-desorption isotherms in Supplementary Figure 7, the sample containing the 1 wt% of magnesium perchlorate presents more “textural porosity”, probably due to alteration of surface charge that may cause disaggregation of the particles and an increase in the number of pores.

²⁷Al and ²⁹Si NMR spectra of samples before and after addition of magnesium perchlorate are superimposable without any difference, as shown in Supplementary Figure 8. The signal centered at 0 ppm in ²⁷Al MAS NMR is attributed to hexa-coordinated aluminium usually present in montmorillonite. The ²⁹Si signal centered around -92 ppm corresponds to Q3 species. For ¹H NMR, as shown in Supplementary Figure 9, a slight difference is observed in the form of the peaks, but overall the signals remain the same and the attribution is as follows: the signal around 4.4 ppm is assigned to the protons of water, the one at 1.8 to the protons in Si-OH, and the one at 0.8 to the protons in Mg-OH. The negative ¹H NMR shifts at -2.85 ppm can be attributed to a strong shielding effect due to the presence of paramagnetic center in the sample.

3.2 Near InfraRed Spectra Pre- and Post- UV Irradiation

Figures 3, 4, 5, 6 and 7 report the NIR spectra pre-irradiation in the 2000-20000 cm⁻¹ spectral region for AMP, GLU, PHE, PHA, and MIX adsorbed on montmorillonite at 5, 1 and 0.1 wt% concentration, along with frequency shifts with respect to the pure molecules in the solid state and tentative assignments based on DFT calculations at anharmonic level for the peaks in the most relevant 2740-8000 cm⁻¹ spectral range.

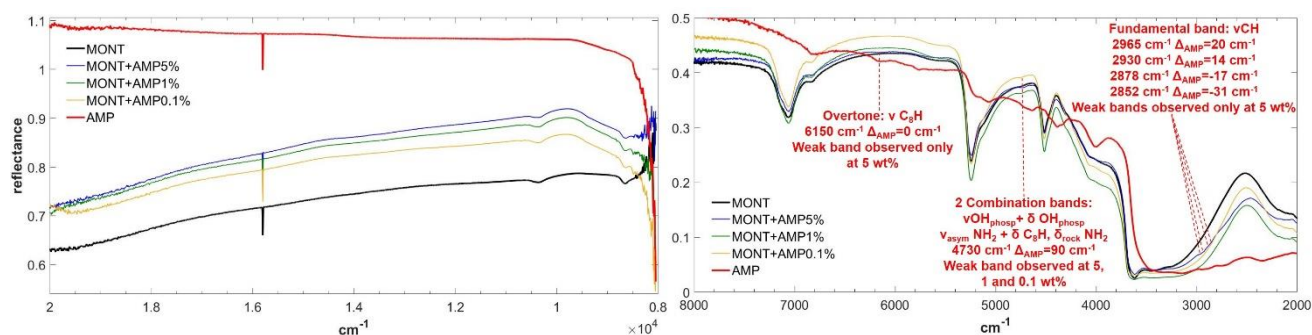


Figure 3. NIR spectra in the 2000-20000 cm^{-1} spectral range of AMP adsorbed on montmorillonite at 5, 1 and 0.1 wt% concentration. Legend: Δ_{AMP} = wavenumber shift with respect to the same peak of the pure molecule in the solid state; phosp = phosphate group; ν = stretching vibrational mode; δ = bending vibrational mode; rock = rocking in-plane vibrational mode; asym = asymmetric vibrational mode.

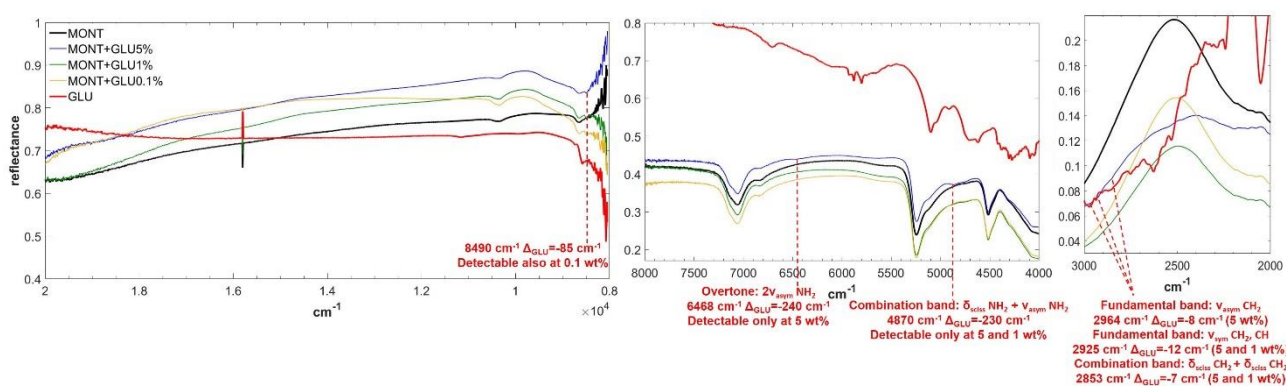


Figure 4. NIR spectra in the 2000-20000 cm^{-1} spectral range of GLU adsorbed on montmorillonite at 5, 1 and 0.1 wt% concentration. Legend: Δ_{GLU} = wavenumber shift with respect to the same peak of the pure molecule in the solid state; ν = stretching vibrational mode; δ = bending vibrational mode; sciss = scissoring in-plane vibrational mode; sym = symmetric vibrational mode; asym = asymmetric vibrational mode.

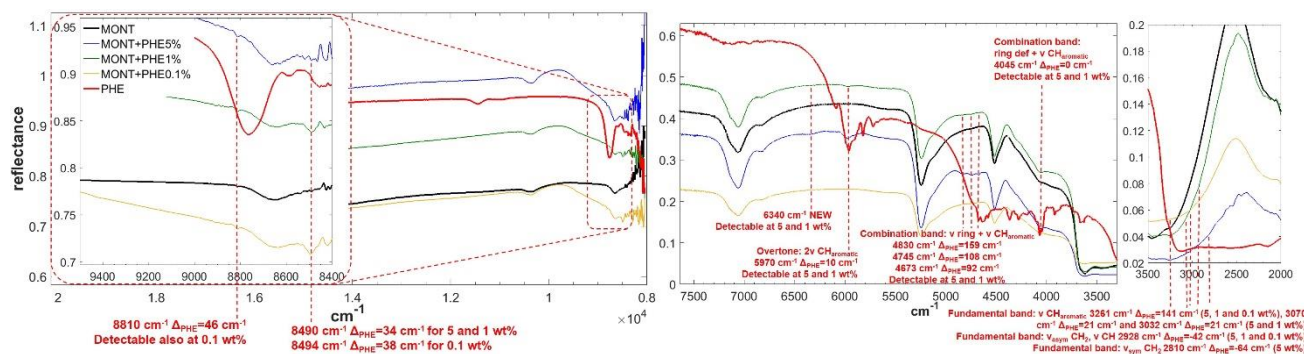


Figure 5. NIR spectra in the 2000-20000 cm^{-1} spectral range of PHE adsorbed on montmorillonite at 5, 1 and 0.1 wt% concentration. Legend: Δ_{PHE} = wavenumber shift with respect to the same peak of the pure molecule in the solid state; ν = stretching vibrational mode; δ = bending vibrational mode; def = deformational vibrational mode.

= deformation vibrational mode; sym = symmetric vibrational mode; asym = asymmetric vibrational mode.

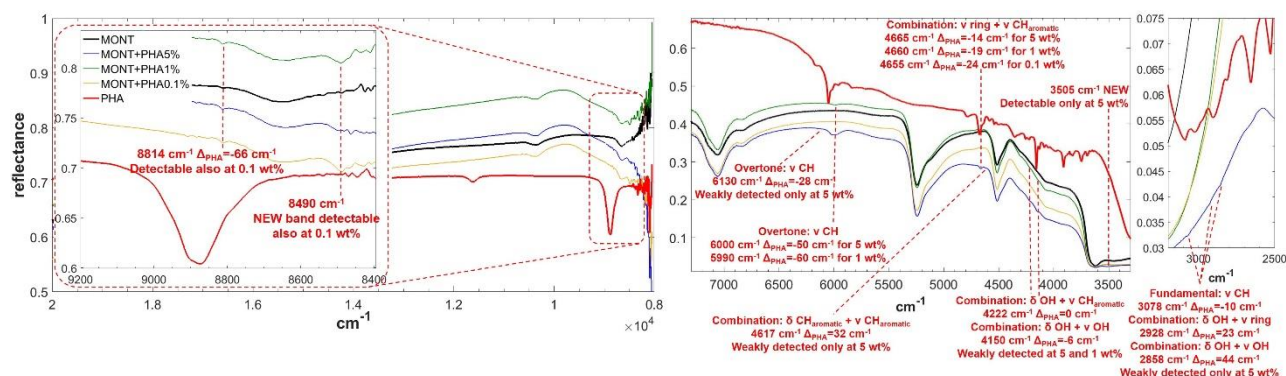


Figure 6. NIR spectra in the 2000-20000 cm^{-1} spectral range of PHA adsorbed on montmorillonite at 5, 1 and 0.1 wt% concentration. Legend: Δ_{PHA} = wavenumber shift with respect to the same peak of the pure molecule in the solid state; ν = stretching vibrational mode; δ = bending vibrational mode.

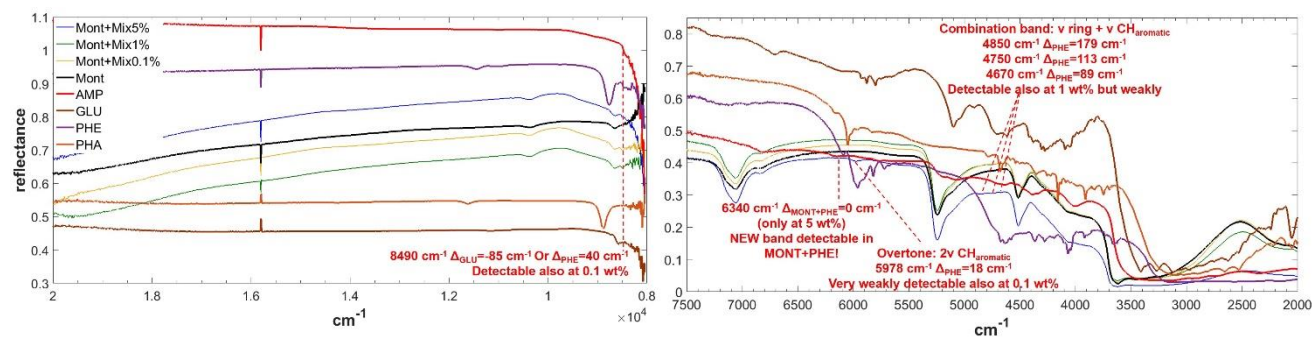


Figure 7. NIR spectra in the 2000-20000 cm^{-1} spectral range of MIX adsorbed on montmorillonite at 5, 1 and 0.1 wt% concentration. Legend: Δ_{PHE} = wavenumber shift with respect to the same peak of the pure molecule in the solid state; $\Delta_{\text{MONT+PHE}}$ = wavenumber shift with respect to the same peak of PHE adsorbed on MONT; ν = stretching vibrational mode.

Table 1 reports the changes of molecular NIR features due to adsorption onto montmorillonite at different concentrations, presence of 1 wt% of magnesium perchlorate and 70 min of UV irradiation, along with the new features forming during UV irradiation. Tentative assignments are based on DFT calculations at anharmonic level, which are reported in detail in Supplementary Table 1.

Table 2 reports the changes of montmorillonite NIR features due to molecular adsorption at different concentrations, presence of 1 wt% of magnesium perchlorate and 70 min of UV irradiation.

Table 1. Changes of molecular NIR features due to adsorption onto montmorillonite at different concentrations (5, 1 and 0.1 wt%), presence of 1 wt% of magnesium perchlorate and 70 min of UV irradiation.

Assignment based on DFT calculations	$\tilde{\nu}$	$\tilde{\nu}$	$\Delta_{\text{mont}}\tilde{\nu}$	$\tilde{\nu}$	$\Delta_{\text{mont}}\tilde{\nu}$	$\tilde{\nu}$	$\Delta_{\text{mont}}\tilde{\nu}$	$\tilde{\nu}$	$\Delta_{\text{mont}}\tilde{\nu}$	$\tilde{\nu}$	$\Delta_{\text{pre-ir}}\tilde{\nu}$	I
	AMP solid state exp.	MONT+AMP5%		MONT+AMP1%		MONT+AMP0.1%		MONT+AMP5%		MONT+AMP5% post-irradiation		
Overtone: $2\nu\text{C}_6\text{H}$	6150 m	6150 w	0					6150 w	0			Too weak to evaluate variations
2 Combinations in 1 band: $\nu\text{OH}_{\text{phosp}} + \delta\text{OH}_{\text{phosp}}, \nu_{\text{asym}}\text{NH}_2 + \delta\text{C}_6\text{H}, \delta_{\text{rock}}\text{NH}_2$	4640 ms	4730 w	90	4730 w	90	4730 w	90	4730 m	90	4730	0	No variations
Fundamental: $\nu_{\text{asym}}\text{CH}$	2945 m	2965 w	20					2945 vvw	0			Too weak to evaluate variations
Fundamental: $\nu_{\text{sym}}\text{CH}_2$	2916 mw	2930 w	14									
Fundamental: νCH	2895 w	2878 vvw	-17									
Fundamental: $\nu_{\text{sym}}\text{CH}$	2883 w	2852 vvw	-31					2883 vvw	0			Too weak to evaluate variations
NEW FEATURES												
												3390
												4070
												4220
												4460
	GLU solid state exp.	MONT+GLU5%		MONT+GLU1%		MONT+GLU0.1%		MONT+GLU5%		MONT+GLU5% post-irradiation		
	8575 vs	8490 m	-85	8490 m	-85	8490 m	-85					
Overtone: $2\nu_{\text{asym}}\text{NH}_2$	6708 m	6468 mw	-240					6468 mw	-240	6450	-18	No variations
Combination: $\delta_{\text{sciss}}\text{NH}_2 + \nu_{\text{asym}}\text{NH}_2$	5100 vs	4870 m	-230	4870 w	-230			4870 m	-230	4850	-20	Too weak to evaluate variations
Fundamental: $\nu_{\text{asym}}\text{CH}_2$	2972 m	2964 w	-8									
Fundamental: $\nu_{\text{sym}}\text{CH}_2\text{CH}$	2937 mw	2925 w	-12	2925 vw	-12							
Combination: $\delta_{\text{sciss}}\text{CH}_2 + \delta_{\text{sciss}}\text{CH}_2$	2860 mw	2853 vw	-7	2853 vw	-7	2853 vvw	-7					
NEW FEATURES												
												4075
												3280
												2343
	PHE solid state exp.	MONT+PHE5%		MONT+PHE1%		MONT+PHE0.1%		MONT+PHE5%		MONT+PHE5% post-irradiation		
	8764 s	8810 vw	46	8810 vw	46	8810 vw	46					
	8456 m	8490 vw	34	8490 m	34	8494 m	38					
		6340 m NEW		6340 vw NEW				6340 m NEW		6340	0	No variations
Overtone: $2\nu\text{CH}_{\text{aromatic}}$	5960 vs	5970 ms	10	5970 mw	10			5970 ms	10	5970	0	Slight intensity decrease (4%)
Combination: $\nu\text{ ring} + \nu\text{CH}_{\text{aromatic}}$	4671 s	4830 m	159	4830 w	159			4830 m	159	4830	0	No variations
Combination: $\nu\text{ ring} + \nu\text{CH}_{\text{aromatic}}$	4637 vs c	4745 m	108	4745 m	108			4745 m	108	4745	0	Significant intensity decrease (43%)
Combination: $\nu\text{ ring} + \nu\text{CH}_{\text{aromatic}}$	4581 s	4673 m	92	4673 m	92			4673 m	92	4673	0	Intensity increase
Combination: ring def + $\nu\text{CH}_{\text{aromatic}}$	4045 s	4045 mw	0	4045 w	0			4045 mw	0	4045	0	Intensity increase
Fundamental: $\nu\text{CH}_{\text{aromatic}}$	3120 s b	3261 w	141	3261 w	141	3261 vw	141	3261 mw	141	3261	0	Intensity increase
Fundamental: $\nu\text{CH}_{\text{aromatic}}$	3049 vvw wb	3070 vvw	21	3070 vvw	21			3070 vw	21	3070	0	Intensity increase
Fundamental: $\nu\text{CH}_{\text{aromatic}}$	3011 vvw wb	3032 vw	21	3032 vvw	21			3032 w	21	3032	0	Intensity increase
Fundamental: $\nu_{\text{asym}}\text{CH}_2, \nu\text{CH}$	2970 m wb	2928 vw	-42	2928 vw	-42	2928 vw	-42	2928 w	-42	2928	0	Too weak to evaluate variations
Fundamental: $\nu_{\text{sym}}\text{CH}_2$	2874 w wb	2810 sh	-64					2810 sh	-64	2810	0	Too weak to evaluate variations
Overtone: δCH	2748 m wb	2670 w	-78	2670 vw	-78			2670 vw	-78	2670	0	Too weak to evaluate variations
NEW FEATURES												
												7140
	PHA solid state exp.	MONT+PHA5%		MONT+PHA1%		MONT+PHA0.1%		MONT+PHA5%		MONT+PHA5% post-irradiation		

	8880 vs	8814 w	-66	8814 w	-66	8814 w	-66						
		8490 m NEW		8490 m NEW		8490 m NEW							
Overtone: vCH	6158 m	6130 vw	-28					6130 vw	-28	6130	0		Too weak to evaluate variations
Overtone: vCH	6050 vs	6000 ms	-50	5990 m	-60			6000 ms	-50	6000	0		No variations
Combination: v ring + vCH _{aromatic}	4679 vs	4665 m	-14	4660 mw	-19	4655 vw	-24	4660 m	-19	4660	0		Intensity increase
Combination: δCH _{aromatic} + vCH _{aromatic}	4585 ms	4617 w	32					4617 w	32	4619	2		Too weak to evaluate variations
Combination: δOH+vCH _{aromatic}	4222 m	4222 sh	0	4222 vw sh	0			4210 sh	-12	4210	0		Too weak to evaluate variations
Combination: δOH+vOH	4156 vs	4150 w	-6	4150 vw	-6			4149 w	-7	4149	0		Intensity increase
Combination: τ molecule + vOH	3910 s							3940 w	30	3940	0		Too weak to evaluate variations
		3505 w NEW						3505 w NEW		3505	0		Too weak to evaluate variations
Fundamental: vCH	3088 s	3078 vw	-10					3069 vw	-19	3071	2		Too weak to evaluate variations
Combination: δOH + v ring	2905 m	2928 vw	23										
Combination: δOH+vOH	2814 m	2858 vw	44										
	Molecule solid state exp.	MONT+MIX5%		MONT+MIX1%		MONT+MIX0.1%		MONT+MIX5%		MONT+MIX5% post-irradiation			
PHE	8450 s		40		40		40						
GLU	8575 vs	8490 m	-85	8490 m	-85	8490 m	-85						
MONT+PHE	6340 m NEW	6340 w NEW	0					6340 w NEW	0	6340	0		Too weak to evaluate variations
PHE Overtone: 2vCH _{aromatic}	5960 vs	5978 m	18	5978 w	18	5978 vw	18	5978 m	18	5978	0		No variations
PHE Combination: v ring + vCH _{aromatic}	4671 s	4850 m	179	4850 vw	179			4850 m	179	4840	-10		Intensity increase
PHE Combination: v ring + vCH _{aromatic}	4637 vs c	4750 m	113	4750 vw	113			4750 m	113	4740	-10		Slight intensity decrease (15%)
PHE Combination: v ring + vCH _{aromatic}	4581 s	4670 m	89	4670 w	89			4670 m	89	4670	0		No variations
NEW FEATURES													
7140													

Legend: $\tilde{\nu}$ = wavenumber in cm^{-1} ; $\Delta_{\text{mol}}\tilde{\nu}$ = wavenumber shift (in cm^{-1}) with respect to the same peak of the pure molecule in the solid state; $\Delta_{\text{pre-ir}}\tilde{\nu}$ = wavenumber shift (in cm^{-1}) with respect to the same peak pre-irradiation; I = Intensity of the band; phosp = phosphate group; v = stretching vibrational mode; δ = bending vibrational mode; τ = torsional vibrational mode; def = deformation vibrational mode; sciss = scissoring in-plane vibrational mode; rock = rocking in-plane vibrational mode; wagg = wagging out-of-plane vibrational mode; sym = symmetric vibrational mode; asym = asymmetric vibrational mode; s = strong band; m = medium band; w = weak band; vw = very weak band; vvw = extremely weak band; c = convolution of peaks; b = broad band; wb = within broad band.

Table 2. Changes of montmorillonite NIR features due to molecular adsorption at different concentrations (5, 1 and 0.1 wt%), presence of 1 wt% of magnesium perchlorate and 70 min of UV irradiation.

Assignment	$\tilde{\nu}$	$\tilde{\nu}$	$\Delta_{\text{MONT}}\tilde{\nu}$	$\tilde{\nu}$	$\Delta_{\text{MONT}}\tilde{\nu}$	$\tilde{\nu}$	$\Delta_{\text{MONT}}\tilde{\nu}$	$\tilde{\nu}$	$\Delta_{\text{MONT}}\tilde{\nu}$	$\tilde{\nu}$	$\Delta_{\text{MONT}}\tilde{\nu}$	$\tilde{\nu}$	$\Delta_{\text{pre-ir}}\tilde{\nu}$	I
	MONT	MONT+AMP5%		MONT+AMP1%		MONT+AMP0.1%		MONT+P		MONT+P+AMP5%		MONT+P+AMP5% post-irradiation		
Overtone: 2vAl ₂ OH ^a	7063 s	7063	0	7063	0	7063	0	7063	0	7063	0	7048	-15	Intensity decrease (29%)
Overtone: 2vH ₂ O ^b	6830 mw	6830	0	6830	0	6830	0	6830	0	6830	0	6830	0	Intensity decrease up to complete disappearance (100%)
Combination : (v+δ)H ₂ O ^c	5244 vs	5244	0	5244	0	5244	0	5244	0	5244	0	5244	0	Significant intensity decrease (53%)
Combination : (v+δ)OH ^b	4515 s	4515	0	4515	0	4515	0	4515	0	4515	0	4515	0	No variations
Fundamental : vOH ^c	3620 s wb	3620	0	3620	0	3620	0	3624	4	3620	-4	3616	-4	Intensity increase
	MONT	MONT+GLU5%		MONT+GLU1%		MONT+GLU0.1%		MONT+P		MONT+P+GLU5%		MONT+P+GLU5% post-irradiation		

Overtone: 2vAl ₂ OH ^a	7063 s	7063	0	7063	0	7063	0	7063	0	7063	0	7063	0	Intensity decrease (28%)
Overtone: 2vH ₂ O ^b	6830 mw	6830	0	6830	0	6830	0	6830	0	6830	0	6830	0	Intensity decrease up to nearly complete disappearance (86%)
Combination : (v+δ)H ₂ O ^c	5244 vs	5244	0	5244	0	5244	0	5244	0	5244	0	5244	0	Significant intensity decrease (56%)
Combination : (v+δ)OH ^b	4515 s	4515	0	4515	0	4515	0	4515	0	4515	0	4515	0	No variations
Fundamental : vOH ^c	3620 s wb	3620	0	3620	0	3620	0	3620	0	3620	0	3620	0	Intensity increase
	MONT	MONT+PHE5%		MONT+PHE1%		MONT+PHE0.1%		MONTP		MONTP+PHE5%		MONTP+PHE5% post-irradiation		
Overtone: 2vAl ₂ OH ^a	7063 s	7063	0	7063	0	7063	0	7063	0	7063	0	7053	-10	Intensity decrease (24%)
Overtone: 2vH ₂ O ^b	6830 mw	6830	0	6830	0	6830	0	6830	0	6830	0	6830	0	Intensity decrease up to nearly complete disappearance (96%)
Combination : (v+δ)H ₂ O ^c	5244 vs	5244	0	5244	0	5244	0	5244	0	5244	0	5236	-8	Significant intensity decrease (61%)
Combination : (v+δ)OH ^b	4515 s	4515	0	4515	0	4515	0	4515	0	4515	0	4515	0	Slight intensity decrease (6%)
Fundamental : vOH ^c	3620 s wb	3620	0	3620	0	3620	0	3620	0	3620	0	3620	0	Intensity increase
	MONT	MONT+PHA5%		MONT+PHA1%		MONT+PHA0.1%		MONTP		MONTP+PHA5%		MONTP+PHA5% post-irradiation		
Overtone: 2vAl ₂ OH ^a	7063 s	7063	0	7063	0	7063	0	7063	0	7063	0	7063	0	Intensity decrease (25%)
Overtone: 2vH ₂ O ^b	6830 mw	6830	0	6830	0	6830	0	6830	0	6830	0	6830	0	Significant intensity decrease (52%)
Combination : (v+δ)H ₂ O ^c	5244 vs	5244	0	5244	0	5244	0	5244	0	5244	0	5235	-9	Significant intensity decrease (55%)
Combination : (v+δ)OH ^b	4515 s	4515	0	4515	0	4515	0	4515	0	4515	0	4515	0	Slight intensity decrease (8%)
Fundamental : vOH ^c	3620 s wb	3620	0	3620	0	3620	0	3620	0	3620	0	3620	0	Intensity increase
	MONT	MONT+MIX5%		MONT+MIX1%		MONT+MIX0.1%		MONTP		MONTP+MIX5%		MONTP+MIX5% post-irradiation		
Overtone: 2vAl ₂ OH ^a	7063 s	7063	0	7063	0	7063	0	7063	0	7063	0	7063	0	Intensity decrease (23%)
Overtone: 2vH ₂ O ^b	6830 mw	6830	0	6830	0	6830	0	6830	0	6830	0	6830	0	Significant intensity decrease (61%)
Combination : (v+δ)H ₂ O ^c	5244 vs	5244	0	5244	0	5244	0	5244	0	5244	0	5240	-4	Significant intensity decrease (47%)
Combination : (v+δ)OH ^b	4515 s	4515	0	4515	0	4515	0	4515	0	4515	0	4515	0	No variations
Fundamental : vOH ^c	3620 s wb	3620	0	3620	0	3620	0	3620	0	3620	0	3620	0	Intensity increase

^a (BISHOP et al., 2002); ^b (Madejová et al., 2006); ^c (Madejová, 2003).

Legend: $\tilde{\nu}$ = wavenumber in cm^{-1} ; $\Delta_{\text{MONT}}\tilde{\nu}$ = wavenumber shift (in cm^{-1}) with respect to the same peak of MONT; $\Delta_{\text{pre-irr}}\tilde{\nu}$ = wavenumber shift (in cm^{-1}) with respect to the same peak pre-irradiation; I = Intensity of the band; v = stretching vibrational mode; δ = bending vibrational mode; s = strong band; m = medium band; w = weak band; vw = very weak band; vvw = extremely weak band; b = broad band; wb = within broad band.

Figures 8, 9, 10, 11 and 12 show the main changes in NIR features after 70 min of UV irradiation for the samples of 5 wt% molecular concentration on montmorillonite containing 1 wt% of Mg-perchlorate.

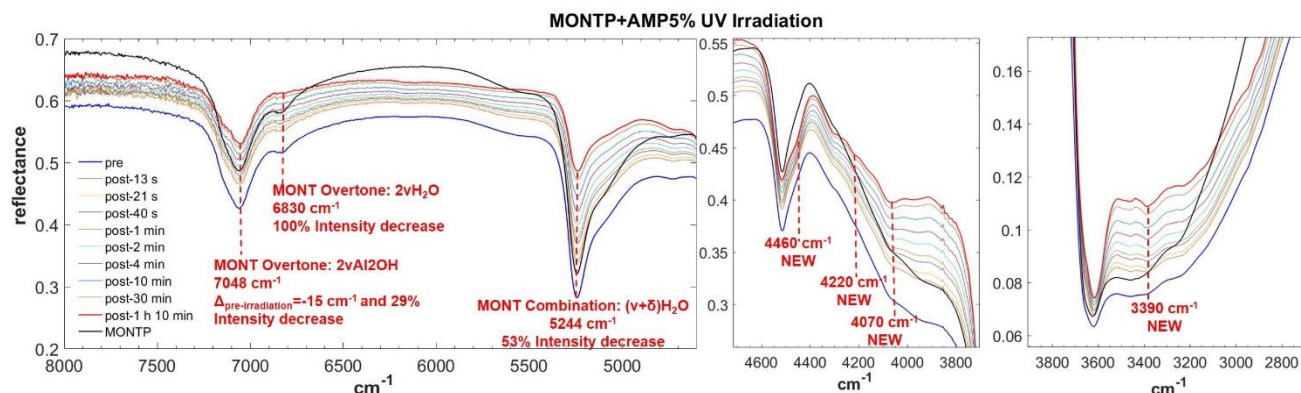


Figure 8. Main changes in NIR features after 70 min of UV irradiation for AMP 5 wt% on montmorillonite containing 1 wt% of Mg-perchlorate. Legend: $\Delta_{\text{pre-irr}}$ = wavenumber shift with respect to the same peak pre-irradiation; ν = stretching vibrational mode; δ = bending vibrational mode.

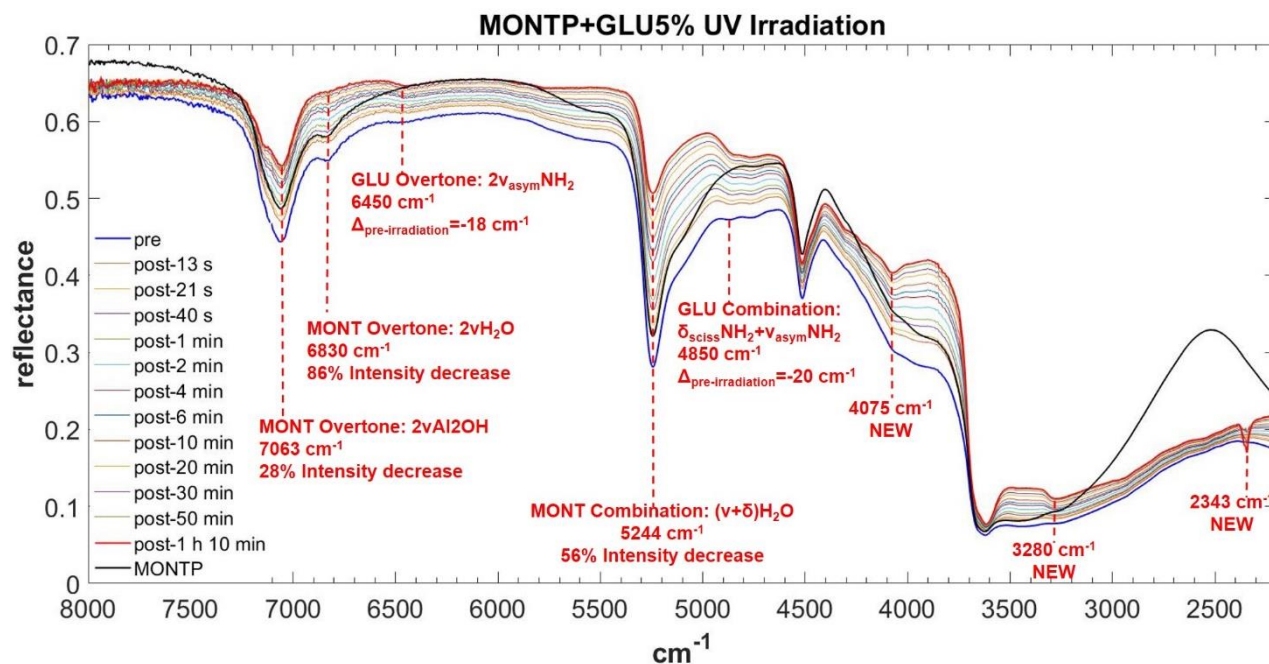


Figure 9. Main changes in NIR features after 70 min of UV irradiation for GLU 5 wt% on montmorillonite containing 1 wt% of Mg-perchlorate. Legend: $\Delta_{\text{pre-irr}}$ = wavenumber shift with respect to the same peak pre-irradiation; ν = stretching vibrational mode; δ = bending vibrational mode; sciss = scissoring in-plane vibrational mode; asym = asymmetric vibrational mode.

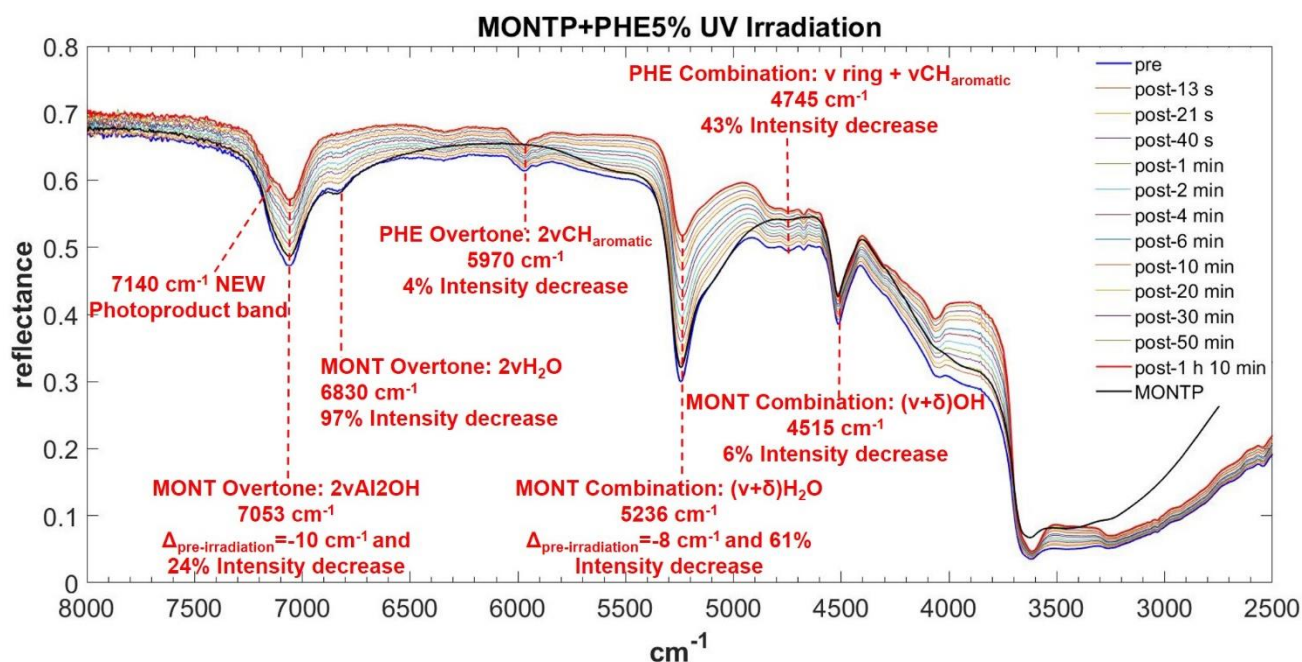


Figure 10. Main changes in NIR features after 70 min of UV irradiation for PHE 5 wt% on montmorillonite containing 1 wt% of Mg-perchlorate. Legend: $\Delta_{\text{pre-irr}}$ = wavenumber shift with respect to the same peak pre-irradiation; ν = stretching vibrational mode; δ = bending vibrational mode.

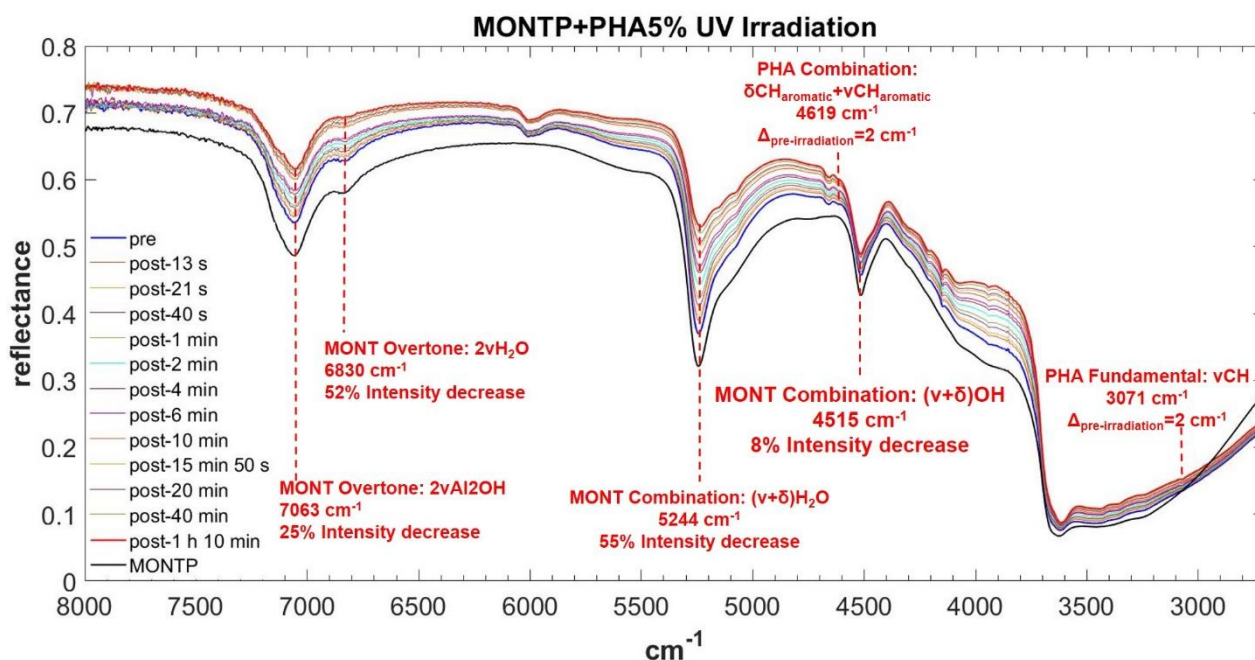


Figure 11. Main changes in NIR features after 70 min of UV irradiation for PHA 5 wt% on montmorillonite containing 1 wt% of Mg-perchlorate. Legend: $\Delta_{\text{pre-irr}}$ = wavenumber shift with respect to the same peak pre-irradiation; ν = stretching vibrational mode; δ = bending vibrational mode.

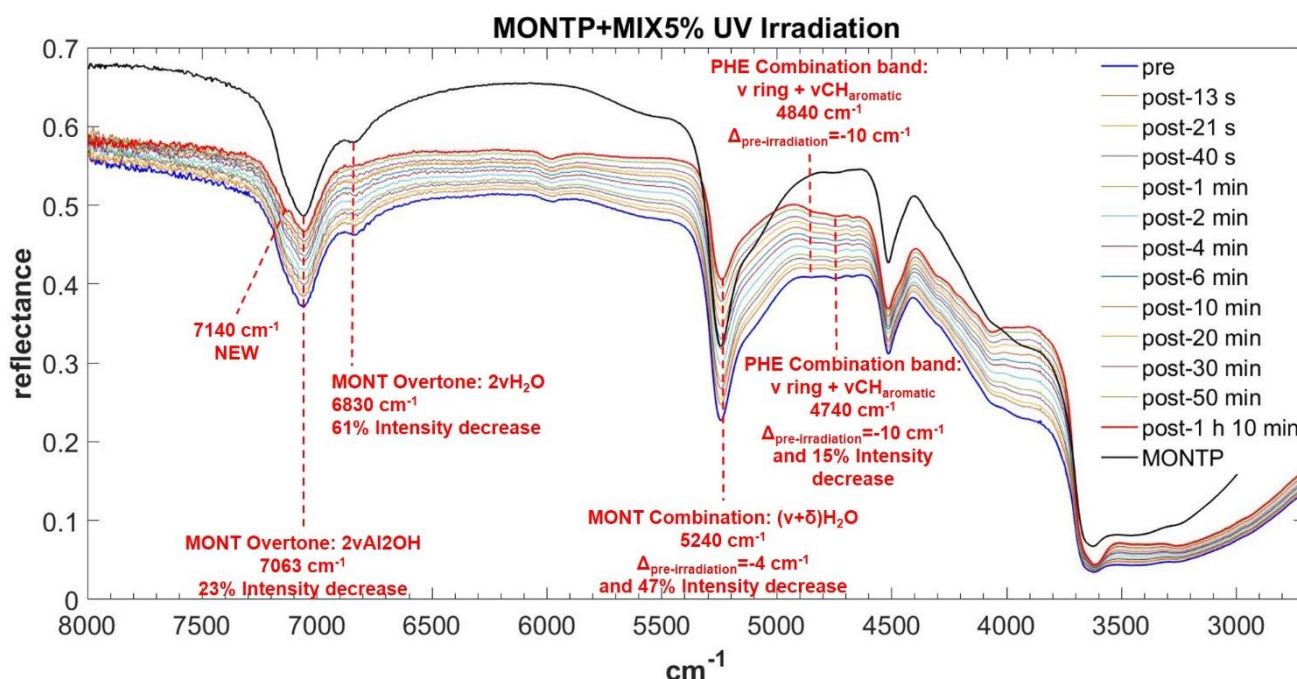


Figure 12. Main changes in NIR features after 70 min of UV irradiation for MIX 5 wt% on montmorillonite containing 1 wt% of Mg-perchlorate. Legend: $\Delta_{\text{pre-irr}}$ = wavenumber shift with respect to the same peak pre-irradiation; ν = stretching vibrational mode; δ = bending vibrational mode.

From these data, AMP appears to be the most difficult molecule to detect in the NIR, because it does not absorb in the 8000-20000 cm^{-1} spectral range and presents only a few weak bands in the 2000-8000 cm^{-1} spectral range detectable mainly at 5 wt%, although a very weak band at 4730 cm^{-1} can still be detected at 1 wt%. Based on DFT calculations at anharmonic level, we assigned this band to two strong combinations: $\nu\text{OH} + \delta\text{OH}_{\text{phosphate}}$ and $\nu_{\text{asym}}\text{NH}_2 + \delta\text{C}_8\text{H}$, $\delta_{\text{rock}}\text{NH}_2$. Due to molecule-mineral interactions, this band undergoes a significant high-frequency shift of 90 cm^{-1} with the respect to the case of pure AMP in the solid state, which indicates that the amino group of the adenine moiety is mainly involved in the interactions with the mineral. The other detectable peaks in the region of C-H stretching, instead, undergo less significant shifts, i.e. 20 cm^{-1} for the peak at 2965 cm^{-1} , 14 cm^{-1} for the peak at 2930 cm^{-1} , -17 cm^{-1} for the peak at 2878 cm^{-1} , -31 cm^{-1} for the peak at 2852 cm^{-1} , and no shift for the peak at 2700 cm^{-1} . In the presence of 1 wt% of magnesium perchlorate, we can observe some differences, i.e. the peaks at 2945 cm^{-1} and 2883 cm^{-1} which are at exactly the same wavenumber of pure AMP, and the peaks at 2930 and 2878 cm^{-1} which are not observable. Unfortunately, most of these peaks are too weak to evaluate intensity variations due to UV irradiation, but new features due to the formation of photoproducts are clearly observed at 3390, 4070, 4220 and 4460 cm^{-1} .

GLU adsorbed on montmorillonite presents remarkable red-shifts with respect to pure GLU in the solid state, of 240 and 230 cm^{-1} for the peaks of highest intensity at 6468 and 4870 cm^{-1} , respectively, attributable to the overtone $2\nu_{\text{asym}}\text{NH}_2$ and the combination $\nu_{\text{asym}}\text{NH}_2 + \delta\text{NH}_2$, which are the same in the presence of 1 wt% of magnesium perchlorate and indicate the involvement of the NH_2 functional group in the interaction with the mineral. New features appear as consequence of UV irradiation, at 4075, 3280, and 2343 cm^{-1} , indicative of the formation of photoproducts.

PHE is the molecule with the most intense bands in the NIR when adsorbed on montmorillonite. Significantly intense bands detectable also at 1 wt% occur at: 5970 cm^{-1} , attributable to the overtone $2\nu\text{CH}_{\text{aromatic}}$, blue-shifted of 10 cm^{-1} with respect to pure PHE in the solid state; 4830 cm^{-1} and 4745 cm^{-1} , attributable to the combinations $\nu\text{ ring} + \nu\text{CH}_{\text{aromatic}}$, blue-shifted of 159 cm^{-1} and 108 cm^{-1} , respectively; 4673 cm^{-1} , attributable to $\nu\text{ ring} + \nu\text{CH}_{\text{aromatic}}$, blue-shifted of 92 cm^{-1} ; and 4045 cm^{-1} , attributable to the combination ring def + $\nu\text{CH}_{\text{aromatic}}$, not shifted with respect to pure PHE in the solid state. Weaker bands can be detected also for the 0.1 wt% concentration at 3261 cm^{-1} , attributable to the fundamental $\nu\text{CH}_{\text{aromatic}}$, blue-shifted of 141 cm^{-1} with respect to pure PHE in the solid state, and 2928 cm^{-1} , attributable to the fundamental $\nu_{\text{asym}}\text{CH}_2 / \nu\text{CH}$, red-shifted of 42 cm^{-1} with respect to pure PHE in the solid state. These variations indicate involvement of the C-H of the aromatic ring in the interaction with the mineral. In addition, in the case of 5 and 1 wt%, a new band due to molecule-mineral interactions is detectable at 6340 cm^{-1} , indicative of chemisorption. The presence of 1 wt% of magnesium perchlorate does not seem to affect the NIR features, while UV irradiation causes intensity decreases for the bands at 5970 cm^{-1} (about 4%), 4745 cm^{-1} (about 43%), 2545 cm^{-1} (about 13%), and 2455 cm^{-1} (about 35%), along with the formation of a peak at 7140 cm^{-1} due to formation of photoproducts.

PHA adsorbed on montmorillonite is also clearly detectable in the NIR thanks to the band at 6000 cm^{-1} , attributable to the overtone $2\nu\text{CH}$, red-shifted by 50 cm^{-1} with respect to pure PHA in the solid state when PHA is adsorbed on montmorillonite at 5 wt%, which undergoes a further red-shift of 10 cm^{-1} at 1 wt%, and the band at 4665 cm^{-1} , attributable to the combination $\nu\text{ ring} + \nu\text{CH}_{\text{aromatic}}$, whose red-shift with respect to pure PHA in the solid state goes from 14 cm^{-1} when PHA is adsorbed on montmorillonite at 5 wt% to 19 cm^{-1} at 1 wt% and 24 at 0.1 wt%. Significant shifts with respect to pure PHA can be observed also for other weaker bands like the combination $\delta\text{CH}_{\text{aromatic}} + \nu\text{CH}_{\text{aromatic}}$ at 4617 cm^{-1} with a blue-frequency shift of 32 cm^{-1} , the fundamental νCH at 3078 cm^{-1} with a red-shift of 10 cm^{-1} , the combination $\delta\text{OH} + \nu\text{ ring}$ at 2905 cm^{-1} with a blue-shift of 23 cm^{-1} , and the combination $\delta\text{OH} + \nu\text{OH}$ at 2814 cm^{-1} with a blue-shift of 44 cm^{-1} . Also in this case, new bands due to molecule-mineral interactions are observed, one at 8490 cm^{-1} at all concentrations, and another at 3505 cm^{-1} only detectable at 5 wt%. In the case of PHA, many peaks are slightly different in the sample spiked with 1 wt% magnesium perchlorate: 4660 cm^{-1} $\Delta_{\text{PHA}}=-19\text{ cm}^{-1}$, 4210 cm^{-1} $\Delta_{\text{PHA}}=-12\text{ cm}^{-1}$, 4149 cm^{-1} $\Delta_{\text{PHA}}=-7\text{ cm}^{-1}$, and 3069 cm^{-1} $\Delta_{\text{PHA}}=-19\text{ cm}^{-1}$. Other differences are the presence of the peak at 3940 cm^{-1} $\Delta_{\text{PHA}}=30\text{ cm}^{-1}$, which is not observable in the analogous sample without magnesium perchlorate, and the absence of the peaks at 2928 and 2858 cm^{-1} , which are detectable in the analogous sample without magnesium perchlorate. The effects of UV irradiation are, instead, negligible.

The NIR spectrum of the mixture of molecules adsorbed on montmorillonite appears dominated by PHE's features. The main peaks of PHE adsorbed on montmorillonite can be detected with small shifts, i.e. 5978 cm^{-1} (blue-shifted of 8 cm^{-1} with respect to PHE adsorbed on montmorillonite), 4850 cm^{-1} (blue-shifted of 20 cm^{-1} with respect to PHE adsorbed on montmorillonite), 4750 cm^{-1} (blue-shifted of 5 cm^{-1} with respect to PHE adsorbed on montmorillonite), and 4670 cm^{-1} (red-shifted of 3 cm^{-1} with respect to PHE adsorbed on montmorillonite). Furthermore, the new peak at 6340 cm^{-1} due to molecule-mineral interactions is the same observed in the case of PHE adsorbed on montmorillonite. Effects of the presence of magnesium perchlorate are not evident, while UV irradiation causes red-shifts of 10 cm^{-1} for the peaks at 4850 and 4750 cm^{-1} , not observed in the case of PHE adsorbed on montmorillonite. The peak at 4740 cm^{-1} also undergoes an intensity decrease, as observed with PHE adsorbed on montmorillonite, but at a much lower extent (15% vs. 43%). Finally, UV irradiation causes the formation of the same photoproduct as in the case of PHE adsorbed on montmorillonite, as indicated by the appearance of the same band at 7140 cm^{-1} .

Regarding NIR features of montmorillonite, as seen in Table 2, they are not affected by molecular interactions, neither by the presence of 1 wt% of magnesium perchlorate. However, UV irradiation causes some changes that depend on the molecule adsorbed.

In the case of AMP, the peak of montmorillonite at 7063 cm^{-1} , attributable to the overtone of the Al_2OH stretching, undergoes a red-shift of 15 cm^{-1} as consequence of 70 min of UV irradiation and a 28% intensity decrease. The peak at 6830 cm^{-1} , due to the overtone of the stretching of the water adsorbed on the mineral, decreases in intensity up to complete disappearance after 70 min of UV irradiation. The peak at 5244 cm^{-1} , due to a combination of stretching and bending of adsorbed water, also undergoes a significant decrease in intensity of 53% in 70 min of UV irradiation.

In the case of GLU, the peak of montmorillonite at 7063 cm^{-1} does not undergo any red-shift as a consequence of 70 min of UV irradiation, but a similar 28% intensity decrease is observed. The peak at 6830 cm^{-1} undergoes a decrease in intensity of 86%, similar to the case of AMP. The peak at 5244 cm^{-1} also undergoes a significant decrease in intensity of 56% in 70 min of UV irradiation.

In the case of PHE, similar to the case of AMP, the peak of montmorillonite at 7063 cm^{-1} undergoes a red-shift of 10 cm^{-1} as consequence of 70 min of UV irradiation and a 24% intensity decrease. The peak at 6830 cm^{-1} undergoes a decrease in intensity of 96% similarly to the case of AMP and GLU. The peak at 5244 cm^{-1} undergoes a red-shift of 8 cm^{-1} and a significant decrease in intensity of 61% in 70 min of UV irradiation, as observed also in the other cases. In contrast to the other examples, the peak at 4515 cm^{-1} undergoes a slight intensity decrease of 6% due to a combination of stretching and bending of the OH group.

For PHA, as for GLU, the montmorillonite peak at 7063 cm^{-1} does not undergo a red-shift as a consequence of 70 min of UV irradiation, but a similar 25% intensity decrease is observed. The peak at 6830 cm^{-1} displays a lower decrease in intensity with respect to the other cases (52%). Similar to PHE, the peak at 5244 cm^{-1} undergoes a red-shift of 9 cm^{-1} and a significant decrease in intensity of 54% in 70 min of UV irradiation. As for PHE, the peak at 4515 cm^{-1} also undergoes a slight intensity decrease of 8%.

In the case of the mixture, as observed for PHA and GLU, the montmorillonite peak at 7063 cm^{-1} does not undergo a red-shift as a consequence of 70 min of UV irradiation but we observe a 23% intensity decrease. As observed with PHA, the 6830 cm^{-1} peak shows a smaller decrease in intensity (61%). Similar to PHE and PHA, the peak at 5244 cm^{-1} undergoes a red-shift of 4 cm^{-1} and a significant decrease in intensity of 47% in 70 min of UV irradiation. No variations of the peak at 4515 cm^{-1} are observed in this case.

4 Discussion

The NIR characterization of Mars soil analog samples obtained by doping montmorillonite (MONT) with the nucleic acid component adenosine 5'-monophosphate (AMP), the amino acids L-glutamic acid (GLU) and L-phenylalanine (PHE), and phthalic acid, which is a plausible precursor of the chlorohydrocarbons detected on Mars by the *Curiosity* rover, has shown that this spectroscopic technique is suitable for detection of these kinds of organic molecules when adsorbed on MONT (Steele et al., 2018). Indeed, some bands can be detected for all molecules adsorbed on MONT, with estimated detection limits of $\leq 250\text{ ppm}$ for PHE, $\leq 1000\text{ ppm}$ for PHA and GLU, and $\leq 50000\text{ ppm}$ for AMP (despite a very weak band at 4730 cm^{-1} can be detected also at 1000 ppm). In the case of samples

with an adsorbed mixture of all four molecules (MIX), PHE's adsorption features predominate, consistent with PHE being the molecule with the lowest detection limit.

We did not observe significant differences in the presence of 1 wt% magnesium perchlorate, except for MONT+PHA, and to smaller extent MONT+AMP, where discrepancies are within 10 cm^{-1} . This finding suggests that such an amount of magnesium perchlorate does not interfere with molecule-mineral interactions.

We did not observe significant frequency shifts at different organic concentrations, but only fewer and less intense bands with lower organic concentrations, as expected, except for PHA, which presents larger red-shifts at lower concentrations. Different molecular peak positions at different surface coverage can be explained considering that molecules may occupy different surface sites at different concentrations, which results in different spectroscopic features.

Very importantly, we observed significant frequency shifts with respect to pure molecules (even greater than 100 cm^{-1}) for some vibrational modes of molecular functional groups involved in the interaction with the mineral and new features due to chemisorption in the case of PHE and PHA, while the mineral features were unaltered. In addition, in the case of samples obtained by adsorbing a mixture of all four molecules on montmorillonite, the bands of PHE detected were shifted with respect to the case of the samples with only PHE adsorbed on montmorillonite, indicating that molecules influence each other's adsorption, consequently affecting spectroscopic features. These results highlight the importance of establishing databases of spectroscopic features of molecule-mineral complexes because the features observed for organic molecules change when they are adsorbed on minerals; therefore, databases of spectroscopic features for pure molecules and pure minerals are not adequate for correct assignments.

Moreover, UV irradiation experiments show that at very short timescales both molecular and mineral NIR features are affected by UV radiation. Regarding molecular features, in most of the cases we do not observe significant intensity decreases, indicating that montmorillonite does not promote molecular degradation, except for the combination band $\nu_{\text{ring}} + \nu_{\text{CH}_{\text{aromatic}}}$ of PHE at 4745 cm^{-1} (43% of loss), which was limited to only 15% loss in the samples for which all four molecules adsorb as a mixture on montmorillonite. This result possibly indicates a photoprotective effect due to interactions among the different molecules. On the other hand, we observed the appearance of new features due to the formation of photoproducts in the case of AMP, PHE and GLU, whose nature is still to be determined. Regarding mineral features, the changes due to UV irradiation are particularly important both for the structural O-H groups of the mineral and the adsorbed water, whose bands undergo significant intensity decreases. Very interestingly, such changes depend on the specific molecule adsorbed. Indeed, in the case of AMP and PHE adsorbed on montmorillonite, we observed red-shifts of the overtone of the Al_2OH stretching, whereas for PHE, PHA and MIX, we observed red-shifts of the combination band of stretching and bending of adsorbed water, indicating a weakening of the respective bonds as consequence of UV irradiation. In addition, the extent of intensity decrease of these bands is different for the different molecules adsorbed. These results indicate that molecule-mineral interactions influence the possible photochemical mechanisms induced by UV radiation.

5 Conclusions

This study reports changes of the NIR spectroscopic features due to molecule-mineral interactions, molecular concentration, presence of oxidants such as magnesium perchlorate, and UV irradiation, for molecules relevant in the context of life detection, such as adenosine 5'-monophosphate, L-glutamic

acid, L-phenylalanine, and phthalic acid, adsorbed on montmorillonite, a clay mineral common on Mars.

We observed that molecule-mineral interactions and UV irradiation cause the most significant changes in NIR spectra.

Specifically, chemisorption determines frequency shifts even greater than 100 cm^{-1} with respect to pure molecules for the vibrational modes of molecular functional groups involved in the interaction with the mineral and the appearance of new features. Very importantly, molecule-mineral interactions strongly depend on many factors, including conditions of preparation of molecule-mineral complexes, specific properties of minerals and molecules, presence of multiple components in the system, and so on. Therefore, to interpret NIR features of martian samples, it is very important to prepare many analog samples obtained with various combinations of relevant molecules and minerals under conditions mimicking the martian environment and establish large libraries of NIR spectra to use as reference.

Based on this study and previous works (Fornaro et al., 2018c, and references therein; Schuerger et al., 2006, 2003), the effects of UV radiation on NIR features occur quite rapidly, showing both variations in intensities and frequencies of molecular and mineral bands, and the appearance of photoproduct marker bands. Such effects concern mostly surficial samples reached by UV radiation, although regolith-bound organics moved around by dust devils and aeolian erosion can be affected as well. Subsurface exploration is therefore necessary to investigate organics in their original form. In this regard, the *ExoMars 2020* mission will offer a great opportunity to inspect the alteration state of organic matter as a function of depth up to 2 m, providing also valuable information for interpreting *Mars 2020* data acquired on more altered surficial samples.

This study highlights the importance of establishing custom databases of spectroscopic features for molecule-mineral complexes under martian-like conditions in order to assist interpretation of data collected by instruments based on NIR spectroscopy, such as SuperCam on board *Mars 2020* space mission, ISEM, *in situ* and MicrOmega on board *ExoMars 2020* space mission. To this end, such investigations provide important information on where to look for molecular spectroscopic features in NIR when molecules are embedded in plausible “martian” mineral matrices, along with assignments of molecular vibrational modes in the NIR spectral region, which are very scarce in the literature.

Furthermore, this work enables us to identify the molecules with the highest detection potential in the NIR when associated with specific minerals and, hence, select the best analog samples to be used for testbed analyses of flight instruments.

6 Conflict of Interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

7 Author Contributions

T.F. prepared and characterized the Mars soil analog samples, analysed all data and wrote the paper. J.B., G.P. and M.A.C. contributed to NIR characterization and UV irradiation experiments. M.B. performed DFT calculations. M.J. carried out mineral characterization. D.F. performed XRF and IRMS measurements. A.S. and R.H. mentored this work and contributed to paper revision.

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11 Supplementary Material

Supplementary Material is available online.