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# The photochemical evolution of polycyclic aromatic hydrocarbons and nontronite clay on early Earth and Mars

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#### ABSTRACT

17		
18	Keywords:	The photochemical evolution of polycyclic aromatic hydrocarbons (PAHs), an abundant form of
19	early Mars	meteoritic organic carbon, is of great interest to early Earth and Mars origin-of-life studies and
20	polycyclic aromatic hydrocarbons	current organic molecule detection efforts on Mars. Fe-rich clay environments were abundant
21	clays	on early Earth and Mars, and may have played a role in prebiotic chemistry, catalyzing the
22	-	breakdown of PAHs and freeing up carbon for subsequent chemical complexification. Current
23		Mars is abundant in clay-rich environments, which are most promising for harboring organic
24		molecules and have comprised the main studied features by the Curiosity rover in search of
25		them. In this work we studied the photocatalytic effects of the Fe-rich clay nontronite on
26		adsorbed PAHs. We tested the effect of ultraviolet radiation on pyrene, fluoranthene, perylene,
27		triphenylene, and coronene adsorbed to nontronite using the spike technique, and in situ diffuse
28		reflectance infrared Fourier transform (DRIFT) spectroscopy in a Mars simulation chamber. We
29		studied the infrared vibrational PAH bands with first order reaction kinetics and observed an
30		extensive decrease of bands of pyrene, fluoranthene, and perylene, accompanied by the formation
31		of PAH cations, while triphenylene and coronene remained preserved. We further analyzed our
32		irradiated samples with nuclear magnetic resonance (NMR). Our study showed certain PAHs
33		to be degraded via the (photo)Fenton mechanism, even under a dry, hypoxic atmosphere. Using
34		solar spectra representative of early Earth, early Mars, and current Mars surface illumination up
35		to 400 nm, the processes occurring in our set up are indicative of the UV-induced photochemistry
36		taking place in Fe-rich clay environments on early Earth and Mars.

#### 1. Introduction 38

Polycyclic aromatic hydrocarbons (PAHs) represent 10-20% of cosmically available carbon (Tielens, 2013). PAHs 39 and PAH clusters are responsible for the 3-15  $\mu$ m infrared emission bands, which have been observed in remarkably 40 similar quantities across the observable universe (Pendleton and Allamandola, 2002). Formed through combustion 41 processes in circumstellar regions, these compounds undergo gas phase processing in the interstellar medium when 42 subjected to ultraviolet (UV) and Lyman- $\alpha$  radiation and shock waves (Frenklach and Feigelson, 1989; Cherchneff 43 et al., 1992).

Once incorporated into ice and rock in small bodies like asteroids and comets, PAHs experience a wholly different 45 chemical evolution than in the gas phase. They can undergo hydrothermal alteration in the interiors of asteroids (Giese 46 et al., 2019) and radiative ice processing at the surfaces of comets (Kofman et al., 2018). PAHs eventually reach 47 planetary surfaces ferried by interplanetary dust particles (IDPs) and (micro)meteorites. Depending on their origin, 48

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the isotopic signatures of PAHs in meteorites may carry information on accretion processes and aqueous alteration conditions (Lecasble et al., 2022).

The further photochemical evolution of PAHs delivered to a planetary surface environment by meteoritic sources is 51 of great interest to early Earth and Mars origin-of-life studies, as well as current organic molecule detection efforts on 52 Mars. Both early Earth and early Mars likely experienced high luminosity in the ultraviolet part of the solar spectrum, 53 the lack of ozone allowing the full range of UV between 200 and 400 nm to reach the surface (Cnossen et al., 2007; 54 Claire et al., 2012; Cockell, 2000). Currently the Martian atmosphere does little to attenuate the UV flux from 200-55 400 nm, with an average mid-day flux at Gale Crater of 34.1 Wm<sup>-2</sup> (Vicente-Retortillo et al., 2020; Razzell Hollis 56 et al., 2021). 57 Of prebiotic interest to both Earth and Mars are clay-rich environments, and the evolution of organic carbon 58

<sup>50</sup> molecules within them. On Mars, clay environments are ency iten environments, and the evolution of organic encoder
 <sup>50</sup> molecules within them. On Mars, clay environments are most promising for harboring organic molecules (Grotzinger
 <sup>61</sup> et al., 2014; Freissinet et al., 2015; Eigenbrode et al., 2018). Clay minerals on early Earth are posited to have played a
 <sup>62</sup> role in prebiotic molecular evolution (Cairns-Smith, 1966; Hartman, 1975). Bernal first speculated on the importance
 <sup>63</sup> of clays in this respect, because of their well-ordered arrangement, their large surface and interlayer adsorption capacity,
 <sup>64</sup> their shielding capacity against ultraviolet radiation, their ability to concentrate organic molecules, and their ability to
 <sup>64</sup> serve as polymerization templates (Bernal, 1949).

Abiogenesis-focused clay experiments have most often used montmorillonite, which has been shown to catalyze 65 a large variety of organic reactions involving small molecules and PAHs (Bujdak et al., 1994; Kumar et al., 2014; 66 Huang and Ferris, 2006; Juntunen et al., 2018), as well as serve in a protective capacity for small molecules in radiative environments (Scappini et al., 2004). However, Fe-rich smectites, and in particular nontronite, are thought to have been 68 significantly more abundant on early Earth and Mars (Kloprogge and Hartman, 2022), and are indeed the dominant form 69 of smectite detected on Mars (Bibring et al., 2005; Murchie et al., 2009). Little research has investigated the catalytic 70 properties of these smectites, and it is unclear if the results obtained for montmorillonite can be directly translated to 71 nontronite and other Fe-rich smectites (Kloprogge and Hartman, 2022). Here we explore the photochemical evolution 72 of PAHs adsorbed to nontronite clay, and speculate on both of their roles in prebiotic carbon chemistry. 73

## **1.1.** Delivery of PAHs to clay environments on early Earth

The annual flux of carbonaceous material (organic and inorganic) delivered by extraterrestrial sources to Earth is 75 currently 300 tonnes per year (Glavin et al., 2018) and could have been as high as 10 000 to 1 million tonnes per year in 76 the first 0.6 Ga of Earth's history (Whittet, 1997; Jenniskens et al., 2000). The significant volume of extraterrestrially 77 delivered organic matter begs the question of whether it was a source of molecules important for prebiotic chemistry. 78 Meteoritic organic matter is thought to be strongly associated with clay minerals in carbonaceous chondrites, wherein 79 complex chemical evolution may already take place (Pearson et al., 2002). Much of the literature has focused on small 80 molecules contained in meteorites, such as amino acids and nucleic acid bases, and their potential as a carbon source 81 for prebiotic chemistry on early Earth (e.g. (Marty et al., 2013)). However, 75% of extraterrestrial organic matter 82 in meteorites is in aromatic form (Sephton, 2002), and is more likely to survive the journey to a planetary surface, 83 during which much of the small molecules can be destroyed (Basiuk and Navarro-González, 1998; Basiuk et al., 1999). 84 PAHs could later be broken down into smaller, more biologically relevant and reactive molecules by photolysis and 85 photocatalysis in the ultraviolet radiation regime of early Earth (Ehrenfreund et al., 2006). 86 Though little is known about Earth's surface during the Hadean and early Archean, it is thought that the crust would 87

have consisted primarily of basalt and komatiite lavas containing peridotite-derived rocks high in Fe and Mg content
(O'Neil et al., 2008; Davies, 1992). The peridotite and komatiite rocks that made up the ocean floor and emerging
plateaus favored the formation of Fe-Mg clay minerals upon interaction with seawater and hydrothermal alteration.
The rocks would first react to trioctahedral phyllosilicates such as talc, kerolite, or stevensite-saponite, until increasing
weathering would cause these to destablize and form dioctahedral Fe-rich clay minerals, with nontronite as the most
abundant species (Kloprogge and Hartman, 2022).

#### **1.2.** Delivery of PAHs to clay environments on Mars

The estimated current global carbon flux on Mars from cometary impacts is 13 tonnes per year within an order of magnitude, and 50 tonnes per year from asteroids (Frantseva et al., 2018). The amount of organic material delivered to the surface of Mars by IDPs and (micro)meteorites is estimated to be of the order of 1000 tonnes per year (Flynn,

<sup>98</sup> 1996; Flynn et al., 2004) and would have been higher in its early history.

On Mars, ancient sedimentary environments have been found with several tens of nanomoles of carbon compounds 99 (Grotzinger et al., 2014; Freissinet et al., 2015; Eigenbrode et al., 2018). Biological, geological, and meteoritic 100 sources are all possible for the refractory aromatic organic material from the Sheepbed lacustrine mudstones detected 101 by the Sample Analysis at Mars (SAM) instrument onboard the Curiosity rover (Eigenbrode et al., 2018). The 102 diverse molecular contributions in the SAM gas chromatography-mass spectrometry (GC-MS) data from Mojave 103 and Confidence Hills are consistent with the complex chemistry of meteoritic and geological organic matter and the 104 interactions that occur during pyrolysis of sediments (Moldoveanu, 2009). Coevolving volatiles analyzed above 500°C 105 in the Mojave and Confidence Hills analyses suggest the pyrolysis of geological refractory organic macromolecules typically found in carbonaceous chondrites (Remusat et al., 2005; Okumura and Mimura, 2011). 107

The diversity of smectites on Mars includes nontronite, saponite, beidellite, and montmorillonite, with Fe/Mg-rich 108 clays dominating having formed by water-restricted alteration of the original basaltic and ultramafic rocks (Kloprogge 109 and Hartman, 2022). Another theory suggests that primordial clays on Mars were formed when Mars' primary crust 110 reacted with a dense steam or supercritical atmosphere of water and carbon dioxide that was outgassed during magma 111 ocean cooling (Cannon et al., 2017). Such a mechanism would have formed a widespread coherent layer of clays on 112 the Martian surface during the pre-Noachian era. 113

The Martian rock record is abundant in preserved Noachian rocks. CheMin data from the Curiosity rover have 114 shown that ancient fluvio-lacustrine rocks in Gale crater contain up to 35 wt.% of clay minerals (Grotzinger et al., 115 2014; Bristow et al., 2015). While the CheMin instrument has detected collapsed clays, i.e., with no interlayer water, 116 these clays were likely hydrated upon their formation on early Mars, before the changing environmental conditions 117 facilitated the desorption of structural water (Tu et al., 2021). The majority of clay minerals detected by orbital imaging 118 spectrometers are Fe/Mg smectites in 4 billion-year-old rocks (Bibring et al., 2005; Murchie et al., 2009). Nontronite 119 spectral signatures have been studied on the flanks of the central mount of Gale Crater (Poulet et al., 2014; Thomson 120 et al., 2011), and have comprised the main studied features by the Curiosity rover in its search for organic molecules. 12:

#### **1.3.** The influence of clay environments on PAHs 122

The widespread abundance of nontronite on early Earth and Mars would have facilitated contact with PAHs after 123 their liberation from their meteoritic parent bodies. Nontronite has been shown to exhibit both photoprotective and 124 photocatalytic properties to adsorbed organic molecules in the presence of UV radiation, facilitating their preservation 125 or eventual removal (Fornaro et al., 2018b; Poch et al., 2015; dos Santos et al., 2016). 126

Fe-rich clays like nontronite exhibit catalytic properties: they are able to create electron-hole pairs where reaction 127 of the holes with adsorbed water molecules create OH radicals that can oxidize organic compounds, as first described 128 for heterogenous catalysis on TiO<sub>2</sub> surfaces (Linsebigler et al., 1995; Fujishima et al., 2000). Electron-hole pairs can 129 also induce oxidation without creating free radicals by involving chemisorbed molecules that serve as electron hole 130 traps (Shkrob and Chemerisov, 2009; Shkrob et al., 2011a,b). 131

Photocatalysis has been shown to increase PAH degradation in aqueous environments under current terrestrial 132 conditions (Wen et al., 2002; Ohno et al., 2003; Dong et al., 2010a). Environmental pollution studies have highlighted 133 the photocatalytic effects of metal oxides and clays that facilitate the degradation of PAHs (Dong et al., 2010a,b). 134 However, PAHs bond to mineral grains via  $\pi$ -cloud interactions (Campisi et al., 2021, 2022). Experiencing only weak 135 bonds or long range interactions with their mineral substrates, the extent of photocatalytic effects on their degradation 136 in a dry, hypoxic environment without abundant (oxi)hydroxide radical production is questioned. 137

In this work we experimentally tested the effect of UV radiation on different PAHs adsorbed to nontronite clay 138 using the spike technique. Analyses were performed with in situ diffuse reflectance infrared Fourier transform (DRIFT) 139 spectroscopy and nuclear magnetic resonance (NMR). 140

#### 2. Materials & Methods 141

We conducted our experiments at the INAF-Astrophysical Observatory of Arcetri, with an experimental setup in 142 which samples were irradiated for 6 hours under ambient temperature and a nitrogen atmosphere. DRIFT spectra were 143 recorded every hour, with more frequent measurements at the beginning of each experiment. The chemical evolution 144 of organic molecules was described with first-order reaction kinetics. We then extracted the organic fraction from the 145 samples and analyzed them with NMR at Utrecht University. 146

#### 147 2.1. Sample preparation

The Fe-rich clay mineral used in our experiments was selected to be representative of both early Earth 148 and Mars and because it exhibits known catalytic properties. Natural nontronite clay was purchased from the 149 Clay Minerals Society's Source Clays Project, where it is designated NAU-2 and is described structurally as 150  $M_{0,72}^+$ [Si<sub>7.55</sub>Al<sub>0.45</sub>][Fe<sub>3.83</sub>Mg<sub>0.05</sub>]O<sub>20</sub>(OH)<sub>4</sub> (Keeling et al., 2000). Prior to the experiments the nontronite sample 151 needed to be crushed to obtain a grain size of  $\sim 2 \mu m$ . Since manual grinding of a clay may introduce torque, 152 preferentially straining the mineral grains in the sample, reducing tensile strength and ultimately introducing structural 153 changes, the nontronite was ground using a Herzog HP-MA automatic pulverizing mill equipped with a tungsten-154 carbide vessel. This mill allows the total pressing force to be program-controlled, and ensures that no particular sheer 155 direction is imposed on the clay lattice (Herzog, 2022). 156

The indigenous organics present in the clay could be removed with organic solvent washing cycles. However, this could also remove the interlayer water. Though on current Mars the clays detected are collapsed, the clays on early Earth and Mars would have likely had water in their interlayer spaces. As we don't have insight into the potential alteration of the nontronite structure due to known clay organic removal techniques (Fornaro et al., 2020) and the readmittance of water with our ensuing spike technique, we chose to use the clay in its natural state. Other studies have also chosen not to remove the indigenous organics from nontronite (dos Santos et al., 2016). We opted instead to track the chemical evolution of these organics throughout the experiment.

The PAHs used for the experiments were pyrene, perylene, fluoranthene, triphenylene, and coronene, which have 164 all been found in meteorites (Sephton, 2002). They were purchased from Sigma Aldrich and had a purity of 98%. 165 We adsorbed pyrene, fluoranthene, and triphenylene individually to the nontronite powder via the incipient wetness 166 impregnation technique (Fornaro et al., 2018a), in which a molecule in solution is added to the mineral sample until 167 it becomes a thick slurry. The volume of the solution is thus equal to the mineral porous volume. To this end we 168 dissolved pyrene, fluoranthene, and triphenylene in toluene, which does not alter the clay mineral structure unlike other solvents such as methanol. The molecules in solution were deposited on nontronite samples to achieve a 1:100 170 molecule to mineral mass ratio. This ratio produced adequate infrared spectra while allowing the PAH to dry in a way 171 that appeared evenly and thinly spread across the mineral surfaces, instead of forming large crystals as was the case 172 when deposited in larger amounts. The samples were left to dry overnight at room temperature in a chemical hood 173 before being measured with DRIFT spectroscopy. 174

Perylene and coronene do not dissolve readily in toluene, and thus they cannot be deposited via the incipient wetness 175 technique. Instead we mechanically ground each PAH individually with the mineral powder, with the assumption that 176 the work introduced by grinding resulted in physisorption of the two species. To obtain adequate infrared spectra 177 (with the same intensity of organic peaks as in the samples treated with the incipient wetness technique), the molecule 178 to mineral ratio needed to be 1:10. It should be noted that for the nontronite samples, the mechanical grinding may 179 introduce structural changes to the mineral. However, this is not apparent in the IR data, and no clear trends between 180 the preparation methods were found in further analyses, thus the nontronite is assumed to be identical in all samples. 181 For pure PAH irradiation experiments, triphenylene, perylene and coronene were introduced into the experimental 182 apparatus directly from the bottle without any further grinding. Pyrene and fluoranthene had a larger crystal size so 183 were first ground and then introduced into the setup. An overview of the PAHs and the experiments performed with 184 each can be seen in Table 1. 185

#### 186 2.2. Experimental setup

The experimental setup at INAF-Astrophysical Observatory of Arcetri allows for the in situ monitoring of infrared 187 spectra of samples subjected to UV irradiation. The apparatus consists of a Bruker Vertex 70v FTIR double pendulum 188 spectrometer with a Harrick *Praying Mantis<sup>TM</sup>* Diffuse Reflection Accessory for DRIFT measurements (Figure 1). The spectrometer is interfaced with a Newport Oriel 300 W Xenon arc discharge lamp with a spectral range of 200-190 930 nm, which is considered a good analog source for the UV that would have reached the surface of early Earth and 101 Mars. The total UV flux of the lamp in the spectral range of 200-400 nm is  $1.94 \times 10^{16}$  photons s<sup>-1</sup>, as measured by 192 a single monochromator Spectro 320 scanning spectrometer (Instrument System) (Fornaro et al., 2018a). We work in 193 this spectral range because here the molecular absorption is higher. The radiation is focused by an optical fiber from the 194 lamp to the spectrometer with a spot size of 800  $\mu$ m onto one of two sample holders: one holder designated 'large' with 195 radius 0.5 cm, and the other holder designated 'small' with a radius of 0.15 cm. For a spectral range of 200-400 nm, this 196 gives a UV flux of 141 Wm<sup>-2</sup> for the large sample holder and 1570 Wm<sup>-2</sup> for the small sample holder, as compared to 197 the average mid-day flux at Gale Crater on Mars of 34.1 Wm<sup>-2</sup>, calculated using the COMIMART radiative transfer 198

#### Table 1

Overview of experiments performed. Samples were irradiated under a 300 W Xe arc lamp in a chamber at ambient temperature under a nitrogen atmosphere.

PAH	pyrene	fluoranthene	triphenylene	perylene	coronene
formula	C <sub>16</sub> H <sub>10</sub>	C <sub>16</sub> H <sub>10</sub>	C <sub>18</sub> H <sub>12</sub>	C <sub>20</sub> H <sub>12</sub>	$C_{24}H_{12}$
nontronite sample	incipient wetness	inc. wetness	inc. wetness	manual	manual
preparation	impregnation	impregnation	impregnation	grinding	grinding
irradiation pure	6 h at 1570 Wm <sup>-2</sup> (200-400 nm)				
irr. on nontronite	6 h at 141 Wm <sup>-2</sup> (200-400 nm)				
analyses FTIR, NMR FTIR, NMR FTIR				FTIR	FTIR

- 199 model with data from the UV sensor on the Rover Environmental Monitoring Station (REMS) on board the Curiosity
- rover (Vicente-Retortillo et al., 2015; Razzell Hollis et al., 2021).



**Figure 1**: Schematic of the experimental setup at INAF-Astrophysical Observatory of Arcetri. The apparatus allows for the *in situ* monitoring of infrared spectra of samples subjected to UV irradiation.

With this apparatus the modification of the infrared spectrum of a sample due to UV irradiation can be monitored at regular time intervals *in situ* without moving the sample itself. The reference spectrum for each PAH, recorded prior to the experiment, can be seen in Figure S1 in Appendix A. DRIFT spectroscopy measurements were taken in steps of increasing seconds for the first five minutes of irradiation, in steps of increasing minutes for the first hour, and every hour for six hours. The evolution of the PAH molecules could thus be seen in real time, with photochemical degradation manifesting itself as the decrease in peak sizes or the formation of new peaks, indicating bond breakage or new species formation, respectively.

#### 208 2.3. FTIR Analysis

PAH infrared spectra contain a cacophony of absorption peaks due to the many vibrational modes: ring deformation,
C-H out-of-plane (oop) bend, C-H in-plane (ip) bend, C-C stretch, C=C stretch, and C-H stretch. In this work the
individual peaks were grouped in these categories for analysis, and the C-H in-plane bend and C-C stretch were
grouped together since many of the bands overlap. Peak assignments were taken from reference spectra from the
Spectral Database of Organic Compounds (SDBS) (SDBS, 2022) and the NASA Ames PAH IR Spectroscopic Database
(Boersma et al., 2014; Bauschlicher et al., 2018; Mattioda et al., 2020a). The peak assignments can be seen in Table
S1 in Appendix A.

The area of each peak can be seen as proportional to the amount of molecules with that vibrational mode. The peak areas were plotted against time, and the destruction or formation rates, the half-lives, and cross sections were calculated following first order kinetics relations (Cottin et al., 2003; ten Kate et al., 2005). First order kinetics assume a one-toone interaction between one UV photon and one broken linkage in a molecule. Not all of the peaks investigated were possible to fit with one exponential function, indicating it might take more than one photon to break the corresponding bond, or multiple pathways take place. Multiphoton processes in which a PAH absorbs a second photon before it has

radiated the energy of the first photon away have a low probability but can be important as well (Tielens, 2008). Such processes were not taken into account in this analysis.

The destruction rate  $\beta$  is calculated fitting the following first order rate equation,

$$\frac{A(t)}{A_0} = Be^{-\beta t} + C \tag{1}$$

where  $A_0$  is the area of the peak at time t = 0,  $\beta$  is the degradation rate of the peak, B is a coefficient describing the fraction of the sample which has interacted with the radiation, and C is a coefficient describing the fraction of

the sample which has not interacted with the radiation. The radiation only penetrates a few microns into the sample,

whereas the IR laser penetrates much further into the sample. B + C should thus equal 1.

The equation for the destruction half-life  $t_{d1/2}$ , or the time it takes for half of the material to be destroyed, is

$$t_{d1/2} = \frac{\ln(2)}{\beta} \tag{2}$$

where  $\beta$  is the degradation rate.

The cross section describes the probability of interaction between the molecule and the UV radiation. Thus the higher the cross section, the higher the chance that there will be an interaction. The destruction cross section is defined as

$$\sigma_d = \frac{\beta}{\Phi} \tag{3}$$

where  $\beta$  is the degradation rate and  $\Phi$  is the flux of the lamp over a specified wavelength range.

Similarly, the formation rate of new species is calculated by fitting the following first order rate equation,

$$\frac{A(t)}{A_{max}} = 1 - e^{-\alpha t} \tag{4}$$

where  $A_{max}$  is the maximum peak area and  $\alpha$  is the formation rate of the peak.

Having obtained the formation rate, we can calculate the formation half-life  $t_{f1/2}$ , or the time it takes for half of the reaction products to form, with  $t_{f1/2} = \frac{\ln(2)}{\alpha}$ . The formation cross-section is then  $\sigma_f = \frac{\alpha}{\Phi}$ .

#### 233 2.4. NMR Analysis

After irradiation we recovered the samples from the setup and stored them in Eppendorf tubes for further proton NMR analysis. This was a qualitative analysis in order to see whether the irradiated samples contained any reaction products, and whether the characteristic PAH peaks showed significant changes.

We extracted the organic fraction from the nontronite samples by incubating the samples in a bath at 60°C in a 3:1 solution of HPLC-grade methanol and chloroform. The samples were then centrifuged at 1500 rpm for 8 min. The supernatant was removed and passed through a syringe filter to eliminate any residual mineral, and subsequently dried under nitrogen overpressure.

PAH standards, pure PAH samples, and extracted samples were suspended in fully deuterated chloroform and analyzed using a 600 Bruker NMR. Deuterated chloroform was used as only non-polar breakdown products were expected from the irradiation of PAHs in a dry, hypoxic environment, where there is little water and oxygen to generate polar compounds. Standard <sup>1</sup>H 1D spectra were acquired using the zg pulse sequence, using 2.7 s acquisition time, 4 sec recycle delay and 256 scans (total measurement time 30 min per 1D). Spectra were processed using exponential line broadening of 0.3 Hz before Fourier transform using Bruker Topspin and plotted in MeRestNova for the paper figures. We performed this analysis only on pyrene, fluoranthene, and triphenylene, as perylene and coronene were not

(1)

#### Table 2

Nontronite infrared bands and assignments, taken from Madejová (2003) and Frost et al. (2002).

vibrational mode	band (cm <sup>-1</sup> )	peak
Si-O-Si bend	475	1
Fe-O oop bend	677	2
Si-O stretch	1200	3
O-H bend	1630	4
O-H stretch	3580	5



#### Figure 2

DRIFT spectrum of nontronite before and after irradiation. Numbered peaks correspond to vibrational modes in Table 2.

sufficiently soluble in chloroform. The NMR spectra of PAHs irradiated on nontronite were compared to the PAHs

irradiated pure, the PAH standards, and the experimental blanks, which consisted of the extraction solvent that had

<sup>250</sup> been passed through a syringe filter, then dried and resuspended in deuterated chloroform.

#### 251 3. Results

#### 252 3.1. Irradiation of nontronite

The infrared spectrum of pure nontronite exhibited some changes after 6 h of irradiation. The Si-O-Si, Fe-O, and 253 Si-O infrared bands (listed in Table 2 and denoted by red numbers in Figure 2), did not show significant changes. The 254 O-H stretching band at 3580 cm<sup>-1</sup> did narrow after irradiation as a consequence of desorption of water physisorbed 255 on the clay, which is responsible for the broader band observed pre-irradiation in the same region and indicated by the 256 purple line in Figure 2. The band at 1630 cm<sup>-1</sup> is attributed to the O-H bending of water in the nontronite interlayer space (Frost et al., 2002). We observe the decrease and shift of this band as some of the interlayer water is desorbed 258 through irradiation. The bands at ~2900 cm<sup>-1</sup> indicate C-H stretching in minor organic contaminants in the clay. These 250 were not removed from the clay prior to the experiment in order to avoid possible alterations of the clay structure that 260 might be caused by treatments usually employed to remove organics. The C-H stretching bands became more prominent 261 post-irradiation as the absorbance of water decreased, revealing the bands of the nontronite spectrum. 262

#### 263 3.2. Irradiation of PAHs

We monitored the irradiation of PAHs *in situ* with DRIFT spectroscopy, with measurements taken in steps of increasing seconds for the first five minutes of irradiation, in steps of increasing minutes for the first hour, and every hour for six hours. All PAHs were irradiated pure and adsorbed to nontronite (Table 1).

Figure 3 shows the FTIR spectrum of fluoranthene adsorbed to nontronite before irradiation (in blue) and after 6 h of irradiation (in orange), where all of its characteristic peaks have decreased or disappeared entirely, revealing the underlying spectrum of nontronite. The nontronite band at 3580 cm<sup>-1</sup> (O-H stretch, marked with a dotted circle in Figure 3), which was obscured by the more prominent fluoranthene spectrum prior to irradiation, is visible postirradiation. We modeled the decay of peaks with Equations 1-4. The decay parameters of fluoranthene bands are listed in Table 3. The C-H in-plane bend and C-C stretch were grouped in one category, since many of the bands overlap. The parameters for the different bands are in a similar range and were averaged to give parameters for the total molecule.

We observed the degradation of perylene when adsorbed to nontronite and exposed to UV for 6 h (Figure 4a), as 274 well as the growth of some peaks (Figure 4b). The perylene bands that decreased did so at similar rates, though many 275 of the perylene bands could not be modeled with Equations 1-4 (Table S1, Figure S6 in Appendix A), and were thus not 276 included in the analysis. The formation of two bands in the C=C range is pointed out with red arrows in Figure 4a, and 277 manifests as the growth and rightward shift of bands at 1543 cm<sup>-1</sup> and 1346 cm<sup>-1</sup> (Figure 4b). The formation kinetics 278 of new bands is shown in Figure 4c, and is consistent with the formation of perylene cations (Szczepanski et al., 1993). 279 However, there should also be a rightward shift and growth of the peak at 1334  $\text{cm}^{-1}$  to 1318  $\text{cm}^{-1}$ , which we do not 280 observe. This could be due to the stifling of certain vibrations because of mineral matrix or PAH crystal effects, which 281

#### Table 3

Degradation kinetics of UV-irradiated fluoranthene ( $C_{16}H_{10}$ ) adsorbed to nontronite. Parameters were calculated with Equations 1-4.

mode	$t_{d1/2}$ (min)	$\sigma_d ~(\times E^{-19} cm^2)$	
ring deformation	83±21	3±1	
C-H oop bend	69±32	$10\pm4$	
C-H ip bend/	62±13	$5\pm1$	
C-C stretch			
C=C stretch	74 <u>+</u> 17	4±1	
total molecule	72±20	6±2	







do not allow the PAHs to vibrate freely as they would in the gas phase. This could also explain the differences in the intensities of the 1543 cm<sup>-1</sup> and 1346 cm<sup>-1</sup> bands between our experiments and those reported in the literature for gas phase or matrix-isolated species. In addition, the area around 1318 cm<sup>-1</sup> is very crowded with intense underlying nontronite bands. This could obscure detection of the 1318 cm<sup>-1</sup>, if it is present.

The formation of PAH ion peaks generally show an initial growth before plateauing off or slightly decreasing due 286 to photobleaching of the ions (Mattioda et al., 2020a; Hudgins and Allamandola, 1997; Hudgins and Sandford, 1998), 287 as seen in our results in Figure 4c. Photoproducts, on the other hand, would increase linearly with time (Mattioda 288 et al., 2020a), and any PAH hydrogenation would be associated with the growth of C-H stretching bands in the 2800-289 3000 cm<sup>-1</sup> region (Cruz-Diaz et al., 2020), which we do not observe as the perylene C-H band in that region is 290 degraded (Table 5). However, the growth of the 1543  $cm^{-1}$  and 1346  $cm^{-1}$  bands could alternatively be indicative of 291 C-H bending bands (Mattioda et al., 2020b), suggesting the hydrogenation of the perylene molecules. Whether we are 292 observing the formation of perylene cations or rather the hydrogenation of perylene molecules could be hinted at by 293 which of the two are more stable on the nontronite surface. Previous work has shown that PAH radical cations can 294 persist on irradiated clay surfaces for several hours in an oxic environment, before they are further broken down by 295 reactive oxygen species (Jia et al., 2019). As the atmosphere in our experiments is largely anoxic, we could expect both 296 perylene cations and hydrogenated perylene to remain stable for the length of the experiment. 297

When pure PAHs were irradiated, only the spectrum of triphenylene exhibited changes (Figure 5a), while no changes were observed in the spectra of pyrene, perylene, fluoranthene, and coronene (Figure S2 in Appendix A). While pure triphenylene manifested peak degradation during irradiation, triphenylene on nontronite exhibited peak growth (Figure 5b) at similar rates for all peaks. This indicates preservation of the molecule: the slight growth of peaks at the beginning of the experiment is probably due to the rearranging of the PAH molecules in the sample due to minimal heating effects. The peak degradation and growth kinetics of triphenylene are compared in Table 4.

The reaction kinetics of all PAHs adsorbed to nontronite are listed in Table 5. Pyrene, fluoranthene, and perylene 304 were degraded in the presence of UV and nontronite, with at least half of the PAHs removed after 6 h of irradiation. 305 Triphenylene and coronene exhibited peak growth in the presence of UV and nontronite, indicating the retention of 306 the PAHs in the samples during 6 h of irradiation. Again, the growth of the IR bands is rather due to the rearranging 307 of the PAH molecules in the sample at the start of the experiment rather than the formation of new products, as there 308 are no new peaks forming. Desorption of PAH molecules by the UV lamp or the IR beam was discounted, based on 300 temperature measurements of the sample area and previous work in the setup, which has not encountered issues with 310 such effects (Potenti et al., 2018; Poggiali et al., 2020; Fornaro et al., 2018a). The spectra pre- and post-irradiation of 311 all UV-irradiated PAHs adsorbed to nontronite are shown in Figure S3 and their reaction kinetics in Figures S4-S6 in 312 Appendix A. 313

The clay-related organic contaminants at  $\sim 3000 \text{ cm}^{-1}$  (Figure 2), which were not removed from the clay, were also subject to the irradiation experiments. The degradation of the band at  $\sim 3000 \text{ cm}^{-1}$  belonging to the contaminants may contribute slightly to the parameters calculated for the C-H stretch of perylene, triphenylene, and coronene. However,



Figure 4: FTIR spectra of perylene adsorbed to nontronite before and after irradiation, compared to the spectrum of nontronite post-irradiation (a), (b) formation of new peaks during irradiation of perylene (indicated by red arrows in (a)), (c) reaction kinetics of new peaks shown in (b).

the overall behavior of the bands is reflecting changes in the PAH molecules, given that their kinetics are in line with the kinetics of the other bands (Table 5). For pyrene and fluoranthene the kinetics of the band at ~3000 cm<sup>-1</sup> were discounted, as they exhibited behavior much different to that of the other bands. This may indicate that the behavior of that band was dominated by the eventual degradation of the organic clay-related contaminants.

#### **321 3.3.** Extractions of PAHs post-irradiation

Pyrene, fluoranthene, and triphenylene samples recovered from the irradiation experiments were further analyzed
 with NMR (Figure 6). The NMR spectra of PAHs irradiated on nontronite were compared to the PAHs irradiated pure,
 the PAH standards, and the experimental blanks. Note that the NMR studies are purely qualitative.

The spectra of pure, irradiated, and clay-irradiated PAHs were superimposed (Figures 6a, c, e) and scaled to match 325 the intensities of the characteristic PAH aromatic peaks. This allowed us to detect changes in relative peak intensities 326 indicating degradation. In the pyrene and fluoranthene clay-irradiated samples (Figure 6a, c), the characteristic PAH 327 peaks decreased strongly in intensity relative to the small aromatic peaks in the region, indicating the degradation 328 of the PAHs. These smaller signals surrounding the PAH peaks are likely from aromatic impurities as they are also 320 present in the pure PAH samples but at much lower intensity to the pure PAH signals. Based on the change in relative 330 intensity, about 30% of pyrene and 40% of fluoranthene were degraded. 331 Further analysis of the spectra showed that a new shoulder formed on the clay-irradiated pyrene peak at 8.18 ppm, 332

indicating the presence of a possible partly degraded PAH intermediate (pointed out with a black arrow in Figure 6a).
 A few more new peaks appeared in the clay-irradiated samples at ~3.5 ppm (the range of alcohols) in the pyrene and

#### Table 4

Reaction kinetics of UV-irradiated triphenylene pure and adsorbed to nontronite. Parameters were calculated with Equations 1-4.

triphenylene	pure		on nontronite	
mode	$t_{d1/2}$ (min)	$\sigma_d \times \mathrm{E}^{-19}(\mathrm{cm}^2)$	t <sub>f1/2</sub> (min)	$\sigma_f \times E^{-19}(cm^2)$
ring deformation	200±120	$0.15 \pm 0.08$	13 <u>+</u> 4	22 <u>+</u> 9
C-H oop bend	230±130	$0.14 \pm 0.06$	17 <u>+</u> 7	17 <u>±</u> 8
C-H ip bend/C-C stretch	97±37	0.28±0.09	8 <u>+</u> 2	62 <u>±</u> 32
C=C stretch	110±33	$0.34 \pm 0.01$	7±3	40±20
C-H stretch	94±29	$0.3 \pm 0.1$	10±4	29±14
total molecule	150±69	$0.23{\pm}0.09$	11±4	34±17
flux (Wm <sup>-2</sup> )		1570	141	

*oop: out-of-plane, ip: in-plane*,  $t_{d1/2}$ : destruction half-life,  $\sigma_d$ : destruction cross section,  $t_{f1/2}$ : formation half-life,  $\sigma_f$ : formation cross section



#### Figure 5

FTIR spectra of triphenylene ( $C_{18}H_{12}$ ) irradiated pure (a) and adsorbed to nontronite (b). The nontronite spectrum post-irradiation is shown in black, and the difference spectra of triphenylene (spectrum at t=0 subtracted from spectrum at t=6) are shown in green.

sus triphenylene samples (indicated by black arrows in Figures 6b, f). Fluoranthene already has this shift in its standard

spectra, and a shift here could also be attributed to indigenous organics within the clay structure. The other new signals

<sup>337</sup> in the clay-irradiated samples are accounted for by the experimental blanks.

## 338 4. Discussion

#### **339 4.1.** Effect of UV on nontronite

Fe-rich clays like nontronite can exhibit catalytic properties in the presence of UV radiation: they can create electron-hole pairs that can react either with adsorbed water molecules creating hydroxyl radicals, or with chemisorbed molecules that serve as electron hole traps (Linsebigler et al., 1995; Fujishima et al., 2000; Shkrob et al., 2011a,b). The presence of water physisorbed on the clay and the structural water of the clay could result in the production of OH radicals that may facilitate the Fenton mechanism (Fenton, 1894). The Fenton reaction results in the oxidation of Fe<sup>2+</sup> to Fe<sup>3+</sup>,

$$Fe^{2+} + H_2O_2 \rightarrow Fe^{3+} + OH^{\bullet} + OH^{-}$$

$$\tag{1}$$

#### Table 5

Reaction kinetics of UV-irradiated PAHs adsorbed to nontronite. Parameters were calculated with Equations 1-4. The destruction half-life  $(t_{d1/2})$ , destruction cross section  $(\sigma_d)$ , formation half-life  $(t_{f1/2})$  and formation cross section  $(\sigma_f)$  are given for each vibrational mode for the corresponding PAH, and averaged for the total molecule. The C-H in-plane bend and C-C stretch are grouped in one category, since many of the bands overlap.

	PAH	pyrene	fluoranthene	perylene	PAH	triphenylene	coronene
mode	degradation	C <sub>16</sub> H <sub>10</sub>	C <sub>16</sub> H <sub>10</sub>	$C_{20}H_{12}$	retention	C <sub>18</sub> H <sub>12</sub>	C <sub>24</sub> H <sub>12</sub>
ring	t <sub>d1/2</sub> (min)	220±65	83±21	n/a	t <sub>f1/2</sub> (min)	13±4	4±3
deformation	$\sigma_{\rm d} \times {\rm E}^{-19}({\rm cm}^2)$	1.4±0.7	3±1	n/a	$\sigma_{\rm f} \times {\rm E}^{-19}({\rm cm}^2)$	22±9	81±87
C-H	t <sub>d1/2</sub> (min)	80±9	69±31	77±33	t <sub>f1/2</sub> (min)	17±7	n/a
oop bend	$\sigma_{\rm d} \times {\rm E}^{-19}({\rm cm}^2)$	3.8±0.8	10 <u>+</u> 4	4±2	$\sigma_{\rm f} \times {\rm E}^{-19}({\rm cm}^2)$	17±8	n/a
C-H in hend	t <sub>d1/2</sub> (min)	180±42	62±13	49±10	t <sub>f1/2</sub> (min)	8±2	n/a
C-C stretch	$\sigma_{\rm d} \times {\rm E}^{-19}({\rm cm}^2)$	1.6±0.5	$5\pm1$	6±2	$\sigma_{\rm f} \times {\rm E}^{-19}({\rm cm}^2)$	62±32	n/a
<u> </u>	t <sub>d1/2</sub> (min)	210±55	74±17	n/a	$t_{f1/2}$ (min)	7±3	2.5±0.6
stretch	$\sigma_{\rm d} \times {\rm E}^{-19}({\rm cm}^2)$	1.4±0.5	4±1	n/a	$\sigma_{\rm f} \times {\rm E}^{-19} ({\rm cm}^2)$	40±20	120±43
C-H	t <sub>d1/2</sub> (min)	n/a	n/a	40±10	t <sub>f1/2</sub> (min)	10±4	7±3
stretch	$\sigma_{\rm d} \times {\rm E}^{-19}({\rm cm}^2)$	n/a	n/a	7±2	$\sigma_{\rm f} \times {\rm E}^{-19} ({\rm cm}^2)$	29±14	39±22
total	t <sub>d1/2</sub> (min)	170±43	72±20	55±17	t <sub>f1/2</sub> (min)	$11\pm4$	4±2
molecule	$\sigma_{\rm d} \times {\rm E}^{-19}({\rm cm}^2)$	2.0±0.6	6±2	6±2	$\sigma_{\rm f} \times {\rm E}^{-19}({\rm cm}^2)$	34±17	79±51

oop: out-of-plane, ip: in-plane

 $t_{d1/2}$ : destruction half-life,  $\sigma_d$ : destruction cross section

 $t_{f1/2}$ : formation half-life,  $\sigma_f$ : formation cross section

The photo-Fenton reaction is enhanced by UV light at wavelengths <350 nm, producing additional OH radicals, and leading to Fe<sup>3+</sup> reduction (Ruppert et al., 1993),

$$Fe^{3+} + OH^{-} \xrightarrow{h\nu(<350 \text{ nm})} Fe^{2+} + OH^{\bullet}$$
(2)

and regeneration of the catalyst,

. . . . .

 $HO^{\bullet} + OH^{\bullet} \rightarrow H_2O_2$ (3)

In our experiments, Reactions 2 and 3 described above can generate Fe<sup>2+</sup> and OH, producing H<sub>2</sub>O<sub>2</sub>, and facilitating the Fenton mechanism leading to the reduction of the clay,

$$Fe(III)smectite + OH^{-} \xrightarrow{h\nu(<350 \text{ nm})} Fe(II)smectite + OH^{\bullet}$$
(4)

The clay may be subject to subsequent reoxidation by H<sub>2</sub>O<sub>2</sub> Jia et al. (2012); Yap et al. (2011),

.

$$Fe(II)smectite + H_2O_2 \rightarrow Fe(III)smectite + OH^{-} + OH^{-}$$
(5)



**Figure 6:** NMR data of irradiated samples. Pure PAHs and PAHs extracted from clay samples are compared to PAH standards and experimental blanks. (a-b) spectra of pyrene, (c-d) spectra of fluoranthene, (e-f) spectra of triphenylene. The data are normalized to the intensity of the PAH peaks. New peaks or shoulders are marked with black arrows. The superimposed spectra were scaled to match the intensities of the characteristic PAH aromatic peaks, revealing potential changes in relative peak intensities.

or in the presence of oxygen,

Fe(II)smectite +  $O_2 \rightarrow Fe(III)$ smectite +  $O_2^-$ 

(6)

forming superoxide radicals (Sỳkora, 1997).

The OH radicals and H<sub>2</sub>O<sub>2</sub> can further catalyze the photo-Fenton reaction and can oxidize PAHs adsorbed to the 341 mineral surface. The hypoxic experimental conditions in our experiments likely inhibit Reaction 6, in part stifling 342 subsequent oxidation of reduced nontronite. Nonetheless, the Fenton mechanism can proceed following the ionization 343 of surface and interlayer water molecules and OH groups by UV radiation. This is supported by the decrease in the O-H 344 bending and stretching bands in the nontronite IR spectrum post irradiation (Figure 2). The removal of interlayer water 345 from the clay is indicative of the processes occurring on Mars, where the CheMin instrument has detected collapsed 346 clays, i.e., with no interlayer water (Tu et al., 2021). The interlayer water was likely removed from these clays over 347 time, facilitated by the changing environmental conditions, particularly the increasingly harsh radiative environment. 348 The catalytic activity of nontronite is excellent over a wide pH range, with nontronite remaining both chemically 349 and mechanically stable, with no measurable Fe leaching, damage to the structure, or noticeable loss of activity (Liu 350 et al., 2014). Reduction in nontronite occurs to the greatest extent in the octahedral sheet iron and oxidation in the 351 tetrahedral sheet iron (Geatches et al., 2012). Reduction of octahedral iron causes a change in specific surface area 352 353 of the smectite sheets, (Lear and Stucki, 1989) which can reduce the interlayer spacing and trap interlayer cations, ultimately changing the reactivity of the clay. 354

Fe<sup>2+</sup> and Fe<sup>3+</sup> act as photocatalysts via the Fenton mechanism at low pHs ( $\leq 4$ ) (Fenton, 1894). Nontronite has the potential to have high surface acidity, due to exchangeable cations in its interlayer space (Laszlo, 1987). This will be more important in a dry environment, with no water molecules to react with to reduce the acidity. Carbocations formed here can further react with interlayer water to produce alcohols and ethers (Theng, 2018; Adams et al., 1979; Nagendrappa et al., 2002). This may explain the appearance of additional peaks at ~3.5 ppm in our NMR spectra of the organic extractions of clay-irradiated samples (Figure 6).

Conversely, Fe-rich minerals can be strong absorbers of UV light, which can provide protection against radiation damage (Pierson et al., 1993; Gauger et al., 2015). Nontronite has been shown to exhibit a photoprotective effect on adsorbed amino acids when irradiated with UV under current Martian surface conditions (Poch et al., 2015; dos Santos et al., 2016). These results suggest that photoprotection by nontronite is not only due to mechanical shielding (elevated because of the clay's high surface area and small pore sizes) (dos Santos et al., 2016), but also caused by stabilizing interactions between the molecules and the mineral surface, which allow absorbed energy to dissipate and photodissociated molecules to recombine (Poch et al., 2015).

Nontronite's high surface area also creates more sites for PAH adsorption to the clay and potential for PAHs to experience the catalytic effects of Fe in the clay. Thus nontronite can exhibit both photoprotective and photocatalytic effects, with the dominant effect seeming to vary per PAH, as described in the next section.

#### **4.2.** Effect of UV on PAHs

The adsorption of PAHs to Fe-rich clay minerals is likely accompanied by the formation of "cation- $\pi$ " interactions at the active sites, inducing electron transfer from PAHs to surface cations (Jia et al., 2018). This would result in oxidation of the PAH,

$$Fe(III)smectite - PAH \xrightarrow{h\nu(<350nm)} Fe(II)smectite - PAH^+$$
(7)

and further oxidation of the PAH by radicals and  $H_2O_2$  produced in Reactions 3-6.

In previous work, the photodegradation of pyrene in solid state on various Fe oxide surfaces had a half-life of 3 to 4 hours depending on the Fe oxide, calculated by extracting and measuring pyrene concentration over irradiation time (Wang et al., 2009). Pyreno was identified as an intermediary reaction product by gas chromatography-mass spectrometry (GC-MS). Similarly, the photodegradation of pyrene on TiO<sub>2</sub> and soil surfaces was studied, and all loss of surfaces was studied to invitation by the Feature mechanism (Zheng et al., 2009, 2010)

of pyrene was attributed to ionization by the Fenton mechanism (Zhang et al., 2008, 2010).

The photodegradation of phenanthrene under visible light (380-780 nm) was found to follow a pseudo-firstorder kinetic model when adsorbed to smectite saturated with  $Fe^{3+}$  cations, with 100% loss after 6 h of irradiation (Jia et al., 2012). They attributed the degradation to the photo-Fenton effect, wherein hydroxyl radicals and singlet

oxygen preferentially attack phenanthrene at positions 9 and 10, eventually leading to the production of 9,10-381 Phenanthrenequinone, phthalate, diisobutyl phthalate and various alkanes, such as esters, alkanoic acids, alkanols, 382 383 alkanes, and dioxanes. Similarly in our experiments, Reaction 4 described above is thought to have generated  $Fe^{2+}$ , producing  $H_2O_2$  and 384 facilitating the Fenton mechanism Jia et al. (2012); Yap et al. (2011), which would have degraded the adsorbed PAHs, 385 along with direct oxidation by OH radicals and water molecules removed from the clay by UV, and the electron transfer 386 in Reaction 7. Interestingly, only pyrene, fluoranthene, and perylene show degradation in their IR spectra and seem to be 387 significantly affected by this mechanism. Though probably experiencing only weak bonding or long range interactions with the nontronite surface Campisi et al. (2021, 2022), in our experiments pyrene, fluoranthene, and perylene were 380 subject to photodegradation under a dry, hypoxic atmosphere. A summary of the results can be seen in Table 6. The 390 averaged total molecule half-lives and cross sections were compared, and scaled with the average mid-day flux at Gale 391 Crater on Mars of 34.1 Wm<sup>-2</sup> (in the 200-400 nm range) Vicente-Retortillo et al. (2015); Razzell Hollis et al. (2021), 392 to reflect their significance on the surface of present Mars. 393 Because of their aromaticity, PAHs are inherently stable molecules with large electronic absorption cross-sections, 394 which makes them strong absorbers of UV light. They are also stable as ions, and are thought to be present both 395 positively and negatively charged in the interstellar medium; when PAHs are ionized the strength of modes in their IR 396 spectra involving C-C stretching vibrations can increase manifold, whereas the C-H stretching and, to a lesser extent, 397 the out-of-plane bending vibrations decrease in strength (Tielens, 2008). In our experiments, the spectra of perylene 398 showed evidence of PAH ions, with the formation of new peaks in the C=C range (Figure 4b, c) and a decrease in

the C-H band. However, we were not able to determine whether perylene was forming cations or rather undergoing
hydrogenation, owing to the difficulty in comparing infrared spectra from our samples with spectra of perylene in the
gas phase. The gaps between our understanding of PAH photochemistry in the gas phase, in ice matrices, and in mineral
matrices warrant a full-scale analytical campaign.

The trends we see in our experiments hold up across the different sample preparation techniques we've used. Our 404 NMR data supports the FTIR data, showing that clay-irradiated pyrene and fluoranthene degraded. The lack of major 405 reaction products in the NMR data could be due to any breakdown products degrading immediately into small volatile 406 hydrocarbons and carbon dioxide (which would not be present in high enough concentrations to cause a change in the 407 spectrum in a DRIFTS type set up). The extraction technique used targeted solid-state products as these are expected to 408 be non-polar given the nature of the parent molecule and the dry, hypoxic experimental conditions. Instead the PAHs 409 may well be degrading directly into CO, CO<sub>2</sub>, C<sub>2</sub>H<sub>2</sub>, CH<sub>3</sub>OH and other small hydrocarbons, which are volatile and 410 readily escape from the system. This is supported by laboratory results of irradiated PAHs adsorbed to forsterite and 411 anatase, where solid state photodegradation of the PAHs resulted in the production of CO<sub>2</sub> (Potenti et al., 2018). 412 Triphenylene and coronene were not degraded by photocatalysis on the surface of nontronite, possibly owing to 413

<sup>413</sup> Triphenylehe and coronene were not degraded by photocatalysis on the surface of nontronite, possibly owing to
<sup>414</sup> different binding geometries and adsorption mechanisms between the PAH and the clay surface. If the PAHs are not
<sup>415</sup> strongly bound to the surface, they may not be affected by the electron transfer reaction and oxidizing radicals produced
<sup>416</sup> by the Fenton mechanism (Reactions 4-7). In general the binding energy increases with the PAH size/surface area, a
<sup>417</sup> trend attributed to the increasing number of contact points (though pyrene is an exception to this, having higher binding
<sup>418</sup> energies than fluoranthene because of its more compact geometry) (Campisi et al., 2022). Conversely triphenylene in
<sup>419</sup> our experiments may have a lower binding energy because of its less compact geometry, facilitating its preservation.
<sup>420</sup> Adsorption energies depend also on the specific geometrical interaction of the PAH and the mineral surface, and might
<sup>421</sup> not have such a straightforward trend, especially with a complex mineral like nontronite.

Triphenylene was shown to degrade when irradiated pure, but was completely preserved when irradiated with nontronite, indicating the clay may be serving in a protective capacity by shielding the PAH from UV radiation. While coronene is stable in pure form, it can be that part of the UV radiation was absorbed by nontronite when they were irradiated together, but its inefficient adsorption to the grains made it immune to the clay's catalytic effects. Alternatively, it is possible a large PAH like coronene with a large electronic absorption cross section absorbed more of the UV radiation, shielding the nontronite and preventing it from forming the required precursors for the Fenton reactions. It is likely the inefficient adsorption of triphenylene and coronene to nontronite grains and/or the large UV absorption capability of coronene are facilitating their deliverance from the photocatalytic effects of nontronite clay.

#### Table 6

Comparing the effect of nontronite on adsorbed irradiated PAHs. Parameters were calculated with Equations 1-4. The halflives were extrapolated to reflect current surface conditions on Mars, where the average mid-day flux at Gale Crater was calculated using the COMIMART radiative transfer model (Vicente-Retortillo et al., 2015; Razzell Hollis et al., 2021).

PAH	pyrene	fluoranthene	perylene	PAH	triphenylene	coronene			
degradation	$C_{16}H_{10}$	$C_{16}H_{10}$	C <sub>20</sub> H <sub>12</sub>	retention	C <sub>18</sub> H <sub>12</sub>	C <sub>24</sub> H <sub>12</sub>			
					$\Rightarrow$				
in the lab where $\Phi = 141 \text{ Wm}^{-2}$ (200-400 nm)									
$t_{d1/2} \text{ (min)}  170\pm43  72\pm20  55\pm17  t_{f1/2} \text{ (min)}  1$						4±2			
$\sigma_{\rm d} \times {\rm E}^{-19}({\rm cm}^2)$	2.0±0.6	6±2	6±2	$\sigma_{\rm f} \times {\rm E}^{-19} ({\rm cm}^2)$	34±17	79±51			
on Mars where $\Phi = 34.1 \text{ Wm}^{-2}$ (200-400 nm)									
t <sub>d1/2</sub> (sol)	$0.5 \pm 0.1$	0.20±0.06	$0.15 \pm 0.05$	$t_{f1/2}(sol)$	0.03±0.01	0.012±0.007			
effect of nontronite	catalytic	catalytic	catalytic		protective	protective			

### 430 5. Conclusion

Our UV irradiation experiments of various polycyclic aromatic hydrocarbons (PAHs) adsorbed to nontronite 431 showed both degradation and retention of PAH molecules (Table 6). The degradation of pyrene and fluoranthene 432 was manifested by the decrease of infrared bands and characteristic NMR aromatic peaks. The formation of new 433 peaks in the perylene spectra indicated the formation of perylene cations or hydrogenation of perylene molecules. The triphenylene and coronene IR bands showed no decrease throughout irradiation, indicating the preservation of 435 the molecules. The degradation of PAHs was attributed to the Fenton mechanism resulting from the photocatalytic 436 activity of the irradiated Fe-rich smectite nontronite. Our results indicate the removal of adsorbed and interlayer water 437 from irradiated nontronite, which facilitates the Fenton mechanism and is indicative of processes on Mars, where the observed clays are collapsed, i.e., with no interlayer water. The interlayer water was likely removed from these clays over 439 time, facilitated by the changing environmental conditions, particularly the increasingly harsh radiative environment. 440 The effect of the Fenton mechanism on PAHs likely depends on their adsorption capacity on nontronite, and 441 their electronic absorption cross section. While nontronite may exhibit both catalytic and protective properties when 442 irradiated with UV, it may also be that PAHs with large electronic cross section can in turn shield nontronite, preventing 443 it from forming the required precursors for the Fenton mechanism. The efficiency of this mechanism thus varies 444 depending on the combination of all these factors, with no straightforward trend. While certain PAHs may have broken 445 down and contributed organic carbon to prebiotic chemistry on clays, others may have remained inert. There exist 446 difficulties in studying PAHs in prebiotic environments, owing to discrepancies in the behavior of infrared spectra 447 between PAHs in experiments like ours and PAHs in the gas phase. The gaps between our understanding of PAH 448 photochemistry in the gas phase, in ice matrices, and in mineral matrices warrant a full-scale analytical campaign. 449 Assuming the simulated solar spectra used in this study are representative of early Earth, early Mars, and current 450 Mars surface illumination up to 400 nm, the processes occurring in our set up are indicative of the UV-induced 451 photochemistry taking place in Fe-rich clay environments on early Earth and Mars. We have demonstrated that 452 453

degradation of PAHs on nontronite via the Fenton mechanism is possible, even under a dry, hypoxic atmosphere. Thus certain meteoritic PAHs were likely degraded into smaller organic molecules on the Martian surface, which should be considered when determining the sources of organic molecules detected by Mars rovers. Future organic molecule irradiation experiments in Martian conditions could explore the reactivity of different kinds of smectites, such as Mg-smectites, and sulfates.

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## **464** Data Availability

- The data supplement to this work is freely accessible at https://public.yoda.uu.nl/geo/UU01/XJOBC1.
- 466 html, https://doi.org/10.24416/UU01-XJOBC1.

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