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Synopsis: Zeolite L was Na- and NH_4 -exchanged and the resulting samples were characterized by structural and chemical analyses. From the perspective of the possible use of zeolite L for rare earth element recovery (in particular Ce) and to unveil the differences in the affinity for Ce dictated by the presence of different counter-cations, the three samples with LTL framework type were Ce³⁺-exchanged and fully characterized.

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Keywords: Ce-exchanged zeolite; structural analysis; synchrotron high-resolution X-ray powder diffraction.

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Cerium exchange by microporous materials, such as zeolites, has important applications in different fields, for example, rare earth element recovery from waste or in catalytic processes. This work investigated the Ce-exchange capacity of zeolite L in three different cationic forms (the as-synthesized K form and Naand NH₄-exchanged ones) from a highly concentrated solution. Chemical analyses and structural investigations allowed determination of the mechanisms involved in the exchanges and give new insights into the interactions occurring among the cations and the zeolite framework. Different cation sites are involved: (i) K present in the original LTL in the cancrinite cage (site KB) cannot be exchanged; (ii) the cations in KD (in the 12-membered ring channel) are always exchanged; while (iii) site KC (in the eight-membered ring channel) is involved only when K⁺ is substituted by NH₄⁺, thus promoting a higher exchange rate for $NH_4^+ \rightarrow K^+$ than for $Na^+ \rightarrow K^+$. In the Ce-exchanged samples, a new site occupied by Ce, in the centre of the main channel, appears, accompanied by a rearrangement of the H₂O molecules and an increase in the number of H₂O molecules [ok as edited?]. In terms of Ce exchange, the three cationic forms behave similarly, from both the chemical and structural point of view (exchanged Ce ranges from 38 to 42% of the pristine cation amount). Beyond the intrinsic structural properties of the zeolite L framework, the Ce exchange seems thus also governed by the water coordination sphere of the cation. Complete Ce recovery from zeolite pores was achieved.

1. Introduction

The scarcity of rare earth elements (REE) in some regions, such as Europe (Massari & Ruberti, 2013; Charalampides et al., 2015), and the well known balance problem (i.e. the balance between natural abundance and market request) (Binnemans et al., 2018; de Boer & Lammertsma, 2013) are pushing the scientific community to find alternative ways to supply REE. The most promising solution is to recycle them from waste, which in the last few years has become an urgent requirement for society (Jowitt et al., 2018; Rademaker et al., 2013; Balaram, 2019). Beyond the economic benefits, this process is also necessary for the ecological preservation of the natural environment (Balaram, 2019). The mining, extraction and refining of these REE? elements are in fact associated with high environmental costs (Charalampides et al., 2016; Haque et al., 2014). Despite the fact that eco-friendly politics is supportive of and encourages the development of recycling solutions, the current REE recovery percentage is still lower than 1%, and is limited to a few and very specific applications such as recovery from magnets and polishing compounds (Chakhmouradian & Wall, 2012; Royen & Fortkamp, 2016). The goal for the near future is, thus, to increase the REE



Figure 1

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(a) LTL framework type; (b) columns constituting the framework, formed by the stacking of cancrinite cages and double six-membered rings.

130 recycling rate by finding an economical and convenient way to recover these elements from waste. Leached liquors obtained 131 132 from waste can be a rich secondary source of REE and many methods have been tested to recover them, i.e. solvent 133 extraction (Meshram & Abhilash, 2020; Deshmane et al., 2020; 134 Ni'Am et al., 2020), precipitation (Zhou et al., 2018; Porvali et 135 al., 2018; Diaz et al., 2016), ion exchange (Royen & Fortkamp, 136 2016). The latter can exploit various materials such as active carbon (Gad & Awwad, 2007), resins (Manos & Kanatzidis, 138 2016) or microporous materials (Kavun et al., 2021; Royen & 139 Fortkamp, 2016) such as zeolites. 140

Zeolites are natural and synthetic microporous materials 141 mainly constituted by a complex interconnection of SiO₄ and 142 AlO₄ tetrahedra which leads to many different types of 143 framework (Baerlocher et al., 2007; Baelocher & McCusker 144 iza-structure database http://www.iza-structure.org/databases/ 145) and pore systems (*i.e.* cages, 1D, 2D and 3D network chan-146 nels). The zeolite pores, thanks to their nanometric sizes, can 147 148 host water molecules (Bryukhanov et al., 2017; Coudert et al., 2009; Saada et al., 2011; Bennett & Smith, 1968), many 149 different types of cations (Arletti et al., 2017 [two Arletti et al. 150 2017 references, please indicate which]; Confalonieri et al., 151 2018, 2020 [three Confalonieri et al. 2020 references, please 152 153 indicate which]; Frising & Leflaive, 2008; Isaac et al., 2020; Simoncic & Armbruster, 2004; Barrer & Meier, 1958; Bennett 154 & Smith, 1968) and/or organic molecules [including CO₂] 155 (Hong et al., 2014; Hudson et al., 2012; Lu et al., 2008; 156 Confalonieri et al., 2020), alcohol (Zhang et al., 2012; Arletti et 157 al., 2017; Confalonieri et al., 2019), various hydrocarbons 158 (Fabbiani et al., 2021; Confalonieri et al., 2020; Santoro et al., 159 2003, 2016; Scelta et al., 2014), dyes (Dejoie et al., 2014) or 160 even amino acids (Boekfa et al., 2008; Krohn & Tsapatsis, 161 2005; Stückenschneider et al., 2013) etc.]. One of the most 162 163 remarkable properties of zeolites is the high cationic exchange capacity, where cations in the pores can be easily exchanged (Pabalan & Bertetti, 2001; Dyer, 2007). Zeolites are green 165 materials and cheap [to produce], and their high cationic 166 exchange capacity could potentially be exploited for REE 167 recovery; however, only a few works have investigated this 169 possibility (Mosai et al., 2019; Faghihian et al., 2005; Duplouy, 2016; Barros et al., 2019), and thus new and extensive studies 170 are desirable. 171

In this paper, we describe a preliminary test of the effectiveness of different cationic forms of zeolite L (LTL framework type) (Baerlocher et al., 2007) for cerium recovery. This framework type has been reported as suitable for REE recovery (Duplouy, 2016), and Ce is one of the most exploited REE in terms of volume (Binnemans et al., 2018; Charalampides et al., 2015), being present in a huge variety of technological applications [i.e. as fluid catalytic cracking (f.c.c.) catalysts, glass polishing, fluorescent lamps, NiMH batteries etc. (Meshram & Abhilash, 2020)]. We examine the use of 172

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synthetic zeolite L in its as-synthesized K form and in the Naand NH₄-exchanged ones. The three cationic forms are tested in experiments of Ce exchange from a highly concentrated solution. The obtained results contribute to our understanding of whether zeolite L is suitable for Ce exchange and recovery, and whether Ce shows a different affinity for the diverse exchanged forms. The study was carried out by a combination of different techniques, including structural refinement of high-resolution X-ray powder diffraction experiments. The exploitation of zeolites as cation exchangers cannot disregard the mechanisms involved in the exchanges and the interaction occurring among extra-framework cations, framework and H₂O molecules. Thus, a structural study unravelling from the atomistic point of view the cation exchange mechanism and the sites involved is fundamental for the choice of a powerful protocol of exchange. Indeed, the determination, from chemical analysis, of the maximum amount of Ce incorporated into the zeolite pores would give only a partial result since it would not shed light on the dynamics of Ce adsorption. Structural analyses reported here are fundamental to determine the interactions between Ce cations and the zeolite framework, and will pave the way to further studies devoted to the investigation of zeolite selectivity towards Ce. This structural interpretation can be exploited for targets other than REE recovery – for example, Ce-exchanged zeolites are very promising catalysts for the selective catalytic reduction of N_xO (van Kooten et al., 1998) or for the sterification of glycerol with acetic acid (Gautam et al., 2020).

2. Materials and methods

2.1. Zeolite L

Zeolite L [K₆Na₃(H₂O)₂₁Al₉Si₂₇O₇₂, a = 18.4, c = 7.52 Å, $\gamma =$ 221 120° (Baerlocher et al., 2007)] (LTL framework type) (Fig. 1) is 222 built by the linking of columns formed by the stacking of 223 cancrinite cages and double six-membered rings (6MR), 224 leading to the formation of a 1D porous system formed from 225 [12-membered ring] 12MR and [eight-membered ring] 8MR 226 channels running along the c axis. The structure is described 227 using the P6/mmm space group. In zeolite L, three sites, 228

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named KB, KC and KD, are occupied by cations (Gigli *et al.*,
2013). KB is placed at the centre of the cancrinite cage, KC at
the centre of the elliptical 8MR channel, while KD is close to
the wall of the 12MR channel.

In the present work zeolite L, purchased from the Tosoh Corporation (Japan) in its K form (from now on termed K-LTL), was initially characterized and used as starting material for the ion-exchange experiments.

2.2. Ion-exchange experiments

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2.2.1. Na⁺ and NH₄⁺ exchange experiments. With the 241 purpose of obtaining different cationic forms, K-LTL was ion-242 exchanged with two solutions, containing Na^+ and NH_4^+ ions. 243 The two solutions were prepared by dissolving NaNO₃ 244 (sodium nitrate 99.5% RPE-ACS, Carlo Erba) and NH₄Cl 245 246 (ammonium chloride 99.5% BAKER ANALYZED ACS, J. T. Baker) in double-distilled water, obtaining a 0.8 M concen-247 tration of sodium and ammonium. The high concentration of 248 the two solutions was chosen to obtain the maximum exchange 249 of the extra-framework cations of K-LTL.

The pristine sample was sieved so as to keep only particles 251 with a uniform mesh size (>2.5 μ m), dried for 24 h at 60°C, and then put in contact with the two solutions under stirring for 5 days; the solution was changed every day to favour the 254 ion-exchange process. The liquid/solid ratio was 16 ml/g, the temperature was set at 25°C and no buffer solution was used to control the pH to avoid interference in the exchange. Exchanged zeolites were rinsed three times with doubledistilled water using an ALC multi-speed centrifuge PK 121 at 5000 rmin^{-1} for 10 min and then dried for 24 h at 60°C. The 260 cationic forms obtained in this way are labelled Na-LTL and 261 262 NH₄-LTL. A second experiment was performed for sodium exchange at 80°C to evaluate the influence of temperature on 263 the ion-exchange efficiency. The obtained sample is labelled 264 Na-LTL-80°C. 265

2.2.2. Ce^{3+} exchange and recovery. A Ce solution 0.48 M 266 was prepared by dissolving Ce(NO₃)₃·6H₂O [cerium(III) 267 nitrate hexahydrate, REacton, 99.5% (REO), Alfa Aesar] in 268 double-distilled water. This concentration was chosen because 269 it allows the number of Ce ions in the solution to be equal to 270 three times the cation content of the zeolite, hence maximizing 271 the Ce incorporation. The pristine K-LTL and samples Na-272 273 LTL and NH₄-LTL were put in contact with the Ce solution (liquid/solid ratio = 20 ml/g) for 5 days at room temperature 274 with no buffer solution. The solution was changed every day to 275 favour the exchange. The samples were rinsed three times with double-distilled water using an ALC multi-speed centrifuge 277 PK 121 at 5000 r min⁻¹ for 10 min and then dried for 24 h at 60°C. The obtained zeolites are labelled K-Ce-LTL, Na-Ce-279 LTL and NH₄-Ce-LTL. For the last sample a further exchange, aimed at Ce recovery from the zeolite pores, was performed. 281 NH_4 -Ce-LTL was put in contact with a solution $[NH_4^+] =$ 0.8 M prepared by dissolving NH₄NO₃ (ammonium nitrate, 99%, Carlo Erba) in double-distilled water; it was left for 24 h 284 under stirring, at room temperature. The obtained zeolite, 285

labelled NH₄-Ce-LTL-rev, was then rinsed as previously reported.

2.3. Analytical methods

Thermogravimetric analysis. Thermogravimetric analyses (TGA) were performed in the temperature range 25–1050°C with a ramp of 10°C min⁻¹ under air flow, using a Seiko SSC/ 5200 thermal analyser. TGA were performed for all phases except Na-LTL-80°C, for which the water content was determined by loss on ignition (LOI).

Elemental analysis. Elemental analyses (EA) were performed for samples NH_4 -LTL and NH_4 -Ce-LTL to evaluate nitrogen content. A Flash 2000 CHNS/MAS200R instrument, equipped with a highly sensitive thermal conductivity (TCD) detector, was used. FLASH dynamic combustion (modified Dumas method) allows a temperature of 1800°C to be reached, in order to release elementary gases from the samples.

X-ray fluorescence analysis. X-ray fluorescence (XRF) analyses of samples K-LTL and Na-LTL were performed using a wavelength-dispersive X-ray fluorescence Philips PW1480. Samples were ground and pressed with boric acid to obtain powder pellets. Data were collected and corrected on the basis of standard materials (Leoni & Saitta, 1976; Franzini *et al.*, 1975) and assuming the water content determined by LOI obtained by heating the powders in a furnace at 1000°C or by TGA.

Scanning electron microscopy with energy-dispersive spectroscopy (SEM-EDS). The chemical composition of samples exchanged with Ce (*i.e.* K-Ce-LTL, Na-Ce-LTL and NH₄-Ce-LTL) and NH₄-LTL was checked using a scanning electron microscope Nova NanoSEM 450 equipped with an X-EDS Bruker QUANTAX-200 detector. Powders were compressed into thin self-supporting discs coated with gold. Chemical data were obtained by averaging the results obtained on five areas for each sample.

High-resolution X-ray powder diffraction (XRPD). Diffraction data were collected at ID22, the high-resolution beamline at the European Synchrotron Radiation Facility (ESRF), Grenoble, France, using the new Extremely Brilliant Source (EBS). The samples were ground and then used to fill borosilicate capillaries. Using a channel-cut Si(111) crystal monochromator the wavelength was set to be equal to 0.354496 Å; diffraction intensities were recorded by a bank of nine detectors, each preceded by a Si(111) crystal analyser.

Structure refinement. Structure refinements were performed using Rietveld methods through the GSAS package (Larson & von Dreele, 1994) with the EXPGUI interface (Toby, 2001). The structure reported by Hirano *et al.* (1992) was used as a starting model. The peak profile was fitted using the Thompson pseudo-Voigt function (Thompson *et al.*, 1987), while the background was refined using a Chebyshev polynomial function with 24 coefficients. Si—O distances were **refined using 'soft constraints' [ok as edited?]**, gradually decreasing the weight of the constraint (up to 10 [**unit?**]) after the initial stages. Extra-framework species, cations and H₂O

Table 1 343

Chemical formulas of the whole sample set. 344

345 The error is calculated as error% = $(Al-K-NH_4-3Ce)/(K+NH_4+3Ce) \times 100$ (Gottardi & Galli, 1985).

Name	Chemical analysis	Chemical formula	Error (%)
K-LTL	XRF. TGA	Ko 02 Nao 41 Size 70 Alo 14 O 72:16.91 H2O	-3.2
Na-LTL	XRF, TGA	$K_{600}Na_{297}Si_{2653}Al_{953}O_{72}\cdot19.51$ H ₂ O	3.2
Na-LTL-80°C	XRF, LOI	$K_{600}Na_{293}Si_{2675}Al_{930}O_{72}\cdot19.60$ H ₂ O	4.3
NH ₄ -LTL	SEM-EDS, TGA, EA	$K_{2,25}(NH_4)_{6,86}Si_{26,91}Al_{9,08}O_{72}$ ·17.07 H ₂ O	-0.3
K-Ce-LTL	SEM-EDS, TGA	$K_{5,26}Na_{0,21}Ce_{1,16}Si_{26,63}Al_{9,50}O_{72} \cdot 20.26 H_2O$	6.1
Na-Ce-LTL	SEM-EDS, TGA	K _{5.24} Na _{0.15} Ce _{1.26} Si _{26.55} Al _{9.55} O ₇₂ ·20.53 H ₂ O	4.2
NH ₄ -Ce-LTL	SEM-EDS, TGA, EA	K _{1.82} (NH ₄) _{3.92} Ce _{1.17} Si _{26.90} Al _{9.05} O ₇₂ ·20.94 H ₂ O	-2.2
NH ₄ -Ce-LTL-rev	SEM-EDS, TGA, EA	K _{1.87} (NH ₄) _{6.81} Ce _{0.09} Si _{27.07} Al _{8.90} O ₇₂ ·17.26 H ₂ O	-0.6

Table 2

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Unit-cell parameters.

Sample	a (Å)	c (Å)	$V(\text{\AA}^3)$
K-LTL	18.37294 (5)	7.52458 (3)	2199.74 (1)
Na-LTL	18.37075 (7)	7.52661 (4)	2199.80 (1)
NH₄-LTL	18.42200 (5)	7.54096 (3)	2216.31 (1)
K-Ce-LTL	18.34930 (5)	7.52420 (4)	2193.97 (1)
Na-Ce-LTL	18.34758 (5)	7.52461 (3)	2193.67 (1)
NH ₄ -Ce-LTL	18.38740 (5)	7.54285 (3)	2208.55 (1)

molecules were located in the zeolite pores by careful inspection of the Fourier difference map of the electronic density; the species were assigned considering the bond distances, the site-occupancy factors and the mutual exclusion rules for distances that are too short [ok as edited?].

3. Results

3.1. Chemical characterization

Fig. 2 shows the TGA for all the exchanged samples and the obtained water content is reported in Table S4 in the supporting information.

The first peak of all the DTG [please define] curves, positioned below 300°C, represents the water loss, while peaks at higher temperatures, observed only for samples NH₄-LTL and NH₄-Ce-LTL, are related to the ammonium content. In particular, the peak near 650°C in the NH₄-LTL DTG (Fig. 2) curve could derive from the condensation of residual H⁺ ions.

Table 1 reports the chemical formulas of the whole sample set, obtained considering: (i) the results obtained from XRF or SEM-EDS (Tables S1 and S2 in the supporting information); (ii) the NH₄ amount determined by elemental analysis (Table S3); (iii) the H₂O content (Table S4) determined by TGA or LOI analyses.

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The chemical compositions reported in Table 1 give a clear idea about the exchange capacity of the samples involved in this study. From the

comparison of the two Na-exchanged samples at different temperatures, it is possible to conclude that this variable is not critical in the improvement of the exchange effectiveness, as the two samples present the same final composition. For this reason, all the other experiments were performed at room temperature. Comparing Na-LTL and NH₄-LTL, we can deduce that the exchange is higher for NH₄ than for Na (2.25 and 6.00 potassium ions are not exchanged from the initial 9.03 in the two samples, respectively). Once the samples are exchanged with Ce³⁺ no clear affinity between Ce and one of the three cationic forms is observed. The three Ce^{3+} exchanged samples show, in fact, a similar amount of Ce (ranging from 1.16 to 1.26 atom p.u.c. [please define]), and also almost the same amount of H₂O molecules.

In order to recover Ce from zeolite pores, NH₄-Ce-LTL was further exchanged with a NH_4^+ solution. The amount of Ce^{3+} , obtained by EA, in the final NH₄-Ce-LTL-rev (0.09 p.u.c.) was compared with that of the NH₄-Ce-LTL (1.17 p.u.c.) proving the almost complete release of Ce and the restoration of the initial amount of ammonium as well as the H₂O molecules.

3.2. Structure analysis

Structure analysis was performed to determine the amount and position of the extra-framework species, in particular of Ce, in the zeolite **pores**, and to understand the guest-guest and host-guest interactions. Cell parameters are reported in Table 2; structural data, as well as refinement details and diffraction



TGA (solid line) and DTG (dotted line) curves for: K-LTL, Na-LTL and NH₄-LTL samples in black, red and green, respectively (left panel), and K-Ce-399 LTL, Na-Ce-LTL and NH₄-Ce-LTL samples in black, red and green, respectively (right panel).

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Figure 3

Zoomed-in view of observed (blue symbols) and calculated (red line) diffraction patterns and final difference curve (grey line) from Rietveld refinement of K-LTL and K-Ce-LTL.



Figure 4

Structure of K-LTL (left panel) and K-Ce-LTL (right panel). Cations are drawn as incomplete spheres as a function of their fractional occupancies. For the H_2O molecules, only the positions actually occupied are shown, rather than all the partially occupied symmetrically equivalent positions.

patterns are reported in Tables S5–S17 and in Figs. S1–S6 in the supporting information. For selected samples (K-LTL and K-Ce-LTL), observed and calculated diffraction patterns

research papers

obtained by Rietveld refinement are reported in Fig. 3. 3.2.1. Na⁺ and NH₄⁺ exchange. K-LTL and Na-LTL show very similar unit-cell parameters (Table 2), while we can observe slightly higher values for NH₄-LTL. Apart from this, the Ce-free cationic forms of zeolite L display similar structures (Tables S6-S11). No new extra-framework sites were located compared with those reported in the literature (see Section 2.1); all the cations are sited in the KB (at the centre of the cancrinite cage), KC (at the centre of the elliptical 8MR channel) and KD (close to the wall of the 12MR channel) sites [see the structure of K-LTL (Fig. 4, left panel)]. KB is bonded to the framework oxygen atoms O3 with six distances of about 2.9 Å and KC is placed at about 2.9 Å from four framework oxygen atoms O5 and at a larger distance from eight O4 atoms. Cations in KD, instead, display a solvation sphere constituted by W5 and W3 H₂O molecules and bonds with the framework oxygen atoms O4 and O6 (Fig. 4, left panel).

 H_2O molecules are all placed in the main channel; they interact with each other, forming a cluster (Fig. 4, left panel). Two W4, at the centre of the 12MR channel, are at bonding distance and are also bonded to W1, W2, W3. Site W5, almost fully occupied in all samples, is located in the external part of the H_2O cluster at bonding distance with KD, W3 and W6. The only interaction between the framework oxygen atoms and H_2O molecules is observed for W2 with O1. In Table 3, the number of H_2O molecules in the six sites W1–W6 is reported for all the samples.

3.2.2. Ce^{3+} exchange. All the zeolites exchanged with Ce show a reduction of cell volume (due to the decrease of the *a* parameter) if compared with their Ce-free counterparts (Table 2). However, as already observed for the three Ce-free zeolites, NH₄-Ce-LTL shows a higher volume. When K, Na and NH₄ cations are exchanged with Ce, similar structural modifications occur in the three samples. All the patterns collected for Ce-exchanged samples show changes in the

relative peak intensities compared with their Ce-free counterparts, especially in the low-angle region, clearly indicating different cation contents (see as an example NH₄-LTL and NH₄-Ce-LTL patterns in Fig. 5).

In the Ce-exchanged forms, in addition to KB, KC and KD cation sites, a new one appears in the middle of the 12MR channel (labelled Ce). In the Ceexchanged samples the occupancy of KD and Ce sites was refined using the Ce scattering factor. The Ce site-occupancy factor ranges from 88 (K-Ce-LTL) to 97% (NH₄-Ce-LTL), indicating that this latter site is filled almost exclusively with Ce in all the exchanged zeolites (Fig. 4, right panel). This new site coordinates with W2 and W3. The

Table 3 571

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 H_2O p.u.c. in each W site for the whole sample set. 572

573 The low amount of H₂O found by XRPD in Ce-exchanged zeolites is explained in Section 4. 574

		K-LTL	Na-LTL	NH ₄ -LTL	K-Ce-LTL	Na-Ce-LTL	NH ₄ -Ce-LTL
W	1	2.0	2.0	2.0			
W	2	1.0	1.0	1.0	3.0	3.0	3.0
W	3	6.0	6.0	6.0	6.0	6.0	6.0
W^2	1	1.2	0.3	0.6			
W_{2}^{*}	5	6.6	6.0	5.7	6.0	6.0	6.0
We	5	1.8	2.5	1.6	1.0	1.0	1.0
То	tal	18.6	17.8	16.9	16	16	16

occupancy of site KD is about 20-22% for all the Ceexchanged samples; thus it is impossible to exclude the presence of various cations in the same position. In the Ceexchanged samples, H2O molecules are positioned only in four sites W2, W3, W5 and W6 (Tables 3 and S12, S14, S16), creating a cluster that interacts slightly with the framework and bonding Ce site.

4. Discussion

4.1. Na⁺ and NH₄⁺ exchange

The K, Na and NH₄ zeolites show similar structures, but the occupancy factors of sites KB, KC and KD are different due to the different types and amount of exchanged extra-framework cations. Combining the information obtained by XRPD and chemical analysis (Table 4), we can assume the cation distribution in the three sites reported in Table 5. Since it is not possible to distinguish between different cations occupying the same site from the XRPD data, the comparison between XRPD and chemical data reported in Table 4 is only based on the number of electrons.

K-LTL presents a certain amount of Na (0.41 p.u.c.) which remains in part in all the exchanged samples, with the



XRPD patterns of NH₄-LTL and NH₄-Ce-LTL.

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exception of the NH_4 -exchanged one (see Table 1) where NH_4 628 is able to remove all the Na. Since the KB site is always 629 occupied by non-exchangeable K, these Na ions should be 630 distributed in KC and KD sites. Na-LTL zeolite shows an 631 exchange Na⁺ \rightarrow K⁺ of about 27%, according to Sato *et al.* 632 (1990), so we hypothesize a prevalent substitution of K ions in 633 the main channel (KD), while the small amount of Na⁺ in KC 634 is that already present in the pristine K-LTL. In sample NH₄-635 LTL, K⁺ ions are found mainly in the KB site and partially in 636 KC, with an exchange $NH_4^+ \rightarrow K^+$ of about 73%, in very good 637 agreement with the results reported by Dyer et al. (1993). The 638 higher value of this last exchange seems related to the ionic 639 radii of the species involved (Dyer et al., 1993). Indeed, NH₄⁺ 640 and K⁺ show a more similar size when compared with the 641 couple Na⁺-K⁺; thus NH₄⁺ can enter in site KD without 642 affecting the framework stability. The presence of NH_4^+ ions 643 both in the 8MR and 12MR channels is also demonstrated by 644 the broad peak (between 290 and 450°C) in DTG data related 645 to the NH_4^+ loss (Fig. 2). Indeed, the two sites occupied by this 646 cation present different interactions with the framework and 647 extra-framework species and hence exhibit different 648 temperatures of release. In addition, in NH₄-LTL, we can 649 notice an enlargement of the cell volume, and that site KD is positioned farther from the framework in comparison with the 651 other samples (*i.e.* KD-O4 = 3.18 and 3.20 Å in K-LTL and 652 Na-LTL, respectively?, and 3.25 in NH_4 -LTL; KD-O6 = 2.98653 and 2.99 Å in K-LTL and Na-LTL, respectively?, and 3.11 in 654 NH₄-LTL). This can be a direct consequence of the slightly 655 larger ionic radius of NH⁴⁺ (1.40 Å) (Sidey, 2016) compared 656 with K⁺ (1.37 Å) (Shannon, 1976) and Na⁺ (0.99 Å) (Shannon, 657 1976) and/or due to the hydrogen bonds occurring between 658 the framework oxygen atoms and this cation. 659

Regarding H₂O molecules, in NH₄-LTL the shift towards lower temperature of the first peak of the DTG curve (Fig. 2), related to H₂O loss, can be caused by longer bonds between W2 and the framework [W2-O1 is 2.65 (6) Å in K-LTL,2.61 (7) Å in Na-LTL and 2.81 (6) Å in NH_4 -LTL]. In general, for all the samples, the total amount of cations and H₂O



Figure 6

Comparison of XRPD patterns of NH₄-LTL (red line) and NH₄-Ce-LTLrev, after Ce recovery (black line) samples.

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Table 4

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Number of electrons (e) for cations (cat.) and H_2O molecules derived from structural (XRPD) (number of atoms $\times Z$ depending on the scattering factor used for the occupancy factor refinement of each cation site) and chemical analyses (number of atoms from the chemical formula \times Z).

The differences between the total number of electrons for cations and H₂O molecules obtained by the two techniques are also reported (e cat. XRPD – e cat. chemical analysis; e H₂O XRPD - e H₂O chemical analysis).

	XRPD						Chemica	al analysis	8					
	e KB	e KC	e KD	e Ce	e cat.	e H ₂ O	e K	e Na	e NH ₄	e Ce	e cat.	e H ₂ O	e Dif. cat.	e Dif. H ₂ O
K-LTL	36.00	51.30	89.64		176.94	148.43	162.54	4.10			166.64	135.28	10.30	13.14
Na-LTL	36.00	50.76	69.12		155.88	142.88	108.00	29.70			137.70	156.00	18.18	-13.12
NH ₄₋ LTL	36.00	28.08	57.24		121.32	135.02	40.50		68.60		109.10	136.56	12.22	-1.53
K-Ce-LTL	36.00	50.22	75.90	48.51	210.63	127.99	94.68	2.10		63.80	160.58	162.08	50.05	-34.08
Na-Ce-LTL	36.00	50.76	75.90	53.90	216.56	128.00	94.32	1.50		69.30	165.12	164.24	51.44	-36.24
NH₄-Ce-LTL	34.92	28.62	66.00	52.80	182.34	127.97	32.76		39.20	64.35	136.31	167.52	46.03	-39.55

Table 5

Cation distribution over the extra-framework sites deduced from structural and chemical results.

		K		Na		NH_4		Ce	
		e	Atom	e	Atom	e	Atom	e	Ator
K-LTL	KB	36.00	2.00						
	KC	51.90	2.88	2.10	0.21				
	KD	74.64	4.15	2.00	0.20				
Na-LTL	KB	36.00	2.00						
	KC	52.50	2.92	1.50	0.15				
	KD	19.50	1.08	28.2	2.82				
NH_4 -LTL	KB	36.00	2.00						
	KC	4.50	0.25			27.50	2.75		
	KD					41.10	4.11		
K-Ce-LTL	KB	36.00	2.00						
	KC	48.12	2.67	2.10	0.21				
	KD	10.56	0.59					15.29	0.28
	Ce	26.00	2 00					48.51	0.88
Na-Ce-LTL	KB	36.00	2.00	1.50	0.15				
	KC	49.26	2.74	1.50	0.15			15 40	0.20
	KD Ca	9.06	0.50					15.40	0.28
NIL CoLTI	VD VD	22.76	1.07					35.90	0.96
NH ₄ -Ce-LIL	KD VC	52.70	1.82			20 62	286		
	KC KD					20.02	2.00	11 55	0.21
	Ce					10.30	1.00	52.80	0.21
	œ							52.80	0.90

molecules obtained by structural refinement is in good agreement with those values resulting from chemical analysis (Table 4). In sample Na-LTL, the number of cations from structure refinement is slightly overestimated, but this could be due to the simultaneous presence of cations and H₂O molecules on the KD site. This is consistent with the finding that the number of H₂O molecules obtained is underestimated with respect to the TGA data. The fact that KD can be shared by the different species (also the case in the other samples) is also suggested by the high value of its thermal parameter.

4.2. Ce³⁺ exchange and recovery

All Ce-exchanged zeolites show a similar amount of Ce 734 p.u.c. (1.1-1.2 atoms p.u.c.) (Table 1) (exchange degree from 735 38% to 42%) and similar cation distribution. On the basis of 736 the cation distribution reported in Table 5, Ce mainly occupies 737 the site Ce and, to a lesser extent, site KD. From Table 4, 738 739 structural analysis always shows an overestimation of the cations, accompanied by an underestimation of the number of 740 H₂O molecules. Thus, we can assume that the cation over-741

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estimation is due to the co-presence of H₂O molecules in sites Ce and KD; these H₂O molecules [ok as edited?] have migrated from the original sites W1 and W4, which are empty in these samples. This hypothesis is confirmed by the high thermal parameters of these two sites, indicating a high degree of structural disorder. In comparison with the pristine materials, Ce-exchanged zeolites show a higher amount of H₂O (Table 1), according to: (i) the bigger solvation sphere of this cation [solvation spheres: $Ce^{3+} = 9.1$ (Lutz *et al.*, 2012), K⁺ = 7.5 (Duignan *et al.*, 2020), NH₄⁺ = 5.8(Chang & Dang, 2003), $Na^+ = 5.6 H_2O$ molecules (Duignan et al., 2020)]; (ii) the larger space available due to the lower amount of trivalent cations present in the main channel. The H₂O content is similar for the three samples, due to the same amount of Ce incorporated, as well as the temperature of its release (see the first two peaks of the

DTG, Fig. 2). It is interesting to observe that these temperatures appear higher than in pristine materials, due to the strong and short bonds occurring between the Ce site and W2 and W3 (respectively, about 2.3 and 2.6 Å). Ce penetration, indeed, leads to a reorganization of the position of the H₂O molecules and their bonds with this element have a different coordination sphere with respect to the original K⁺, Na⁺ and NH₄⁺ cations. This, beyond the intrinsic properties of zeolite L, can play an important role in the exchange capacity.

The complete recovery of cerium from the NH₄-Ce-LTL was achieved through a further exchange with a NH₄ solution. Chemical results demonstrated complete recovery of the initial amount of ammonium and water molecule in the zeolite with an almost complete release of cerium. As further demonstration, the two XRPD patterns, reported in Fig. 6, of the initial NH₄-LTL and the NH₄-Ce-LTL-rev after cerium recovery show the same features, suggesting the same structure and the same type and number of cations.

799 5. Conclusions

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The aim of this work was the evaluation of the Ce-exchange capacity of three cationic forms of LTL zeolites (the assynthesized K form and Na- and NH₄- exchanged ones) using a highly concentrated solution of Ce. The effect of the Ce exchange in the different cationic forms was evaluated through chemical analyses and XRPD structural investigations. Indeed, structural studies can give new insight into the sites involved in the exchange, and this is fundamental for the choice of a powerful protocol of exchange for future application such as Ce recovery from diluted solution. The results reported here also have important implications for catalytic applications, since many zeolite catalysts are prepared through Ce exchange.

813 The obtained results show a $Na^+ \rightarrow K^+$ substitution of 814 about 27%, while $NH_4^+ \rightarrow K^+$ reached 73%, because the 815 cations in the extra-framework site KC - in the 8MR channel -816 are exchanged only if K^+ is replaced by NH_4^+ ions due to their 817 similar ionic size. The cation-exchange degree for Ce ranges 818 from 38 to 42% and the LTL cation-exchange capacity for Ce 819 is not influenced by the nature of the cations in the pristine sample. In contrast to what was observed for Na and NH₄ 821 exchange, in which the cations occupy the pre-existing 822 potassium positions, the Ce cations occupy a new site in the 823 12MR site and induce a reorganization of the H₂O molecules. 824 This may be due to the higher ionic potential of Ce atoms, 825 which tend to form a stronger solvation sphere that hinders 826 cation mobility. The Ce-exchange capacity of zeolite L and its 827 affinity with this element can thus be driven not only by the 828 intrinsic property of the LTL framework, but also by the coordination sphere of Ce compared with the original K⁺, Na⁺ 830 and NH₄⁺ cations. With the aim of possible REE recovery 831 from zeolite pores, the Ce-exchanged NH₄-LTL was tested 832 through exchange with a NH_4^+ solution, leading to the 100% 833 recovery of this element. The results presented here are 834 promising for future Ce-recovery experiments from very 835 diluted solutions. 836

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References

- Arletti, R., Fois, E., Gigli, L., Vezzalini, G., Quartieri, S. & Tabacchi, G. (2017). Angew. Chem. Int. Ed. 56, 2105–2109.
- Arletti, R., Giacobbe, C., Quartieri, S. & Vezzalini, G. (2017). Minerals, 7, 18.
- Baerlocher, C., McCusker, L. B. & Olson, D. H. (2007). Atlas of Zeolite Framework Types, 6th ed. Amsterdam, The Netherlands: Elsevier.
- Balaram, V. (2019). Geosci. Front. 10, 1285–1303.
- Barrer, R. M. & Meier, W. M. (1958). J. Chem. Soc. p. 299.
- Barros, O., Costa, L., Costa, F., Lago, A., Rocha, V., Vipotnik, Z.,
 Silva, B. & Tavares, T. (2019). *Molecules*, 24, 1005.
- 855 Bennett, J. M. & Smith, J. V. (1968). *Mater. Res. Bull.* 3, 633–642.

Files: j/vb5024/vb5024.3d j/vb5024/vb5024.sgml **VB5024** FA IU-2110/43(11)11 2113/42(8)11 ()

Binnemans, K., Jones, P. T., Müller, T. & Yurramendi, L. (2018). J. Sustain. Met. 4, 126–146.

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Boekfa, B., Pantu, P. & Limtrakul, J. (2008). J. Mol. Struct. 889, 81-88.

- Boer, M. A. de & Lammertsma, K. (2013). Chemsuschem, 6, 2045-
- 2055. Bryukhanov, I. A., Rybakov, A. A., Larin, A. V., Trubnikov, D. N. &
- Vercauteren, D. P. (2017). J. Mol. Model. 23, 68. Chakhmouradian, A. R. & Wall, F. (2012). Elements, 8, 333–340.
- Chang, T. M. & Dang, L. X. (2003). J. Chem. Phys. 118, 8813–8820.
- Charalampides, G., Vatalis, K., Karayannis, V. & Baklavaridis, A. (2016). *IOP Conf. Ser.: Mater. Sci. Eng.* **161**, 012069.
- Charalampides, G., Vatalis, K. I., Apostoplos, B. & Ploutarch-Nikolas, B. (2015). *Procedia Econ. Financ.* **24**, 126–135.
- Confalonieri, G., Fabbiani, M., Arletti, R., Quartieri, S., Di Renzo, F., Haines, J., Tabacchi, G., Fois, E., Vezzalini, G., Martra, G. & Santoro, M. (2020). *Microporous Mesoporous Mater.* **300**, 110163.
- Confalonieri, G., Grand, J., Arletti, R., Barrier, N. & Mintova, S. (2020). *Microporous Mesoporous Mater.* **306**, 110394.
- Confalonieri, G., Quartieri, S., Vezzalini, G., Tabacchi, G., Fois, E., Daou, T. J. & Arletti, R. (2019). *Microporous Mesoporous Mater*. 284, 161–169.
- Confalonieri, G., Ryzhikov, A., Arletti, R., Nouali, H., Quartieri, S., Daou, T. J. & Patarin, J. (2018). *J. Phys. Chem. C*, **122**, 28001–28012.
- Confalonieri, G., Ryzhikov, A., Arletti, R., Quartieri, S., Vezzalini, G., Isaac, C., Paillaud, J. L., Nouali, H. & Daou, T. J. (2020). *Phys. Chem. Chem. Phys.* **22**, 5178–5187.
- Coudert, F. X., Cailliez, F., Vuilleumier, R., Fuchs, A. H. & Boutin, A. (2009). *Faraday Discuss.* **141**, 377–398.
- Dejoie, C., Martinetto, P., Tamura, N., Kunz, M., Porcher, F., Bordat, P., Brown, R., Dooryhée, E., Anne, M. & McCusker, L. B. (2014). *J. Phys. Chem. C*, **118**, 28032–28042.
- Deshmane, V. G., Islam, S. Z. & Bhave, R. R. (2020). Environ. Sci. Technol. 54, 550–558.
- Diaz, L. A., Lister, T. E., Parkman, J. A. & Clark, G. G. (2016). J. Clean. Prod. 125, 236–244.
- Duignan, T. T., Schenter, G. K., Fulton, J. L., Huthwelker, T., Balasubramanian, M., Galib, M., Baer, M. D., Wilhelm, J., Hutter, J., Del Ben, M., Zhao, X. S. & Mundy, C. J. (2020). *Phys. Chem. Chem. Phys.* 22, 10641–10652.
- Duplouy, L. (2016). PhD thesis, University of Helsinki, Université de Lille.
- Dyer, A. (2007). Studies in Surface Science and Catalysis, Vol. 168, edited by J. Čejka, H. van Bekkum, A. Corma & F. Schüth, pp. 525– 553. <u>City of publication?</u>: Elsevier.
- Dyer, A., Amini, S., Enamy, H., El-Naggar, H. A. & Anderson, M. W. (1993). *Zeolites*, **13**, 281–290.
- Fabbiani, M., Confalonieri, G., Morandi, S., Arletti, R., Quartieri, S., Santoro, M., Di Renzo, F., Haines, J., Fantini, R., Tabacchi, G., Fois, E., Vezzalini, G., Ricchiardi, G. & Martra, G. (2021). *Microporous Mesoporous Mater.* **311**, 110728.
- Faghihian, H., Amini, M. K. & Nezamzadeh, A. R. (2005). J. Radioanal. Nucl. Chem. 264, 577–582.
- Franzini, M., Leoni, L. & Saitta, M. (1975). Rendiconti Societa' Italiana di Mineralogia e Petrologia, XXXI, 365–378.
- Frising, T. & Leflaive, P. (2008). *Microporous Mesoporous Mater.* **114**, 27–63.
- Gad, H. M. H. & Awwad, N. S. (2007). Sep. Sci. Technol. 42, 3657–3680.
- Gautam, P., Barman, S. & Ali, A. (2020). Int. J. Chem. React. Eng. 18, 20200081.
- Gigli, L., Arletti, R., Quartieri, S., Di Renzo, F. & Vezzalini, G. (2013). *Microporous Mesoporous Mater.* **177**, 8–16.
- Gottardi, G. & Galli, E. (1985). *Natural Zeolites*. Berlin, Heidelberg: Springer-Verlag .
- Haque, N., Hughes, A., Lim, S. & Vernon, C. (2014). *Resources*, **3**, 614–635.
- Hirano, M., Kato, M., Asada, E., Tsutsumi, K. & Shiraishi, A. (1992). *Adv. X-ray Chem. Anal. Jpn*, **23**, 101–110.

35024 PROOFS J:FA:2022:53

- 913 Hong, S. H., Jang, M. S., Cho, S. J. & Ahn, W. S. (2014). Chem.
 914 Commun. 50, 4927–4930.
- Hudson, M. R., Queen, W. L., Mason, J. A., Fickel, D. W., Lobo, R. F. & Brown, C. M. (2012). J. Am. Chem. Soc. 134, 1970–1973.
- Isaac, C., Confalonieri, G., Nouali, H., Paillaud, J. L., Arletti, R.,
 Daou, T. J. & Ryzhikov, A. (2020). *Microporous Mesoporous Mater.* 298, 110047.
- 919 Jowitt, S. M., Werner, T. T., Weng, Z. H. & Mudd, G. M. (2018). Curr. Opin. Green Sustain. Chem. 13, 1–7.
- Kavun, V., van der Veen, M. A. & Repo, E. (2021). Microporous Mesoporous Mater. 312, 110747.
- Weight Kooten, W. E. J. van, Calis, H. P. A. & van den Bleek, C. M. (1998).
 Catal. Automot. Pollut. Contr. IV, **116**, 357–366.
- 924 Krohn, J. E. & Tsapatsis, M. (2005). Langmuir, 21, 8743–8750.
- Larson, A. C. & Von Dreele, R. B. (1994). Reference details?
- Leoni, L. & Saitta, M. (1976). *Rendiconti Societa' Italiana di* Mineralogia e Petrologia, **32**, 497–510.
- Lu, C. Y., Bai, H. L., Wu, B. L., Su, F. S. & Hwang, J. F. (2008). Energy
 Fuels, 22, 3050–3056.
- Lutz, O. M. D., Hofer, T. S., Randolf, B. R. & Rode, B. M. (2012).
 Chem. Phys. Lett. 539–540, 50–53.
 - Manos, M. J. & Kanatzidis, M. G. (2016). Chem. Sci. 7, 4804–4824.
- ⁹³¹ Massari, S. & Ruberti, M. (2013). *Resour. Policy*, **38**, 36–43.
- Meshram, P. & Abhilash (2020). *Miner. Process. Extr. Metall. Rev.* 41, 279–310.
- Mosai, A. K., Chimuka, L., Cukrowska, E. M., Kotzé, I. A. & Tutu, H. (2019). Water Air Soil Pollut. 230, 188.
- Ni'Am, A. C., Wang, Y. F., Chen, S. W., Chang, G. M. & You, S. J. (2020). Chem. Eng. Process. 148, 107831.
- Pabalan, R. T. & Bertetti, F. P. (2001). *Rev. Mineral. Geochem.* 45, 453–518.

- Porvali, A., Wilson, B. P. & Lundström, M. (2018). *Waste Manage.* (*Oxford*), **71**, 381–389.
- Rademaker, J. H., Kleijn, R. & Yang, Y. X. (2013). *Environ. Sci. Technol.* **47**, 10129–10136.
- Royen, H. & Fortkamp, U. (2016). Report of IVL Swedish Environmental Research Institute 2016.
- Saada, M. A., Soulard, M., Marler, B., Gies, H. & Patarin, J. (2011). *J. Phys. Chem. C*, **115**, 425–430.
- Santoro, M., Ciabini, L., Bini, R. & Schettino, V. (2003). J. Raman Spectrosc. 34, 557–566.
- Santoro, M., Scelta, D., Dziubek, K., Ceppatelli, M., Gorelli, F. A., Bini, R., Garbarino, G., Thibaud, J. M., Di Renzo, F., Cambon, O., Hermet, P., Rouquette, J., van der Lee, A. & Haines, J. (2016). *Chem. Mater.* 28, 4065–4071.
- Sato, M., Morikawa, K. & Kurosawa, S. (1990). *Eur. J. Mineral.* **2**, 851–860.
- Scelta, D., Ceppatelli, M., Santoro, M., Bini, R., Gorelli, F. A., Perucchi, A., Mezouar, M., van der Lee, A. & Haines, J. (2014). *Chem. Mater.* 26, 2249–2255.
- Shannon, R. D. (1976). Acta Cryst. A32, 751-767.
- Sidey, V. (2016). Acta Cryst. B72, 626-633.
- Simoncic, P. & Armbruster, T. (2004). Am. Mineral. 89, 421-431.
- Stückenschneider, K., Merz, J., Hanke, F., Rozyczko, P., Milman, V. & Schembecker, G. (2013). J. Phys. Chem. C, 117, 18927–18935.
- Thompson, P., Cox, D. E. & Hastings, J. B. (1987). J. Appl. Cryst. 20, 79–83.
- Toby, B. H. (2001). J. Appl. Cryst. 34, 210-213.
- Zhang, K., Lively, R. P., Noel, J. D., Dose, M. E., McCool, B. A., Chance, R. R. & Koros, W. J. (2012). *Langmuir*, **28**, 8664–8673.
- Zhou, H. Y., Wang, Y. L., Guo, X. G., Dong, Y. M., Su, X. & Sun, X. Q. (2018). *J. Mol. Liq.* **254**, 414–420.

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