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Combined effect of citrate and fluoride ions on hydroxyapatite nanoparticles

Lorenzo Degli Esposti^{a, *}, Alessio Adamiano^a, Anna Tampieri^a, Gloria Belen Ramirez-Rodriguez^b, Dritan Siliqi^c, Cinzia Giannini^c, Pavlo Ivanchenko^d, Gianmario Martra^d, Feng-Huei Lin^{e,f}, José Manuel Delgado-López^b, Michele Iafisco^{a,*}

^aInstitute of Science and Technology for Ceramics (ISTEC), National Research Council (CNR), Via Granarolo 64, 48018 Faenza (Italy)

^bDepartment of Inorganic Chemistry, University of Granada, Av. Fuente Nueva, s/n, 18071, Granada (Spain)

^cInstitute of Crystallography (IC), National Research Council (CNR), Via Amendola 122/0, 70126 Bari (Italy)

^dDepartment of Chemistry and Interdepartmental Centre "Nanostructured Interfaces and Surfaces-NIS", University of Torino, Via P. Giuria 7, 10125, Torino (Italy)

^eDepartment of Biomedical Engineering, National Taiwan University, 10617 Taipei (Taiwan)

^fInstitute of Biomedical Engineering and Nanomedicine, National Health Research Institutes, Keyan Road 35, 35053 Miaoli (Taiwan)

Authors for correspondence:Dr. Michele Iafisco, Ph. D.Institute of Science and Technology for Ceramics (ISTEC)National Research Council (CNR),Via Granarolo 64, 48018 Faenza (RA), Italy.E-mail: michele.iafisco@istec.cnr.it

Lorenzo Degli Esposti, M. Sc. Institute of Science and Technology for Ceramics (ISTEC) National Research Council (CNR), Via Granarolo 64, 48018 Faenza (RA), Italy. E-mail: lorenzo.degliesposti@istec.cnr.it **FT-IR spectra of citrate-HA.** The IR-ATR spectra of the citrate-HA samples are reported in Figure S1B. All samples displayed a main broad band at 1030 cm⁻¹ with shoulders at 1046 and 1075 cm⁻¹ due to the triply degenerated anti-symmetric stretching mode of the apatitic PO₄ groups (v_3PO_4). Other features emerge at 961 cm⁻¹ (symmetric stretching mode of the apatitic PO₄ groups, v_1PO_4) and at 603, 576 (as a shoulder) and 565 cm⁻¹ (triply degenerated bending mode of the same groups, v_4PO_4). Apatitic hydroxyl groups are evinced by a band at 631 cm⁻¹ that is due to OH⁻ librational modes [1]. The presence of traces amounts of carbonate ions is revealed by the presence of a very weak B-type carbonate substitution (CO₃ occupying PO₄ sites) band at 873 cm⁻¹ [2].



Figure S1. (A) PXRD diffractograms of citrate-HA 5m, citrate-HA 30m, and citrate-HA 4h. (B) FT-IR spectra of citrate-HA 5m, citrate-HA 30m, and citrate-HA 4h.



Figure S2. Plot as a function of the maturation time of (A) *a* cell axis, (B) *c* cell axis of citrate-FHA and citrate-HA.

Sample	<i>a-b</i> cell axes (Å)	c cell axis (Å)	D ₍₀₀₀₂₎ (nm)	D ₍₃₁₋₄₀₎ (nm)	D ₍₀₀₀₂₎ / D ₍₃₁₋₄₀₎
Citrate-HA 5m	9,499	6,898	21,6±0,5	5.5±0,4	3,9
Citrate-HA 30m	9,472	6,895	28,6±0,5	6,3±0,7	4,3
Citrate-HA 4h	9,462	6,891	42,9±0,5	7.3±0,6	5,9

Table S1. Cell parameters and crystal domain of citrate-HA samples.



Figure S3. Plot as a function of the maturation time of (A) $D_{(0002)}$ crystal domain, (B) $D_{(31-40)}$ crystal domain and (C) $D_{(0002)}/D_{(31-40)}$ ratio of citrate-FHA and citrate-HA.

Sample	Ca/P ^a (mol)	F ^b (% wt)	Citrate ^c (% wt)	Carbonate ^c (% wt)	ζ-Potential (mV)
Citrate-HA 5m	1,54±0,02	-	4,8±0,3	1,4±0,2	-13.1 ± 0.4
Citrate-HA 30m	1,54±0,02	-	3,2±0,3	0,8±0,1	-12.4 ± 0.4
Citrate-HA 4h	1,53±0,01	-	2,6±0,2	1,2±0,1	-9.3 ± 0.4

Table S2. Chemical composition of citrate-HA samples.

^(a)Quantified by ICP-OES; ^(b)Quantified by fluoride ion electrode; ^(c)Quantified by TGA.

Thermogravimetric analysis. Thermogravimetric curves and their first derivatives of all the samples mainly shows four weight losses (Figure S4): (i) from room temperature to 200°C due to the adsorbed water, (ii) from 200°C to 400°C related to structural water, (iii) from 400 to 600°C due to the citrate, and (iv) from 600 to 1000°C corresponding to the carbonate ions [3].



Figure S4. TGA and DTG curve of citrate-FHA 4h. The TGA curves of the other samples have the same profile.

Raman spectroscopy. Figure S5A-B shows the Raman spectra of the samples. The most intense peak appears at 960 cm⁻¹, which corresponds to v_1PO_4 mode. Other features from apatitic PO₄ group emerge at 1042 cm⁻¹ (v_3PO_4), 586 (v_4PO_4) and 428 cm⁻¹ (v_2PO_4). In the spectrum of both citrate-FHA and citrate-HA nanoparticles, very weak B-type carbonate bands appeared at 1070 cm⁻¹ (v_1CO_3) and at 1430 cm⁻¹ (v_2CO_3), confirming the data of IR-ATR spectra and chemical analysis [2]. Moreover, the Raman spectrum of citrate-HA samples exhibit an intense peak at 3570 cm⁻¹ (Figure S5C-D) associated to the apatitic vOH mode. The intensity of this peak increases with maturation time. On the contrary, this band is not present in the spectrum collected for the citrate-FHA nanoparticles, further confirming that OH-ions were completely replaced by fluoride ions. The bands at 2930 and 845 cm⁻¹ are related to the vCH₂ and the δ OCO modes of citrate [4], respectively, and their relative intensity decreases with maturation time as observed in the IR-ATR spectra.



Figure S5. Raman spectra of (A) of citrate-FHA 5m, citrate-FHA 30m, citrate-FHA 4h and (B) citrate-HA 5m, citrate-HA 30m, and citrate-HA 4h. (C) and (D) show an enlarged view of the OH and CH stretching modes spectral region for citrate-FHA and citrate-HA, respectively.

FT-IR spectroscopy in controlled atmosphere mode (Figure S6). The v _{asym}COO mode of citrates falling in 1750-1500 cm⁻¹ range overlaps with δ H₂O mode of adsorbed water (centered at ca. 1645 cm⁻¹). Thus, the detailed analysis of citrate profile requires the complete removal of the adsorbed water. To this aim, the sample in contact with H₂O at 20 mbar (curve a) was outgassed at b.t. for 60 min (curve b). Further, surface-accessible water along with OH species was exchanged with D₂O (δ D₂O is located at 1200 cm⁻¹) by 10 cycles of contacting the sample with 20 mbar of D₂O for 5 min followed by 5 min outgassing with subsequent 60 min outgassing at b.t. (curve c). The resulting profile is 'cleaned' from the contribution of water vibrations in the 1750-1500 cm⁻¹ range.



Figure S6. FT-IR spectra of citrate-FHA 4h. Curve (a) in contact with H₂O vapor at 20 mbar; curve (b) after 60 min outgassing at b.t.; curve (c) after exchange with D₂O and subsequent 60 min outgassing at b.t. Inset: zoomed view of 1750-1620 cm⁻¹ range. *small feature due to non-compensated atmospheric CO₂.



Figure S7. Plot as a function of the maturation time of (A) length, (B) width, and (C) aspect ratio of citrate-FHA nanoparticles evaluated by TEM. A comparison with the respective crystal domains calculated by PXRD patterns is reported in each panel.



Figure S8. FEG-SEM micrographs of (A) citrate-FHA 4h, and (B) citrate-HA 4h.



Figure S9. SAXS curves and data fitting with lamellar model for (A) of citrate-FHA 5m, (B) citrate-FHA 30m, (C) citrate-FHA 4h, (D) of citrate-HA 5m, (E) citrate-HA 30m, and (F) citrate-HA 4h.

Table S3. Mean thickness and polydispersity of citrate-HA samples extracted as lamellar model fitting SAXS data.

Sample	Thickness (nm)	Polydispersity
Citrate-HA 5m	6,8	0,47
Citrate-HA 30m	7,1	0,70
Citrate-HA 4h	5,5	0,30

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