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The effect of type-B carbonate content on the elasticity of fluorapatite

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Abstract

The mechanical behavior of carbonate-bearing fluorapatite (CFAP) (with up to 5.5 wt% CO₃) was investigated at high pressure up to 7 GPa. The incorporation of carbonate in CFAP samples was investigated by FTIR spectroscopy. The chemical formulae and cell parameters are Ca_{4.90}Fe_{0.04} $(PO_4)_{2.87} (CO_3)_{0.13} F_{1.23}$ and $a = 9.3527(1), c = 6.8752(1) \text{ Å}, V = 520.83(1) \text{ Å}^3$ for the FOW CFAP (Fowey Consols area, UK), and Ca4.97Sr0.03 (PO4)2.55 $(CO_3)_{0.45}$ F_{1.42} and a = 9.3330(1), c = 6.8984(1) Å, V = 520.38(1) Å³ for the FRA CFAP (Framont region, France). Preliminary characterization at ambient conditions was done by single-crystal X-ray diffraction study. The structure refinements, in space group $P6_3/m$, confirm a type-B substitution of the phosphate $(PO_4)^{3-}$ group by the carbonate ion $(CO_3)^{2-}$. The site occupancies for the C atom are 0.04 for FOW and 0.11 for FRA CFAP, in quite good agreement with the 1.6 and 5.5 wt% CO3 amount obtained by analytical methods. Single-crystal high-pressure XRD study on the two type-B CFAP samples was performed. The FOW and FRA crystals were mounted concurrently in a ETH-type DAC and cell parameters were determined at 26 different pressures up to 6.86 GPa at room T. The variation with pressure of the unit-cell parameters and volume shows no discontinuity that could be related to any possible phase transition in the P range investigated. The linear compressibility coefficients are $\beta_a = 3.63 \times 10^{-3}$ GPa⁻¹ and $\beta_c = 2.47 \times 10^{-3}$ GPa⁻¹ for FOW, and $\beta_a = 3.67 \times 10^{-3}$ GPa⁻¹ and $\beta_c = 2.65 \times 10^{-3}$ GPa⁻¹ for FRA, giving an axial anisotropy of β_a : $\beta_c = 1.47$:1 and 1.38:1, respectively. The P-V data were fitted by a second-order Birch-Murnaghan EoS and the resulting BM2-EoS coefficients are $V_0 = 519.81(7)$ Å³, $K_{T0} = 92.1(3)$ GPa for FOW, and $V_0 = 518.95(9)$ Å³, $K_{T0} = 89.1(4)$ GPa for FRA CFAP. The results obtained indicate that a 5.5 wt% CO₃ content (type-B) reduces the isothermal bulk modulus by about 9%.

Keywords

Carbonate-rich fluorapatite FTIR spectra HP SC-XRD Crystal structure Bulk modulus EoS

Electronic supplementary material

The online version of this article (https://doi.org/10.1007/s00269-018-0962-1) contains supplementary material, which is available to authorized users.

Introduction

Apatite is an important mineral occurring in igneous, metamorphic and hydrothermal rocks and the knowledge of its mineral physics helps to assess the geological implications of its stability field. Understanding its stability at the Earth's interior is essential to grasp its role as a possible repository of phosphorous at mantle depths. Beside its potential application to mantle mineralogy, knowledge of the apatite Equation of State (EoS) is also relevant to metamorphic petrologic applications such as interpreting pressure-temperaturetime paths (Brunet et al. 1999; Ashley et al. 2017). Apatite has many geological uses, including dating techniques and studies of the variation of rare-earth elements in rocks, and is also widely used in material science (Hughes 2015 and references therein). Rock phosphates (microcrystalline apatites), mostly of biological origin, are the starting material for the manufacture of phosphate fertilizer and a source of phosphorus for the chemical industry (Elliott et al. 2002; Rakovan and Pasteris 2015). In the biomedical field, its mechanical properties play a major role in its technological applications, as apatite is the main constituent of dental enamel and human bone (Elliott 1994, 2002; Pasteris 2016).

The apatite group has three calcium phosphate mineral species, fluorapatite (FAP), chlorapatite (CAP), and hydroxylapatite (HAP) with the general formula of Ca_{5} -(PO₄)₃-(F,Cl,OH) (Pasero et al. 2010). Recent attention has focused on a more complete understanding of the apatite atomic arrangement and its properties, both in inorganic and biogenic apatite, and knowledge of the details of its atomic arrangement is increasing (Hughes et al. 1989, 1990, 2014; Hughes and Rakovan 2002, 2015; McCubbin et al. 2008; Hughes 2015). It is wellestablished that the incorporation of carbonate into the apatite structure has a destabilizing effect, resulting in increased solubility (LeGeros 1991). Whereas the structural details of FAP, CAP, and HAP are generally well-established, those of "carbonated" apatite are not yet very clear (Liu et al. 2011; Yi et al. 2013). Due to the importance of carbonate-bearing HAP in biomineralization (Elliott 2002), the incorporation of carbonate into apatite and its resulting physicalchemical changes have been investigated extensively and yet are still not well understood (McClellan 1980; Perdikatsis 1991; Elliott 1994; Regnier et al. 1994; Nathan 1996; Wilson et al. 1999, 2004; Comodi and Liu 2000; Leventouri et al. 2000; Ivanova et al. 2001; Elliott et al. 2002; Pan and Fleet 2002; Fleet and Liu 2004, 2007, 2008a, b; Yi et al. 2013). Incorporation of carbonate into the apatite structure can occur by substitution for both channel anion (type-A) and phosphate ion (type-B), as first suggested by McConnell (1938). Several authors (LeGeros 1965; McClellan and Lehr 1969; McClellan and Van Kauwenbergh

1990) have shown a correlation between increased carbonate and decreased phosphate, with different effects on the unit-cell parameters (Knudsen and Gunter 2002). The response of the unit-cell parameters of apatite to accommodation of the carbonate ion is complicated (Elliott 2002; Fleet et al. 2004; Liu et al. 2011). Substitution of hydroxyl by type-A carbonate in the channels results in a progressive increase in a values and a decreasing trend for c values, whereas substitution of phosphate by type-B carbonate gives origin to inverse trends where a decreases and c increases. However, many of the type-B carbonate apatites have somewhat increased a dimensions due mainly to the vacancies in oxygen sites coordinating tetrahedra and the possible presence of Ca substituted by H₂O groups (Ivanova et al. 2001).

With regard to the elastic behavior of apatites, several compression experiments are done (Brunet et al. 1999; Comodi et al. 2001; Matsukage et al. 2004; Schouwink et al. 2010; Liu et al. 2011) in Table 1 to investigate the compressibility of synthetic and natural apatite. Literature results are listed in Table 1 to compare the bulk modulus and the volume variations between the previous studies for apatite species and our results for carbonate-bearing fluorapatite (CFAP). Allan et al. (1996) and Brunet et al. (1999) studied the compressibility of three synthetic crystals of FAP, HAP, and CAP up to maximum pressures of 18, 20 and 52 GPa, respectively, at ambient temperature, by high-pressure synchrotron X-ray powder diffraction. Regardless of the large P-range, only the unit-cell volumes measured to about 10 GPa were used to derive their EoS, because higher-pressure data were affected by non-hydrostatic effects. A second-order Birch-Murnaghan EoS was fitted to the data giving values of $K_0 = 97.9$, 97.5, and 93.1 GPa for FAP, HAP, and CAP, respectively (Table 1). Because the three bulk moduli are identical within one estimated standard deviation, the halogen content of apatite was considered to not have a significant effect on volume compressibility, whereas the compressibility of a and c parameters seems more sensitive to halogen composition. Consistently, for each species, the *a*-axis is more compressible than the *c*-axis.

TableNote. In Tables 1,3,4,5,6, the (esd) values are separated by means of a space by their numbervalues are generally attacked, is this a new format of the magazine?

I mean: 522.4 (4) instead of 522.4(4)

Since you have separated them all in all the tables, you can also leave them like this, so as not to make you as you like, thank you $1\,$

Compression data for apatite in the P range: $0.0001 - P_{max}$

Reference Co	Composition	Diffraction techniques	P _{max} (GPa)	EoS	V_0 (Å ³)	K _T (G
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Reference	Composition	Diffraction techniques	P (GPa)	EoS	V (Å)	K (G
Brunet et al. (1999)	FAP (synthetic)	Powder	10	BM2	522.4 (4)	97 (1.
Brunet et al. (1999)	HAP (synthetic)	Powder	10	BM2	526.9 (5)	97 (1.
Brunet et al. (1999)	CAP (synthetic)	Powder	10	BM2	539.3(1.4)	93 (4.
Comodi et	FAP	Single-	()	BM2	_	97 (1.
al. (2001)	(synthetic)	crystal	6.9	BM3	524.21 (8)	93
Matsukage			7.1	BM2	524.2 (2)	91 (1.
et al. (2004)	FAP ₉₄ CAP ₆	Powder	/.1	BM3	524.2 (3)	91. (3.
Schouwink et al. (2010)	FAP ₉₅ CAP ₅	Single- crystal	9.2	BM3	525.89 (4)	88. (7)
Liu et al.	CHAP (5 wt% CO ₃)	Powder	10	BM2 <i>BM3</i> please split the words BM2 <i>BM3</i> and wrap:	529.16 (8)	89 (5)
(2011)	(synthetic)			BM2 BM3	529.17 (9)	89
Liu et al.	CHAP (11 wt%	Powder	10	BM2	527.1 (4)	84
(2011)	CO ₃) (synthetic)		10	BM3	527.5 (2)	73
FOW (this study)	CFAP (1.6 wt% CO ₃)	Single- crystal	6.9	BM2	519.81 (7)	92 (3)
FRA (this study)	CFAP (5.5 wt% CO ₃)	Single- crystal	6.9	BM2	518.95 (9)	89 (4)

The parameters of the Birch–Murnaghan Equation of State (BM EoS) are: $V_0 =$ zero-pres volume, K_{T0} isothermal bulk modulus, K' first pressure derivative of K_{T0} (esd in parenth *BM3* values are in italic

FAP fluorapatite, *HAP* hydroxylapatite, *CAP* chlorapatite, *CHAP* carbonate-bearing hydr *CFAP* carbonate-bearing fluorapatite

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Comodi et al. (2001), using single-crystal XRD, determined the compressibility up to about 7 GPa of a synthetic fluorapatite with a = 9.375(2), c = 6.887(1) Å,

and V = 524.23(5) Å³ cell parameters. The compressibility along *a* was greater compared to the one measured along *c*. Unit-cell parameters showed a linear variation with *P* and their calculated linear compressibility coefficients give $\beta_a:\beta_c = 1.38:1$. Volume compressibility data fitted with BM3 EoS gave a bulk modulus of $K_0 = 93$ (Table 1), whereas data fitted with BM2 EoS gave a bulk modulus of $K_0 = 97.8$ GPa, in good accord with the value obtained by Brunet et al. (1999) in their fluorapatite, although the latter contained 0.025 apfu hydroxyl.

A fluorapatite of composition $\text{FAP}_{94}\text{CAP}_6$ was studied by Matsukage et al. (2004) by in-situ synchrotron X-ray powder diffraction up to 7.12 GPa at ambient temperature. Both unit-cell parameters and volume decrease systematically with increasing pressure-up to 7.12 GPa; a larger compression along *a* compared to the one along *c*, as previously observed in synthetic fluorapatite by Comodi et al. (2001), was observed. The BM3 EoS fitted to the volume–pressure data gave the isothermal bulk modulus $K_T = 91.5(3.8)$ GPa, and the error became smaller with *K'* fixed at 4 (BM2) (Table 1). The bulk modulus of Matsukage et al. (2004) for fluorapatite is about 6% lower compared to the values of Brunet et al. (1999) and Comodi et al. (2001). The unit-cell axial moduli were fitted by a BM2 EoS and gave $Ka_T = 80(1)$ and $Kc_T = 126(1)$ GPa for *a* and *c* axis, respectively, confirming that fluorapatite is elastically anisotropic. The anisotropy of the axial moduli was estimated to be about 58%.

Schouwink et al. (2010), studying ion tracks in apatite, performed high-pressure single-crystal XRD up to 10 GPa on irradiated and non-irradiated crystals of fluorapatite ($FAP_{95}CAP_5$), coming from Durango (Mexico). The values of bulk modulus are about 3–5% lower than the results reported by Matsukage et al. (2004) and Comodi et al. (2001) (Table 1). Regarding the axial compressibilities, the Authors confirm the previously determined elastic anisotropy with the *c*-axis being less compressible relative to the *a*-axis.

More recently, Liu et al. (2011) investigated the mechanical behavior of "carbonated" hydroxylapatite (CHAP) and studied the effect of carbonate ion on the compressibility of HAP. The incorporation of the carbonate oxy-anion (for both types A-B) into the structure of HAP results in the formation of vacancies, oxygen-loss and disorder, with consequent changes in the physical–chemical properties. Liu et al. (2011) results were obtained using a diamond-anvil cell and synchrotron X-ray powder diffraction up to 10 GPa on two synthetic CHAP samples with up to 11 wt% CO_3 . The results of their BM3 EoS data-fitting, reported in Table 1, suggest that carbonate content significantly reduces the isothermal bulk modulus of CHAP (up to about 15%).

To clarify the influence on the elastic properties of the incorporation of carbonate into the structure of FAP, we have examined two type-B carbonatebearing fluorapatites (CFAP) by HP single-crystal X-ray diffraction. The purpose of our study is to determine the Equation of State (Eos) and to understand the compressional properties of type-B CFAP.

Experimental methods

The two samples of carbonate-bearing fluorapatite (CFAP) are from the National History Museum of Natural Science of Paris (France). One is labeled "francolite" MNHN-63.116 (hereafter FOW), and comes from Fowey Consols area, St Austell District, Cornwall, England, UK (one of the deepest, richest and most important of the Cornish copper mines), and the other is labeled "francolite" MNHN-96.1356 (hereafter FRA) from the Framont-Grandfontaine region in Alsace, France (an ancient iron-mining area). "Francolite" is a carbonate-rich variety of fluorapatite, characterized by an excess amount of F, with carbonate groups $(CO_3)^{2-}$ partly substituting the phosphate $(PO_4)^{3-}$ ones. The name "francolite" comes from Wheal Franco (Tavistock District, Devon, England, UK) (Henry 1850; Sandell et al. 1939) and was used to identify apatite containing at least 1% fluorine and appreciable CO_3 . The name "francolite" was afterwards discredited by IMA and replaced with "carbonate-rich" or "carbonate-bearing" fluorapatite (Pasero et al. 2010).

The fluorapatite samples were investigated by FTIR spectroscopy using a Bruker Hyperion 2000 IR microscope equipped with a liquid-nitrogen-cooled MCT detector. Spectra over the range 4000–650 cm⁻¹ were obtained by averaging 100 scans with a resolution of 4 cm⁻¹. Base-line correction was done using the OPUS spectroscopic software (Bruker Optic GmbH). The spectra were collected on thin films having a similar thickness; to compare the IR intensity (absorbance) of the carbonate bands (proportional to concentration) for the two samples, the sizes of the selected crystal fragments used to obtain the films (via a diamond press) were very similar.

The chemical compositions of the crystals were obtained using a Cameca SX-100 electron microprobe operating in wavelength-dispersive mode with an accelerating voltage of 15 kV, a specimen current of 20 nA, and a beam diameter of 10 μ m. The following standards were used: Si: diopside; F, Ca and P: apatite; Sr: SrTiO₃; Fe: fayalite. The elements Si, Na, La, Ce, Pr, Nd, Sm, Y, Mg, Cl and S were sought but were below detection limits. The data were reduced and corrected by the PAP method of Pouchou and Pichoir (1985). The CO₂ content (presence of CO₃ groups confirmed by IR spectroscopy, see below) was calculated from the results of the crystal-structure refinement and on the basis of

P + C = 3 apfu, as the absence of CO₃ and OH in the channels was confirmed by IR spectroscopy. The results are reported in Table 2. The chemical formula obtained on the basis of 13 (O + F) pfu are $Ca_{4.90} Fe_{0.04} (PO_4)_{2.87} (CO_3)_{0.13} F_{1.23}$ (1.6 wt% CO₃) for FOW type-B CFAP, and $Ca_{4.97} Sr_{0.03} (PO_4)_{2.55} (CO_3)_{0.45} F_{1.42}$ (5.5 wt% CO₃) for FRA type-B CFAP. Both formulae show an excess of F > 1 pfu, which reasonably correlates with the CO₃ content.

Table 2

Electron microprobe analyses (mean of 10 analytical points, in wt%) of FOW and FRA samples, estimated standard deviation (esd) and relative formulae on the basis of 13 (O + F)

Sample Please split the words "Sample oxide" and	FOW		FRA			
wrap: Sample Oxide Note: "Sample" is referred to the two samples FOW and FRA. oxide	Average	ESD lowercase: esd	Range	Average	ESD lowercase: esd	Rai
FeO	0.59	0.13	0.36 - 0.80	0.01	0.01	0.0
CaO	54.99	0.20	54.67– 55.30	56.29	0.15	55. <u>1</u> 56.4
SrO	0.04	0.02	0.03- 0.09	0.65	0.11	0.3 0.7
P ₂ O ₅	40.71	0.33	40.03– 41.06	36.51	0.28	36. 36.
CO ₂ ^a	1.14			4.00		
F	4.69	0.10	4.55– 4.86	5.43	0.20	5.21 5.71
Total	102.16	0.15	102.11 - 102.56	102.89	0.19	102 103
O=F	1.98	0.04	1.92 - 2.05	2.28	0.08	2.19 2.4

Calculated by stoichiometry

22/	3/2	20	18

100.18	0.15	100.09– 100.55	100.61	0.18	100 101
2.868			2.549		
0.130			0.450		
2.998			2.999		
0.041			0.001		
4.903			4.974		
0.002			0.031		
4.946			5.006		
1.235			1.415		
	100.18 2.868 0.130 2.998 0.041 4.903 0.002 4.946 1.235	100.18 0.15 2.868 0.130 0.130 2.998 0.041 4.903 0.002 0.002 4.946 1.235	100.18 0.15 100.09-100.55 2.868 0.130 2.998 0.041 4.903 0.002 1.235	100.180.15100.09- 100.55100.612.8682.5492.5490.1300.4500.4502.9982.9992.9990.0410.0010.0014.9034.9740.0310.0020.0315.0061.2351.415	100.180.15100.09- 100.55100.610.182.8682.8682.5490.1300.1300.4500.4500.4502.9982.9992.9990.0010.0410.0010.0010.0014.9034.9740.0310.0310.0020.0315.0061.415

Two crystals of these carbonate-bearing fluorapatites (FOW size: $135 \times 110 \times 50$ μ m and FRA size: 100 × 80 × 60 μ m) were selected on the basis of their optical features for single-crystal investigation. A preliminary characterization at room conditions was done using an Oxford Gemini R Ultra single-crystal X-ray diffractometer (CrisDi Centre, University of Torino), equipped with a CCD area detector, with graphite-monochromatized MoK α radiation ($\lambda = 0.7107$ Å), and the tube operating at 50 kV and 40 mA. Diffraction data were collected in the 2θ range 0-82°. A total of 27 runs with 2806 frames (for FOW) and 2264 frames (for FRA) were collected, covering the whole accessible reciprocal space; 66,135 reflections of with 1209 unique were measured for FOW, and 66,143 of with 1210 unique for FRA. Omega-rotation frames (frame width 1°, exposure time 8 s, sample-to-detector distance 53 mm) were processed with the CrysAlisPro software (Agilent Technologies) and intensities were corrected for Lorentz and polarization effects. Data were corrected for empirical absorption. Scattering curves were chosen according to the calculated chemical formulae. All reflections with $I_0 > 2\sigma(I)$ were considered as observed during weighted fullmatrix least-squares refinement. Crystal-structure refinements were done in the space group $P6_2/m$, starting from the fractional coordinates of Comodi et al. (2001). Refinement of site scattering at the P sites showed a deficit. The maximum content of 15% CO₃ substitution at the P-sites did not produce evidence of site splitting in FOW CFAP (as it represents an electron density < $0.3 \text{ e}^{-}\text{Å}^{-3}$), whereas a maximum was clearly observed for FRA CFAP and was accordingly added to the model. The excess of fluoride was placed at the O3 oxygen site (the F3 site). Structural details were refined using the SHELX-97 package (Sheldrick 2008). Unit-cell parameters at room conditions are a =

9.3527(1), c = 6.8752(1) Å, V = 520.83(1) Å³ for FOW CFAP, and a = 9.3330(1), c = 6.8984(1) Å, V = 520.38(1) Å³ for FRA CFAP, respectively. Anisotropic refinement with 45 parameters converged at R = 0.018 for 1170 reflections with $F_o \ge 4\sigma(F)$ and at R = 0.020 for all 1209 data for FOW CFAP. The refinement with 47 parameters converged at R = 0.039 for 1161 reflections with $F_o \ge 4\sigma(F)$ and at R = 0.042 for all 1210 data for FRA CFAP. Details of data collection and refinement for FOW and Table 3. FRA CFAP are listed in Table 3. Atom coordinates, displacement parameters and site occupancies are given in Table 4. Selected bond lengths are shown in Table 5 (CIF files deposited of Electronic supplementary material).

Table 3

Single-crystal data for FOW and FRA CFAP samples at ambient conditions (esd in parentheses)

	FOW	FRA
Space group	<i>P</i> 6 ₃ / <i>m</i>	<i>P</i> 6 ₃ / <i>m</i>
<i>a</i> (Å)	9.352-7 (1) please correct: 9.3527 (1)	9.3330 (1)
<i>c</i> (Å)	6.8752 (1)	6.8984 (1)
V (Å3)	520.83 (1)	520.38 (1)
Density (g/cm ³)	3.31	3.29
Refl. unique	1209	1210
Refl. F_0 please replace "zero" with the letter "o" replace F_0 with $F_0 \ge 4\sigma(F)$	1170	1161
<i>R</i> (%) <i>all</i>	2.01	4.24
$\begin{array}{cc} R(\%) & \underline{F_0} & \text{please replace "zero" with the letter} \\ \text{"o"} & \text{replace } F_0 & \text{with } F_0 & \geq 4\sigma(F) \end{array}$	1.84	3.96
Weight	0.05	0.09
Goodness of fit	1.15	2.53

Table 4

Atom coordinates, displacement parameters ($Å^2$) and site occupancies of FOW and FRA CF.

FOW						FRA	
Site	x	Y lowercase:	Z lowercase:	U _{eq} ^a	s.o.f. ^b	Site	x
Ca1	2/3	1/3	0.00113 (3)	0.01055 (6)	1.00 (1)	Cal	2
Ca2	- 0.00730 (2)	0.24139 (2)	1⁄4	0.00883 (5)	0.99 (1)	Ca2	- (
Р	0.36903 (3)	0.39827 (3)	1⁄4	0.00628 (7)	0.96 (1)	Р	0 (
01	0.48506 (10)	0.32724 (11)	1⁄4	0.01188 (16)	1	01	0 (
$\begin{array}{c c} O & O2 \text{ is} \\ correct \\ please \\ substitute \\ O_2 \text{ with} \\ O2 & 2 & O2 \\ \text{is correct} \\ please \\ substitute & O_2 \\ \text{with } O2 \\ \end{array}$	0.46687 (11)	0.58793 (10)	1/4	0.01459 (17)	1	$\begin{array}{c c} O & O2 \text{ is} \\ correct \\ please \\ substitute \\ O_2 \text{ with} \\ O2 & 2 & O2 \\ \text{is correct} \\ please \\ substitute & O_2 \\ \text{with } O2 \\ \end{array}$	0 (
03	0.25709 (8)	0.34152 (9)	0.07053 (9)	0.01636 (14)	1.00 (2)	03	0 (
						F3	0(
F	0	0	1/4	0.0207 (3)	1	F	0
С	Not include	ed into the refin	ement			С	0

All atoms except for C were refined with anisotropic-displacement parameters. Values w during the refinement. The combined occupancies of the C–P and O3–F3 pairs of atoms

 $^{\rm a}U_{\rm eq}$ defined as one-third of the trace of the orthogonalized $U_{\rm ij}$ tensor

^bs.o.f. = site occupancy factor; ${}^{c}U_{iso}$ (the isotropic displacement exponent) takes the form

∢

	FOW	FRA
Ca1-O1 (x3)	2.3914 (6)	2.3983 (11)
Ca1–O2 (x3)	2.4501 (6)	2.4561 (13)
Ca1-O3 (x3)	2.7983 (7)	2.7855 (15)
<cal-o></cal-o>	2.547	2.547
Ca2–O1	2.6882 (9)	2.6816 (18)
Ca2–O2	2.3687 (9)	2.3709 (16)
Ca2–O3 (x2)	2.3461 (6)	2.3523 (12)
Ca2–O3′(x2)	2.4897 (7)	2.4896 (13)
<ca2–o></ca2–o>	2.455	2.456
Ca2–F	2.2925 (2)	2.2819 (4)
Ca2–Ca2	3.9708 (3)	3.9523 (7)
P-O1	1.5298 (8)	1.5241 (16)
$\begin{array}{c} P-O O2 \text{ is correct} \\ please \ substitute \ O_2 \ with \ O2 \\ \hline please \ substitute \ O_2 \ with \ O2 \\ \hline \end{array} \begin{array}{c} 2 O2 \ \text{is correct} \\ \hline \end{array}$	1.5364 (9)	1.5250 (17)
P-O3 (x2)	1.5312 (6)	1.5352 (13)
<p=0></p=0>	1.532	1.530
C-O1	_	1.48 (2)
C-O O2 is correct please substitute O ₂ with O2 2 O2 is correct please substitute O ₂ with O2	_	1.45 (2)
C-O3	_	1.40 (3)
<c-0></c-0>		1.446

Selected bond distances (Å) of FOW and FRA CFAP (esd in parentheses)

After characterization at room conditions, the two crystals of FOW and FRA CFAP were loaded concurrently into a ETH-type diamond-anvil cell (DAC, Miletich et al. 2000) for the high-pressure XRD investigations. A foil of T301 steel 250- μ m thick was used as a gasket to hold the crystal, and it was pre-indented to 110 μ m before drilling a hole by spark erosion (Ø 250 μ m). A 16:3:1 mixture of methanol:ethanol:water was used as hydrostatic pressure-transmitting

medium (Angel et al. 2007; Klotz et al. 2009). The internal pressure and the uncertainty on P were calculated using a single crystal of quartz as an internal pressure calibrant (using 12 reflections) (Angel et al. 1997). The unit-cell parameters were please remove the dot and do not wrap $\frac{1}{2}$

Table 6 <u>d</u> ATTENTION: the sentence is attached to the previous one _____etermined at room temperature in the *P* range 0.0001–6.86 GPa (Table 6) from the setting angles of 19–21 reflections centered in eight positions (King and Finger 1979; Angel et al. 2000), using a Siemens P4 four-circle X-ray diffractometer (Department of Earth Sciences, University of Torino) with a point detector (graphite-monochromatized MoKa radiation, θ -2 θ scan mode, variable scanning speed 2.4–30°/min); the SINGLE Control Program software (Angel and Finger 2011) was used.

Table 6

Unit-cell parameters of FOW and FRA CFAP as a function of pressure (esd in parentheses)

Р	FOW			FRA		
(GPa)	a (Å)	c (Å)	↓ - <i>V</i> (Å ³)	a (Å)	c (Å)	₩ - <i>V</i> (Å ³)
0.0001	9.3549 (8)	6.8739 (14)	520.97 (14)	9.330 (1)	6.8980 (14)	519.97(15)
0.02 (2)	9.3467 (8)	6.8704 (6)	519.79 (9)	9.3210 (11)	6.8976 (19)	518.98 (15)
0.25 (1)	9.3379 (8)	6.8665 (6)	518.52 (9)	9.310 (2)	6.895 (4)	517.5 (3)
0.72 (2)	9.3188 (8)	6.8590 (6)	515.84 (9)	9.2926 (14)	6.885 (2)	514.89 (19)
1.40 (2)	9.2940 (7)	6.8459 (6)	512.11 (9)	9.2699 (12)	6.869 (2)	511.19 (17)
1.62 (2)	9.2867 (8)	6.8429 (6)	511.09 (9)	9.2612 (11)	6.865 (2)	509.95 (16)
1.87 (2)	9.2788 (8)	6.8396 (6)	509.97 (9)	9.2527 (13)	6.861 (2)	508.69 (17)
2.18 (2)	9.2670 (8)	6.8336 (6)	508.23 (9)	9.2399 (12)	6.857 (2)	506.99 (16)
2.56 (2)	9.2530 (8)	6.8269 (6)	506.20 (10)	9.2282 (11)	6.8487 (19)	505.10 (15)
3.11 (3)	9.2336 (7)	6.8163 (5)	503.29 (8)	9.2088 (13)	6.836 (2)	502.03 (17)

Data collected during decompression

Р	FOW			FRA		
(GPa)	a (Å)	c (Å)	₩ - <i>V</i> (Å)	a (Å)	c (Å)	₩ - <i>V</i> (Å)
3.31 (2)	9.2277 (7)	6.8132 (5)	502.42 (8)	9.2019 (12)	6.835 (2)	501.19 (16)
3.63 (2)	9.2183 (6)	6.8089 (5)	501.08 (7)	9.1925 (11)	6.8278 (19)	499.66 (15)
4.05 (2)	9.2051 (8)	6.8022 (6)	499.16 (9)	9.1778 (11)	6.818 (2)	497.38 (16)
4.41 (2)	9.1946 (7)	6.7959 (6)	497.55 (9)	9.1667 (11)	6.812 (2)	495.75 (15)
4.94 (3)	9.175 (4)	6.791 (4)	495.1 (5)	9.1527 (11)	6.805 (2)	493.71 (15)
5.25 (4)	9.1704 (10)	6.7843 (8)	494.10 (12)	9.1435 (11)	6.8015 (19)	492.44 (15)
5.64 (3)	9.1570 (10)	6.7771 (7)	492.14 (11)	9.1314 (11)	6.7953 (19)	490.70 (15)
5.91 (6)	9.1487 (7)	6.7727 (5)	490.91 (8)	9.1225 (10)	6.7885 (17)	489.26 (13)
6.36 (4)	9.1373 (8)	6.7660 (6)	489.22 (10)	9.1110 (10)	6.7824 (17)	487.58 (13)
6.55 (3)	9.1319 (7)	6.7640 (5)	488.49 (8)	9.1051 (13)	6.780 (2)	486.80 (17)
6.86 (4)	9.1220 (9)	6.7574 (7)	486.96 (10)	9.0954 (11)	6.7724 (19)	485.20 (15)
6.78 (4) a	9.1249 (8)	6.7589 (6)	487.37 (9)	9.0966 (12)	6.773 (2)	485.36 (15)
6.69 (5) a	9.1269 (8)	6.7596 (6)	487.64 (9)	9.1001 (10)	6.7764 (18)	485.98 (14)
6.52 (4) ^a	9.1324 (9)	6.7628 (7)	488.46 (10)	9.1048 (11)	6.779 (2)	486.64 (15)
5.83 (3) ^a	9.1500 (8)	6.7721 (6)	491.01 (9)	9.1247 (13)	6.789 (2)	489.52 (17)
5.38 (3) ^a	9.1635 (8)	6.7792 (6)	492.98 (10)	9.1365 (12)	6.797 (2)	491.35 (16)
3.31 (3) ^a	9.2247 (15)	6.8132 (12)	502.10 (18)	9.2020 (12)	6.832 (2)	501.01 (17)

Results and discussion

FTIR spectroscopy

The incorporation of carbonate in fluorapatite was investigated by Fourier transform infrared (FTIR) spectroscopy. Figure 1 shows the FTIR spectra (2000-700 cm⁻¹) for FOW (Fig. 1a) and FRA (Fig. 1b) fluorapatite samples. The spectra display the absorption bands characteristic of carbonate-bearing apatites (Elliott 2002). The prominent absorption band at around 1070 cm^{-1} and the weak peak at 965 cm^{-1} are due to stretching vibrations of the PO₄ groups. Both spectra of the CFAP samples exhibit the typical absorption bands of CO₃ groups in the spectral region 1400–1600 cm⁻¹. The asymmetric-stretching bands of CO_3 (v_3) are observed at 1455 and 1430 cm⁻¹, and are characteristic of a type-B substitution, whereas the out-of-plane bending mode of $CO_3(v_2)$ is observed at ~ 865 cm⁻¹, in agreement with those reported in the literature (Elliott 1994; Wilson et al. 2004; Fleet 2009; Yi et al. 2013). The carbonate-band intensities (absorbance) in the FRA spectrum (Fig. 1b) are higher than those in the FOW spectrum (Fig. 1a), indicating a larger amount of carbonate in the FRA sample, in accord with the carbonate contents determined by electron microprobe. There is no evidence of $(CO_3)^{2-}$ ions substituting for OH⁻ ions (type-A substitution), as the characteristic absorption at ~ 1545 cm⁻¹ was not observed. In addition, no sharp peaks are observed in the OH-stretching region $(3700-3100 \text{ cm}^{-1}, \text{ not})$ shown in Fig. 1) which indicates the absence of OH⁻ ions in the structure of the apatite samples examined.

Fig. 1

The Fourier-transform infrared (FTIR) spectra for **a** FOW and **b** FRA carbonatebearing fluorapatite in the region 2000–700 cm⁻¹



Structure refinement at room temperature

The chemical formulae of CFAP samples, obtained from electron microprobe analysis (Table 2) and FTIR spectra, indicate the presence of carbonate in both samples and show an excess of F. In their analysis of fluor-chlorapatite, Mackie and Young (1974) noted (F + Cl) > 2 halogen atoms per formula unit (apfu) for their two samples, both by chemical analysis (2.06, 2.25 halogen apfu for two samples) and by X-ray site refinement (2.09, 2.19 apfu). They proposed the presence of an additional Cl site at (0,0,0) that they could not asses from their results. The presence of this site was unassailable in the accurate work of Hughes et al. (2014) who found an 8% occupancy of Cl at (0,0,0). Yet their microprobe analyses, with their coincidence with the stoichiometry of the reactants, demonstrate that there are no excess halogens in the F–Cl anion column of their samples. Our refinement shows no residual electron density along (0,0,z). Therefore, we believe that F excess in our samples is related to type-B CO₃ in our samples.

The estimated amount of carbonate is 1.6 wt% for FOW CFAP and 5.5 wt% for FRA CFAP. The higher content of carbonate in FRA with respect to FOW is consistent with the observed *a* unit-cell parameter reduction from 9.3527(1) to 9.3330(1) Å and the significant increase of the *c* parameter from 6.8752(1) to 6.8984(1) Å (Table 3). The results of the structure refinement at room temperature (Table 4) show that, within FRA CFAP, the B carbonate ion is located close to the substituted phosphate group and the C atoms occupies a

position (occupancy = 0.11) similar to the one observed by Wilson et al. (2004) in a sodium carbonate-bearing apatite and by Yi et al. (2013) in CFAP. In FOW CFAP, it was not possible to determine the position of the C atoms because of the low carbonate content.

As it is well known, within the type-B apatite the effect of the replacement of PO_4 ions by CO_3 ions on the structure is a reduction of the PO_4 tetrahedron volume together with a reduction of the P occupancy. Several studies (Elliott 1994; Wilson et al. 1999, 2004; Elliott et al. 2002) have noted that the occupancy of the phosphorus position gives the most reliable estimate of the extent of substitution by type-B carbonate. The occupancies of P in FOW and FRA CFAP (Table 4) become significantly less than unity, and these data are consistent with a substitution of the $(PO_4)^{3-}$ phosphate group by the $(CO_3)^{2-}$ carbonate ion. Within the FOW sample, the observed P occupancy is 0.96 (Table 4), equivalent to 2.88 P apfu, whereas, for the FRA sample, the observed P occupancy is 0.89, equivalent to 2.67 P apfu, i.e. substantially less than 3. The amounts of C atoms calculated directly from the P occupancies (C occ. = 0.04and 0.11) are 0.12 and 0.33 apfu, respectively, in agreement with the chemical analyses (0.13 and 0.45). For the FRA sample, the occupancy of the O3 site (Table 4) becomes less than unity (0.96) whereas some F (occ. = 0.04) is present within the same site, which corresponds to 0.22 F apfu, lower than the amount found by electron probe analysis. With regard to the P-O distances, Table 5 shows that the replacement of P^{5+} ion by C^{4+} ion causes a deformation of the PO₄ tetrahedron, with changes in the P-O bond lengths as the CO₃ content increases. As expected, the P-O1 and P-O2 distances became shorter whereas the P-O3 distances increase slightly. This configuration is in agreement with the findings of Yi et al. (2013), who concluded from FTIR, ¹⁹F DQ-SQ MAS NRM, and ${}^{13}C{}^{19}F{}$ frequency-selective REDOR experiments, that the planar carbonate oxyanion replacing the tetrahedral phosphate one, lies in the sloping face of the tetrahedron opposite to the fluoride ion occupying the remaining vertex. Yi et al. (2013) also propose a "francolite-type" diagnostic narrow IR absorption band at 864 cm⁻¹ that is indeed present in our FTIR spectrum and its absorbance is proportional to the amount of this "francolite-type" defect (i.e. O by F substitution at the O3 site).

Elastic behavior and equation of state

The FOW and FRA unit-cell parameters determined up to P = 6.86 GPa are reported in Table 6 and the variation of *a* and c parameters and volume with pressure are shown in Figs. 2 and 3, respectively. No discontinuities that could be related to any possible phase transition are present up to 6.86 GPa. In the *P* range investigated, both *a* and *c* parameters decrease with increasing pressure

(Fig. 2). The linear compressibility coefficients are $\beta_a = 3.63 \times 10^{-3}$ GPa⁻¹ and $\beta_c = 2.47 \times 10^{-3}$ GPa⁻¹ for FOW, and $\beta_a = 3.67 \times 10^{-3}$ GPa⁻¹ and $\beta_c = 2.65 \times 10^{-3}$ GPa⁻¹ for FRA CFAP, giving an axial anisotropy of β_a : $\beta_c = 1.47$:1 and 1.38:1, respectively. As previously seen in Table 3, the higher content of carbonate in FRA CFAP causes the reduction of the *a* parameter and an increase of the *c* one with respect to FOW. The average compressibility of the cell volume is $\beta_V = 9.5 \times 10^{-3}$ GPa⁻¹ for FOW and $\beta_V = 9.7 \times 10^{-3}$ GPa⁻¹ for FRA.

Fig. 2

Variation of the *a* and *c* unit-cell parameters with pressure in FOW (filled circles) and FRA (open circles) CFAP. Lines are the EoS fit to the linear data (solid line: FOW; dashed line: FRA). Experimental uncertainties (bars) for almost all points are smaller than symbols



Variation in unit-cell volume as a function of pressure for FOW and FRA CFAP. The lines show the second-order Birch–Murnaghan eEquation of sState (BM2-EoS) in the P range 0.0001–6.86 GPa. Symbols as in Fig. 2. Experimental uncertainties are smaller than symbols



The *P-V* data (Fig. 3) were fitted by a third-order Birch–Murnaghan Equation of State (BM3-EoS) using 26 pressure points. The resulting EoS coefficients, calculated using the EosFit7 suite software (Angel et al. 2014; Gonzalez-Platas et al. 2016), are $V_0 = 519.93(8)$ Å³, $K_{T0} = 89.9(1.1)$ GPa, K' = 4.7(3) for FOW and $V_0 = 519.08(13)$ Å³, $K_{T0} = 87.2(1.5)$ GPa, K' = 4.6(4) for FRA CFAP. The isothermal bulk modulus of FRA is lower than that of FOW, indicating that carbonate content influences the compressibility of CFAP.

The confidence ellipses in K_{T0} and K' for the BM3 EoS fit are reported in Fig. 4 and confirm that the compressibility of FRA CFAP (dashed line) is higher than that of FOW.

Fig. 4

Confidence ellipses in K_{T0} and K' for the fit of the third-order Birch–Murnaghan EoS to the fluorapatite *P*-*V* data at 95.4% level (2 σ) (solid line: FOW; dashed line: FRA)



When our *P-V* data are reformulated as a normalized pressure —- finite strain plot ($F_E - f_E$ plot, Angel 2000) (Fig. 5), the values lie on a horizontal line of constant F_E (K' = 4), indicating that the data are adequately described by a BM2-EoS. Therefore, the data can be fitted by a second-order truncation of the Birch– Murnaghan Equation of State. Fitting the *P-V* data to a BM2-EoS gives the resulting coefficients of $V_0 = 519.81(7)$ Å³, $K_{T0} = 92.1(3)$ GPa for FOW and of $V_0 = 518.95(9)$ Å³, $K_{T0} = 89.1(4)$ GPa for FRA samples (Table 1). BM2-EoS gives higher bulk moduli for both samples with respect to BM3-EoS, and confirms that type-B carbonate substitution reduces the isothermal bulk modulus of FAP.

Fig. 5

Normalized pressure $F_{\rm E}$ versus Eulerian finite strain $f_{\rm E}$ in FOW and FRA CFAP $(F_{\rm E} = P/3 \times f_{\rm E} \times (1+2 f_{\rm E})^{5/2}$ and $f_{\rm E} = [(Vo/V)^{2/3} -1]/2$, Angel 2000). The horizontal lines of constant $F_{\rm E}$ indicate that the *P*-*V* data could be fitted by a second-order truncation of the BM-EoS for both apatites. Symbols as in Fig. 2. Bars indicate experimental uncertainties



Figure 6 shows the variation of both *a* and *c* unit-cell parameters for FOW and FRA CFAP as function of pressure, normalized to their room-pressure values. Compressibilities along unit-cell axes were determined by fitting a BM2-EoS to *a* and *c* parameters in the range 0.0001–6.86 GPa. The coefficients obtained are $a_0 = 9.3461(5)$ Å, $1/3 M_i = 81.2(3)$ GPa; $c_0 = 6.8716(4)$ Å, $1/3 M_i = 124.2(7)$ GPa for FOW, and $a_0 = 9.3204(7)$ Å, $1/3 M_i = 80.7(4)$ GPa; $c_0 = 6.8981(10)$ Å, $1/3 M_i = 111.9(1.4)$ GPa for FRA, respectively (with i = *a*, *c* and the linear moduli $M_i = \beta_i^{-1}$, Angel et al. 2014). The linear moduli indicate axial-compression anisotropy with $\beta_a > \beta_c$ and confirm that the *a*-axis direction is more compressible than *c*-axis direction for both samples. As the elastic anisotropy $(M_c:M_a)$ ratios for of FOW and FRA are a little different (1.53 and 1.39, respectively), the two samples of CFAP show quite different elastic anisotropy, stronger on the compressibility along the *c*-axis.

Fig. 6

Variation of *a* and *c* unit-cell parameters of FOW and FRA CFAP as a function of pressure, normalized to their room-pressure values. Symbols as in Fig. 2. Lines correspond to linear fits in the *P* range 0.0001–6.86 GPa with linear moduli $M_i = 244(1)$ and 372(2) GPa for FOW and 242(1) and 336(4) GPa for FRA (with i = a, c and $M_i = \beta_i^{-1}$, Angel et al. 2014). Experimental uncertainties (bars) for almost all points are smaller than symbols



Figure 7 compares the bulk modulus, as a function of carbonate content within both FOW and FRA CFAP, with the values obtained by the compressibility experiments on apatite. In Fig. 7a the bulk moduli for FOW and FRA type-B carbonate-bearing FAP samples, along with those obtained on synthetic carbonate-free fluorapatites (Brunet et al. 1999; Comodi et al. 2001) are showed. Considering the values of K_{T0} obtained on FAP (Table 1), our results show that a 5.5 wt% carbonate content has a considerable effect on the elastic properties and significantly reduces the bulk modulus of FAP by about 9%. The very low values obtained by Matsukage et al. (2004) and Schouwink et al. (2010) in their FPA (Table 1) disagree with our results, probably because of the 5–6 mol% of chlorapatite content of their samples. Our results on the compressibility of CFAP are in good agreement with those obtained by Liu et al. (2011) on synthetic type A-B CHAP. In fact, the results of their high-pressure experimental investigation up to 10 GPa (LM006 and LM005 samples in Fig. 7b) show that the isothermal bulk modulus in hydroxylapatite is significantly reduced (about 15%) by a carbonate content up to 11 wt%.

Fig. 7

Bulk modulus (K_{T0}) as a function of carbonate content (wt% CO₃) for **a** fluorapatite and **b** hydroxylapatite. **a** Square: carbonate-free fluorapatite (Brunet et al. 1999); triangle: carbonate-free fluorapatite (Comodi et al. 2001); filled circle: FOW and open circle: FRA carbonate-bearing fluorapatites (this work). **b** Diamond: carbonate-free hydroxylapatite (Brunet et al. 1999); asterisks: carbonate-rich hydroxylapatite (LM006 and LM005 samples, Liu et al. 2011)



Conclusion

The substitution of the carbonate ion for either the monovalent channel anion (type-A substitution) or for the oxyanion ZO_4^{3-} (type-B substitution) is of particular biological and mineralogical interest. The HAP is the mineral species most similar to the inorganic component of bones; the carbonate is present within bone in a range of concentrations from ~5 to 8 wt%, and ~2 to 4 wt% in enamel, substituting primarily for phosphate (Pasteris 2016). Apatite is the necessary hard phase in natural bone and, therefore, its elastic properties are particularly important for the design of bioapatite substitutes (prosthesis). Powdered FAP is used in dental applications. In fact, fluorine hosted by CHAP is the important anticaries component of the dental enamel; there is only ~ 100 ppm F in the interior of dental enamel (Elliott 2002) but up to 1 wt% in the outermost micrometer-scale layer (Brudevold et al. 1956). Moreover, FAP is a potential host for the containment of high-level nuclear waste, as indicated by its occurrence in the natural reactor at Oklo, Gabon (Bros et al. 1996).

Our study confirms the anisotropic elastic behavior of apatite, even in the presence of less common components as the CO_3 . This has to be considered

when elasticity of apatite is used for pressure determination like apatite-ingarnet barometer used by Ashley et al. (2017) because as discussed by Angel et al. (2015) the anisotropic behavior of the inclusion can modify the elastic response of the inclusion. In fact, the anisotropy could be treated as a deviatoric effect relative to the isotropic case (Angel et al. 2015).

Whereas there has been discussion on the carbonate substitution in fluorapatite, CFAP with only B carbonate has not been studied so far by single-crystal X-ray structure refinement (Fleet and Liu 2008a). Additionally, prior to the present study, there were no data available on the elastic properties of CFAP. The results of this study show that the replacement of P by $(CO_3)^{2-}$ (type-B substitution) involves a softening of the bulk modulus, as previously observed in CHAP (Liu et al. 2011). Whereas data on CHAP cannot discern between the role of type-A and type-B substitutions (as the samples studied by Liu et al. 2011 contained both types of substitution), the results of our study clearly reinforce the prominent role of type-B substitution in the softening.

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Electronic supplementary material

Below is the link to the electronic supplementary material.

Supplementary material 1 (CIF 56 KB)

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