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Original

Novel multifunctional strontium-copper co-substituted mesoporous bioactive particles / Bari, A., Molino, G., Fiorilli, S.L., VITALE BROVARONE, C.. - In: MATERIALS LETTERS. - ISSN 0167-577X. - ELETTRONICO. - 233:(2018), pp. 37-40. [10.1016/j.matlet.2018.04.006]

Availability:

This version is available at: 11583/2707715 since: 2018-05-18T18:45:01Z

Publisher:

elsevier

Published

DOI:10.1016/j.matlet.2018.04.006

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Featured Letter

Novel multifunctional strontium-copper co-substituted mesoporous bioactive particles



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ARTICLE INFO

Article history:

Received 26 January 2018

Received in revised form 28 March 2018

Accepted 2 April 2018

Available online 3 April 2018

Keywords:

Mesoporous bioactive glasses

Therapeutic ions

Ion co-substitution

Biomaterials

Sol-gel preparation

ABSTRACT

Sr-Cu co-substituted mesoporous bioactive glasses were synthesized by two different sol-gel approaches: an ultrasound-assisted base-catalyzed sol-gel procedure and an aerosol-assisted spray-drying method. The produced spherical shaped particles showed high specific surface area values and tunable pore size. The two different routes allowed obtaining samples with different morphology and size, characterized by the ability to induce hydroxyapatite deposition, to incorporate a specific amount of therapeutic ions (strontium with the aim to improve bone formation and copper for its antibacterial and pro-angiogenic properties) and to release them. The obtained multifunctional biomaterials synergistically combine the ability to promote bone formation and angiogenesis and to impart an antibacterial effect.

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1. Introduction

Mesoporous bioactive glasses (MBGs) have been extensively studied for bone repair [1] and, more recently, for soft tissue applications [2]. The incorporation of metal ions has gained increasing interest to impart different biological functions and enhance their therapeutic potential [3,4].

Among these ions, strontium is one of the most attractive for its role in enhancing bone cell activity and inhibiting osteoclast differentiation [5]. An essential and complementary role in tissue regeneration could be performed by copper due to its capability to provide antibacterial properties [6] and to promote angiogenesis [7]. In the literature, the therapeutic effect associated to the single ion substitution is well established [5–7], and with the aim to combine different biological effects, the joint presence of two different therapeutic ions (i.e. Sr/Co and Cu/Zn) have been investigated for hydroxyapatites [8,9] and bioactive glasses [10,11]. Huang and co-workers [9] fabricated a Sr-Cu co-substituted hydroxyapatite coating and demonstrated the potential to combine in a single formulation the antimicrobial property and the osteogenic differentiation stimulus. In this contribution, for the first time to the best of authors' knowledge, novel multifunctional Sr-Cu-containing MBGs, with binary SiO₂-CaO composition, have been synthesized by two different approaches: an ultra-sound assisted sol-gel technique (US) [6] and an aerosol-assisted spray-drying method (SD) [12].

These two routes allow to obtain MBG particles with adjustable features (i.e. size, surface area, pore size), able to co-release different therapeutic ions and to potentially incorporate and delivery drugs and/or biomolecules.

2. Materials and methods

2.1. SrCu_MBG_2%_US and SrCu_MBG_2%_SD synthesis

Sr-Cu-containing MBGs (molar ratio Sr/Cu/Ca/Si = 1/1/13/85) were produced by modifying an ultra-sound assisted base catalyzed sol-gel procedure previously reported by the authors [6]. First, CTAB was dissolved by stirring in absolute methanol, whose pH was adjusted to 12.5 through addition of NH₄OH. Into this solution, Ca(NO₃)₂·4H₂O, CuCl₂ and SrCl₂ were dissolved. Then TEOS, firstly dissolved in absolute methanol, was added to the first solution with the application of ultrasounds (150 W in a 10 s on/10 s off cycle for 20 min). After 24 h of stirring, the precipitate was washed by centrifugation and dried at 70 °C for 12 h and the final precipitate (named hereafter SrCu_MBG_2%_US) was calcined at 600 °C in air (5 h, 1 °C/min).

The spray-drying Sr-Cu substituted MBGs (molar ratio Sr/Cu/Ca/Si = 1/1/13/85) were produced following the procedure reported by Molino et al. [12]. Briefly, Pluronic P123 was dissolved in bidistilled water by stirring. A solution of TEOS, pre-hydrolyzed for 1 h in diluted HCl (pH 2), was poured in the previous one and stirred for 20 min. SrCl₂ and CuCl₂ were simultaneously added to this batch. Ca(NO₃)₂·4H₂O was added to the solution 15 min before

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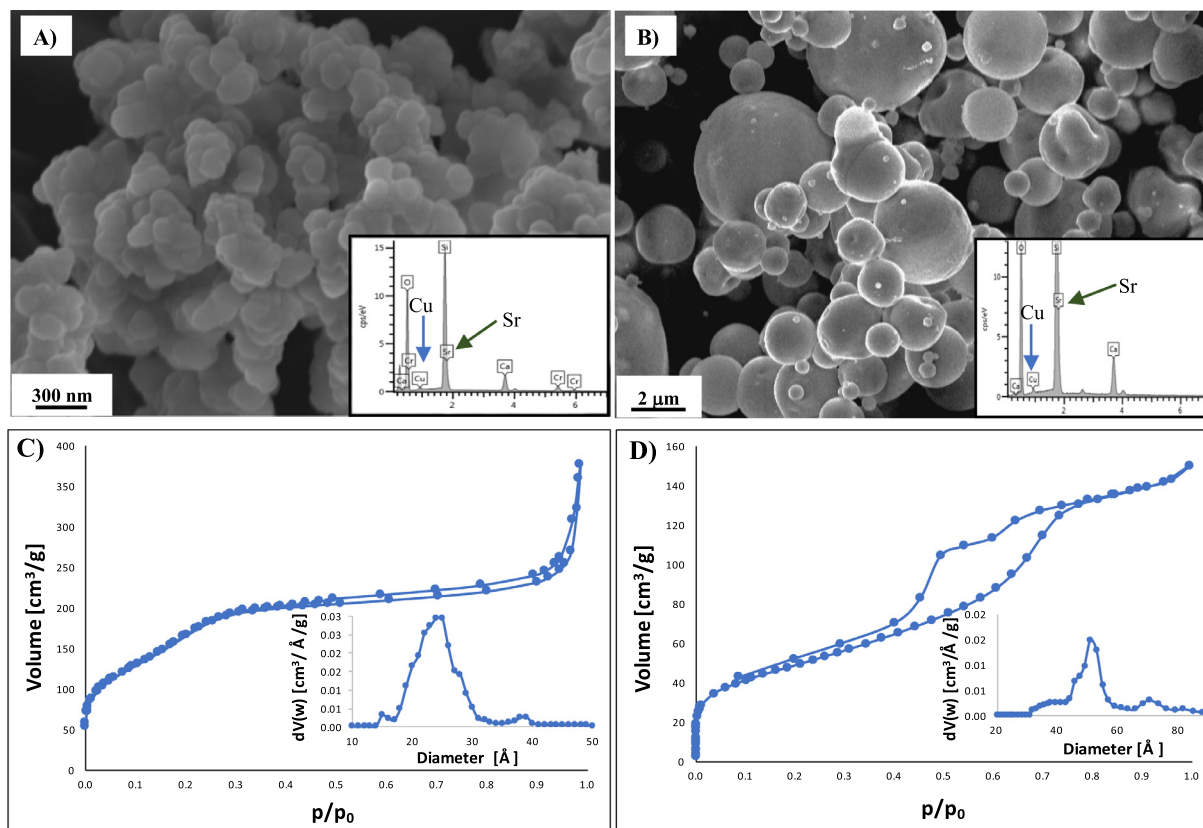


Fig. 1. FE-SEM image of SrCu_MBG_2%_US (A) and SrCu_MBG_2%_SD (B) with related EDS spectra (inset); isotherm curve of SrCu_MBG_2%_US (C) and SrCu_MBG_2%_SD (D) with related pore size distributions (inset).

spraying using a Mini Spray-Dryer B-290 (Buchi). The collected powder (named hereafter SrCu_MBG_2%_SD) was calcined in air at 600 °C (5 h, 1 °C/min).

2.2. Characterization

The particle morphology was analyzed using a FE-SEM microscope (Philips 525 M) equipped with EDS analyzer (Philips Edax 9100). The crystalline nature of HA deposits was investigated by wide-angle XRD analysis (XRD, X'Pert PRO, PANalytical). N₂ adsorption-desorption analysis was performed on powders outgassed at 423 °K for 5 h by using a Quantachrome Autosorb 1. The BET (Brunauer–Emmett–Teller) equation was adopted to calculate the specific surface area (SSA_{BET}) in the relative pressure range of 0.04–0.1. The pore size distribution was studied through DFT (Density Functional Theory) using the NLDFT equilibrium model. To evaluate the sample bioactivity, 30 mg of powder were kept immersed up to 14 days in simulated body fluid (SBF) with a concentration of 1 mg/mL inside an orbital shaker (37 °C, 150 rpm) [13].

Ion release was assessed by soaking powder in SBF with a concentration of 1 mg/mL for 3 h, 1, 3 and 7 days. At each time point, 1 mL of supernatant was collected after centrifugation and analysed by ICP (ICPMS Thermoscientific ICAPO spectrometer).

3. Results and discussion

SrCu_MBG_2%_US particles were spheroidal, slightly aggregated, with a size of about 100 nm (Fig. 1A). SrCu_MBG_2%_SD showed spherical particles ranging between 500 nm and 5 μ m. For both samples, EDS analysis (inset in Fig. 1A and B) revealed that the incorporation successfully occurred for both ions.

Table 1

Textural properties of samples.

Samples	SSA_{BET} (m ² /g)	Pore Volume (cm ³ /g)	DFT Pore Size (nm)
SrCu_MBG_2%_US	470	0.23	2.5
SrCu_MBG_2%_SD	176	0.21	5

N₂ adsorption-desorption measurement showed IV type isotherms for both samples (Fig. 1C and D), confirming their mesoporous structure. SrCu_MBG_2%_US curve exhibited a well-defined step between 0.1 and 0.2 p/p_0 , indicative of the filling of regular mesopores. Pore size distribution in Fig. 1C (inset) revealed an average pore diameter of 2.5 nm. SrCu_MBG_2%_SD isotherm showed a hysteresis loop corresponding to the filling of mesopores and pore size distribution revealed an average pore diameter of 5 nm (Fig. 1D – inset). Although for spray-dried sample the specific surface area is lower (Table 1), the value is still remarkably higher compared to that of not-templated sol-gel glasses [14]. These data highlighted that the ion co-substitution did not significantly affect the textural properties of the final powder that resulted only slightly different from Sr- or Cu-containing MBGs [6,12].

In vitro bioactivity assessment highlighted that partial replacing of Ca with Cu and Sr ions did not significantly hamper the hydroxyapatite nucleation and deposition. After 1 day of soaking, the Ca/P ratios revealed by EDS analysis were 2.89 and 1.63 for SrCu_MBG_2%_US and for SrCu_MBG_2%_SD, respectively. After 7 days of soaking in SBF the samples were fully covered by a layer of needle-like crystals (Fig. 2) and, for both samples, the Ca/P ratio was in good accordance to the stoichiometric value of hydroxyapatite (1.67). The formation of hydroxyapatite layer was confirmed by wide-angle XRD analysis (Fig. S1).

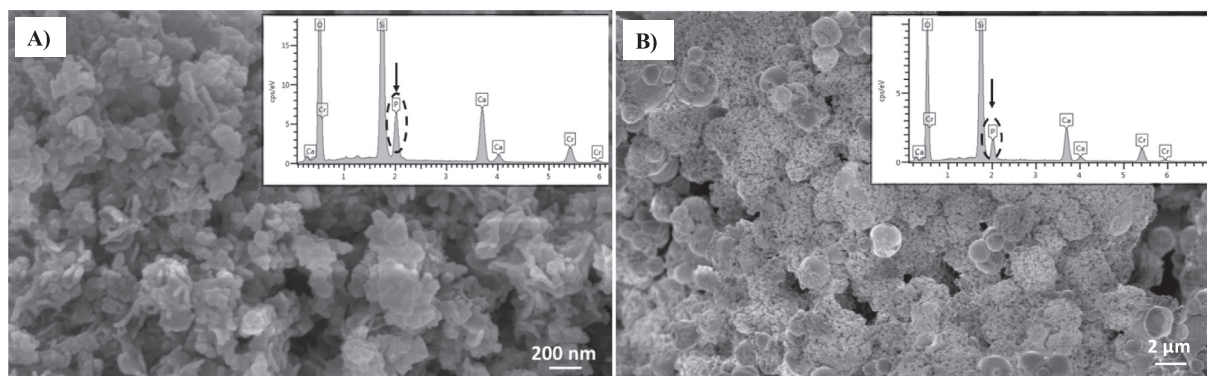


Fig. 2. (A) SrCu_MBG_2%_US and (B) SrCu_MBG_2%_SD after 7 days of soaking in SBF with EDS spectra (inset).

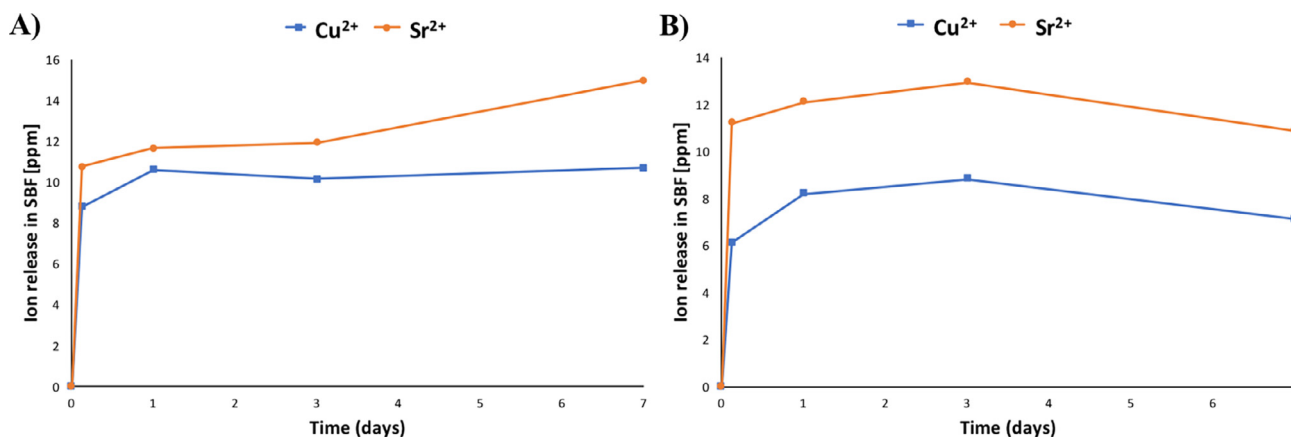


Fig. 3. Ion release curve of SrCu_MBG_2%_US (A) and SrCu_MBG_2%_SD (B).

The pH value of SBF was below 7.8 during the entire test, which is considered an optimal value for osteoblasts [15].

The incorporated Cu²⁺ and Sr²⁺ were successfully co-released in SBF by both types of MBGs (Fig. 3A and B). SrCu_MBG_2%_US showed a burst release of both Cu²⁺ and Sr²⁺ ions within the first 3 h of incubation (9 and 11 ppm respectively). After 1 day of soaking, a lower diffusion rate was observed and could be ascribed to the progressive occlusion of mesopores due to the dissolution of silica framework and its re-precipitation in the form of silica gel at the pore entrance [16].

Similarly, a burst release within the first hours was observed for SrCu_MBG_2%_SD (6 and 11 ppm respectively) followed by a slower increase in the released concentration up to 3 days (9 and 13 ppm respectively). After this time point, the released concentration was constant suggesting that the precipitation of a hydroxyapatite layer could partially hinder further ion release. The increasing trend of Sr²⁺ release profile after 7 days (Fig. 3A) suggests the capability of SrCu_MBG_2%_US to release a further amount of this ion specie, differently from SrCu_MBG_2%_SD.

The higher release of Sr²⁺ compared to Cu²⁺ suggests a higher accessibility and ion-exchange ability of Sr²⁺ within the MBG framework compared to incorporated copper species, most likely due to its similarity with calcium as network modifier [5].

4. Conclusions

For the first time, strontium-copper co-substituted SiO₂/CaO based mesoporous particles were synthesized by two different routes. Both procedures allowed to obtain powders with excellent

bioactive behavior, due to the high exposed specific surface area and open porous structure, and able to simultaneously co-release Sr²⁺ and Cu²⁺ in order to impart an antibacterial effect against different bacteria strains [6] and to enhance bone re-modelling [17]. Based on these results, co-substituted MBGs deserve further scrutiny as candidates for treating bone related pathologies in the presence of bacterial infection.

Acknowledgements

This project has received funding from the European Research Council (ERC) under the European Union's Horizon 2020 research and innovation programme (grant agreement No 681798-BOOST) (www.ercprojectboost.eu).

Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at <https://doi.org/10.1016/j.matlet.2018.04.006>.

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