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New Insights in the Production of Simulated Moon Agglutinates: the Use of Natural Zeolite-Bearing Rocks

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New insight in the production of simulated Moon agglutinates: the use of natural zeolites bearing rocks

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Complete List of Authors:	Manzoli, Maela; University of Turin, Drug Science and Technology Tammaro, Olimpia; Politecnico di Torino Facoltà di Ingegneria Marocco, Antonello; Università degli Studi di Cassino e del Lazio Meridionale, Department of Civil and Mechanical Engineering Bonelli, Barbara; Politecnico di Torino, Materials Science & Chemical Engineering Barrera, Gabriele; INRIM, Electromagnetism Department Tiberto, Paola; INRIM Allia, Paolo; Politecnico di Torino Velez, Jean-Charles; ONERA Midi-Pyrénées Toulouse Centre roggero, aurelien; Institut National des Sciences Appliquées de Lyon Ingénierie des Matériaux Polymères Dantras, Eric; Université Toulouse III Paul Sabatier, Physique des Polymères Arletti, Rossella; Università degli Studi di Modena e Reggio Emilia, Dipartimento di Scienze Chimiche e Geologiche Pansini, Michele; Università degli Studi di Cassino e del Lazio Meridionale, Ingegneria Meccanica, Strutture, Ambiente e Territorio ESPOSITO, Serena; Politecnico di Torino, Department of Applied Science and Technology

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11 Maela, Manzoli¹, Olimpia Tammaro², Antonello Marocco³, Barbara Bonelli², Gabriele Barrera⁴,
12 Paola Tiberto⁴, Paolo Allia^{2,4}, Jean-Charles Matéo-Vélez⁵, Aurélien Roggero^{6,7}, Eric Dantras⁷,
13 Rossella Arletti⁸, Michele Pansini³, Serena Esposito^{2*}
14
15

16
17
18 ¹*Department of Drug Science and Technology, NIS Interdepartmental Centre and INSTM Unit of Torino - University of*
19 *Torino, Via Giuria 9, Torino, 10125, Italy*
20

21
22 ²*Department of Applied Science and Technology and INSTM Unit of Torino – Politecnico, Politecnico di Torino, Corso*
23 *Duca degli Abruzzi 24, 10129, Torino, Italy.*
24

25
26 ³*Department of Civil and Mechanical Engineering and INSTM Research Unit, Università degli Studi di Cassino e del*
27 *Lazio Meridionale, Via G. Di Biasio 43, Cassino, FR 03043, Italy*
28

29 ⁴*INRiM Torino, Advanced Materials for Metrology and Life Sciences, Strada delle Cacce 91, 10143 Torino, Italy*
30

31 ⁵*ONERA, The French Aerospace Lab, 2 Avenue Edouard Belin, Toulouse, France*
32

33 ⁶*Université de Lyon, CNRS, UMR 5223, Ingénierie des Matériaux Polymères, INSA Lyon F-69621 Villeurbanne,*
34 *France*
35

36 ⁷*CIRIMAT-Physique des Polymères, Université Toulouse 3 - Paul Sabatier, 118 route de Narbonne, Toulouse, France*
37

38 ⁸*Department of Chemical and Geological Sciences, University of Modena e Reggio Emilia, Via Campi 103, 41125,*
39 *Modena, Italy*
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47 *properties; electric properties.*
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56 Corresponding Author: Serena Esposito (serena_esposito@polito.it)
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Abstract

Two natural zeolite bearing rocks (one containing clinoptilolite, the other chabazite, phillipsite and analcime) were Fe-exchanged and thermally treated in a reducing atmosphere at 750 °C for 2 h. Two nanocomposites, formed by a dispersion of Fe nanoparticles in a ceramic matrix, were obtained. The prepared lunar dust simulants also contain Na⁺, K⁺, Ca²⁺, and Mg²⁺ and other mineral phases originally present in the starting materials. The samples were fully characterized by different techniques such as atomic absorption spectrometry, X-ray powder diffraction followed by Rietveld analysis, transmission electron microscopy, N₂ adsorption/desorption isotherms at 77 K, measurements of grain size distribution, magnetic properties measurements, broadband dielectric spectroscopy, and DC conductivity measurements. The results of this characterization showed that the obtained metal-ceramic nanocomposites exhibit a chemical and mineralogical composition and electrical and magnetic properties similar to real moon dust and, thus, appear valid moon dust simulants.

1. INTRODUCTION

In the last decade a continuously increasing interest toward manned space missions to the Moon, both from public institutions, such as NASA and ESA, and the private sector, was triggered by the perspective to use it as a test bed either for missions to Mars or other airless planetary bodies and the possibility of field-applications of remote sensing instrumentation.¹⁻³

The surface of airless planetary bodies is generally covered by a thick blanket of regolith, which is the result of space weathering processes, especially comminution, cementation and chemical modification by means of (micro)meteorites bombardment, solar wind, irradiation and sputtering of cosmic and galactic rays.¹ Such a thick blanket of regolith is composed by particles having dimension below 1 cm.^{4,5} Moreover, its finest part (about 20 wt.%) exhibits dusty-like features and, in particular, consists of i) a coarse fraction of breathable lunar dust (2.5-10 μm) which can be deposited inside trachea and bronchiole ducts; ii) a fine fraction (0.1-2.5 μm) which can penetrate the respiratory zone and iii) an ultra-fine fraction (less than 0.1 μm) which can interact with mucous membranes or be trapped by macrophages and give rise to various serious toxic effects.⁶⁻⁸ Thicknesses of typical dusty layers going from few up to 15 meters were recorded in various regions of the Moon.^{8,9}

Furthermore, it should be taken into account that the harsh conditions to which all the lunar regolith fractions are subjected may result into their partial melting.^{3,10,11} In particular, the molten

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3 phase successively quenches to a glass that welds the mineral and lithic particles into aggregate
4 grains, called “agglutinates”.¹² Such harsh conditions can confer unique properties to the lunar
5 agglutinates, namely such as electrostatic charge and high superficial reactivity. Moreover, such
6 agglutinitic glass, with dimension ranging from a few nm to several μm , contains abundant iron
7 particles,^{1,13,14} occurring either as “globules” with mean size of $120 \pm 20 \text{ nm}$ ¹⁵ or metallic nanoscale
8 particles (np-Fe⁰) having dimensions between 3-33 nm with an average size of 7 nm.¹⁶ It must be
9 noted that in agglutinitic glass, iron occurs as Fe⁰ or Fe²⁺ species, and not as Fe³⁺ species, like it
10 occurs, instead, in terrestrial minerals.¹² Such Fe⁰ and Fe²⁺ species may produce reactive oxygen
11 species dangerous to human cells,¹⁷ like hydrogen peroxide (H₂O₂) and hydroxyl radical (OH•).¹⁸
12 Thus, the agglutinitic glass, bearing many metallic iron “globules” and nanoparticles can dissolve
13 into body fluids and reduce hemoglobin Fe³⁺ species according to the Fenton reaction.^{6,7,19} Finally,
14 the review of the main features of the lunar rocks reports that agglutinitic glass is mainly formed by
15 plagioclases and pyroxenes, with smaller amounts of pyroclastic volcanic glass beads, ilmenite and
16 olivine minerals.¹ Its chemical, structural and textural features strongly affect the lunar dust
17 properties, such as the particles adherence, which is also modified by both the reduced Moon gravity
18 (1/6 terrestrial gravity) and by the barbed shape of the dust grains.⁶

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37 The previous considerations accounted for the need of studying in deep the lunar dust properties.
38 However, the poor availability of the actual Moon soil has determined the manufactory of several
39 lunar dust simulants, which were carefully reviewed in our previous work.²⁰ From this review it was
40 found that, despite the efforts of many skillful researchers, all the produced Moon dust simulants did
41 not satisfactory mimic some of the features of the real lunar dust.²¹ On the basis of these
42 considerations it was thought that the patented process,^{22,23} proposed by some of us, aiming at
43 obtaining **metal-ceramic (where ceramic means inorganic, non-metallic)** magnetic nanocomposites
44 from commercial zeolites, could be used to produce reliable Moon dust simulants. The process
45 envisages two steps, *i.e.* a heavy metal (Fe, Ni, or Co) cation exchange of a commercial zeolite and a
46 thermal treatment at relatively mild temperatures (**500-850 °C** range) under reducing atmosphere (2.0
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3 vol.% H₂ in Ar).^{24–29} The nanocomposites were already successfully used for other biochemical and
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5 environmental application and the same^{30–33} efficient, inexpensive and scalable synthesis method was
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7 used to obtain two nanocomposites containing Fe⁰ nanoparticles embedded in a glassy matrix starting
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9 from two commercial zeolites, namely zeolite X and A.²⁰
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11 The so-obtained nanocomposites, referred to as SMA_A and SMA_X (i.e. Simulated Moon
12 Agglutinate from zeolite A and X, respectively), showed proper physico-chemical properties and
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14 magnetic and electric behavior to be considered as suitable materials to simulate the agglutinate
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16 fraction of regolith, lacking in most of lunar simulants.²⁰ However, with respect to real Moon dust,
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18 the two nanocomposites obtained in ref.²⁰ exhibited higher a Fe⁰ content; lower Si/Al ratio and the
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20 presence of only Na⁺ ions (whereas the main cations present in real Moon dust are Na⁺, K⁺, Ca²⁺, and
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22 Mg²⁺).²⁰ Luckily, the versatility and tuneability of the proposed preparation method left room to
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24 further improve the obtained results as^{22,24–29} the Fe⁰ content may be reduced by iterating a lower
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26 number of times the exchange procedures, a higher Si/Al ratio may be obtained by using a natural
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28 zeolite bearing rock as raw material and natural zeolites in their original composition contain Na⁺,
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30 K⁺, Ca²⁺, and Mg²⁺ together with, possibly, traces of other cations.³⁴
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35 On the basis of the above considerations, a clinoptilolite bearing Epiclastite from Northern Sardinia
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37 (Italy) and a chabazite and phillipsite bearing Ignimbrite from Campania (Southern Italy) (labeled as
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39 LacBen and Cab70, respectively, for the sake of uniformity with previous works) were initially
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41 subjected to just one iteration of the Fe²⁺ exchange procedure and subsequently thermally treated at
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43 relatively mild temperatures (500–850 °C range) under reducing atmosphere (2.0 vol.% H₂ in Ar).
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22,24,33,25–32 These natural zeolite bearing rocks, which were already fully characterized,^{35–37} were
selected on the basis of their high natural zeolite content, and huge potentiality, which gives rise to a
very low cost.^{38–40} Moreover, these natural zeolites bearing rocks were already well known as they
were already tested in environment protection studies,^{41–45} for ceramization purposes aiming at tiles
production,^{46–48} for producing lightweight aggregates to be used for concrete manufacture,^{49–51} in

oenological refining processes,⁵² in the immobilization of radionuclides,⁵³ as additive in animal diet,⁵⁴ as drug carriers,⁵⁵ and in solar energy storage applications.⁵⁶

The so-obtained np-Fe⁰ containing nanocomposites were characterized by means of physico-chemical, magnetic and electric techniques, in order assess whether they have the proper features to be considered as simulants of lunar regolith agglutinates.

2. MATERIALS AND METHODS

2.1 Chemicals and Materials Synthesis

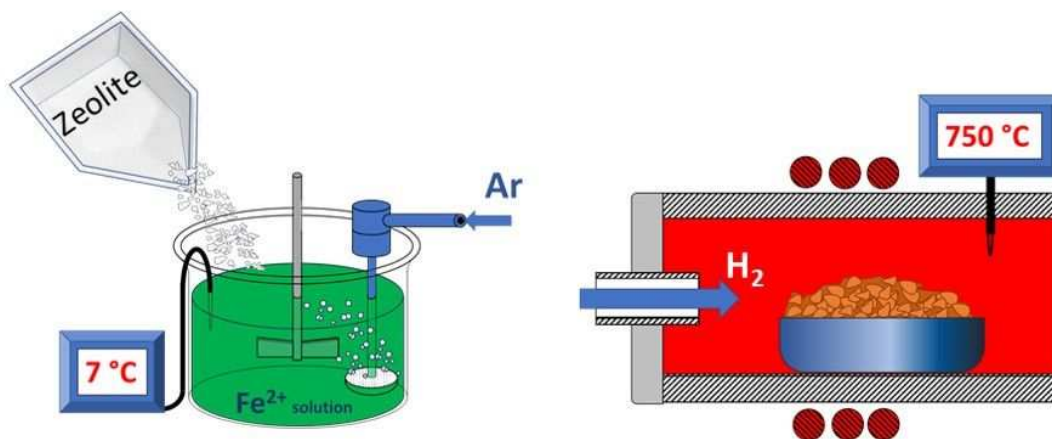
The sample of Campanian Ignimbrite used in this work is a product traded by Italiana Zeoliti s. r. l. (Pigneto – Modena) under the trade-name Cab70, whereas the sample of Epiclestite (LacBen) comes from a deposit sited in Bortivuille (Sassari, Northern Sardinia). The Campanian Ignimbrite is the product of a huge eruption of the Campi Flegrei volcanic area (37,000 years b.p.). This trachytic pyroclastite is mainly constituted by scoriae and pumice set in a cineritic matrix. Two different lithofacies characterize this formation: a zeolite-bearing yellow one and a grey one, with epigenetic feldspars. The sum of chabazite (framework type CHA),⁵⁷ phillipsite (framework type PHI)⁵⁷ and analcime (framework type ANA)⁵⁷ content, generally close to 50 wt. %, in some deposits can even reach about 80 wt. %.^{35–37} Overall the mineralogical analyses performed underlined that Campanian Ignimbrite is strongly heterogeneous.^{39,40}

The Sardinian samples (LacBen) belong to a volcano-sedimentary succession generally resting on the Palaeozoic basement and linked to a calcalkaline eruptive activity, oligo-aquitanean in age. The circulation of hydrothermal fluids through a faults system caused the transformation of the rhyolitic glassy fraction of epiclastic and unwelded pyroclastic deposits in clinoptilolite (framework type CLI)⁵⁷, smectite and opal-CT.^{35–37}

Bulk chemical analysis of the parent zeolites (reported in Table 1) were performed by X-Ray Fluorescence (XRFS, Philips PW1400). Analytical procedures were carried out according to refs,^{58–61}

Table 1: Composition of Cab70 and LacBen as obtained by XRFS

Sample	LacBen	Cab70
	Chemical composition (wt %)	
SiO ₂	65.63	53.71
TiO ₂	0.20	0.04
Al ₂ O ₃	13.50	16.10
Fe ₂ O ₃	1.06	4.01
MnO	0.01	0.02
MgO	1.04	1.01
CaO	2.49	4.01
Na ₂ O	2.46	2.06
K ₂ O	1.11	7.03
P ₂ O ₅	tr.	0.01
H ₂ O	12.50	12.00



Scheme 1. Sketch of the preparation procedure of both SMA_Cab70 and SMA_LacBen by Fe²⁺ cation exchange of Cab70 and LacBen zeolite followed by reductive thermal treatment at 750 °C.

The natural zeolite bearing materials Cab70 and LacBen were subjected to the adopted preparation procedure shown in Scheme 1: the natural zeolite bearing material was contacted with a 0.1 M Fe²⁺ solution (pH = 3.8), prepared by dissolving FeSO₄·7H₂O (Aldrich, 99.5 wt. %) in doubly distilled water (solid/liquid weight ratio = 1/50 g/g; contact time $t = 2$ h). To prevent Fe²⁺ oxidation, during the ionic exchange the temperature was kept at about 7 °C and Ar was continuously bubbled into the solution.⁶² The solid was then recovered by filtration, washed with doubly distilled water, dried for about one day at 80 °C and eventually stored for at least 3 days in a 50% relative humidity environment (created by a saturated Ca(NO₃)₂ aqueous solution), to allow water saturation of the zeolite(s). Afterwards, the Fe²⁺ exchanged natural zeolite bearing materials (hereafter referred to as Fe-Cab70 and Fe-LacBen) were treated in a Pt crucible at 750 °C for 2 h (heating rate 15 degrees

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3 min⁻¹) by flowing a 2 vol.% H₂ in Ar mixture inside an Al₂O₃ tubular furnace (inner diameter = 6.9
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5 cm, height = 91 cm). Afterwards, the furnace was switched off and the samples were left to cool down
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7 to room temperature (r.t.).
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9 The two nanocomposites obtained from the Cab70 and LacBen natural zeolite bearing materials will
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11 be hereafter referred to as SMA_Cab70 and SMA_LacBen, respectively (SMA stands for Simulated
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13 Moon Agglutinate).
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18 2.2 Characterization Methods

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21 The content of Fe²⁺ and (residual) Na⁺, K⁺, Ca²⁺, and Mg²⁺ in both Fe-Cab70 and Fe-LacBen
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23 samples was determined by atomic absorption spectrophotometry (AAS, Perkin-Elmer Analyst 100
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25 apparatus) after dissolving the solids in a 40 wt.% HF and 14 wt.% HClO₄ aqueous solution.^{63,64}
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28 Quantitative phase analyses (QPA) were carried out on the parent materials (LacBen, Cab70),
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30 on the Fe-exchanged materials (Fe-LacBen, Fe-Cab70) and on the nanocomposites (SMA_LacBen,
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32 SMA_Cab70) by exploiting synchrotron radiation: the synchrotron X-ray powder diffraction (XRPD)
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34 patterns were collected on the XRD1 beamline at Elettra Sincrotrone Trieste (Italy) with a fixed
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36 wavelength of 0.70 Å. The powders were packed along with Al₂O₃ (10 wt.%) as internal standard in
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38 a boron capillary and spun under the beam. The diffraction patterns were collected using a Dectris
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40 Pilatus 2M detector. The combined RIR-Rietveld method, which enables the QPA and the calculation
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42 of both the crystalline and amorphous fractions,⁶⁵ was performed by using the GSAS package⁶⁶ with
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44 the EXPGUI interface.⁶⁷ The results of QPA on the six analysed samples is reported in table 2.
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48 N₂ adsorption/desorption isotherms at 77 K were measured on samples previously outgassed
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50 at 200 °C for 3h to remove water and other atmospheric contaminants (Micrometrics ASAP
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52 2020Plus). The samples specific surface area was calculated according to the Brunauer–Emmett–
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54 Teller (BET) method (S_{BET}); total pore volume (V_p) and micropore volume (V_{mp}) were determined
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56 from the amount of adsorbed N₂ at P/P₀ = 0.9 and according to the *t*-plot method, respectively.
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3 Transmission electron microscopy (TEM) characterization was performed by using a side
4 entry JEOL 3010-UHR microscope operating at 300 kV, equipped with a LaB6 filament, a (2k×2k)-
5 pixel Gatan US1000 CCD camera and an OXFORD INCA EDS instrument for atomic recognition
6 via energy dispersive spectroscopy (EDS). To prepare the samples, the powders were either briefly
7 contacted with lacey carbon Cu grids, which resulted in the mere electrostatic adhesion of some
8 particles to the sample holder. Particle size distribution and mean particle diameter (d_m) of np-Fe⁰
9 were obtained by considering a statistically representative number of particles (200-400
10 nanoparticles) on several TEM images (at least 30 different images). In particular, the d_m was
11 calculated using the following equation:
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$$d_m = \Sigma d_i n_i / \Sigma n_i \quad (1)$$

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25 where n_i is the number of particles of diameter d_i . It is worth noting that both exchanged Fe-Cab70
26 and Fe-LacBen samples were stable to prolonged exposition under the electron beam of the
27 instrument (no metal coalescence, nor modification of the zeolitic framework were observed during
28 the measurements.
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34 Differential centrifugal sedimentation method was applied to calculate the particle size
35 distribution on a disc centrifuge (CPS DC24000). Before the measurements, a defined aliquot of
36 samples was dispersed in water and sonicated for 5 min. A particle density of 2.0 mg ml⁻¹ and a
37 refractive index of 1.48 have been used for all the measurement with a non-sphericity factor of 1
38 (assuming a spherical form). The particle absorption values have been varied because of the different
39 color of the samples, namely 0.1 for LacBen, Fe-LacBen and CAB70, 0.4 for Fe-CAB70 and 1.0 for
40 SMA_Cab70 and SMA_LacBen.
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51 **2.3 Magnetic and electrical properties measurement**

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53 The magnetic hysteresis loops of the SMA_Cab70 and SMA_LacBen nanocomposites were
54 studied between 100 K and room temperature on a SQUID magnetometer operating in the 0-70 kOe
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range. In order to extract further information on the Fe⁰ nanoparticles, the FC/ZFC curves of both samples were measured in the temperature interval 10 - 300 K under a field of 50 Oe.

The electrical properties of the nanocomposites were studied using Broadband Dielectric Spectroscopy (BDS) and DC conductivity measurements techniques.^{68,69} In both types of measurement, samples made of dust layers were approximately 200 μm thick and confined between two metallic electrodes. Isotherm dielectric spectroscopy measurements were carried out under dry N₂ flow at atmospheric pressure at the CIRIMAT laboratory (Toulouse, France) in the 133 - 423 K temperature range and in the 10⁻² -10⁶ Hz frequency range. By steps of 283 K, sinusoidal voltages U* (amplitude 1V, 10 points per frequency decade) were isothermally applied to the sample. The measurements of both the induced current (I*) and its phase shift relative to the applied voltage yielded the complex impedance (Z*) values as a function of temperature and frequency. The complex dielectric permittivity ε* (eq. 2) and electrical conductivity σ* (eq. 3) formalisms were used to represent the BDS data.

$$\varepsilon^*(\omega) = \varepsilon'(\omega) - i\varepsilon''(\omega) = \frac{1}{i\omega C_0 Z^*(\omega)} \quad (2.1)$$

$$C_0 = \frac{\varepsilon_0 A}{l} \quad (2.2)$$

$$\sigma^*(\omega) = \sigma'(\omega) + i\sigma''(\omega) = i\omega\varepsilon_0\varepsilon^*(\omega) \quad (3)$$

where ε' and ε'' are the real and imaginary parts of ε*, ω the angular frequency, and C₀ the capacitance of the vacuum-filled capacitor formed by the two electrodes of area A separated by the sample thickness l (ε₀ is the vacuum permittivity), where σ' and σ'' are the real and imaginary parts of σ*.

DC conductivity measurements were carried out under secondary vacuum (< 10⁻⁶ mbar) at the ONERA laboratory (Toulouse, France) with a constant voltage between the electrodes and with temperature from 298 to 423 K. Dust layer density was around 0.6 g cm⁻³.

3. Results and Discussion

3.1 Physico-chemical characterization

The QPA results obtained with the parent materials, reported in Table 2, confirm the literature data of samples coming from the same sites. The Cab70 sample contains three zeolitic phases, phillipsite being the most abundant one (30.4 wt. %) followed by chabazite (6.3 wt. %) and analcime (5.3 wt. %). Along with the zeolites, the sample also contains K-Feldspar (22.9 wt.%), phyllosilicates (6.3 wt.% Muscovite/illite) and a relatively high amount of amorphous phase (28.9 wt.%). The LacBen sample is a clinoptilolite zeolite (54.2 wt. %) containing a relatively high amount of K-Feldspar (22.3 wt.%), quartz (9.7wt.%), low levels of phyllosilicates (<1wt% Muscovite/illite) and a moderate amount of amorphous phase (12.8 wt.%).

As reported in Table 2, the relative phase amounts are not affected by the Fe-exchange and only minor variations in the phase wt.% are observed, probably due to minor inhomogeneity of the parent powders (e.g. the increase of the phyllosilicates and zeolitic phase in sample LacBen). Moreover, it must be considered that the contact with the acidic Fe²⁺ exchange solution might have dissolved minor amount of some constituents of the parent materials. The appearance of a new weak peak (at 5.25° 2θ) is noteworthy, as it corresponds to an iron oxide hydrate phase (Fe₂O₃·H₂O), probably precipitated from the exchange solution. The peak indicates that the phase is present in very low percentage and with the Fe-LacBen sample is even not possible to quantify it from Rietveld method.

Table 2: Results of the QPA as obtained by applying the Rietveld method to synchrotron XRPD.

Phase	LacBen (wt %)	Fe-LacBen (wt %)	SMA_LacBen (wt %)
Clinoptilolite	54.2	60.2	13.5
K-feldspar	22.3	21.5	17.9
Quartz	9.7	7.2	7.9
Cristobalite	-	-	2.9
Muscovite/Illite	0.9	2.5	1.9

Fe₂O₃·H₂O	-	present	-
Fe⁰	-	-	0.6
Amorphous	12.8	8.6	55.2
Total	99.9	99.9	99.9

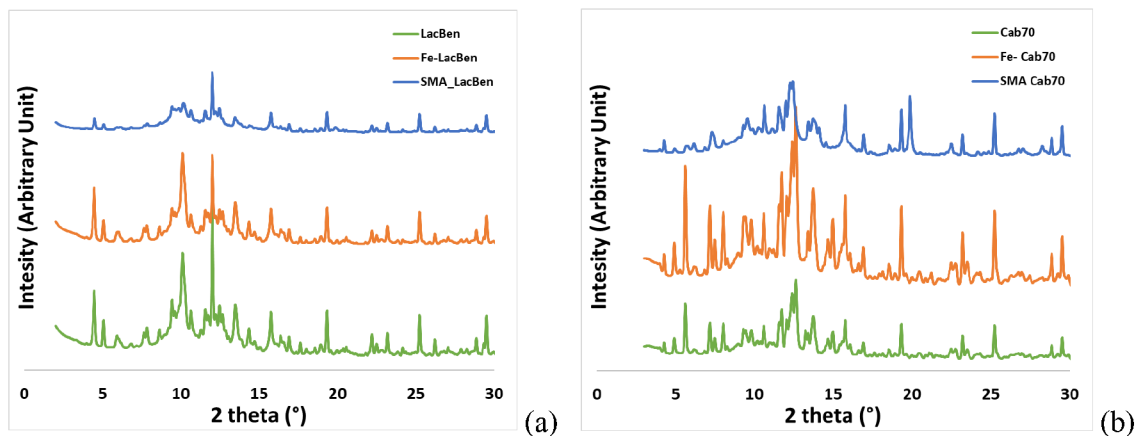
Phase	Cab70 (wt %)	Fe-Cab70 (wt %)	SMA_Cab70 (wt %)
Chabazite	6.3	5.7	5.0
K-Feldspar	22.9	25.0	27.9
Phillipsite	30.4	27.9	5.0
Analcime	5.3	6.4	4.1
Muscovite/Illite	6.3	3.9	4.9
Fe₂O₃·H₂O	-	0.4	-
Fe⁰	-	-	2.7
Amorphous	28.9	30.7	50.5
Total	100.0	100.0	100.0

As expected, the thermal treatment induces strong modification on both the nanocomposites (Figure 1), the strongest effect being related to the SMA_Cab70 sample, where the content of phillipsite drops from 30.4 to 5.0 wt. %, whereas the analcime and chabazite contents seem to be less affected by the thermal treatment, being stable up to about 800 °C.⁷⁰ The clinoptilolite content in sample LacBen strongly decreases after the thermal treatment: such a behavior, somehow slightly inconsistent with the thermal behavior of clinoptilolite (stable up to 900-1000 °C),⁷⁰ has to be ascribed to the Fe-exchange procedure that leads to a more unstable zeolite. It has been indeed demonstrated that the type of extra-framework cation occurring in the clinoptilolite porosities strongly affects its thermal behavior.³⁷ **It must be noticed that the amount of phases other than zeolites, present in the parent materials, do not show large variations owing to the thermal treatment as they exhibit a thermal stability higher than the considered zeolites.³⁵⁻³⁷**

Moreover, it must be borne in mind that the overall mineralogical composition of the two Moon dust simulants appears similar to the one of real Moon dust as reported by Loftus et al.⁴

Both the nanocomposites showed the presence of some Fe⁰, the highest amount (2.7 wt. %) being related to the SMA_Cab70 sample. The different amount of Fe found in the two samples could

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3 be, however, due to the dimension of the particles, being X-ray powder diffraction not very effective
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5 in quantifying nano-phases (if the diffraction domains are too small).
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24 **Figure 1.** XRPD patterns collected of the investigated samples: (a) LacBen, Fe-LacBen and
25 SMA_LacBen; (b) Cab70, Fe-Cab70 and SMA_Cab70

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27 According to AAS analysis, the Fe-Cab70 and Fe-LacBen samples contain 1.79 and 0.89 meq
28 g^{-1} Fe^{2+} , respectively, due to partial exchange of Na^+ , K^+ , Ca^{2+} , and Mg^{2+} present in the framework
29 of the zeolitic phases. Apparently, the content of alkali- and alkali-metal cations after Fe^{2+} exchange
30 turns out lower than the one present in the parent Cab70 and LacBen, but remains sizable. However,
31 the amount of Na^+ , K^+ , Ca^{2+} , and Mg^{2+} present in the framework Fe-exchanged Cab70 and LacBen
32 remains considerable as: 1) the amount of exchanged Fe^{2+} is not sufficient to exchange for all the
33 cation exchange capacity of zeolites;^{34–37} 2) in Cab70 and LacBen samples there are also other phases
34 that do not exhibit cation exchange properties and, thus, keep their original cation composition and
35 3) a portion of the iron introduced in the solution for the cation exchange crystallizes as iron oxide
36 hydrate ($\text{Fe}_2\text{O}_3 \cdot \text{H}_2\text{O}$) and thus, does not contribute to the Na^+ , K^+ , Ca^{2+} , and Mg^{2+} cation exchange
37 (see above).
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51 From these data, the overall iron wt. % in SMA_Cab70 and SMA_LacBen, was calculated
52 considering the nanocomposites as completely dehydrated materials, resulting 5.7 and 2.9 wt. %,
53 respectively. The different iron content in the two Fe^{2+} -exchanged zeolites is probably determined
54 by their different cation exchange capacity, and kinetics of exchange. It must be borne in mind that
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3 these percentages of iron encompass also the Fe present in Cab70 and LacBen samples in their
4 original cation composition (see Table 1), i.e. before the Fe-exchange. However, such Fe is likely
5 present in the various crystalline and amorphous phases of Cab70 and LacBen samples as trivalent
6 iron. Clearly, such trivalent iron does not appear prone to undergo reduction as it is not a cation bound
7 to the negative framework of a zeolite by an ionic bond weakened by the dielectric action of water
8 molecules shielding it.
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16 The normalized grain size distribution curves and the relative cumulative curves, in the 0.1-
17 30 μm range, obtained by CPS are reported in Figure 2. Only one main peak is observed in all the
18 samples but the effect of the iron exchange and subsequent heat treatment is different in the two
19 series. Indeed, in the case of Cab70 series (Fig. 2b), the peak position is very close in all samples.
20 The peak value of Cab70 is about $5\mu\text{m}$ with 50 wt % grain smaller than $3.3\mu\text{m}$ (dotted green curve);
21 small changes are made by the exchange and Fe-Cab70 shows 50 wt. % grain smaller than $3.8\mu\text{m}$,
22 (dotted orange curve). The grain size distribution curve shifts to slightly higher values for the sample
23 heat treated at $750\text{ }^\circ\text{C}$, SMA_Cab70, (50 wt. % grain smaller than $4.5\mu\text{m}$, dotted blue curve). Instead,
24 for the LacBen series a shift in the peak position is clearly visible (Fig. 2a). The pristine material
25 LacBen shows a peak at $7.3\mu\text{m}$ with a shoulder at $1\mu\text{m}$ and 50 wt. % grain size smaller than $5.5\mu\text{m}$,
26 dotted green curve. The finest fraction of the powder is lost with the exchange operations and the
27 maximum of the grain size distribution curve of the sample Fe-LacBen moves to $10.6\mu\text{m}$ with 50 wt%
28 grain smaller than $7.7\mu\text{m}$, dotted orange curve. The Moon dust simulant prepared starting from
29 LacBen, SMA_LacBen, shows the main peak at $6.3\mu\text{m}$ with 50 wt% grain smaller than $5\mu\text{m}$, dotted
30 blue curve, suggesting that the heat treatment causes a finer fraction to be formed.
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48 The grain size distribution curves of Fig. 2 allow the following considerations:
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50 1) The parent materials Cab 70 and Lac Ben used in this work to manufacture the moon
51 dust simulants exhibits a grain size distribution slightly finer than the one of the zeolite A used in our
52 previous work.^{20,60}
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2) The grain size distribution of SMA_LacBen and SMA_Cab70 appears slightly finer than the one SMA_A and similar to the one of SMA_X of our previous work.²⁰

3) SMA_LacBen and SMA_Cab70, like SMA_A and SMA_X, exhibit a grain size distribution similar to that of lunar soil coarse fraction (2.5-10 μm) but still larger than the one of fine (0.1-2.5 μm) and ultrafine (< 0.1 μm) fractions. However, for studies requiring smaller grain size, proper ball milling procedures applied to the present nanocomposites or to their parent materials could allow decreasing the grain size down to 300 nm, thus overcoming this limit.⁷¹

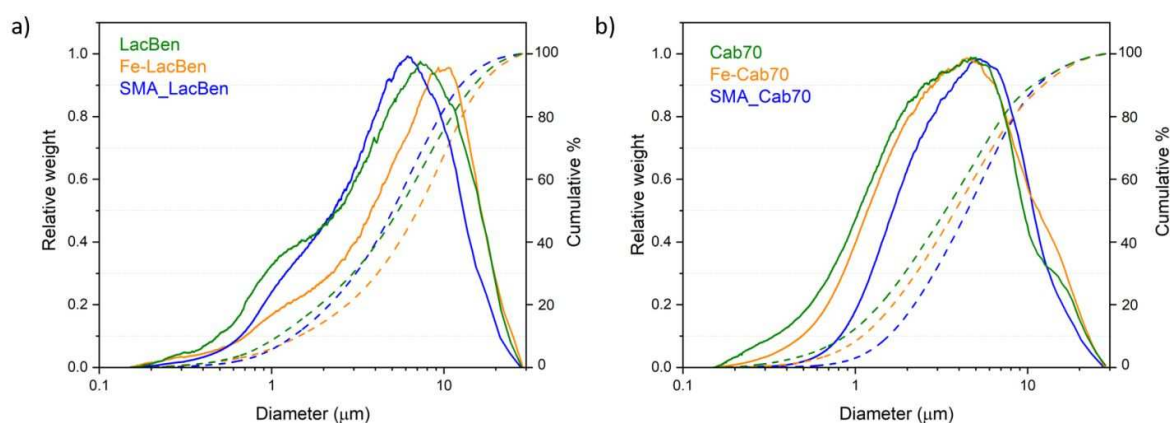


Figure 2. Normalized grain size distribution plots (solid curves) and cumulative curves (dotted curves) for Cab70 (a) and LacBen samples (b).

The N_2 adsorption/desorption isotherms of SMA_Cab70 and SMA_LacBen are reported in Fig. 3, whereas those of the parent Cab70 and LacBen samples and of the exchanged samples, Fe-Cab70 and Fe-LacBen are reported in Figure SI-1. The shape of the isotherms corresponds to type II according to the (IUPAC) classification characterized by no saturation at P/P^0 values close to 1. The classification of the isotherms is not straightforward because although the shape resembles a type II isotherm, the presence of a hysteresis would indicate a type IV isotherm. This type II isotherm commonly observed in non-porous adsorbents or adsorbents possessing relatively large pores, was in accordance with the literature data for the current zeolite phases.^{72,73} The hysteresis loops are

classified as Type H3 suggesting that the porosity is caused by the aggregates of the plate-like particles, which bring about slit-shaped pores.

According to Sing's work, the isotherms can be classified as pseudo-type II, describing delayed capillary condensation due to the small rigidity of the aggregate structure of the adsorbent.⁷⁴

The textural properties derived by the isotherms, i.e. the values of surface area (S_{BET}), total pore volume (V_{p}) and micropore volume (V_{mp}) are reported in Table 3. The trend of the surface area values for both sets of samples may initially appear unexpected. Indeed, the surface area increases after iron exchange, and then decreases after the thermal treatment. The second phenomenon is the result of the expected material densification in the nanocomposites, whereas the increase of area after Fe exchange is less straightforward. A possible explanation can be found in the partial dissolution of some mineral phases contained in the starting samples, Table 2, caused by the acid pH of the exchange solutions (3.8), with a consequent increase in the percentage of porous fraction.

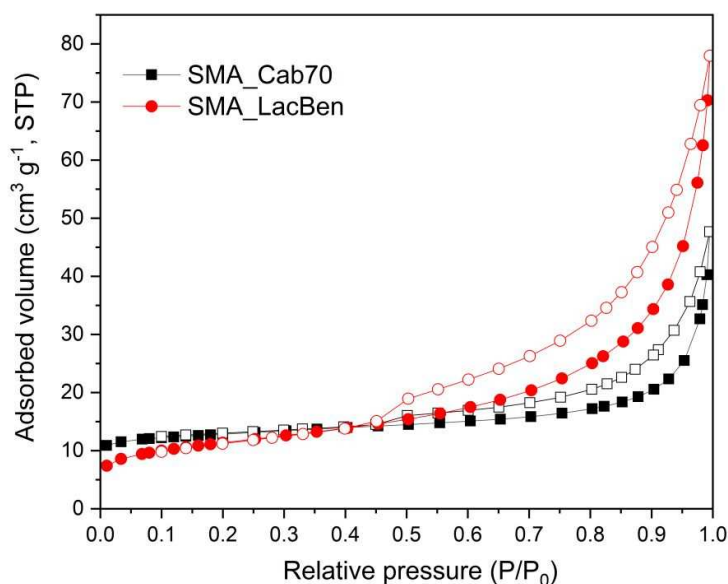


Figure 3. N_2 adsorption-desorption isotherms at 77 K of SMA_Cab70 and SMA_LacBen materials.

Table 3: BET Specific surface area (S_{BET}); total pore volume (V_{p}); micropore volume (V_{mp}), as obtained by N_2 isotherms at 77 K.

Sample	S_{BET} ($\text{m}^2 \text{g}^{-1}$)	V_{p} ($\text{cm}^3 \text{g}^{-1}$)	V_{mp} ($\text{cm}^3 \text{g}^{-1}$)
Cab70	38.45	0.064	0.013
Fe_Cab70	45.31	0.077	0.013

SMA_Cab70	20.49	0.066	5×10^{-3}
Lac-Ben	39.94	0.120	3×10^{-3}
Fe_Lac-Ben	85.52	0.230	8×10^{-3}
SMA_Lac-Ben	27.44	0.102	-

A detailed TEM and HR-TEM characterization has been carried out on the Fe-LacBen and Fe-Cab70 samples and on the nanocomposites (SMA_LacBen and SMA_Cab70, with the aim of following the morphological and structural evolution of the materials during the formation of the Fe-containing nanocomposites. In addition, particular attention was devoted to the determination of the dispersion of Fe. The results of the characterization performed on Fe-LacBen and Fe-Cab70 are shown in Figure 4 and Figure 5, respectively. The former sample appears composed by large zeolitic particles with globular shape alternate to elongated ones (Figure 4, panels a-c). In addition, highly dispersed nanoparticles, possibly iron oxide hydrate ($\text{Fe}_2\text{O}_3 \cdot \text{H}_2\text{O}$), appearing with dark contrast with respect to the zeolitic framework and with mean diameter $d_m = 2.7 \pm 1.0$ nm (Figure 4, inset of panel e) were easily recognizable as can be appreciated in panels b and d of Figure 4 and in Figure SI-2, in which an image at high magnification is reported. According to the particle size distribution, the majority of these nanoparticles has size between 2 and 3 nm. The EDS analysis confirmed the presence of iron-containing nanoparticles (Figure 4, panel e) as well as its quite uniform relative spatial distribution sample, as measured in several regions of the material (an example is shown in Figure SI-3). Moreover, the nanoparticles tend also to form agglomerates with spherical shape and with size in the 10–20 nm range, as those shown in panel c of Figure 4, therefore lowering the metal exposed surface area. Finally, EDS mapping also revealed the presence of residual cations, which did not contribute to the exchange with iron, in agreement with the results of the AAS analysis.

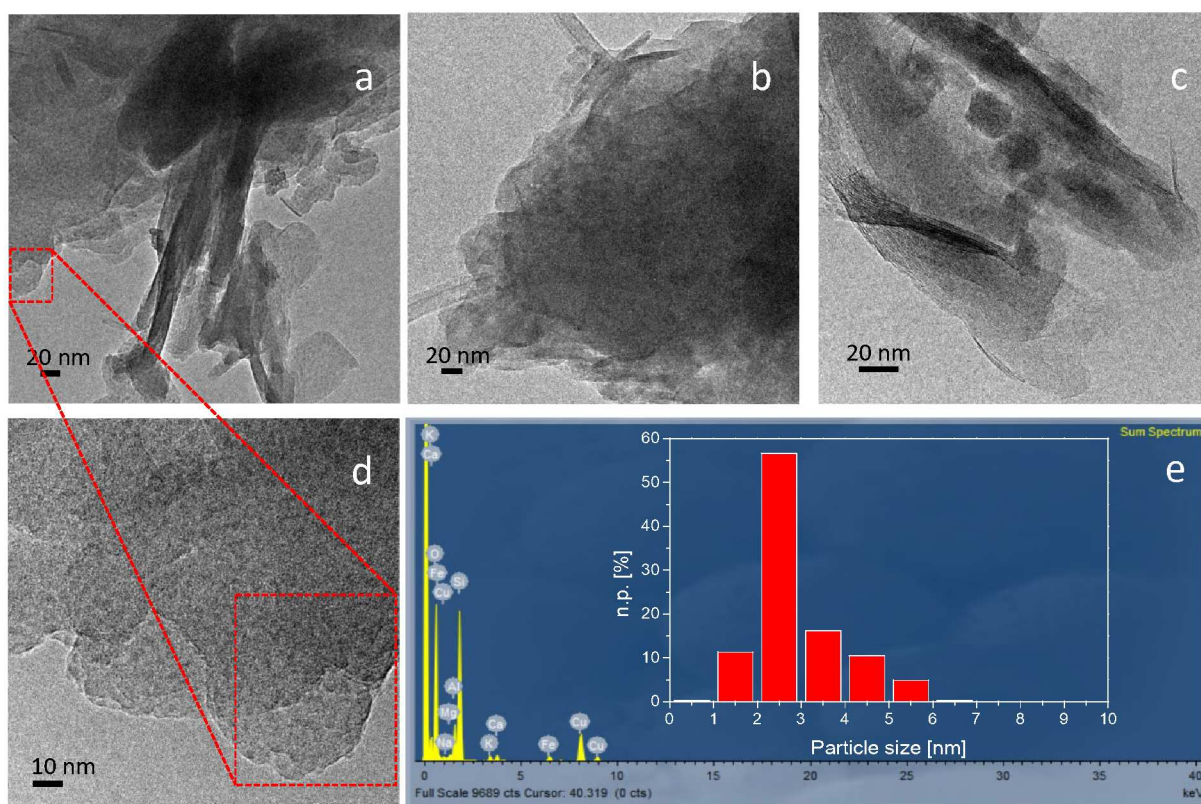


Figure 4. Representative TEM images of Fe-LacBen sample (a-c). Zoom on the region shown in a highlighted by the red box (d). EDS spectrum collected on the region shown in panel d (e) and Fe particle size distribution (inset). Instrumental magnification: 50000 \times (a and b), 100000 \times (c) and 150000 \times (d).

Overall, the morphology of the Fe-Cab70 material is similar to that observed for the exchanged Fe-LacBen sample, but in this case the Fe-containing nanoparticles have average diameter $d_m = 3.1 \pm 0.7$ nm (inset in panel c of Figure 5) and appeared more abundant than on the Fe-LacBen material, according to the results of the EDS analysis that indicated that the relative amount of iron present on this sample is higher than that found on the LacBen material.

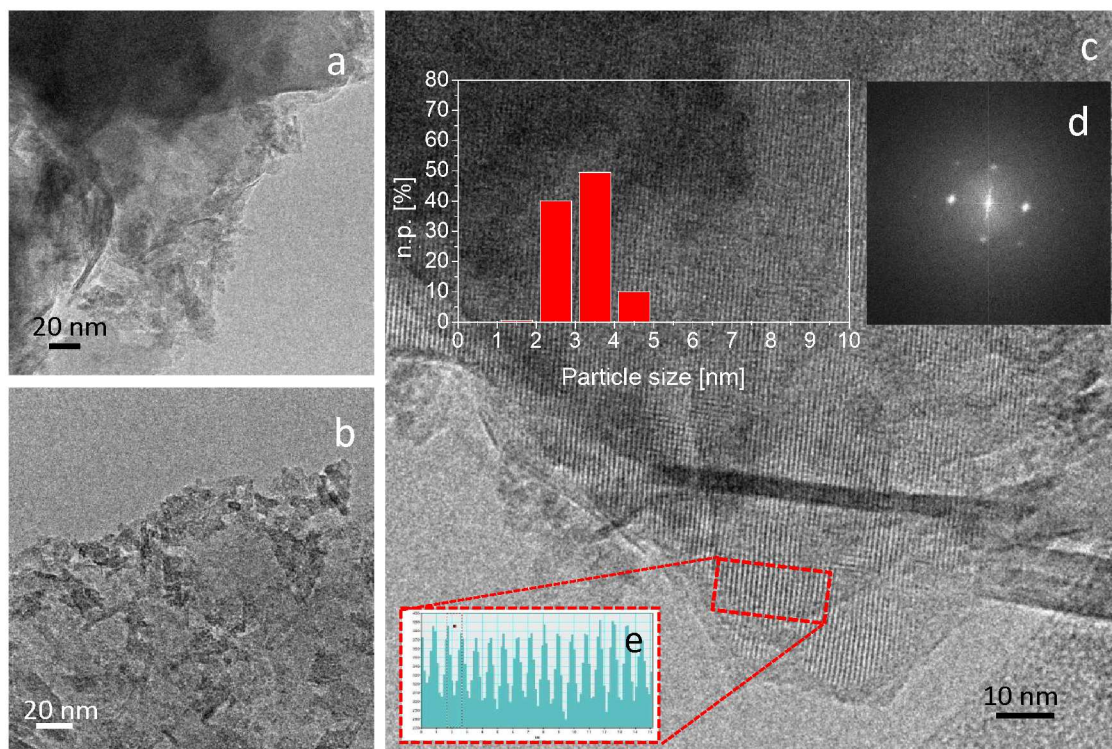


Figure 5. Representative TEM (a and b) and HR-TEM (c) images of the Fe-Cab70 sample. Inset in panel c: Fe particle size distribution. Fourier Transform of the HR-TEM image reported in panel c (d) and evaluation of the distance among the diffraction fringes (e). Instrumental magnification: 80000 \times (a), 100000 \times (b) and 150000 \times (c).

In addition, analogously to what previously observed, the Fe-containing nanoparticles were organized as small agglomerates (Figure 5, panels a-c and Figure SI-4). Nevertheless, such nanoparticles appear homogeneously distributed within the zeolitic matrix as demonstrated by EDS mapping reported in Figure SI-6. Many regions of the sample appear crystalline, as shown in Figure 5 (panel c) in which diffraction fringes with spacing of 9.3 Å that can be attributed to the (101) plane of rhombohedral chabazite (JCPDS file number 00-002-0062) have been measured, according to the corresponding FT and to the measure of the spacing shown in panel d and e, respectively.

Upon thermal treatment under reducing atmosphere, the Fe-containing nanoparticles previously observed on the exchanged Lachen sample undergo coalescence giving rise to a nanocomposite made up by a reduced metallic phase that appears embedded in the zeolitic framework, as shown in Figures 6 and SI-6. Particularly, an evolution to small rounded Fe nanoparticles signalled by red arrows in the Figure and with average diameter of 4.4 ± 1.7 nm (panel a), and to large Fe

particles with rounded globular shape and size ranging between about 15 and 75 nm has been observed (panels b and c).

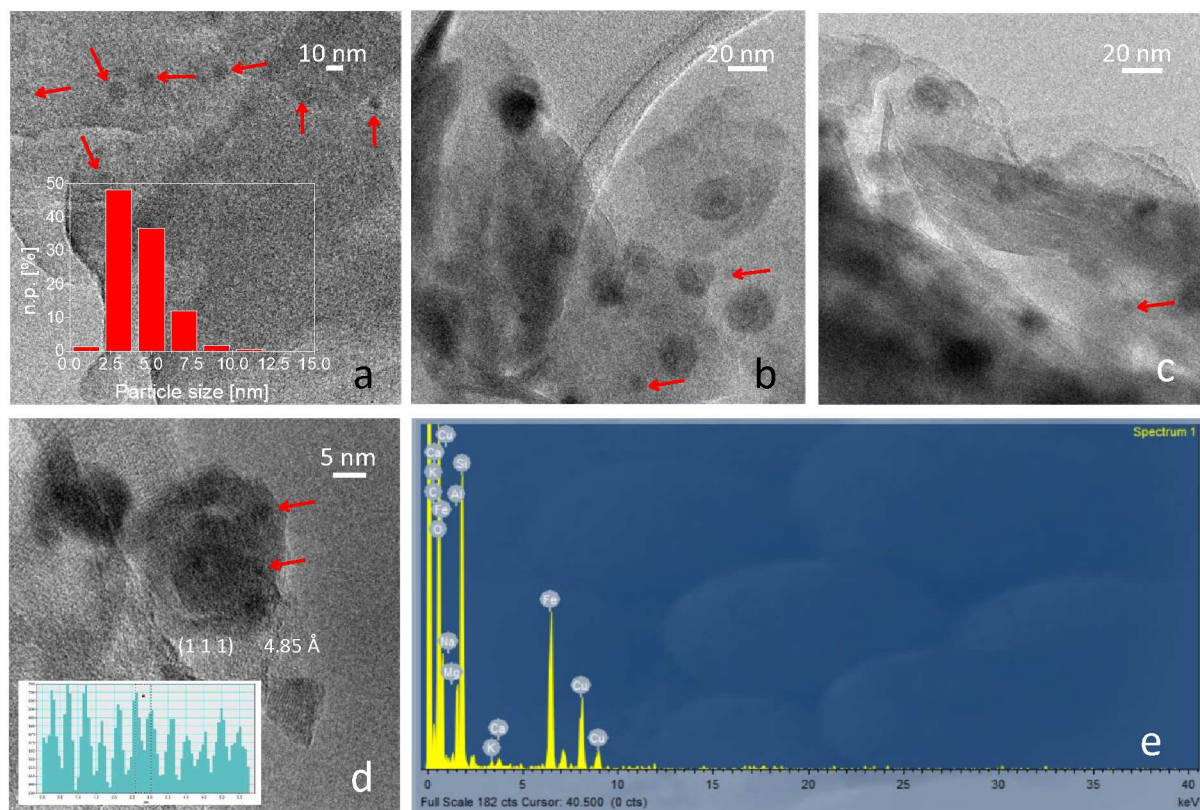
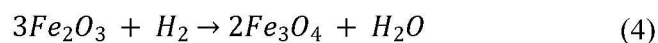


Figure 6. Representative TEM (a-c), HR-TEM (d) images, Fe particle size distribution (inset in panel a) and evaluation of the distance among the diffraction fringes (inset in d) of the SMA_Lacben sample. EDS spectrum collected on the big Fe particle shown in panel d (e). The small Fe nanoparticles are indicated by red arrows. Instrumental magnification: 150000 \times (a, b and c), and 300000 \times (d).

Moreover, these larger Fe nanoparticles display a core-shell-like morphology (panels b and c of Figure 6 and Figure SI-7), in which the core displays a darker contrast with respect to the shell and the EDS analysis revealed an intense peak related to Fe (Figure SI-8). Diffraction fringes with spacing of 4.85 Å, related to the (1 1 1) face of Fe₃O₄ in the cubic phase (JCPDS file number 00-001-1111) were observed on the shell at higher magnification. Magnetite could have been produced from the partial reduction of the iron-containing clusters with hydrogen at high temperature according to the following reaction:

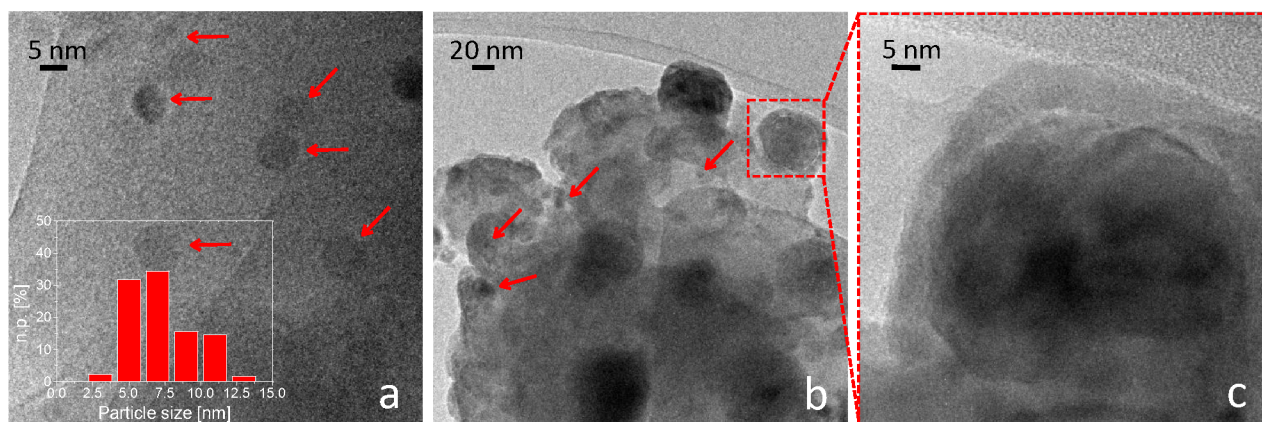


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5 Basing on these findings, a core-shell-like morphology in which a metallic Fe^0 core is
6 surrounded by a Fe_3O_4 shell can be hypothesized after thermal treatment.
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9 It is worth noting that HR-TEM revealed presence of Fe_3O_4 in contrast QFA results. This can
10 be explained by assuming that Fe_3O_4 is mainly not crystalline and/or that the amount of crystalline
11 Fe_3O_4 is below the XRD detection limit.
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15 Also, for the Fe-Cab70 sample, the reducing thermal led to the formation of both small and
16 larger Fe particles within the zeolitic matrix (Figures 7, SI-8, SI-9 and SI-10). Indeed, the
17 SMA_Cab70 material display a less dispersed metallic phase, with average size of 7.3 ± 2.0 nm, as
18 revealed by the particle size distribution reported in panel a of Figure 7. In addition, rounded big Fe
19 particles with core-shell morphology and size around 40-80 nm were observed also in this case.
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21 Overall, the dispersion of the metallic phase observed on the SMA_Cab70 sample is lower than that
22 of the SMA_LacBen sample.
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31 It appears noteworthy that the contemporary presence of Fe^0 and Fe oxides is reported also in
32 all real Moon dust samples.⁴ Thus, it seems another point in which the obtained Moon dust simulants
33 faithfully mimic the features of real Moon regolith.
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54 **Figure 7.** TEM representative images of the SMA_Cab70 sample (a, b). Zoom on the region shown
55 in b highlighted by the red box (c). Fe particle size distribution (inset of panel a). Instrumental
56 magnification: $250000\times$ (a), $50000\times$ (b) and $300000\times$ (c).
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3.2 Magnetic properties

The hysteresis loops of the two Fe-exchanged samples and of the two nanocomposites measured in the interval 100-300 K are shown in Figure 8 (panels *a,b*). The lower value corresponds to the minimum temperature typically measured at midnight on the Moon surface. In both materials, a significant magnetic signal is observed, consisting of a ferromagnetic-like contribution from iron particles followed by a paramagnetic behavior at high fields, to be ascribed to the presence of a fraction of exchanged Fe^{2+} ions still dispersed in the amorphous phase, as previously observed in similar materials^{26,29} and/or to the constituents of the natural zeolite bearing materials which naturally contain Fe ions, such as the illite.⁷⁵

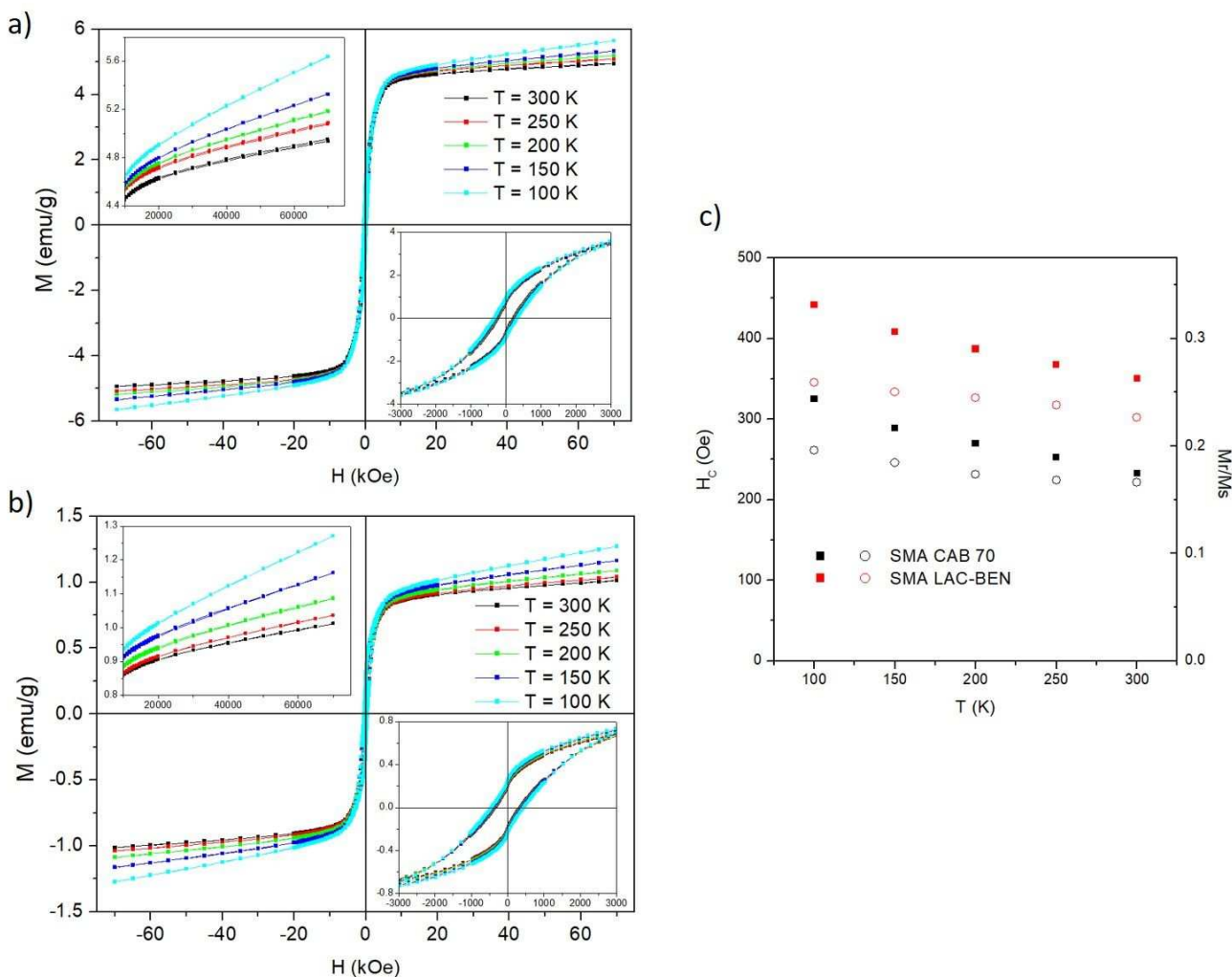


Figure 8. (a,b): hysteresis loops between $T = 100$ K and $T = 300$ K for samples SMA_Cab70 and SMA_LacBen, respectively. The insets put in evidence the paramagnetic contribution at high fields

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3 and the shape of the loops around $H = 0$. (c): temperature behavior of the coercive field measured in
4 the two samples (full symbols) and of the remanence-to saturation ratio (open symbols).
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7 The intensity of the ferromagnetic-like signal is much larger in sample SMA_Cab70. The
8 insets in the panels show the detail of the paramagnetic contribution (displaying the usual increase in
9 the slope of the $M(H)$ line with decreasing temperature) and the shape of the hysteresis loops close to
10 $H = 0$, respectively. Neither the loop shape nor the loop area shows important changes in the
11 considered temperature interval. In particular, the coercive field H_c is a weak function of temperature,
12 as shown in panel *c*; a high value of H_c is still measured at room temperature. This suggests that a
13 fraction of magnetic nanoparticles is still blocked at $T = 300$ K. The ratio M_r/M_s of remanent
14 magnetization M_r to the saturation value of the ferromagnetic component is also reported in panel (c).
15 Even this property turns out to be weakly dependent of temperature. Remarkably, the magnetic
16 properties of both the nanocomposites are in good general agreement with the ones of actual lunar
17 soils, as published elsewhere.²⁰
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31 Using the information on the content of the Fe^0 phase after thermal treatment of samples given
32 by the QPA (rightmost column in Table I), and exploiting the asymptotic value of the room-
33 temperature $M(H)$ curves at high field (after proper subtraction of the paramagnetic contribution), we
34 obtain an intrinsic saturation magnetization M_s of the particle phase of about 160 emu/g and about
35 175 emu/g for the samples SMA_LacBen and SMA_Cab70, respectively. These values of M_s are
36 somewhat smaller than the ones found in the literature for Fe^0 nanoparticles, which are peaked around
37 $M_s = 200$ emu/g,^{76–78} and are consistent with the existence of a fraction of oxidized Fe (Fe_3O_4) in the
38 outer shell of the bigger core-shell particles, as observed by TEM in both samples.
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48 The FC/ZFC curves obtained on both materials are shown in Figure 9. While both FC curves
49 are almost featureless, as often observed in nanoparticle systems where the individual magnetic units
50 (i.e., single or magnetically correlated particles) are characterized by a wide distribution of blocking
51 temperatures,⁷⁹ the ZFC curves exhibit peculiar features: in SMA_Cab70 the ZFC magnetization
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increases monotonically, whereas in sample SMA_LacBen a very broad maximum centered at about 97 K is observed (see upper inset in the Figure).

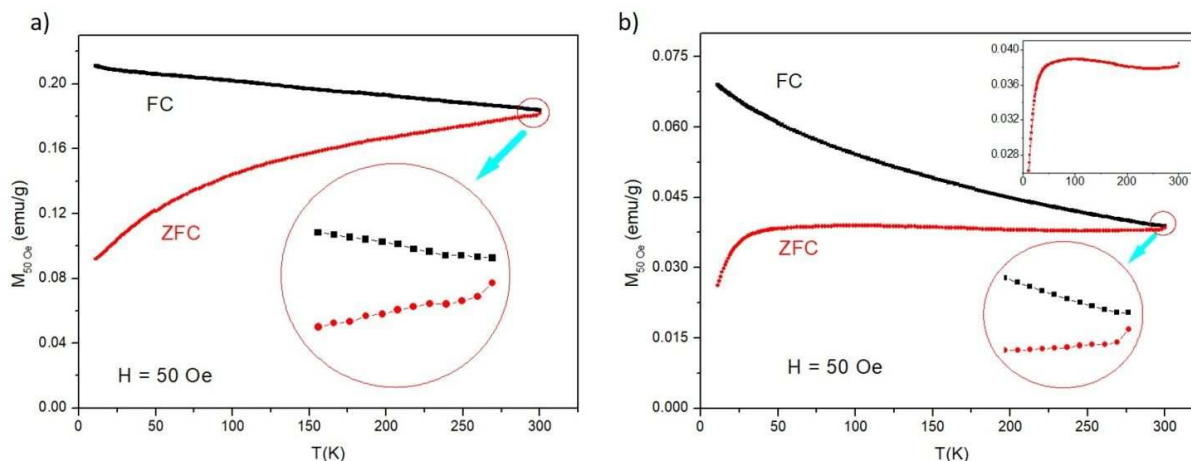


Figure 9. (a): Field-Cooled and Zero-Field-cooled magnetization curves under a field of 50 Oe for sample SMA_Cab70. The inset shows that the two curves do not touch at the starting temperature ($T = 300$ K); (b): the same for sample SMA_LacBen; the upper inset put in evidence the broad maximum of the ZFC curve.

As a common feature, the FC/ZFC curves are not coincident at the starting temperature of the measurement procedure ($T = 300$ K), as shown in the two lower insets. All these features^{29,80} concur to show that a considerable fraction of individual magnetic units are still blocked at r.t., in agreement with the remarkable value of the coercive field (see above).

The difference between the shape of the ZFC curves can be ascribed to a different distribution in size of the magnetic units which individually respond to the magnetic field. Information on the particle size distribution is obtained by exploiting a standard procedure involving the derivative of the difference between ZFC and FC curves.^{81,82}

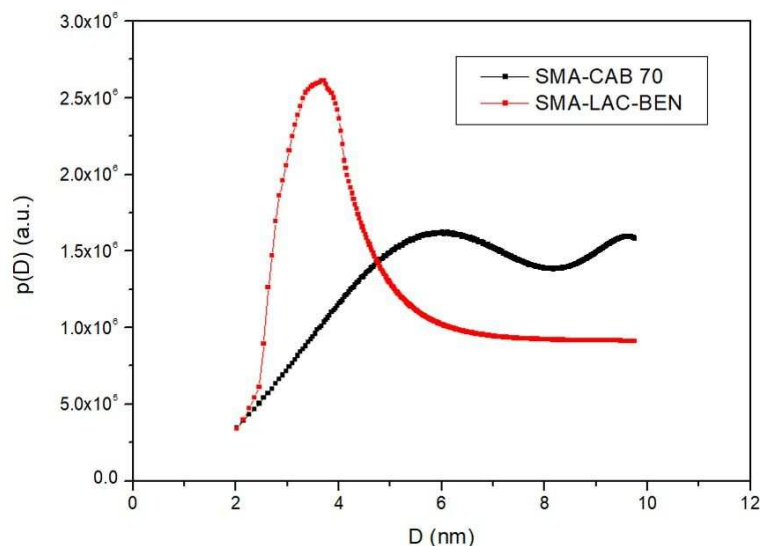


Figure 10: Size probability density $p(D)$ (normalized to the range of existence of the two curves) for the two samples. Particles (or particle aggregates) having diameter values larger than about 10 nm are still blocked at r.t. and therefore cannot be studied by this method of analysis.

The results of this analysis are reported in Figure 10 assuming that the effective magnetic anisotropy constant of the individual Fe^0 nanoparticles takes a value $K_{\text{eff}} = 2 \times 10^6 \text{ erg/cm}^3$ over the whole temperature interval, in agreement with the literature, where K_{eff} has been reported to typically range between 2×10^6 and $3.5 \times 10^6 \text{ erg/cm}^3$.^{83–86} In sample SMA_Cab70, the size distribution density $p(D)$ monotonically increases at low D and reaches a kind of a plateau with weak oscillations; on the contrary, in sample SMA_LacBen the distribution exhibits a sharp peak at $D \cong 3.5 \text{ nm}$ followed by a plateau in the 6-10 nm interval. In both cases, the distributions show no traces of being reduced at high D values.

The $p(D)$ curves obtained from magnetic measurements are in good general agreement with the histograms referring to the small rounded Fe^0 particles observed by HRTEM in both materials (Figures 6 and 7). Instead, the larger globular particles with core-shell structure put in evidence by the same technique are beyond the size interval covered by the analysis, the upper limit of particle sizes ($D_{\text{max}} \approx 10 \text{ nm}$) being determined by the maximum temperature of FC/ZFC measurements (in this case, r.t.).

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3 In the light of the results of HRTEM analysis reported in Section 3.1 (Figures 4-7), it is possible to
4 draw the following picture on the basis of the magnetic properties of Fe⁰ particles measured in both
5 samples: in SMA_LacBen the sharp maximum of $p(D)$ observed at small D values is related to the
6 individual response of small Fe⁰ particles such as the ones put in evidence in Figure 6 (*a-d*, indicated
7 by red arrows), which can be mostly considered as weakly interacting and responding individually
8 to the action of the magnetic field. On the other hand, the constant tail of the $p(D)$ function in the
9 interval 6-10 nm, i.e., in a size region where the histogram distribution obtained by TEM analysis is
10 already decreasing to zero (see inset in panel *a* of Figure 6), is ascribed to the presence of a magnetic
11 correlation among some of the small Fe⁰ particles, which makes them to respond to the magnetic field
12 as larger individual units. In fact, pairs or groups of small particles characterized by a lower than
13 average interparticle distance d can behave collectively when the dipolar magnetic interaction, locally
14 enhanced by the small value of d , is no longer negligible. These interacting groups are viewed by the
15 present analysis as magnetically individual particles of slightly larger size.

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31 On the other hand, in sample SMA_Cab70 no definite peak of the $p(D)$ function emerges at low D .
32 This result is in agreement with the larger histogram distribution obtained by TEM (see inset in panel
33 *a*) of Figure 7). Moreover, the absence of a definite peak in $p(D)$ can be related as well to a stronger
34 dipolar interaction among Fe⁰ particles in this sample with respect to SMA_LacBen. In fact, the larger
35 average diameter of the individual particles determined by HRTEM (7.3 nm in this case against 4.4
36 nm in SMA_LacBen) and their higher concentration and lower dispersion (see Section 3.1) concur in
37 increasing their dipolar interaction energy (which increases as the cube of D and as the reciprocal of
38 the cube of d). The larger dipolar interaction can be instrumental in making more probable the
39 formation of correlated pairs or clusters of particles acting as single magnetic units. This explains
40 why a broader, less structured $p(D)$ function is observed over the entire interval of diameters explored
41 by the FC/ZFC curve analysis.

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55 The absence of a decreasing trend of both $p(D)$ curves for $D \rightarrow D_{max}$ points to the existence of a
56 conspicuous fraction of individual magnetic units (i.e. correlated particles) whose size is larger than
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3 10 nm. These units appear as magnetically blocked over the entire interval of investigated
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5 temperatures.

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7 The magnetic properties of the investigated materials bear close similarities with the results of other
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9 measurements done both on actual lunar soils, simulants and synthetic materials.^{14,20} All
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11 measurements point to the coexistence of various magnetic contributions in these materials, including
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13 para-, antiferro-, ferro- and superparamagnetic phases, as evidenced in the present case.¹⁴ A
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15 comparison with a few similar materials can be found in a previous paper.²⁰ The saturation
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17 magnetization of the ferromagnetic phase is very close to the values reported elsewhere.^{14,20}
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19 However, the coercive field values (in the 250 - 370 Oe interval around room temperature) make the
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21 present materials more similar to simulants made from a terrestrial volcanic tuff or to synthetic
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23 materials than to actual lunar soils, characterized by much lower values of H_c (less than 50 Oe).¹⁴
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25 This difference can be ascribed to a smaller nanoparticle size in actual lunar soils; the size analysis
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27 performed here (Figure 10) points to the existence of magnetically blocked nanoparticles having sizes
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29 much larger than 10 nm, whilst in lunar soils the low measured coercive field is coherent with the
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31 presence of almost superparamagnetic nanoparticles; however, a quantitative estimate of the size of
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33 magnetic nanoparticles is not given in the existing literature.
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39 3.3 Analysis of SMA_LacBen and SMA_Cab70 by broad dielectric spectroscopy

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43 **Dielectric relaxations maps ϵ'' (f, T)**
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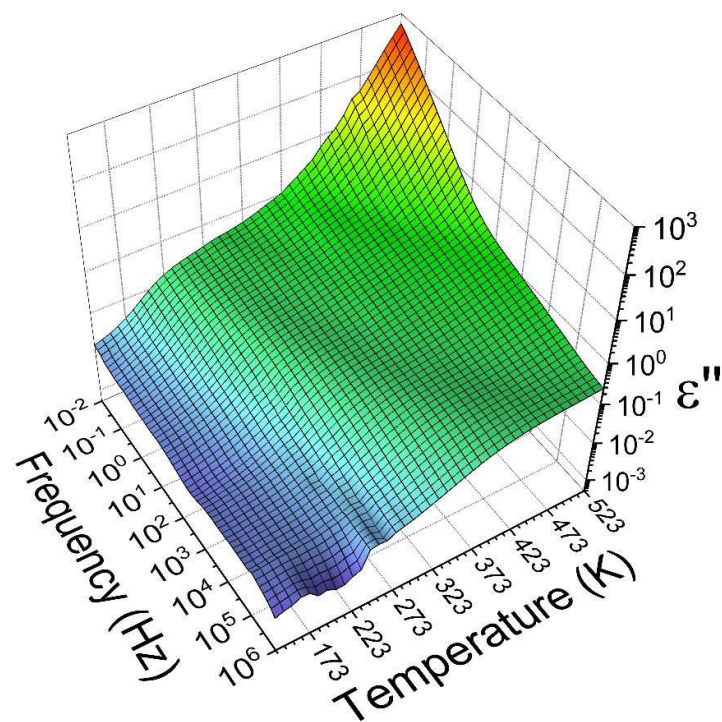


Figure 11 – Dielectric relaxation map – SMA_LacBen.

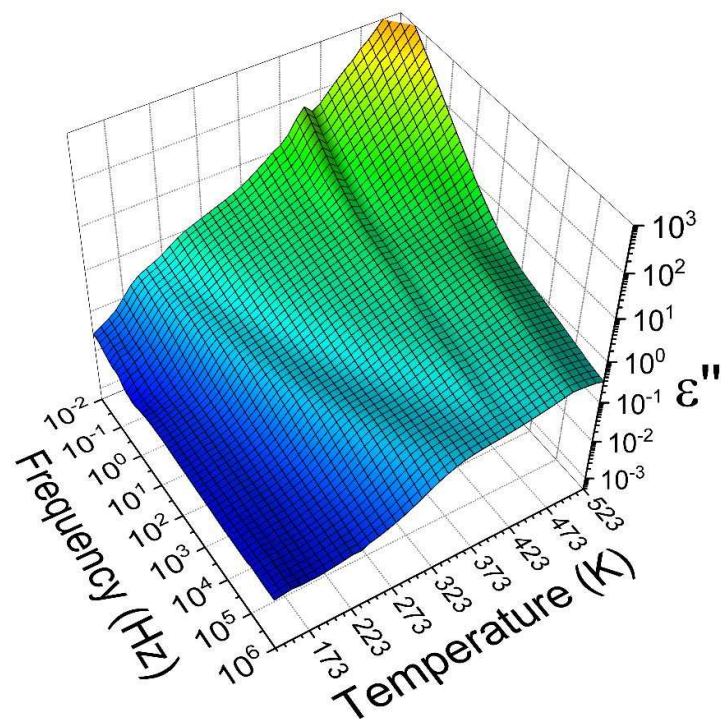


Figure 12 – Dielectric relaxation map SMA_Cab70.

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3 The BDS map of the SMA_LacBen sample (Figure 11) shows one relaxation mode in the
4 vicinity of 223 K at 10^{-2} Hz, that shifts towards higher frequencies as the temperature increases (in
5 the 106 Hz range at 373 K). This mode could have various different physical origins. On one hand, it
6 could be the manifestation of interfacial polarization phenomena, of the Maxwell-Wagner-Sillars
7 (MWS) type, at the particle-void interfaces in the compacted powder sample. On the other hand, the
8 mode could arise from ionic dipole-relaxations involving back and forth ion transfer between
9 potential energy wells (by a hopping mechanism). Such relaxations have been reported for various
10 zeolites.⁸⁷ Due to the loosely controlled packing of the particles, there is insufficient data to ascribe
11 the dielectric mode to either of the two phenomena.
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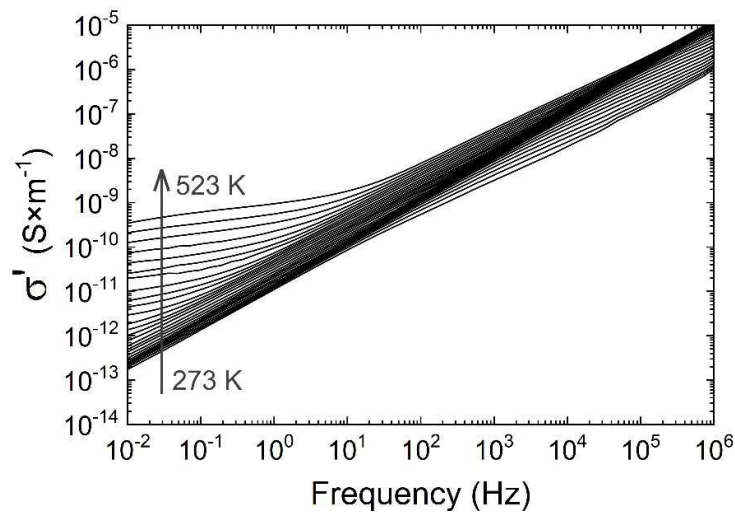
22 The BDS map of SMA_Cab70 (Figure 12) shows a relaxation mode similar to the one
23 observed in SMA_LacBen, but followed by a second mode (shifted by approximately 50 K towards
24 higher temperatures). The same discussion as for the SMA_LacBen case remains regarding the
25 physical origins of these two modes. Because the SMA_Cab70 sample did not display enough
26 conductivity in the range 133-423 K, additional measurements were performed in the range 413-523
27 K (the next day). Those measurements do not strictly overlap with the first series, probably due to
28 water desorption during the first run, resulting in a small isothermal event visible at 413 K in Figure
29 12. The measurements on LacBen were performed in one run between 133 and 523 K and therefore
30 do not display this small event.
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41 At high temperatures and low frequencies, both moon dust simulants display a conductivity
42 front associated with large scale transport of charge carriers across the sample.
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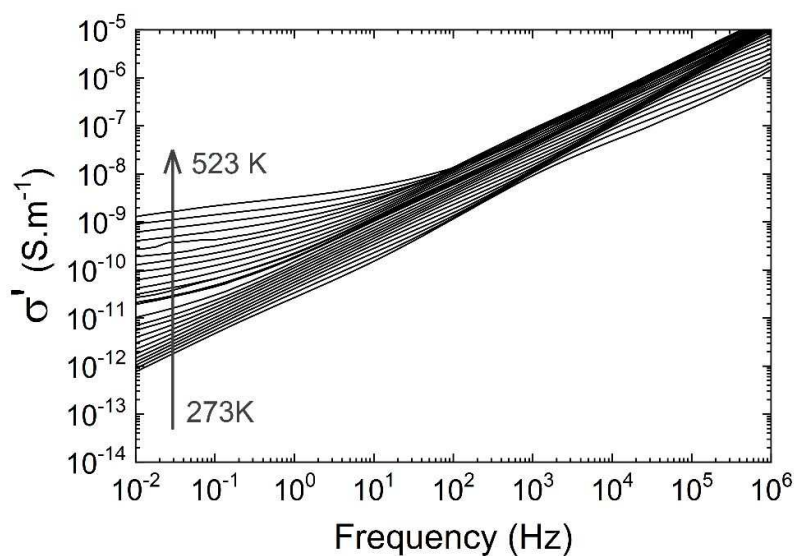
48 **3.4 Isothermal real conductivity spectra, $\sigma'(f)$, and Arrhenius diagrams of the real conductivity** 49 **values at 10⁻² Hz, $\sigma'(1000/T)$** 50

51 The isothermal real conductivity spectra are displayed in the temperature range 273-523 K for
52 SMA_LacBen (Figure 13) and SMA_Cab70 (Figure 14). This representation emphasizes low-
53 frequency charge transport processes which are expected to appear as frequency-independent
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3 plateaus. In both cases, at sufficiently high temperature, such plateaus are observed even though they
4 are slightly frequency dependent. The temperature dependence of the low-frequency conductivity
5 (taken at 10^{-2} Hz) is visible in the Arrhenius diagram of Figure 15.
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27 **Figure 13** – Isothermal real conductivity spectra between 273 and 523 K of sample SMA_LacBen.
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48 **Figure 14** – Isothermal real conductivity spectra between 273 and 523 K of sample SMA_Cab70
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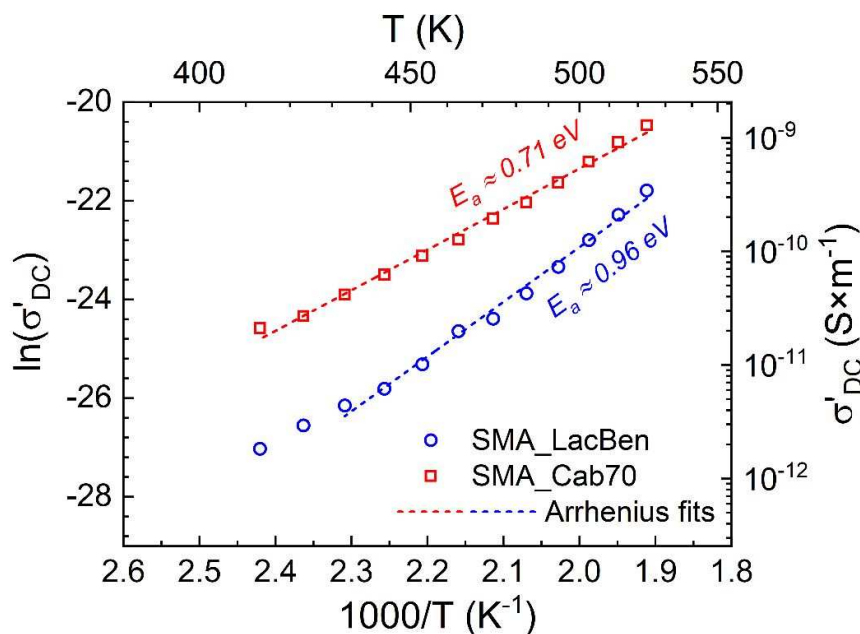


Figure 15 – Arrhenius diagram of the real conductivity at 10⁻² Hz for SMA_LacBen and SMA_Cab70, with the corresponding Arrhenius fits and activation energies

The conductivity values of the SMA_Cab70 sample is systematically higher than that of SMA_LacBen in the temperature range 413-523 K, as can be seen in Figure 15. However, the one-decade difference between the two samples at 423 K decreases to a factor of 4 at 523 K, due to different values of the activation energy for charge transport (0.71 eV for SMA_Cab70 and 0.96 eV for SMA_LacBen).

When compared to the previously studied moon dust simulants (SMA_A and SMA_X), the SMA_LacBen and SMA_Cab70 samples are overall less conductive (sometimes by several orders of magnitude). This is why frequency independent plateaus are not observed below approximately 412 K for SMA_LacBen and SMA_Cab70 samples (as seen in Figure 13 and Figure 14) and the temperature range had to be extended up to 523 K.²⁰ The activation energy is 0.71 eV and 0.96 eV for SMA_Cab70 and SMA_LacBen, respectively, slightly larger than the 0.32 eV and 0.58 eV for Apollo lunar samples 15301,38 and 14613,131, respectively. The DC conductivity at 200 °C is about 3e⁻¹⁰ and 2e⁻¹¹ S/m for SMA_Cab70 and SMA_LacBen, respectively, which is closer to the 7e⁻¹³ and 4e⁻¹¹ S/m of 15301,38 and 14613,131, respectively, than DC conductivity of SMA_A and SMA_X.

CONCLUSIONS

The already good results obtained in our previous work with Moon agglutinates simulants obtained by thermally transforming, under a reducing atmosphere, Fe-exchanged zeolite A and X in metal-ceramic nanocomposites have been further improved in this work by starting from natural zeolite samples.

The following improvements were obtained in in this work:

- 1) A chemical composition decidedly closer to the real Moon dust was obtained.
- 2) A mineralogical composition decidedly closer to the real Moon dust was obtained.
- 3) As a consequence of obtaining an improved chemical and mineralogical composition, also the electric and magnetic behavior of the magnetic metal-ceramic nanocomposites of this work are decidedly similar to those of the real Moon dust.

Moreover, the obtained metal-ceramic nanocomposites do not appear prone to weathering as ceramic phase are well known to be stable and most of the Fe⁰ nanoparticles are embedded into the ceramic matrix which protects them from oxidation.

However, this work has suggested possible further improvements of the, already good, results obtained. Actually, the Fe-exchange natural zeolite bearing material to be transformed in the metal-ceramic nanocomposite simulating the Moon dust, could be split in two portions of about equal entity. The first portion should be thermally treated under a reducing atmosphere at 750 °C, as it was done in this work, and the second portion should be thermally treated, under the same reducing atmosphere, at a decidedly higher temperature, such as 1000 °C. The thermal treatment at this temperature would result in a partial melting of the ceramic matrix which could properly mimic the effects of the meteorite bombardment on the real moon dust. Finally, the two portions of the metal-ceramic nanocomposites obtained should be mixed together to form a lone metal-ceramic nanocomposite. This resulting material should still better mimic the features of the real moon dust as also the effects of the meteorite bombardment on moon dust would be, at least partially, reproduced.

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3 Acknowledgements

4 Authors thank Federico Mondino (Department of Environment, Land and Infrastructure Engineering,
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References

- (1) Liu, Y.; Taylor, L. A. Characterization of Lunar Dust and a Synopsis of Available Lunar Simulants. *Planet. Space Sci.* **2011**, *59* (14), 1769–1783.
- (2) WALLACE, W. T.; TAYLOR, L. A.; LIU, Y.; COOPER, B. L.; McKAY, D. S.; CHEN, B.; JEEVARAJAN, A. S. Lunar Dust and Lunar Simulant Activation and Monitoring. *Meteorit. Planet. Sci.* **2009**, *44* (7), 961–970. <https://doi.org/10.1111/j.1945-5100.2009.tb00781.x>.
- (3) Carpenter, J. D.; Crawford, I. A.; Cockell, C.; Koschny, D.; Jaumann, R.; Wieczorek, M. Scientific Preparations for Lunar Exploration. *Planetary and Space Science*. December 2012, pp 1–2. <https://doi.org/10.1016/j.pss.2012.03.001>.
- (4) Loftus, D. J.; Tranfield, E. M.; Rask, J. C.; McCrossin, C. The Chemical Reactivity of Lunar Dust Relevant to Human Exploration of the Moon. *NASA Ames Research Center* <https://www.lpi.usra.edu/decadal/leag/DavidJLoftus.pdf>
- (5) Carrier, W. D. Particle Size Distribution of Lunar Soil. *J. Geotech. Geoenvironmental Eng.* **2003**, *129* (10), 956–959. [https://doi.org/10.1061/\(asce\)1090-0241\(2003\)129:10\(956\)](https://doi.org/10.1061/(asce)1090-0241(2003)129:10(956)).
- (6) Liu, Y.; Park, J.; Schnare, D. W.; Hill, E.; Taylor, L. A. . Characterization of Lunar Dust for Toxicological Studies. II: Texture and Shape Characteristic. *J. Aerosp. Eng.* **2008**, *21* (4), 272–279.
- (7) Park, J.; Liu, Y.; Kihm, K. D.; Taylor, L. A. Characterization of Lunar Dust for Toxicological Studies. I: Particle Size Distribution. *J. Aerosp. Eng.* **2008**, *21* (4), 266–271.
- (8) Khan-Mayberry, N. The Lunar Environment: Determining the Health Effects of Exposure to Moon Dusts. *Acta Astronaut.* **2008**, *63* (7), 1006–1014.
- (9) Zhang, J.; Yang, W.; Hu, S.; Lin, Y.; Fang, G.; Li, C.; Peng, W.; Zhu, S.; He, Z.; Zhou, B.; Lin, H.; Yang, J.; Liu, E.; Xu, Y.; Wang, J.; Yao, Z.; Zou, Y.; Yan, J.; Ouyang, Z. Volcanic History of the Imbrium Basin: A Close-up View from the Lunar Rover Yutu. *Proc. Natl. Acad. Sci.* **2015**, *112* (17), 5342–5347. <https://doi.org/10.1073/pnas.1503082112>.
- (10) Colwell, J. E.; Batiste, S.; Horányi, M.; Robertson, S.; Sture, S. Lunar Surface: Dust Dynamics and Regolith Mechanics. *Rev. Geophys* **2007**, *45* (2), 2006. <https://doi.org/10.1029/2005RG000184>.
- (11) Caston, R.; Luc, K.; Hendrix, D.; Hurowitz, J. A.; Demple, B. Assessing Toxicity and Nuclear and Mitochondrial DNA Damage Caused by Exposure of Mammalian Cells to Lunar Regolith Simulants. *Wiley Online Libr.* **2018**, *2* (4), 139–148. <https://doi.org/10.1002/2017GH000125>.
- (12) McKay, D. S. .; Blacic, J. D.; Center, N. J. S. Workshop on Production and Uses of Simulated Lunar Materials, LPI Technical Report 91-01. In *Workshop on Production and Uses of Simulated Lunar Materials. A Lunar and Planetary Institute Workshop*; McKay, D. S. ., Blacic, J. D., Eds.; Lunar and Planetary Institute, 3303 NASA Road 1, Houston, TX 77058: The Lunar and Planetary Institute, in Houston, Texas, 1989; Vol. 1, p 83.
- (13) Wentworth, S. J.; Keller, L. P.; McKay, D. S.; Morris, R. V. Space Weathering on the Moon: Patina on Apollo 17 Samples 75075 and 76015. *Meteorit. Planet. Sci.* **1999**, *34* (4), 593–603.
- (14) Liu, Y.; Taylor, L. A.; Thompson, J. R.; Schnare, D. W.; Park, J.-S. Unique Properties of Lunar Impact Glass: Nanophase Metallic Fe Synthesis. *Am. Mineral.* **2007**, *92* (8–9), 1420–

- 1
2
3 1427. <https://doi.org/10.2138/am.2007.2333>.
- 4
5 (15) James, C. L. .; Letsinger, S. L. .; Basu, A. .; Wentworth, S. J. .; McKay, D. S. Size
6 Distribution of Fe₀ Globules in Lunar Agglutinitic Glass. *33rd Annu. Lunar Planet. Sci.*
7 *Conf.* **2002**.
- 8
9 (16) Wallace, W. T.; Phillips, C. J.; Jeevarajan, A. S.; Chen, B.; Taylor, L. A. Nanophase Iron-
10 Enhanced Chemical Reactivity of Ground Lunar Soil. *Earth Planet. Sci. Lett.* **2010**, *295*,
11 571–577. <https://doi.org/http://dx.doi.org/10.1016/j.epsl.2010.04.042>.
- 12
13 (17) Fubini, B.; Hubbard, A. Reactive Oxygen Species (ROS) and Reactive Nitrogen Species
14 (RNS) Generation by Silica in Inflammation and Fibrosis. *Free Radic. Biol. Med.* **2003**, *34*
15 (12), 1507–1516.
- 16
17 (18) Hurowitz, J. A.; Tosca, N. J.; McLennan, S. M.; Schoonen, M. A. A. Production of
18 Hydrogen Peroxide in Martian and Lunar Soils. *Earth Planet. Sci. Lett.* **2007**, *255* (1–2), 41–
19 52.
- 20
21 (19) Horwell, C. J.; Fenoglio, I.; Fubini, B. Iron-Induced Hydroxyl Radical Generation from
22 Basaltic Volcanic Ash. *Earth Planet. Sci. Lett.* **2007**, *261* (3–4), 662–669.
- 23
24 (20) Freyria, F. S.; Marocco, A.; Esposito, S.; Bonelli, B.; Barrera, G.; Tiberto, P.; Allia, P.;
25 Oudayer, P.; Roggero, A.; Matéo-Vélez, J. C.; Dantras, E.; Pansini, M. Simulated Moon
26 Agglutinates Obtained from Zeolite Precursor by Means of a Low-Cost and Scalable
27 Synthesis Method. *ACS Earth Sp. Chem.* **2019**, *3* (9), 1884–1895.
28 <https://doi.org/10.1021/acsearthspacechem.9b00042>.
- 29
30 (21) Witze, A. Why Planetary Scientists Want Better Fake Space Dirt. *Nature* **2017**, *547*, 146–
31 147.
- 32
33 (22) Esposito, S.; Marocco, A.; Bonelli, B.; Pansini, M. Produzione Di Materiali Compositi
34 Metallo-Ceramici Nano Strutturati Da Precursori Zeolitici. MI 2014, A000522.
- 35
36 (23) Esposito, S.; Marocco, A.; Bonelli, B.; Pansini, M. Production of Magnetic Metal
37 Nanoparticles Embedded in a Silica-Alumina Matrix. World Patent WO/ 145230 A1, 2015.
- 38
39 (24) Marocco, A.; Dell’Agli, G.; Esposito, S.; Pansini, M. Metal-Ceramic Composite Materials
40 from Zeolite Precursor. *Solid State Sci.* **2012**, *14* (3), 394–400.
41 <https://doi.org/10.1016/j.solidstatesciences.2012.01.006>.
- 42
43 (25) Ronchetti, S.; Turcato, E. A.; Delmastro, A.; Esposito, S.; Ferone, C.; Pansini, M.; Onida, B.;
44 Mazza, D. Study of the Thermal Transformations of Co- and Fe-Exchanged Zeolites A and X
45 by “in Situ” XRD under Reducing Atmosphere. *Mater. Res. Bull.* **2010**, *45* (6), 744–750.
46 <https://doi.org/10.1016/j.materresbull.2010.02.006>.
- 47
48 (26) Barrera, G.; Tiberto, P.; Esposito, S.; Marocco, A.; Bonelli, B.; Pansini, M.; Manzoli, M.;
49 Allia, P. Magnetic Clustering of Ni²⁺ Ions in Metal-Ceramic Nanocomposites Obtained from
50 Ni-Exchanged Zeolite Precursors. *Ceram. Int.* **2018**, *44* (14), 17240–17250.
51 <https://doi.org/10.1016/j.ceramint.2018.06.182>.
- 52
53 (27) Esposito, S.; Dell’Agli, G.; Marocco, A.; Bonelli, B.; Allia, P.; Tiberto, P.; Barrera, G.;
54 Manzoli, M.; Arletti, R.; Pansini, M. Magnetic Metal-Ceramic Nanocomposites Obtained
55 from Cation-Exchanged Zeolite by Heat Treatment in Reducing Atmosphere. *Microporous*
56 *Mesoporous Mater.* **2018**, *268*, 131–143. <https://doi.org/10.1016/j.micromeso.2018.04.024>.
- 57
58
59
60

- 1
2
3 (28) Barrera, G.; Tiberto, P.; Allia, P.; Bonelli, B.; Esposito, S.; Marocco, A.; Pansini, M.;
4 Leterrier, Y. Magnetic Properties of Nanocomposites. *Appl. Sci.* **2019**, *9* (2), 212.
5 <https://doi.org/10.3390/app9020212>.
6
7 (29) Barrera, G.; Allia, P.; Bonelli, B.; Esposito, S.; Freyria, F. S.; Pansini, M.; Marocco, A.;
8 Confalonieri, G.; Arletti, R.; Tiberto, P. Magnetic Behavior of Ni Nanoparticles and Ni²⁺
9 Ions in Weakly Loaded Zeolitic Structures. *J. Alloys Compd.* **2020**, *817*, 152776.
10 <https://doi.org/10.1016/j.jallcom.2019.152776>.
11
12 (30) Pansini, M.; Dell'Agli, G.; Marocco, A.; Netti, P. A.; Battista, E.; Lettera, V.; Vergara, P.;
13 Allia, P.; Bonelli, B.; Tiberto, P.; Barrera, G.; Alberto, G.; Martra, G.; Arletti, R.; Esposito,
14 S. Preparation and Characterization of Magnetic and Porous Metal-Ceramic Nanocomposites
15 from a Zeolite Precursor and Their Application for DNA Separation. *J. Biomed.*
16 *Nanotechnol.* **2017**, *13* (3), 337–348. <https://doi.org/10.1166/jbn.2017.2345>.
17
18 (31) Pansini, M.; Sannino, F.; Marocco, A.; Allia, P.; Tiberto, P.; Barrera, G.; Polisi, M.; Battista,
19 E.; Netti, P. A.; Esposito, S. Novel Process to Prepare Magnetic Metal-Ceramic
20 Nanocomposites from Zeolite Precursor and Their Use as Adsorbent of Agrochemicals from
21 Water. *J. Environ. Chem. Eng.* **2018**, *6* (1), 527–538.
22 <https://doi.org/10.1016/j.jece.2017.12.030>.
23
24 (32) Marocco, A.; Dell'Agli, G.; Sannino, F.; Esposito, S.; Bonelli, B.; Allia, P.; Tiberto, P.;
25 Barrera, G.; Pansini, M. Removal of Agrochemicals from Waters by Adsorption: A Critical
26 Comparison among Humic-Like Substances, Zeolites, Porous Oxides, and Magnetic
27 Nanocomposites. *Processes* **2020**, *8* (2), 141. <https://doi.org/10.3390/pr8020141>.
28
29 (33) Esposito, S.; Marocco, A.; Dell'Agli, G.; Bonelli, B.; Mannu, F.; Allia, P.; Tiberto, P.;
30 Barrera, G.; Pansini, M. Separation of Biological Entities from Human Blood by Using
31 Magnetic Nanocomposites Obtained from Zeolite Precursors. *Molecules* **2020**, *25* (8), 1803.
32 <https://doi.org/10.3390/molecules25081803>.
33
34 (34) Esposito, S.; Marocco, A.; Dell'Agli, G.; De Gennaro, B.; Pansini, M. Relationships between
35 the Water Content of Zeolites and Their Cation Population. *Microporous Mesoporous Mater.*
36 **2015**, *202*, 36–43. <https://doi.org/10.1016/j.micromeso.2014.09.041>.
37
38 (35) Cappelletti, P.; Langella, A.; Cruciani, G. Crystal-Chemistry and Synchrotron Rietveld
39 Refinement of Two Different Clinoptilolites from Volcanoclastites of North-Western
40 Sardinia. *Eur. J. Mineral.* **1999**, *11* (6), 1051–1060. <https://doi.org/10.1127/ejm/11/6/1051>.
41
42 (36) Cerri, G.; Langella, A.; Pansini, M.; Cappelletti, P. Methods of Determining Cation
43 Exchange Capacities for Clinoptilolite-Rich Rocks of the Logudoro Region in Northern
44 Sardinia, Italy. *Clays Clay Miner.* **2002**, *50* (1), 127–135.
45 <https://doi.org/10.1346/000986002761002739>.
46
47 (37) Langella, A.; Pansini, M.; Cerri, G.; Cappelletti, P.; De'Gennaro, M. Thermal Behavior of
48 Natural and Cation-Exchanged Clinoptilolite from Sardinia (Italy). *Clays Clay Miner.* **2003**,
49 *51* (6), 625–633. <https://doi.org/10.1346/CCMN.2003.0510605>.
50
51 (38) Cerri, G.; Cappelletti, P.; Langella, A.; De'Gennaro, M. Zeolitization of Oligo-Miocene
52 Volcaniclastic Rocks from Logudoro (Northern Sardinia, Italy). *Contrib. to Mineral. Petrol.*
53 **2001**, *140* (4), 404–421. <https://doi.org/10.1007/s004100000196>.
54
55 (39) Cappelletti, P.; Cerri, G.; Colella, A.; de'Gennaro, M.; Langella, A.; Perrotta, A.; Scarpati, C.
56 Post-Eruptive Processes in the Campanian Ignimbrite. *Mineral. Petrol.* **2003**, *79* (1–2), 79–
57
58
59

- 1
2
3 97. <https://doi.org/10.1007/s00710-003-0003-7>.
- 4
5 (40) Langella, A.; Bish, D. L.; Cappelletti, P.; Cerri, G.; Colella, A.; de Gennaro, R.; Graziano, S.
6 F.; Perrotta, A.; Scarpati, C.; de Gennaro, M. New Insights into the Mineralogical Facies
7 Distribution of Campanian Ignimbrite, a Relevant Italian Industrial Material. *Appl. Clay Sci.*
8 **2013**, *72*, 55–73. <https://doi.org/10.1016/j.clay.2013.01.008>.
- 9
10 (41) Colella, C.; De' Gennaro, M.; Langella, A.; Pansini, M. Evaluation of Natural Phillipsite and
11 Chabazite as Cation Exchangers for Copper and Zinc. *Sep. Sci. Technol.* **1998**, *33* (4), 467–
12 481. <https://doi.org/10.1080/01496399808544991>.
- 13
14 (42) Pansini, M.; Colella, C. Dynamic Data on Lead Uptake from Water by Chabazite.
15 *Desalination* **1990**, *78* (2), 287–295. [https://doi.org/10.1016/0011-9164\(90\)80048-G](https://doi.org/10.1016/0011-9164(90)80048-G).
- 16
17 (43) Albino, V.; Cioffi, R.; Pansini, M.; Colella, C. Disposal of Lead-Containing Zeolite Sludges
18 in Cement Matrix. *Environ. Technol. (United Kingdom)* **1995**, *16* (2), 147–156.
19 <https://doi.org/10.1080/09593331608616255>.
- 20
21 (44) Colella, C.; Pansini, M. Lead Removal from Wastewaters Using Chabazite Tuff. In *ACS*
22 *Symposium Series*; Publ by ACS, 1988; pp 500–510. [https://doi.org/10.1021/bk-1988-](https://doi.org/10.1021/bk-1988-0368.ch032)
23 [0368.ch032](https://doi.org/10.1021/bk-1988-0368.ch032).
- 24
25 (45) Cioffi, R.; Pansini, M.; Caputo, D.; Colella, C. Evaluation of Mechanical and Leaching
26 Properties of Cement-Based Solidified Materials Encapsulating Cd-Exchanged Natural
27 Zeolites. *Environ. Technol. (United Kingdom)* **1996**, *17* (11), 1215–1224.
28 <https://doi.org/10.1080/09593331708616491>.
- 29
30 (46) Pansini, M.; De Gennaro, R.; Parlato, L.; De'Gennaro, M.; Langella, A.; Marocco, A.;
31 Cappelletti, P.; Mercurio, M. Use of Sawing Waste from Zeolitic Tuffs in the Manufacture of
32 Ceramics. *Adv. Mater. Sci. Eng.* **2010**, *2010*. <https://doi.org/10.1155/2010/820541>.
- 33
34 (47) de' Gennaro, R.; Cappelletti, P.; Cerri, G.; de' Gennaro, M.; Dondi, M.; Guarini, G.;
35 Langella, A.; Naimo, D. Influence of Zeolites on the Sintering and Technological Properties
36 of Porcelain Stoneware Tiles. *J. Eur. Ceram. Soc.* **2003**, *23* (13), 2237–2245.
37 [https://doi.org/10.1016/S0955-2219\(03\)00086-4](https://doi.org/10.1016/S0955-2219(03)00086-4).
- 38
39 (48) de Gennaro, R.; Dondi, M.; Cappelletti, P.; Cerri, G.; de' Gennaro, M.; Guarini, G.; Langella,
40 A.; Parlato, L.; Zanelli, C. Zeolite-Feldspar Epiclastic Rocks as Flux in Ceramic Tile
41 Manufacturing. *Microporous Mesoporous Mater.* **2007**, *105* (3), 273–278.
42 <https://doi.org/10.1016/j.micromeso.2007.04.023>.
- 43
44 (49) de Gennaro, R.; Langella, A.; D'Amore, M.; Dondi, M.; Colella, A.; Cappelletti, P.; de'
45 Gennaro, M. Use of Zeolite-Rich Rocks and Waste Materials for the Production of Structural
46 Lightweight Concretes. *Appl. Clay Sci.* **2008**, *41* (1–2), 61–72.
47 <https://doi.org/10.1016/j.clay.2007.09.008>.
- 48
49 (50) de Gennaro, R.; Cappelletti, P.; Cerri, G.; de' Gennaro, M.; Dondi, M.; Graziano, S. F.;
50 Langella, A. Campanian Ignimbrite as Raw Material for Lightweight Aggregates. *Appl. Clay*
51 *Sci.* **2007**, *37* (1–2), 115–126. <https://doi.org/10.1016/j.clay.2006.11.004>.
- 52
53 (51) de' Gennaro, R.; Cappelletti, P.; Cerri, G.; de' Gennaro, M.; Dondi, M.; Langella, A. Zeolitic
54 Tuffs as Raw Materials for Lightweight Aggregates. *Appl. Clay Sci.* **2004**, *25* (1–2), 71–81.
55 <https://doi.org/10.1016/j.clay.2003.08.005>.
- 56
57 (52) Mercurio, M.; Bish, D. L.; Cappelletti, P.; de Gennaro, B.; de Gennaro, M.; Grifa, C.; Izzo,
58
59
60

- F.; Mercurio, V.; Morra, V.; Langella, A. The Combined Use of Steam-Treated Bentonites and Natural Zeolites in the Oenological Refining Process. *Mineral. Mag.* **2016**, *80* (2), 347–362. <https://doi.org/10.1180/minmag.2016.080.004>.
- (53) Cappelletti, P.; Rapisardo, G.; De Gennaro, B.; Colella, A.; Langella, A.; Graziano, S. F.; Bish, D. L.; De Gennaro, M. Immobilization of Cs and Sr in Aluminosilicate Matrices Derived from Natural Zeolites. *J. Nucl. Mater.* **2011**, *414* (3), 451–457. <https://doi.org/10.1016/j.jnucmat.2011.05.032>.
- (54) Mercurio, M. ; Langella, A. ; Cappelletti, P. ; de Gennaro, B. ; Monetti, V. ; de Gennaro M. May the Use of Italian Volcanic Zeolite-Rich Tuffs as Additives in Animal Diet Represent a Risk for the Human Health? . *Period. di Mineral.* **2012**, *81* (3), 393–407.
- (55) Cappelletti, P.; Colella, A.; Langella, A.; Mercurio, M.; Catalanotti, L.; Monetti, V.; de Gennaro, B. Use of Surface Modified Natural Zeolite (SMNZ) in Pharmaceutical Preparations Part I. Mineralogical and Technological Characterization of Some Industrial Zeolite-Rich Rocks. *Microporous Mesoporous Mater.* **2017**, *250*, 232–244. <https://doi.org/10.1016/j.micromeso.2015.05.048>.
- (56) Basile, A.; Cacciola, G.; Colella, C.; Mercadante, L.; Pansini, M. Thermal Conductivity of Natural Zeolite-PTFE Composites. *Heat Recover. Syst. CHP* **1992**, *12* (6), 497–503. [https://doi.org/10.1016/0890-4332\(92\)90018-D](https://doi.org/10.1016/0890-4332(92)90018-D).
- (57) Baerlocher, C. ; Meier, W. M. ; Olson, D. H. *Atlas of Zeolite Framework Types* ; Elsevier, 2001.
- (58) Dell’Agli, G.; Ferone, C.; Mascolo, G.; Pansini, M. Crystallization of Monoclinic Zirconia from Metastable Phases. *Solid State Ionics* **2000**, *127* (3), 223–230. [https://doi.org/10.1016/S0167-2738\(99\)00279-9](https://doi.org/10.1016/S0167-2738(99)00279-9).
- (59) Clayden, N. J.; Esposito, S.; Ferone, C.; Pansini, M. ²⁹Si and ²⁷Al NMR Study of the Thermal Transformation of Barium Exchanged Zeolite-A to Celsian. *J. Mater. Chem.* **2003**, *13* (7), 1681–1685. <https://doi.org/10.1039/b212717b>.
- (60) Marocco, A.; Liguori, B.; Dell’Agli, G.; Pansini, M. Sintering Behaviour of Celsian Based Ceramics Obtained from the Thermal Conversion of (Ba, Sr)-Exchanged Zeolite A. *J. Eur. Ceram. Soc.* **2011**, *31* (11), 1965–1973. <https://doi.org/10.1016/j.jeurceramsoc.2011.04.028>.
- (61) Ferone, C.; Liguori, B.; Marocco, A.; Anaclerio, S.; Pansini, M.; Colella, C. Monoclinic (Ba, Sr)-Celsian by Thermal Treatment of (Ba, Sr)-Exchanged Zeolite A. *Microporous Mesoporous Mater.* **2010**, *134* (1–3), 65–71. <https://doi.org/10.1016/j.micromeso.2010.05.008>.
- (62) Weidenthaler, C.; Zibrowius, B.; Schimanke, J.; Mao, Y.; Mienert, B.; Bill, E.; Schmidt, W. Oxidation Behavior of Ferrous Cations during Ion Exchange into Zeolites under Atmospheric Conditions. *Microporous Mesoporous Mater.* **2005**, *84* (1–3), 302–317. <https://doi.org/10.1016/j.micromeso.2005.04.022>.
- (63) Colantuono, A.; Dal Vecchio, S.; Mascolo, G.; Pansini, M. Thermal Shrinkage of Various Cation Forms of Zeolite A. *Thermochim. Acta* **1997**, *296* (1–2), 59–66. [https://doi.org/10.1016/s0040-6031\(97\)00065-8](https://doi.org/10.1016/s0040-6031(97)00065-8).
- (64) Marocco, A.; Dell’Agli, G.; Spiridigliozzi, L.; Esposito, S.; Pansini, M. The Multifarious Aspects of the Thermal Conversion of Ba-Exchanged Zeolite A to Monoclinic Celsian. *Microporous and Mesoporous Materials*. Elsevier B.V. January 15, 2018, pp 235–250.

- 1
2
3 <https://doi.org/10.1016/j.micromeso.2017.08.018>.
- 4
5 (65) Gualtieri, A. F. Accuracy of XRPD QPA Using the Combined Rietveld-RIR Method. *J. Appl. Crystallogr.* **2000**, *33* (2), 267–278. <https://doi.org/10.1107/S002188989901643X>.
- 6
7 (66) Larson, A. C. and Von Dreele, R. B. GSAS General Structure Analysis System, LANSCE
8 ,MS-H80. *Los Alamos Natl. Lab.* **1998**, No. Los Alamos, NM 87545, USA,.
- 9
10 (67) Toby, B. H. EXPGUI, a Graphical User Interface for GSAS. *J. Appl. Crystallogr.* **2001**, *34*
11 (2), 210–213. <https://doi.org/10.1107/S0021889801002242>.
- 12
13 (68) Strangway, D. W.; Chapman, W. B.; Olhoeft, G. R.; Carnes, J. Electrical Properties of Lunar
14 Soil Dependence on Frequency, Temperature and Moisture. *Earth Planet. Sci. Lett.* **1972**, *16*
15 (2), 275–281. [https://doi.org/10.1016/0012-821X\(72\)90203-8](https://doi.org/10.1016/0012-821X(72)90203-8).
- 16
17 (69) Alvarez, R. Lunar Powder Simulator under Lunarlike Conditions: Dielectric Properties. *J.*
18 *Geophys. Res.* **1973**, *78* (29), 6833–6844. <https://doi.org/10.1029/JB078i029p06833>.
- 19
20 (70) Cruciani, G. Zeolites upon Heating: Factors Governing Their Thermal Stability and
21 Structural Changes. *J. Phys. Chem. Solids* **2006**, *67* (9–10), 1973–1994.
22 <https://doi.org/10.1016/j.jpics.2006.05.057>.
- 23
24 (71) Charkhi, A.; Kazemian, H.; Kazemeini, M. Optimized Experimental Design for Natural
25 Clinoptilolite Zeolite Ball Milling to Produce Nano Powders. *Powder Technol.* **2010**, *203*
26 (2), 389–396. <https://doi.org/10.1016/j.powtec.2010.05.034>.
- 27
28 (72) Budnyak, T. M.; Yanovska, E. S.; Kichkiruk, O. Y.; Sternik, D.; Tertykh, V. A. Natural
29 Minerals Coated by Biopolymer Chitosan: Synthesis, Physicochemical, and Adsorption
30 Properties. *Nanoscale Res. Lett.* **2016**, *11* (1), 492. [https://doi.org/10.1186/s11671-016-1696-](https://doi.org/10.1186/s11671-016-1696-y)
31 [y](https://doi.org/10.1186/s11671-016-1696-y).
- 32
33 (73) García, J. E.; González, M. M.; Notario, J. S. Phenol Adsorption on Natural Phillipsite.
34 *React. Polym.* **1993**, *21* (3), 171–176. [https://doi.org/10.1016/0923-1137\(93\)90119-Z](https://doi.org/10.1016/0923-1137(93)90119-Z).
- 35
36 (74) Sing, K. S. W.; Williams, R. T. Physisorption Hysteresis Loops and the Characterization of
37 Nanoporous Materials. *Adsorpt. Sci. Technol.* **2004**, *22* (10), 773–782.
38 <https://doi.org/10.1260/0263617053499032>.
- 39
40 (75) Potter, D. K.; Corbett, P. W. M.; Barclay, S. A.; Haszeldine, R. S. Quantification of Illite
41 Content in Sedimentary Rocks Using Magnetic Susceptibility - A Rapid Complement or
42 Alternative to X-Ray Diffraction. *J. Sediment. Res.* **2004**, *74* (5), 730–735.
43 <https://doi.org/10.1306/021304740730>.
- 44
45 (76) Kandapallil, B.; Colborn, R. E.; Bonitatibus, P. J.; Johnson, F. Synthesis of High
46 Magnetization Fe and FeCo Nanoparticles by High Temperature Chemical Reduction. *J.*
47 *Magn. Magn. Mater.* **2015**, *378*, 535–538. <https://doi.org/10.1016/j.jmmm.2014.11.074>.
- 48
49 (77) Kin, M.; Kura, H.; Tanaka, M.; Hayashi, Y.; Hasaegawa, J.; Ogawa, T. Improvement of
50 Saturation Magnetization of Fe Nanoparticles by Post-Annealing in a Hydrogen Gas
51 Atmosphere. *J. Appl. Phys.* **2015**, *117* (17), 17E714. <https://doi.org/10.1063/1.4919050>.
- 52
53 (78) Margeat, O.; Respaud, M.; Amiens, C.; Lecante, P.; Chaudret, B. Ultrafine Metallic Fe
54 Nanoparticles: Synthesis, Structure and Magnetism. *Beilstein J. Nanotechnol.* **2010**, *1* (1),
55 108–118. <https://doi.org/10.3762/bjnano.1.13>.
- 56
57
58
59
60

- 1
2
3 (79) Barrera, G.; Sciancalepore, C.; Messori, M.; Allia, P.; Tiberto, P.; Bondioli, F. Magnetite-
4 Epoxy Nanocomposites Obtained by the Reactive Suspension Method: Microstructural,
5 Thermo-Mechanical and Magnetic Properties. *Eur. Polym. J.* **2017**, *94*, 354–365.
6 <https://doi.org/10.1016/j.eurpolymj.2017.07.022>.
7
- 8 (80) Allia, P.; Barrera, G.; Tiberto, P. Linearized Rate-Equation Approach for Double-Well
9 Systems: Cooling- and Temperature-Dependent Low-Field Magnetization of Magnetic
10 Nanoparticles. *Phys. Rev. B* **2018**, *98* (13), 134423.
11 <https://doi.org/10.1103/PhysRevB.98.134423>.
12
- 13 (81) Mamiya, H.; Ohnuma, M.; Nakatani, I.; Furubayashim, T. Extraction of Blocking
14 Temperature Distribution from Zero-Field-Cooled and Field-Cooled Magnetization Curves.
15 In *IEEE Transactions on Magnetics*; 2005; Vol. 41, pp 3394–3396.
16 <https://doi.org/10.1109/TMAG.2005.855205>.
17
- 18 (82) Sciancalepore, C.; Bondioli, F.; Messori, M.; Barrera, G.; Tiberto, P.; Allia, P. Epoxy
19 Nanocomposites Functionalized with in Situ Generated Magnetite Nanocrystals:
20 Microstructure, Magnetic Properties, Interaction among Magnetic Particles. *Polymer*
21 *(Guildf)*. **2015**, *59*, 278–289. <https://doi.org/10.1016/j.polymer.2014.12.047>.
22
- 23 (83) Ibusuki, T.; Kojima, S.; Kitakami, O.; Shimada, Y. Magnetic Anisotropy and Behaviors of
24 Fe Nanoparticles. *IEEE Trans. Magn.* **2001**, *37* (4), 2223–2225.
25 <https://doi.org/10.1109/20.951130>.
26
- 27 (84) Shah, L. R.; Ali, B.; Hasanain, S. K.; Mumtaz, A.; Baker, C.; Shah, S. I. Effective Magnetic
28 Anisotropy and Coercivity in Fe Nanoparticles Prepared by Inert Gas Condensation. *Int. J.*
29 *Mod. Phys. B* **2006**, *20* (1), 37–47. <https://doi.org/10.1142/S0217979206033097>.
30
- 31 (85) Mehdaoui, B.; Meffre, A.; Lacroix, L. M.; Carrey, J.; Lachaize, S.; Respaud, M.; Gougeon,
32 M.; Chaudret, B. Magnetic Anisotropy Determination and Magnetic Hyperthermia Properties
33 of Small Fe Nanoparticles in the Superparamagnetic Regime. *J. Appl. Phys.* **2010**, *107* (9),
34 25. <https://doi.org/10.1063/1.3348795>.
35
- 36 (86) Peddis, D.; Qureshi, M. T.; Baker, S. H.; Binns, C.; Roy, M.; Laureti, S.; Fiorani, D.;
37 Nordblad, P.; Mathieu, R. Magnetic Anisotropy and Magnetization Dynamics of Fe
38 Nanoparticles Embedded in Cr and Ag Matrices. *Philos. Mag.* **2015**, *95* (33), 3798–3807.
39 <https://doi.org/10.1080/14786435.2015.1090640>.
40
- 41 (87) Kalogeras, I. M.; Vassilikou-Dova, A. Electrical Properties of Zeolitic Catalysts. *Defect*
42 *Diffus. Forum* **1998**, *164* (164), 1–36. <https://doi.org/10.4028/www.scientific.net/ddf.164.1>.
43
44
45
46
47
48
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51
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