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2	<b>Rb</b> <sup>+</sup> , and <b>Cs</b> <sup>+</sup> as extra-framework cations
3	by
4	Mihye Kong, Yongmoon Lee, G. Diego Gatta, Yongjae Lee
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6 7	Corresponding author: <b>Yongjae Lee</b> , Department of Earth System sciences, Yonsei University, Seoul 03722, Republic of Korea - E-Mail: yongjaelee@yonsei.ac.kr
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25	Mihye Kong <sup>1</sup> , Yongmoon Lee <sup>2</sup> , G. Diego Gatta <sup>3</sup> , Yongjae Lee <sup>1, 2, *</sup>
26	
	Department of Forth System sciences, Vensei University, Second 02722, Depublic of Keree
27	<sup>1</sup> Department of Earth System sciences, Yonsei University, Seoul 03722, Republic of Korea
28	<sup>2</sup> Center for High Pressure Science and Technology Advanced Research, Shanghai 201203,
29	China
30	<sup>3</sup> Dipartimento di Scienze della Terra, Università degli Studi di Milano, Via Botticelli 23, I-
31	20133 Milano, Italy
32	
33	Abstract
34	The high-pressure behavior of monovalent-cation-exchanged chabazites was
35	investigated by means of <i>in-situ</i> synchrotron X-ray powder diffraction with a diamond anvil
36	cell, and using water as penetrating pressure-transmitting medium, up to 5.5 GPa at room
37	temperature. In all cases, except for Na-containing chabazites, a phase transition from the
38	original rhombohedral $(R\overline{3}m)$ to triclinic symmetry (likely $P\overline{1}$ ) was observed in the range
39	between 3.0 GPa and 5.0 GPa. The phase transition is accompanied by an abrupt decrease of
40	the unit-cell volume by up to 10 %. Evidence of pressure-induced hydration (PIH), i.e., P-
41	induced penetration of H2O molecules through the zeolitic cavities, was observed, as
42	reflected by the incompressibility of the cation-exchanged chabazites, which is governed by
43	the distribution of the extra-framework cations. The reversibility of the PIH and P-induced
44	phase transitions in the high-pressure behavior of the cation-exchanged chabazites are
45	discussed in the context of the role played by the chemical nature and bonding configuration

of the extra-framework cations, along with that of the H<sub>2</sub>O content at room conditions.

- 48 Keywords: Chabazite, compressibility, high pressure, pressure-induced hydration,
- 49 synchrotron diffraction

## Introduction

52 There is a growing interest in understanding the behavior of microporous materials at non-ambient conditions and, in particular, at high pressure (e.g., Bish and Carey 2001; Alberti 53 and Martucci 2005; Cruciani 2006; Gatta and Lee 2014; Gatta et al. 2017 and references 54 55 therein). Pressure can cause important structural changes in microporous materials, 56 modifying their physical-chemical properties and hence affecting their potential technological utilizations. Pressure-induced hydration (PIH) or pressure-induced insertion (PII), i.e., P-57 58 induced penetration of external molecules through the zeolitic sub-nanocavities at moderate pressure ( $\leq 1$  GPa), is one of the most fascinating discovery in material science over the last 59 60 decade, with potential technological and geological implications, recently reviewed by Gatta et al. (2017), promoting new routes for creating hybrid host-guest composite materials or for 61 understanding the stability of clathrates or the role played by zeolites as carrier of H<sub>2</sub>O or 62 63 CO<sub>2</sub> in subduction zones (e.g., Lee et al. 2011; Seoung et al. 2013; Seoung et al. 2014; Seoung et al. 2015; Im et al. 2015). Framework topology and extra-framework content are the 64 65 key factors that govern the structural deformations at high pressure (e.g., Gatta et al. 2005; Gatta 2010; Danisi et al. 2015). Previous studies showed that the pressure-induced 66 67 deformation of the tetrahedral framework in zeolites can be described in terms of tilting of quasi-rigid tetrahedra (e.g., Gatta 2008; 2010; Gatta and Lee 2014). There has not, however, 68 been any systematic study on how the framework distortion in response to the applied 69 70 pressure is influenced by the nature and distribution of the extra-framework cations. Only the "fibrous zeolites group", which was extensively investigated at high pressure, provided a 71 72 preliminary model to describe the effect of the extra-framework population on the elastic behavior of isotypic materials (e.g., Gatta 2005; Gatta et al. 2005; Seoung et al. 2013; Seoung 73 et al. 2015). 74

75 Chabazite (ideally  $|(Ca_{0.5},K,Na)_x(H_2O)_{12}|[Al_xSi_{12-x}O_{24}]$ , with x = 2.4 - 5.0, 76 http://www.iza-online.org/natural/Datasheets/Chabazite/chabazite.htm) is one of the most 77 widespread natural zeolites with excellent ion-exchange properties (*e.g.*, Barrer et al. 1969;

Shang et al. 2012). Its framework is built up by double 6-membered rings (D6R), stacked in 78 an ABC sequence and linked together through single 4-membered rings (S4R) (e.g., 79 Calligaris et al. 1982; Zema et al. 2008). As a result, the framework contains large ellipsoidal 80 cavities (*i.e.*, the CHA cage) with apertures of about  $6.7 \times 10$  Å, which are accessible through 81 single 8-rings (S8R) (Breck 1974). The largest opening of the S8R has a dimension of  $3.8 \times$ 82 3.8 Å and is located in the direction normal to the (001) crystal plane (Smith et al. 2001; 83 Shang et al. 2012). Chabazites crystallizes with rhombohedral symmetry (space group  $R\overline{3}m$ ), 84 with only one independent tetrahedral framework site, populated by Al and Si with a 85 statistically disordered distribution (Dent and Smith 1958). Exchangeable extra-framework 86 cations and H<sub>2</sub>O molecules are distributed over the D6R, S8R, and CHA cages with various 87 occupancies (e.g., Fialips et al. 2005). A recent structural study of our group on various 88 monovalent cation-exchanged chabazites revealed the systematic interplay between the 89 framework and the extra-framework cations, i.e., the unit-cell volume of monovalent-cation-90 exchanged chabazites varies in response to the ion selectivity, in the order of  $Cs^+ \ge K^+ > Ag^+$ 91  $> Rb^+ > Na^+ > Li^+$  (Kong et al. 2016). 92

93 The aim of this study is the description of the comparative compressional behavior of these monovalent cation-exchanged chabazites (Kong et al. 2016) and the potential crystal-94 fluid interactions in response to the applied hydrostatic pressure. We have performed in-situ 95 high-pressure (at room temperature) synchrotron X-ray powder diffraction experiments on 96 97 Li-, Na-, Ag-, K-, Rb-, and Cs-exchanged chabazites, using a diamond-anvil cell and pure water as a nominally pore-penetrating pressure-transmitting medium, in order to emulate the 98 same conditions generated in industrial processes, or occurring in nature, in which water is 99 the dominant P-fluid. 100

101

#### 102 **Experimental methods**

103

A natural chabazite (hereafter ORI-CHA, Ca1.6Na0.5Si8.4Al3.6O24·14.3H2O, space

group  $R\overline{3}m$ , a = 9.405(5) Å,  $\alpha = 94.22(2)^{\circ}$ ) from Rubendorfel, Bohemia, was used in this 104 study. Cation exchange was performed by stirring a mixture of ground ORI-CHA and the 105 106 respective nitrate solution of Li, Na, Ag, K, Rb, and Cs, in a 1:100 weight ratio, in a closed system at 80°C for 72h. The final product was filtered, washed with distilled water, and air-107 dried. Elemental analysis (by X-ray fluorescence with energy-dispersive system detector) 108 revealed that a complete ion-exchange was achieved, with the respective aforementioned 109 cations. Further details pertaining to the ion-exchange protocols and cristallochemical 110 characterization of the natural and final products are reported by Kong et al. (2016). 111

In-situ high-pressure (HP) synchrotron X-ray powder diffraction experiments on the 112 113 as-prepared cation-forms of chabazites were performed at beamline 10-2 at the Stanford Synchrotron Radiation Lightsource (SSRL) at the SLAC National Accelerator Laboratory. 114 At the beamline 10-2, the synchrotron radiation from the wiggler insertion device impinges 115 on a Si(111) crystal followed by two pinholes in order to generate an approximately 200 µm 116 diameter beam of monochromatic X-rays with a wavelength of 0.61992(5) Å. A Pilatus 117 118 300K-w Si-diode CMOS detector, manufactured by DECTRIS, was used to collect the powder diffraction data. The detector, held at a distance of 1032(2) mm from the sample, was 119 stepped to produce scattering angle coverage in  $2\theta$  up to ca.  $40^{\circ}$ . The position of the incident 120 beam, sample to detector distance, and detector tilt were determined using LaB<sub>6</sub> (SRM 660) 121 as a standard polycrystalline material. 122

123 A modified Merrill-Bassett diamond anvil cell (DAC), with two opposing diamonds supported by tungsten-carbide plates, was used for the high-pressure X-ray diffraction 124 measurements. A stainless-steel foil of 250 µm thickness was pre-indented to a thickness of 125 about 100 µm, and a 300 µm hole was obtained by electro-spark erosion. The powdered 126 samples of Li-, Na-, Ag-, K-, Rb-, and Cs-exchanged chabazites were placed in the gasket 127 hole together with a few ruby chips (~20 µm in diameter) for pressure measurements by the 128 ruby-fluorescence method (following the protocol of Mao et al. 1986; error:  $\pm 0.05$  GPa). 129 Ambient pressure data were collected first on the dry zeolite powder sample inside the DAC. 130

Subsequently, pure water was added into the gasket hole as a (hydrostatic, at  $P \le 1$  GPa) P-131 transmitting medium (PTM), and the second ambient pressure data were collected using the 132 'wet' sample. The pressure was then increased and, at any pressure point, the sample was 133 equilibrated for about 10 minutes before collecting the X-ray diffraction data. Water 134 transforms to a solid phase at  $P \ge 1$  GPa (and room temperature), and the diffraction peaks of 135 ice VI and VII were observed at pressure in excess of 1 GPa. The experiments were 136 deliberately performed under non-hydrostatic conditions at P > 1 GPa, in order to emulate the 137 conditions of natural or industrial processes. 138

Pressure-dependent changes of the unit-cell lengths and volumes were derived from a series of Le Bail profile fittings (Le Bail et al. 1988) using the GSAS-EXPGUI suite of programs (Larson and Von Dreele 2004; Toby 2001). The background was fitted with a Chebyshev polynomial (with  $\leq$  24 coefficients), and the pseudo-Voigt profile function of Thompson et al. (1987) was used to model the Bragg peaks shape. Unfortunately, any attempt to perform Rietveld structure refinements (Rietveld 1969) was unsuccessful.

The (isothermal) bulk compressibility of the (low-*P*) rhombohedral polymorphs of Li<sup>+</sup>-, Na<sup>+</sup>-, Ag<sup>+</sup>-, K<sup>+</sup>-, Rb<sup>+</sup>- and Cs<sup>+</sup>-chabazites is here described by the bulk modulus  $K_0$  ( $K_0 =$ 1/ $\beta = -V \cdot \partial P / \partial V$ , where  $\beta$  is the isothermal compressibility coefficient), obtained by a secondorder Birch-Murnaghan Equation of State (II-BM-EoS) fit (Birch 1947), using the EOS-fit V7.0 program (Angel et al. 2014) and the data weighted by the uncertainties in *P* and *V*.

150

# 151 **Results**

Synchrotron X-ray powder diffraction patterns collected at high pressure, using pure water as PTM, are shown in Fig. 1. A visual examination of the diffraction patterns reveals that, upon increasing pressure, the diffraction peaks exhibit gradual broadening. The broadening effect can be due to a number of factors, such as an increase in the long-range structural disorder and the growth of microstrains in response to the non-hydrostatic

conditions at P > 1 GPa (e.g., Yamanaka et al. 1997; Weidner et al. 1998; Fei and Wang 2000). 157 Similar effects have been observed for the other isotypic CHA materials (i.e., SAPO-34, 158 ALPO-34) by Leardini et al. (2010, 2013). After pressure release back to ambient conditions, 159 the peak positions, widths, and intensities revert back to those before compression, indicating 160 the reversibility of the P-induced deformation mechanisms in all the cation-exchanged 161 chabazites within the *P*-range investigated (Fig. 1). At P > 3 GPa, phase transitions from 162 rhombohedral to triclinic symmetry are observed in chabazites exchanged with Li<sup>+</sup>, K<sup>+</sup>, Ag<sup>+</sup>, 163 Rb<sup>+</sup> and Cs<sup>+</sup>, whereas the natural chabazite and the Na-form do not experience any transition 164 (Figs. 1 and 2 and Table 1). These phase transitions are driven by an abrupt decrease of the 165 unit-cell volume in the range between 2.0 and 10 % (Fig. 2). 166

The compressional of chabazite 167 pattern the natural (ORI-CHA, Ca1.6Na0.5Si8.4Al3.6O24·14.3H2O) in water PTM shows a monotonic trend, though with a 168 softening which is more pronounced at P > 2 GPa (Figs. 1 and 2, Table 1). The refined bulk 169 modulus (deduced on the basis of the low-P data pre-softening) is  $K_0$  (ORI-CHA) = 88(3) 170 171 GPa, while the measured unit-cell volume at ambient pressure is  $V_0$  (ORI-CHA) = 824.9(9) ų. 172

When Li-CHA (Li<sub>2.9</sub>Si<sub>8.6</sub>Al<sub>3.4</sub>O<sub>24</sub>·13.2H<sub>2</sub>O) is compressed in water PTM from  $P_{amb}$  to 173 174 5.5 GPa, the unit-cell volume decreases steadily below 3.0 GPa. Above this pressure, the rhombohedral structure transforms into a triclinic one (Figs. 1 and 2, Table 1), accompanied 175 by abrupt and anisotropic contraction of the unit-cell edges by ca. 0.8 %, 2.0 %, and 4.5 % for 176 the a-, b-, and c-edge lengths, respectively, of the high-P triclinic polymorph (Fig. 2). This 177 leads to an overall volume reduction by ca. 3.0 %. Bulk modulus at ambient pressure, 178 calculated for the low-P rhombohedral polymorph of Li-CHA, is  $K_0$  (Li-CHA) = 202(2) GPa 179 with the measured  $V_0$  (Li-CHA) of 819.9(9) Å<sup>3</sup>. The bulk modulus of Li-CHA is the highest 180 amongst the studied cation-exchanged chabazites (hence with the lowest compressibility), 181 whereas its volume at ambient pressure is the smallest. 182

In the case of Na-CHA (Na<sub>3.4</sub>Si<sub>8.6</sub>Al<sub>3.4</sub>O<sub>24</sub>·11.4H<sub>2</sub>O), compression in water PTM up to 5.3 GPa leads to a steady decrease of unit-cell volume without phase transition, though with a modest volume expansion at very low-*P* (0.5 GPa, Table 1) and softening at *P* > 2 GPa (Figs. 1 and 2, Table 1). The refined bulk modulus at ambient pressure (deduced on the basis of the low-*P* data pre-softening) is  $K_0$  (Na-CHA) = 114(9) GPa with the measured  $V_0$  (Na-CHA) of 824.9(9) Å<sup>3</sup>.

In Ag-CHA(Ag<sub>3.5</sub>Si<sub>8.5</sub>Al<sub>3.5</sub>O<sub>24</sub>·15.9H<sub>2</sub>O), the steady initial contraction of the unit-cell edges in water PTM is followed by a transition to a triclinic structure above ca. 5.7 GPa, accompanying abrupt and anisotropic contractions of the *a*-, *b*-, and *c*-edge lengths, of the triclinic polymorph, by ca. 0.4 %, 3.3 %, and 8.7 %, respectively (Figs. 1 and 2, Table 1). This leads to an overall volume reduction by ca. 10.0 %. The refined bulk modulus at ambient pressure, calculated for the low-*P* rhombohedral polymorph of Ag-CHA, is  $K_0$  (Ag-CHA) = 116(2) GPa with the measured  $V_0$  (Ag-CHA) of 829.2(2) Å<sup>3</sup>.

Similar transition from rhombohedral to triclinic structure is observed in K-CHA 196 (K<sub>3.2</sub>Si<sub>8.7</sub>Al<sub>3.3</sub>O<sub>24</sub>·10.7H<sub>2</sub>O) compressed in water at ca. 5.1 GPa (Figs. 1 and 2, Table 1). Also 197 in this case, the transition is accompanied by abrupt and anisotropic contraction of the unit-198 cell edges by ca. 1.5 %, 1.5 %, and 6.5 % for the a-, b-, and c-edge lengths, respectively (Fig. 199 200 2), which leads to an overall volume reduction of the high-P triclinic polymorph by ca. 6.0 %. The refined bulk modulus of the low-P rhombohedral polymorph of K-CHA is  $K_0$  (K-CHA) = 201 93(1) GPa, the lowest value amongst the ion-exchanged chabazites of this study, whereas the 202 measured unit-cell volume at ambient pressure is  $V_0$  (K-CHA) = 830.8(8) Å<sup>3</sup>. 203

Compression of Rb-CHA (Rb4.1Si7.9Al4.1O24·6.5H2O) in water PTM to 6.0 GPa shows a modest volume expansion at very low-*P* (0.5 GPa, Table 1) and then a gradual monotonic decrease of the unit-cell volume up to ca. 4.9 GPa, followed by abrupt contraction by ca. 5.0 % in response to the rhombohedral-to-triclinic phase transition (Figs. 1 and 2, Table 1). This transition is also driven by anisotropic contraction of the unit-cell edges, of the triclinic polymorph, by ca. 1.9 %, 0.7 %, and 4.8 % for the *a*-, *b*-, and *c*-edge lengths, respectively (Fig. 2). The refined bulk modulus of low-*P* rhombohedral Rb-CHA is the second largest after Li-CHA:  $K_0$  (Rb-CHA) = 149(5) GPa, with a measured  $V_0$  (Rb-CHA) = 826.0(1) Å<sup>3</sup>.

For Cs-CHA (Cs<sub>3.4</sub>Si<sub>8.6</sub>Al<sub>3.4</sub>O<sub>24</sub>·6.4H<sub>2</sub>O), a modest volume expansion at very low-P 212 213 (0.5 GPa, Table 1) followed by a monotonic compression is also observed (Figs. 1 and 2, Table 1). The degree of volume contraction during the rhombohedral-to-triclinic transition, 214 between 3 and 4 GPa, is modulated to ca. 2.0% with anisotropic reduction of the unit-cell 215 edges by ca. 1.4 %, 1.2 %, and 1.1 % for the a-, b-, and c-edges lengths, respectively, of the 216 triclinic form (Figs. 1 and 2, Table 1). Bulk modulus and (measured) unit-cell volume at 217 218 ambient pressure for the low-P rhombohedral polymorph are:  $K_0$  (Cs-CHA) = 137(1) GPa and  $V_0$  (Cs-CHA) = 830.4(4) Å<sup>3</sup>, respectively. 219

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# **Discussion and Implications**

222 The experimental findings of this study, in which a nominally penetrating Ptransmitting fluid is used (sensu Gatta 2008), allow first of all a comparison between the 223 compressional behavior of a natural chabazite in penetrating and non-penetrating media. 224 Leardini et al. (2010, 2013) reported the behavior of two natural chabazites, with slightly 225 different compositions, compressed in silicone oil (a non-penetrating P-medium) and showed: 226 227 a change of the compressional behavior at 1.4 GPa in one of the samples, with chemical formula (K1.36Ca1.04Na0.28Sr0.4Ba0.06Mg0.02)[Si7.17Al4.87O24]·13.16H2O, with an estimated bulk 228 modulus of 35(5) GPa at P < 1.4 GPa and 62(1) at P > 1.4 GPa (Leardini et al. 2010); a 229 rhombohedral-to-triclinic phase transition at 2.1 GPa in the second chabazite sample, with 230 chemical formula (Ca1.32K0.45Na0.13Sr0.10)[Si8.55Al3.45O24]·11.30H2O, with an estimated bulk 231 232 modulus of 54(3) GPa for the low-P polymorph. Further HP-experiments on the synthetic ALPO-34 and SAPO-34, isotypic materials with CHA framework topology, were performed 233 using non-penetrating fluids: the bulk modulus of the ALPO-34 was reported to be 54(3) 234

(Leardini et al. 2012) and that of SAPO-34 of 29(1) GPa (Leardini et al. 2010). ALPO-34 and 235 SAPO-34 are materials with a nominally neutral framework, and thus with no extra-236 237 framework cations. If we consider all the data available in the open literature, the "expected" bulk modulus (at room conditions) of a natural (rhombohedral) chabazite is 50±15 GPa. In 238 our study, the bulk modulus of the natural chabazite compressed in water, a nominally 239 penetrating fluid, leads to a bulk modulus of about 90 GPa. This value is, in general, unusual 240 for zeolites (i.e., too high, Gatta and Lee 2014) and, in this specific case, suggests that the 241 H<sub>2</sub>O molecules penetrate through the zeolitic cavities in response to the applied pressure. The 242 continuous penetration of the extra H<sub>2</sub>O molecules would lead to more efficient stuffing of 243 the pores by extra-framework species, making the zeolite structure less compressible. This 244 can explain the higher bulk modulus observed in this study if compared to those obtained in 245 previous experiments with non-penetrating P-fluids, in which the inherent compressibility is 246 obtained. A similar effect was previously observed in several HP-experiments on zeolites 247 (compressed in penetrating and non-penetrating fluids) and provides "indirect" evidence of 248 PIH in our experiment, useful when "direct" evidence are missed due to the lack of abrupt 249 structural changes and/or structural models (i.e., impossibility to perform Rietveld structure 250 251 refinements).

Without data at atomic scale obtained by structure refinements, it is not sure if the 252 penetration of extra H<sub>2</sub>O molecules occurs entirely at very low-P ( $\leq 0.5$  GPa), as suggested 253 254 by the modest volume expansion in Na-, Rb- and Cs-CHA (Table 1) and as observed for several zeolites (Gatta 2008; Gatta and Lee 2014 and references therein), or it is a continuous 255 process within the P-range investigated. In the second case, the bulk modulus value does not 256 have a robust physical meaning, because the composition of the zeolite changes with 257 increasing pressure (i.e., the system is "open"). However, the "apparent" compressibility, 258 259 through the bulk modulus, remains a useful measure for a comparative analysis (e.g., the same zeolite compressed in different fluids; zeolites with the same framework topology and 260 different extra-framework population compressed in the same fluid). 261

The compressional behavior of all the cation-exchanged chabazites of this study allow us to make the following observations and considerations:

264 1) Our results indicate an inverse relationship between the onset pressure of the rhombohedral-to-triclinic transition and the radius of extra-framework cation in 265 266 chabazite, above the ca. 1.0 Å threshold (Fig. 3). Similar trend is observed between the degree of volume contraction and the radius of extra-framework cation, which 267 appears to be mainly driven by the c-edge length contraction of the triclinic 268 polymorph (Fig. 3, Table 1). The largest contraction along the *c*-axis is ca. 8.7% in 269 270 Ag-CHA, whereas in K-CHA, Rb-CHA, and Cs-CHA, the contractions are by ca. 6.5, 271 4.8, and 1.4%, respectively (Fig. 3). The different volume contraction, in response to the phase transition, might be partly related to the initial H<sub>2</sub>O content at ambient 272 conditions. In Ag-CHA there are ca. 15.9 H<sub>2</sub>O molecules per formula unit (p.f.u.), 273 which decrease to ca. 10.7, 6.5, and 6.4 in K-CHA, Rb-CHA, and Cs-CHA, 274 275 respectively (Fig. 3). On the other hand, there are ca. 13.2 H<sub>2</sub>O p.f.u. in Li-CHA, 276 which exhibits lower transition pressure and volume contraction than Ag-CHA (Fig. 3): Li-CHA appears to be an outlier in the contraction vs. cation radius relationship 277 and needs further structural investigation. 278

2) There is an additional experimental finding about a potential relation between the 279 observed bulk modulus and the distribution of extra-framework cations over the 280 281 different segments forming the chabazite cavities, *i.e.*, D6R, S8R, and CHA-cage (Fig. 4). The highest bulk modulus of 202(2) GPa is observed for the rhombohedral low-P 282 polymorph of Li-CHA, where Li-cations fill all the three cavities (D6R, S8R, and 283 284 CHA-cage) at ambient conditions. More compressible than Li-CHA are Rb-CHA and Cs-CHA with bulk moduli of 149(5) and 137(1) GPa, respectively. In these chabazites, 285 286 the extra-framework cations populate the S8R and CHA-cages only (i.e., no D6R). The most compressible forms are then Ag-CHA, Na-CHA, and K-CHA with bulk 287 moduli of 116(2), 114(9), and 93(1) GPa, respectively. In these compounds, extra-288

framework cations are only located in the largest CHA-cages (*i.e.*, no D6R or S8R).

290 291  All the high-P deformation mechanisms and penetration phenomena are reversible, as proved by the diffraction data collected at room conditions after decompression (Fig.

292 1, Table 1).

293 Overall, it appears that:

1) PIH occurs in the natural and in all the cation-exchanged chabazites of this study, and 294 295 it is reversible. This is true even in the case of Na-CHA, which does not experience any *P*-induced phase transition but reacts, in response to the applied pressure, with a 296 bulk modulus of 114(9) GPa, not realistic for a zeolite without any crystal-fluid 297 interaction (Gatta 2008, Gatta and Lee 2014). At this stage, it is unknown why the 298 ORI-CHA and Na-CHA do not experience the P-induced phase transition observed 299 for the other cation-exchanged forms of this study. Likely, the higher number of 300 independent extra-framework sites in these two chabazites (i.e., ORI-CHA: 4Ca + 301 3Na + 5OW; Na-CHA: 4Na + 7OW; Li-CHA: 4Li + 5OW; K-CHA: 3K + 5OW; Rb-302 CHA: 2Rb + 2OW; Cs-CHA: 2Cs + 2OW; Ag-CHA: 2Ag + 2OW; Kong et al. 2016) 303 makes their structures more "flexible", with higher degrees of freedom 304 accommodating the P-induced deformation effects. 305

2) The degree of PIH is someway controlled by the distribution of the extra-framework 306 cations (which, in turn, reflects their ionic radius and charge) and how these can 307 coordinate extra H<sub>2</sub>O molecules. Li, for example, is a small ion and its coordination 308 polyhedra leaves room in the cavities for additional H<sub>2</sub>O molecules, which can be 309 further coordinated by Li or can be H-bonded to the framework oxygens. However, 310 the different number (and location inside the cavities) of independent cation sites and 311 H<sub>2</sub>O molecules in the cation-exchanged chabazites of this study does not allow to 312 define a universal and unambiguous model to explain the behavior of all the cation-313 exchanged chabazites. 314

We can draw geological implications of our experimental findings as follows. Our 316 results demonstrate that small molecules (in kinetic diameters), like H<sub>2</sub>O, CO<sub>2</sub>, CH<sub>4</sub> or H<sub>2</sub>S, 317 318 can potentially penetrate into the CHA-type zeolites in response to the applied pressure. Such a penetration phenomenon is likely to be active even at very low pressures (kilobar level or 319 even lower). Geological fluids can, therefore, interact efficiently with this zeolite with a 320 significant fluid-to-crystal mass transfer. In other words, the ability of zeolites, as 321 microporous materials, to act as geochemical traps of small molecules can be drastically 322 enhanced at moderate pressures even at room temperature; it is highly likely that the 323 324 combined effect of pressure and temperature would improve the magnitude of the PIH and PII, as previously observed in other zeolites (Gatta and Lee 2014 and references therein). 325

326 The technological implications of our results are even more relevant. Our experimental findings demonstrate that it is possible to modulate the elastic behavior of a 327 given zeolite simply by cation-exchange and using a penetrating P-transmitting fluid. A 328 combined  $[A^+-CHA + H_2O]$  system (with  $A^+ = Li$ , Na, Ag, K, Rb, Cs) can behave like a low-329 330 compressibility "spring": the bulk modulus of the Li-CHA in H<sub>2</sub>O (*i.e.*, 202(2) GPa) is higher, in certain P-range, than those of garnets (~190 GPa, Hazen et al. 1994), mullites (~ 170 GPa, 331 Gatta et al. 2010, 2013) or topaz (~160 GPa, Gatta et al. 2006, 2014). With different cations, 332 it is possible to generate hybrid softer systems with modulated bulk moduli targeting certain 333 solids such as olivines (~ 120-130 GPa, Smyth et al. 2000), pyroxenes (~ 90-130 GPa, 334 335 McCarthy et al. 2008) or feldspars (~ 50-80 GPa, Angel 2004). This is surprising if we consider that zeolites are microporous materials and intuitively considered as soft compounds. 336 PIH observed in this study for the natural and for all the cation-exchanged chabazites, is a 337 reversible phenomenon and cannot be used to generate super-hydrated zeolites which remain 338 metastable at room conditions after decompression. However, it would be different for other 339 340 small molecules and/or mixed cation chabazites. In this light, further studies are in progress in order to expand the number of small molecules able to penetrate the CHA-cavities at high 341 342 pressure.

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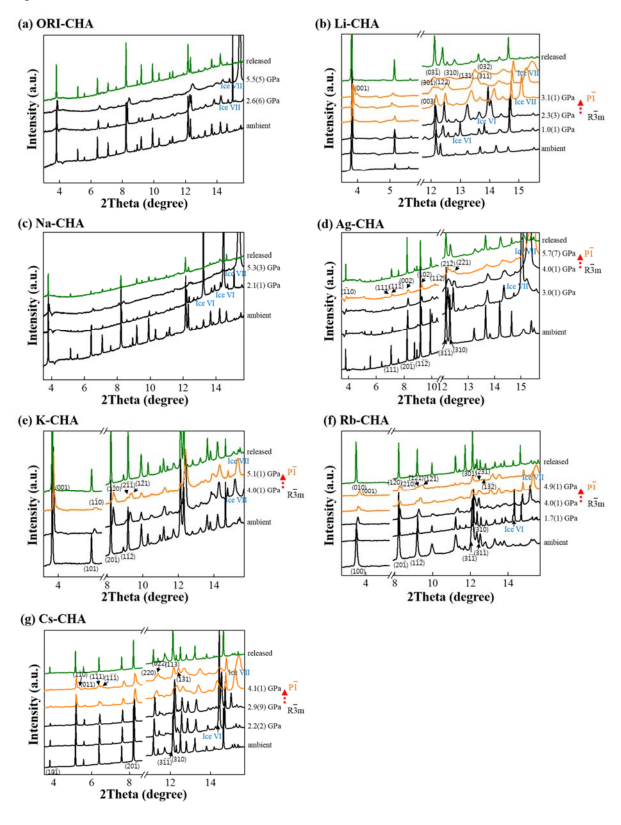
## 470 Figure captions

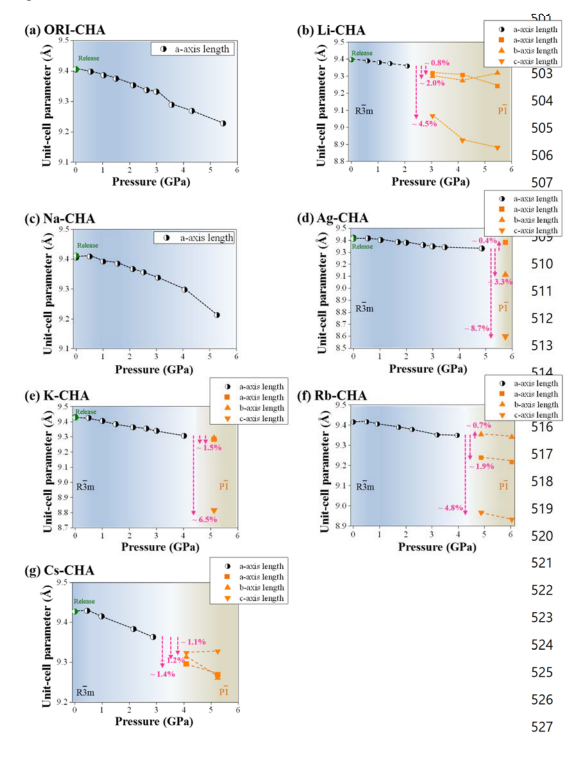
Figure 1. Synchrotron X-ray powder diffraction patterns as a function of hydrostatic
pressure mediated by pure water as *P*-transmitting medium for (a) ORI-CHA, (b) Li-CHA, (c)
Na-CHA, (d) Ag-CHA, (e) K-CHA, (f) Rb-CHA, and (g) Cs-CHA. Some of the new peak
positions due to symmetry lowering are indicated with Miller indices.

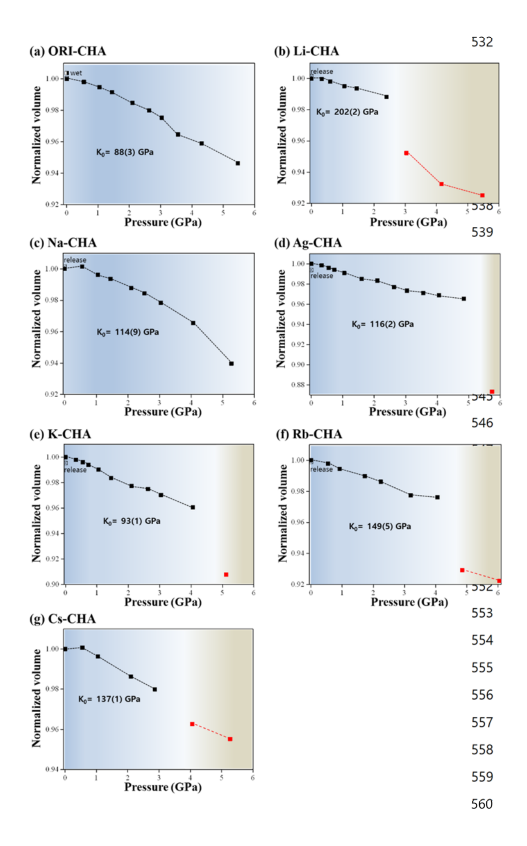
Figure 2. Evolution of the unit-cell edges lengths (Å) and volume (Å<sup>3</sup>) with *P*, using pure
water as *P*-transmitting medium, for (a) ORI-CHA, (b) Li-CHA, (c) Na-CHA, (d) Ag-CHA,
(e) K-CHA, (f) Rb-CHA, and (g) Cs-CHA. The errors associated with the cell parameters are
smaller than the symbols. The dashed lines represent only a guide for eyes. For the unit-cell
volume, the red symbols indicate the triclinic high-*P* polymorphs.

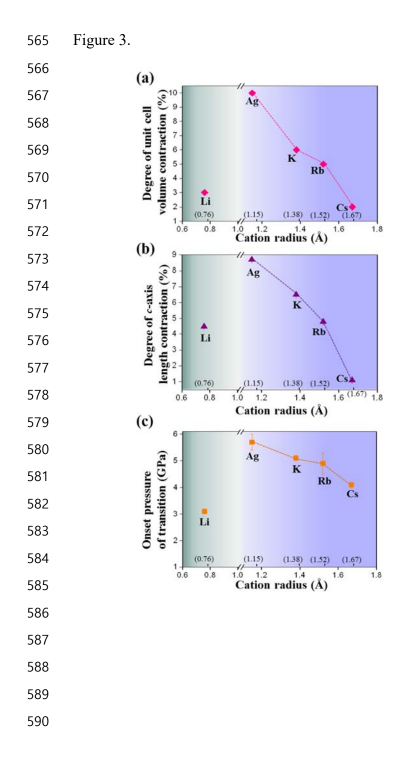
Figure 3. Changes in the (a) unit-cell volume, (b) *c*-edge length, and (c) onset pressure of the rhombohedral-to-triclinic transition as a function of the ionic radius of the extraframework cation in the alkali-metal-exchanged chabazites.

Figure 4. (a) Site distribution and (b) occupancy of the extra-framework cations, and (c)
initial H<sub>2</sub>O molecular contents per formula unit in the alkali-metal-exchanged chabazites at
ambient conditions. (d) "Observed" bulk moduli plotted as a function of cation radius.

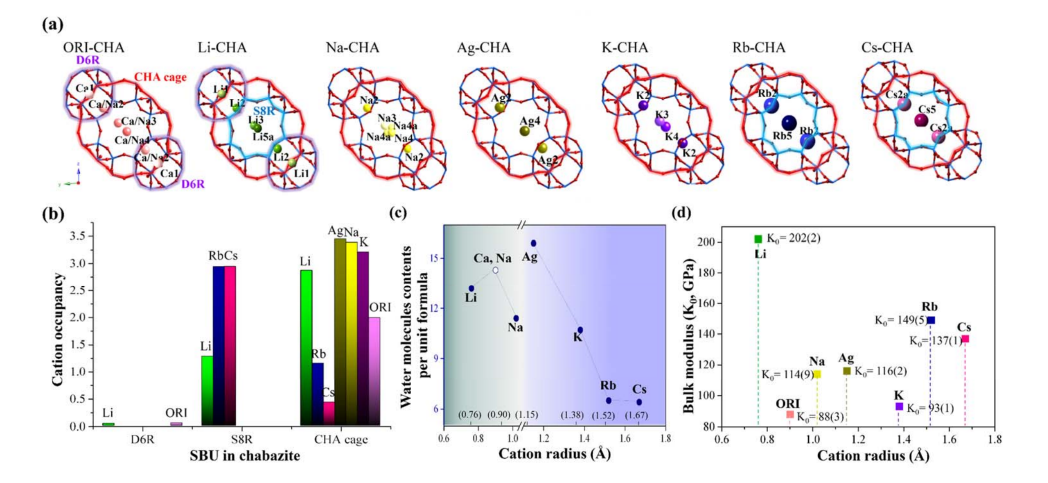








591 Figure 4.



ORI- CHA		Ambient	GPa	1.05(1) GPa	1.45(1) GPa	2.10(1) GPa	2.63(1) GPa	3.04(1) GPa	3.56(1) GPa	4.31(1) GPa	5.47(1) GPa	Released
	S.G.	$R\overline{3}m$	R∃m	R3m	R∃m	R∃m	R∃m	R∃m	$R\overline{3}m$	R∃m	R∃m	R∃m
	R <sub>wp</sub> (%)	1.56	1.80	1.52	1.71	2.1	1.92	1.58	1.63	1.42	1.53	1.78
	$\chi^2$	0.12	0.14	0.10	0.13	0.15	0.14	0.10	0.11	0.10	0.10	0.14
	$\tilde{a}$ (Å)	9.405(5)	9.398(8)	9.386(6)	9.375(5)	9.353(3)	9.337(7)	9.332(2)	9.289(9)	9.269(1)	9.228(1)	9.406(6
	α (°)		94.13(3)	94.02(2)	93.93(3)	93.85(5)	93.83(3)	93.83(3)	93.88(1)	93.73(1)	93.76(2)	94.21(1
	$V(\text{\AA}^3)$	824.9(9)	823.3(3)	820.5(5)	817.8(8)	812.3(3)	808.4(4)	804.4(1)	795.7(1)	791.0(4)	780.6(3)	825.1(1)
Na-		Ambient	0.51(1)	1.01(1)	1.55(1)	2.12(1)	2.52(1)	3.06(1)		4.06(1)	5.27(1)	Release
СНА	S.G.	R3m	GPa R3m	GPa R3m	GPa R3m	GPa R3m	GPa R3m	GPa R3m		GPa R3m	GPa R3m	R3m
	$R_{\rm wp}(\%)$		1.30	1.56	1.35	2.00	1.8	1.79		1.33	1.25	1.40
	2	0.16										
	χ		0.10	0.10	0.10	0.15	0.12	0.12		0.10	0.10	0.10
	a(Å)		9.409(1)	9.392(1)	9.385(5)	9.367(1)	9.356(1)	9.338(1)		9.298(8)	9.213(2)	9.412(2)
	$\alpha$ (°)		94.14(1)	94.09(1)	94.17(1)	94.19(1)	94.22(1)	94.25(1)		94.32(1)	94.28(4)	94.32(1)
	$V(\text{\AA}^3)$		826.1(1)	821.8(1)	819.7(1)	814.9(1)	812.1(1)	807.2(1)		796.6(1)	775.2(4)	826.2(1)
Ag- CHA	_	Ambient	0.55(1) GPa	1.03(1) GPa	1.68(1) GPa	1.99(1) GPa	2.63(1) GPa	2.99(1) GPa	3.45(1) GPa	4.85(1) GPa	5.74(1) GPa	Release
	S.G.	$R\overline{3}m$	R3m	R3m	R3m	R3m	R3m	R3m	R∃m	R3m	$P\overline{1}$	$R\overline{3}m$
	$R_{wp}(\%)$	4.11	2.03	2.27	1.75	1.97	2.38	1.74	1.69	1.52	1.06	2.05
	$\chi^2$	1.61	0.36	0.45	0.26	0.32	0.47	0.25	0.24	0.19	0.10	0.35
	a(Å) b(Å)		9.417(7)	9.402(2)	9.385(5)	9.38(8)	9.36(6)	9.349(9)	9.342(2)	9.332(1)	9.382(4) 9.112(2)	9.412(2)
	c (Å) α (°) β (°)	94.17(7)	94.21(1)	94.25(5)	94.39(9)	94.4(4)	94.4(4)	94.43(3)	94.45(5)	94.51(1)	8.598(2) 86.29(3) 93.00(2)	94.31(1)
	$\gamma$ (°) V (Å <sup>3</sup> )	829.2(2)	828.1(1)	823.9(1)	819.1(1)	817.6(6)	812.3(1)	809.3(1)	807.4(1)	804.6(3)	97.77(2) 726.0(3)	826.4(1)
Li-CHA		Ambient	0.61(1) GPa	0.98(1) GPa	1.47(1) GPa	2.26(1) GPa		3.06(1) GPa		4.16(1) GPa	5.48(1) GPa	Released
	S.G.	R3m	R3m	R3m	R3m	R3m		$P\overline{1}$		$P\overline{1}$	$P\overline{1}$	R∃m
	$R_{wp}(\%)$	3.11	2.92	2.84	2.35	3.53		3.41		3.27	2.03	3.49
	$\chi^2$	0.47	0.38	0.36	0.24	0.55		0.46		0.43	0.19	0.55
	a (Å) b (Å)		9.389(9)	9.38(8)	9.374(4)	9.359(9)		9.317(1) 9.299(2)		9.307(3) 9.273(2)	9.242(2) 9.319(3)	9.399(2)
	$c (Å)  \alpha (°)  \beta (°)  \gamma (°) $	94.88(8)	94.86(6)	94.86(6)	94.75(5)	94.89(9)		9.067(2) 91.00(3) 92.48(1) 95.72(2)		8.924(2) 90.60(3) 92.47(4) 96.53(2)	8.881(6) 91.79(4) 92.46(5) 96.61(3)	94.8(8)
	$V(\text{\AA}^3)$	819.9(9)	818.3(1)	815.9(1)	814.6(6)	810.2(1)		780.7(3)		764.4(2)	758.5(5)	821.1(1)
К-СНА		Ambient	0.49(1) GPa	1.00(1) GPa	1.49(1) GPa	2.13(1) GPa	2.62(1) GPa	3.00(1) GPa		4.01(1) GPa	5.12(1) GPa	Release
	S.G.	R3m	R3m	R3m	R3m	R3m	R3m	R3m		R3m	$P\overline{1}$	R3m
	R <sub>wp</sub> (%)	2.63	3.52	3.76	4.01	3.32	3.40	3.41		2.81	2.44	3.91
	2	0.36	0.62	0.73	0.81	0.51	0.55	0.57		0.37	0.27	0.73
	χ a (Å) b (Å)	9.43(3)	9.425(5)	9.405(5)	9.384(4)	9.363(3)	9.355(5)	9.34(4)		9.307(1)	9.285(1) 9.291(3)	9.427(7)
	c (Å) α (°)	94.39(9)	93.97(7)	93.85(5)	93.86(6)	93.8(8)	93.74(4)	93.71(1)		93.59(1)	8.816(2) 90.77(3) 93.95(2) 93.47(2)	94.3(3)
	β(°) γ(°)											
	γ(°)	830.8(8)	830.8(8)	826.1(1)	820.5(5)	815.3(1)	813.3(1)	809.4(1)		801.4(2)	757.3(2)	830.3(3
Rb- CHA		830.8(8) Ambient	0.51(1)	826.1(1) 0.92(1) GPa	820.5(5) 1.73(1) GPa	815.3(1) 2.24(1) GPa	813.3(1)	809.4(1) 3.20(1) GPa	3.96(1) GPa	801.4(2) 4.87(1) GPa	757.3(2) 6.04(1) GPa	830.3(3) Release
Rb- CHA	γ(°)		0.51(1)	0.92(1)	1.73(1)	2.24(1)	813.3(1)	3.20(1)		4.87(1)	6.04(1)	

Table 1. Changes in the unit-cell edge lengths and volume of the cation-exchanged chabazites
 with *P*, compressed in pure water as pore-penetrating pressure transmitting medium.

	$R_{wp}(\%)$	1.84	2.54	2.36	2.16	1.56	2.11	1.52	2.02	1.87	4.88
	$\chi^2$	0.47	0.50	0.44	0.36	0.18	0.34	0.17	0.32	0.26	1.84
	a (Å) b (Å)	9.416(6)	9.417(7)	9.406(6)	9.39(9)	9.379(9)	9.352(2)	9.35(5)	9.24(4) 9.352(3)	9.218(2) 9.342(4)	9.416(1)
	$c (Å)  \alpha (°)  \beta (°)  (°)$	94.69(9)	94.45(5)	94.42(2)	94.36(6)	94.36(6)	94.42(2)	94.56(1)	8.967(2) 91.51(2) 92.72(2)	8.933(5) 91.9(9) 92.86(4)	94.58(1)
	$\gamma$ (°) V (Å <sup>3</sup> )	826.0(1)	827.2(2)	824.3(3)	820.5(1)	817.5(5)	810.2(1)	809.2(3)	95.01(2) 770.2(2)	95.3(3) 764.5(4)	826.4(2)
Cs- CHA		Ambient	0.45(1) GPa	0.98(1) GPa		2.15(1) GPa	2.87(1) GPa		4.09(1) GPa	5.24(1) GPa	Released
	S.G.	R3m	R3m	R3m		R3m	R3m		$P\overline{1}$	$P\overline{1}$	R3m
	$R_{wp}(\%)$	2.23	3.44	5.00		3.49	3.57		1.96	1.87	3.52
	$\chi^2$	0.25	0.58	1.23		0.57	0.57		0.17	0.16	0.58
	a (Å) b (Å) c (Å)	9.427(7)	9.429(9)	9.415(5)		9.383(3)	9.363(3)		9.296(2) 9.313(1) 9.324(1)	9.269(1) 9.261(2) 9.328(1)	9.427(1)
	α (°) β (°) γ (°)	94.25(5)	94.26(6)	94.26(6)		94.21(1)	94.22(2)		94.53(1) 95.27(1) 93.5(5)	94.78(1) 95.19(2) 92.96(2)	94.24(4)
	$V(\text{\AA}^3)$	820 A(A)	831.0(1)	827.3(3)		819.1(1)	813.7(1)		799.4(2)	793.2(2)	830.5(1)