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1. Introduction

Aerogels were invented in the early 1930's by Kistler^{1,2} and are defined as solid colloidal or polymeric networks of nanoparticles expanded throughout their entire volume by a gas.^{3,4} Early emphasis was based on silica and other inorganic aerogels, $1,2,5$ while organic aerogels, based on organic polymers⁶⁻¹² or biopolymers,¹³⁻¹⁶ and hybrid organic-inorganic aerogels¹⁷⁻²¹ were developed later.

Alginates are polysaccharides obtained from brown algae. They are block copolymers of two diastereomeric monomers, b- $(1\rightarrow 4)$ -linked D-mannuronic acid (M) and α - $(1\rightarrow 4)$ -linked Lguluronic acid (G; Fig. 1). Alginate aerogels are among the most well-studied biopolymer-based aerogels, as their synthesis is simple, inexpensive and environmentally friendly²²⁻²⁷ and they can also be carbonized in good yields.²⁸ Alginate hydrogels can

Patrina Paraskevopoulou, D^{*a} Irina Smirnova, ^{*b} Ta[ma](http://orcid.org/0000-0002-9012-2495)ra Athamneh, *i*^b Maria Papastergiou[,](http://orcid.org/0000-0002-0528-3930)^a Despoina Chriti,^a Gregor Mali, ^{D c} Tomaž Čendak, D ^c Grigorios Raptopoulos^a and Pa[v](http://orcid.org/0000-0003-0598-243X)el Gurikov D^{*d}

Polyurea-crosslinked calcium alginate (X-Ca-alginate) and chitosan (X-chitosan) aerogel beads have been prepared via reaction of an aliphatic triisocyanate (Desmodur N3300) with the –OH groups of calcium alginate or the $-NH₂$ groups of chitosan preformed wet gels, forming surface urethane or urea groups, respectively, and the subsequent formation of a polyurea conformal coating of the network via reaction of the triisocyanate with adsorbed water retained on the inner surfaces of the wet gels. That conformal coating is evident by SEM, which showed that all aerogels, native and crosslinked, were fibrous and their micromorphology did not change after crosslinking. The chemical identity of the new materials was confirmed with ATR-FTIR spectroscopy and solid-state 13 C and 15 N CPMAS NMR spectroscopy. The percent uptake of polyurea in X-Ca-alginate aerogels (43–66%) and X-chitosan aerogels (31–51%) was calculated from skeletal density data. This work expands the methodology for polymer-crosslinked alginate aerogels from monoliths to beads, and from alginates to chitosan, establishing its generality for all biopolymer aerogels. Possible applications of these materials include environmental remediation and conversion to carbon aerogels. PAPER

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be prepared *via* gelation of sodium alginate in aqueous solutions using divalent (e.g., Ca^{2+} , Ba^{2+} , Co^{2+} , Ni^{2+} , Cu^{2+} , or Zn^{2+}) or trivalent (e.g., Fe³⁺ or Al³⁺) cations, which act by binding to the carboxylate groups of different alginate strands.^{26,29,30}

Chitosan is the deacetylated derivative of chitin; it is also a linear polysaccharide consisting of randomly distributed β -(1– 4)-linked D-glucosamine (usually >80% w/w/; deacetylated unit) and N-acetyl-D-glucosamine (acetylated unit; Fig. 1). Chitosan is soluble in slightly acidic aqueous solutions and can be gelled either by increasing the pH of the solution, 31 or by introducing ionic metal species.³²⁻³⁴

Alginate and chitosan hydrogels can be converted into the corresponding aerogels via solvent exchange with ethanol or acetone, and drying from supercritical fluid (SCF) $CO₂$. Both biopolymer aerogels have several desirable features: (a) they come from natural products; (b) the sol–gel process is carried out in water; (c) they bear a large number of functional groups available for coordination to a wide array of metal ions; and, (d) they can be converted pyrolytically to carbon aerogels with ultra-high open porosities and surface areas.²⁸ In addition to classical aerogel applications in thermal insulation, catalysis and adsorption, both alginate and chitosan aerogels are biocompatible, biodegradable and non-toxic materials, and can be used in medical applications, e.g., in drug delivery, as antimicrobial and wound healing biomaterials, and in cosmetics.15,35,36

The main disadvantage of most biopolymer aerogels, including those of alginate and chitosan, is that they are mechanically weak materials. This has been rectified recently,

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a Inorganic Chemistry Laboratory, Department of Chemistry, National and Kapodistrian University of Athens, Panepistimiopolis Zografou, 15771, Athens, Greece. E-mail: paraskevopoulou@chem.uoa.gr

^bInstitute of Thermal Separation Processes, Hamburg University of Technology, Eißendorfer Straße 38, 21073 Hamburg, Germany. E-mail: irina.smirnova@tuhh.de c National Institute of Chemistry, Hajdrihova 19, 1000 Ljubljana, Slovenia

^dLaboratory for Development and Modelling of Novel Nanoporous Materials, Hamburg University of Technology, Eißendorfer Straße 38, 21073 Hamburg, Germany. E-mail: pavel.gurikov@tuhh.de

[†] Electronic supplementary information (ESI) available: Size distributions of aerogel beads, TGA plots, N_2 -sorption diagrams, SEM images. See DOI: 10.1039/d0ra07337g

[‡] Current address: Jadara University, P.O. Box 733, Irbid 21110, Jordan.

Fig. 1 The structures of calcium alginate, chitosan and Desmodur N3300.

by synthesizing polyurea/polyurethane-crosslinked alginate (X-alginate) aerogels, 37 via reaction of a preformed calcium alginate network with an aliphatic triisocyanate that was introduced, post-gelation, to the pores of the wet gel via diffusion, in analogy to the crosslinking of silica and other inorganic aerogels.18,19,38–⁴⁰ The reaction formed urethane groups on the surface of the alginate framework, and a polyurea (PUA) network connecting those urethane groups, resulting in a conformal nano-thin film of PUA around the entire alginate network. Although X-alginate aerogels are essentially copolymers, they are different from literature materials based on: (a) mixtures, blends, or composites of polyurethanes and alginates, (b) polyurethane/alginate copolymers, or (c) products from urethane bond formation between alginates and isocyanates.⁴¹–⁴⁵ This is because the relative topology of the alginate network and the crosslinker is defined at the nanoscopic scale rather than at the molecular level. Xalginate aerogels are mechanically-strong materials, in contrast to their native counterparts, which are extremely weak. Compared to various organic aerogels from the literature, X-alginate aerogels can be as stiff as many other polymeric aerogels, with two- or three-times higher densities than those of X-alginate aerogels.³⁷ PSC Advances

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In this work we have prepared X-alginate aerogels in the form of beads, which have several advantages compared to monoliths, for example fewer defects,⁴⁶ easier processing via accelerated solvent exchange and continuous supercritical drying,⁴⁷ and closer proximity to applications.⁴⁸⁻⁵⁴ Indeed, although isolated cases of aerogel beads based on inorganic materials, 51,54-56 and organic polymers¹⁰ have also been reported, almost all biopolymer aerogels in the literature are prepared in the form of beads, because, in addition to the above advantages, fast gelation renders the procedure of making biopolymer aerogel beads more straightforward than making monoliths, and easily scalable.48,49,57,58 Finally, this new class of materials has been expanded to include crosslinked chitosan (X-chitosan) aerogel beads, thus suggesting the generality of this method for other biopolymer aerogels.

2. Experimental section

2.1 Materials and methods

Two different sodium alginate sources with different guluronic acid (G) content were used: PROTANAL LF 240 D, G 30–35% (referred to as G35) and PROTANAL LF 200 S, G 65–75% (referred to as G75). Chitosan (200–500 mPa s, 0.5% in 0.5%

acetic acid at 20 °C; degree of deacetylation $> 80\%$) was purchased from TCI chemicals. $CaCl₂$ and NaOH were purchased from Sigma. Desmodur N3300 (trimer of hexamethylene diisocyanate, an aliphatic triisocyanate) was generously provided by Covestro AG. MeCN (HPLC grade) and acetone were purchased from Fisher, and glacial acetic acid was purchased from Panreac, and they were used as received.

SCF drying was carried out in an autoclave (E3100, Quorum Technologies, East Sussex, UK). Wet-gels were placed in the autoclave at 12 °C and were covered with acetone. Liquid $CO₂$ was allowed in the autoclave; acetone was drained out as it was being displaced by liquid $CO₂$ (five times, 30 min each). Afterwards, the temperature of the autoclave was raised to 45 $^{\circ}$ C and was maintained for 1 h. Finally, the pressure was gradually released, allowing SCF $CO₂$ to escape as a gas, leaving dry-gels (aerogels).

 13 C and 15 N Cross-Polarization Magic Angle Spinning (CPMAS) NMR spectra were obtained with a 600 MHz Varian spectrometer (Varian, Palo Alto, CA, USA) operating at 150.73 MHz for 13 C and 60.74 MHz for 15 N nuclei. In 13 C CPMAS measurements samples were spun with the MAS frequency of 20 kHz , whereas in $15N$ CPMAS measurements the samples were spun at 10 kHz. In all CPMAS experiments an amplitude ramp during the CP block and high-power decoupling during acquisition were employed on the proton channel. Chemical shifts in $13¹³C$ NMR spectra are given vs. tetramethylsilane and in $15¹⁵N$ NMR spectra *vs.* liquid NH_3 .

ATR-FTIR spectra were obtained with a Perkin Elmer Spectrum 100 Spectrometer.

The thermal stability of the materials was studied by thermogravimetric analysis (TGA) employing a Mettler-Toledo TGA/ DSC1 instrument (Schwerzenbach, Switzerland). Samples were placed in platinum crucibles. An empty platinum crucible was used as a reference. Samples were heated from ambient temperatures to 800 °C in a 50 mL min⁻¹ flow of N₂ at a heating rate of 10 $^{\circ}$ C min⁻¹.

N₂-sorption measurements were made on a Micromeritics Tristar II 3020 surface area and porosity analyzer (Micromeritics, Norcross, GA, USA). Native biopolymer aerogels were degassed at 50 \degree C for 24 h and X-biopolymer aerogels were degassed at 80 °C for 24 h. Skeletal densities (ρ_s) were determined by He pycnometry, using a Micromeritics AccuPyc II 1340 pycnometer (Micromeritics, Norcross, GA, USA). Bulk densities (ρ_b) of the samples were calculated from their weight and natural dimensions.

Scanning Electron Microscopy (SEM) characterization was conducted with Pt-coated samples adhered onto a conductive double-sided adhesive carbon tape, using a high resolution FESEM JEOL JSM 7401f.

2.2 Synthesis of biopolymer (Ca-alginate and chitosan) wetgel and aerogel beads

Ca-alginate and chitosan hydrogels were prepared by slight modifications of literature procedures.^{25,30,31}

A solution of sodium alginate in $H₂O$ (1 or 2% w/w) was prepared by dissolving sodium alginate (1.00 g or 2.00 g; G35 or G75) in H₂O (c) at 25 °C. The solution was added dropwise, using a 25 mL-burette, to a 0.2 M solution of CaCl₂ under mild magnetic stirring. Spherical hydrogel Ca-alginate beads were formed on contact with the receiving solution (instantly) and were left to age for 18 h. Afterwards, they were either used for the synthesis of crosslinked Ca-alginate wet-gel and aerogel beads (as described below), or they were stepwise solvent-exchanged with acetone/H₂O mixtures (30, 60, 90% v/v) and then with dry acetone $(4\times)$, and were dried with SCF CO₂ to produce the corresponding aerogels (referred to as native Ca-alginate aerogel beads).

A solution of chitosan (0.5 or 1% w/w) was prepared by dissolving chitosan (0.50 or 1.00 g) in aqueous acetic acid (0.9 M; 99.5 or 99.00 g) at 50 \degree C. The solution was added dropwise, using a 25 mL burette, to a 1 M solution of NaOH under mild magnetic stirring. Spherical hydrogel chitosan beads were formed on contact with the receiving solution (instantly) and were left to age for 18 h. Afterwards, they were either used for the synthesis of crosslinked chitosan wet-gel and aerogel beads (as described below), or they were stepwise solvent-exchanged with acetone/ $H₂O$ mixtures (30, 60, 90% v/v) and then with dry acetone $(4\times)$, and were dried with SCF CO₂ to produce the corresponding aerogels (referred to as native chitosan aerogel beads).

2.3 Synthesis of crosslinked biopolymer (X-Ca-alginate and X-chitosan) wet-gel and aerogel beads

X-Ca-alginate and X-chitosan wet-gels and aerogels were

Fig. 2 Synthesis of native and crosslinked biopolymer (X-biopolymer) wet-gel and aerogel beads.

Fig. 3 Optical photographs and size distributions of X-Ca-alginate and X-chitosan aerogel beads, as indicated (diameters measured with ImageJ; histograms were calculated using OriginPro 9.0). Mean diameter and sample size (N) are shown on the figure. "1%" refers to the initial concentration of the sodium alginate or the chitosan solution that was 1% w/w.

prepared following a literature procedure.³⁷ In brief, a solution of the triisocyanate in MeCN was prepared by dissolving Desmodur N3300 (41.00 g, 0.081 mol) in dry MeCN (100 mL). Caalginate or chitosan hydrogel beads from Section 2.2 were solvent-exchanged with MeCN/H₂O mixtures (30, 60, 90% v/v) and then with MeCN $(4\times)$. Afterwards, they were immersed in the solution of the triisocyanate (volume of solution was $4 \times$ the volume of the beads) and they were kept into that solution for 24 h, allowing the triisocyanate to diffuse into the wet-gel beads. Subsequently, the system was transferred into the oven at 70 $\mathrm{^{\circ}C}$ for 24 for the crosslinking reaction to be complete. After that time period, crosslinked Ca-alginate or chitosan wet-gel beads were solvent-exchanged with acetone $(3\times)$ and were dried with SCF $CO₂$ to produce the corresponding crosslinked aerogels (referred to as X-biopolymer – or X-Ca-alginate or X-chitosan – aerogel beads).

Fig. 4 The reactions of crosslinking biopolymer (top: calcium alginate; bottom: chitosan) wet gels with a triisocyanate

3. Results and discussion

3.1 Preparation of crosslinked biopolymer (X-biopolymer) aerogel beads

The synthetic protocol for the preparation of biopolymer hydrogel and aerogel beads is presented in Fig. 2. Ca-alginate hydrogels were prepared by dropwise addition of an aqueous sodium alginate solution into an aqueous solution of $CaCl₂$. Chitosan hydrogels were prepared by dropwise addition of an aqueous chitosan solution in aqueous acetic acid into an aqueous solution of NaOH. In either case, gelation was observed on contact with the receiving solution (instantly) and wellshaped spherical beads were formed. After aging, beads were used to produce the corresponding native aerogels (referred to as native biopolymer aerogels), or they were used for crosslinking with Desmodur N3300 (Fig. 1) to produce the corresponding crosslinked aerogels (referred to as X-biopolymer, X-Ca-alginate or X-chitosan aerogels), according to wellestablished procedures.^{37,39,59} Representative optical photographs of X-biopolymer aerogel beads are shown in Fig. 3 together with their mean diameter and size distributions. Mean diameters and size distributions of all aerogel beads (both native and crosslinked) of this study are shown in Fig. S1–S3.†

The crosslinking reactions are described in Fig. 4. As has been recently reported,³⁷ the triisocyanate is being "attached" to the alginate backbone through urethane linkages that are formed from the reaction of the –OH groups of the alginate with one of the –NCO groups of the triisocyanate. In the case of chitosan, the triisocyanate is being "attached" to the chitosan backbone through urea linkages that are formed from the reaction of the $-NH₂$ groups of the chitosan with one of the –NCO groups of the triisocyanate. The reaction of isocyanates with amines is faster than their reaction with alcohols, therefore no urethane linkages are expected (or observed) in this case. In either case, the remaining –NCO groups of the triisocyanate are hydrolyzed by water adsorbed on the surface of the biopolymer wet-gel network and form amines, which react with free triisocyanate that is dissolved in the solution filling the pores of the wet gels, and form urea groups. Hydrolysis of the new dangling –NCO groups continues, followed by reaction with new triisocyanate, until the biopolymer network is coated by PUA.

Fig. 5 ATR-FTIR spectra of native Ca-alginate and X-Ca-alginate aerogels (left), and native chitosan and X-chitosan aerogels (right).

Fig. 6 13 C CPMAS NMR spectra of native Ca-alginate and X-Ca-alginate aerogels, as indicated.

3.2 Chemical characterization of native and crosslinked biopolymer aerogels

Native biopolymer and X-biopolymer aerogels were chemically characterized with ATR-FTIR (Fig. 5), 13 C and 15 N CPMAS NMR (Fig. 6–8), and TGA (Fig. 9 and S4†). Ca-alginate beads were crosslinked with Desmodur N3300, providing materials with a chemical composition analogous to the X-Ca-alginate monoliths published previously.³⁷

ATR-FTIR spectra of native biopolymer and X-biopolymer aerogels are shown in Fig. 5. In agreement with the literature, $20,37$ the spectra of both native and X-Ca-alginate aerogels (Fig. 5 left) show the characteristic asymmetric and symmetric stretching vibrations of the carboxylate groups coordinated to Ca^{2+} ions at 1603 cm^{-1} and 1419 cm^{-1} , the corresponding stretching vibrations of the C–O–C groups on the sugar ring at 1082 $\mathrm{cm^{-1}}$ and 1033 $\mathrm{cm^{-1}}$, respectively. Similarly, the spectra of both native and X-chitosan aerogels (Fig. 5 right) show the stretching vibration of carbonyl groups (amide I) at 1655 cm^{-1} and the bending vibration of N–H groups at 1554 cm^{-1} (amide II), due to the N-acetyl groups, the scissoring vibration of $-NH_2$ groups at 1599 cm^{-1} and the wagging vibrations of –NH₂ groups

Fig. 7^{13} C CPMAS NMR spectra of native chitosan and X-chitosan aerogels, as indicated.

Fig. $8⁻¹⁵N$ CPMAS NMR spectra of native chitosan and X-chitosan aerogels, as indicated.

at 948 and 897 $\rm cm^{-1}$, the stretching vibration of C–OH groups at 1150 cm^{-1} and the asymmetric and symmetric stretching vibrations of the C–O–C groups on the sugar ring at 1066 cm^{-1} and 1031 cm^{-1} , respectively.⁶⁰ The spectra of the X-biopolymer aerogels show all characteristic peaks expected for PUA derived

Fig. 9 Weight loss with temperature for native Ca-alginate and X-Caalginate aerogels (left; 1% w/w initial sodium alginate concentration), and for native chitosan and X-chitosan aerogels (right; 1% w/w initial chitosan concentration).

Table 1 Selected material properties of native (Ca-alginate and chitosan) and crosslinked (X-Ca-alginate and X-chitosan) biopolymer aerogel beads Selected material properties of native (Ca-alginate and chitosan) and crosslinked (X-Ca-alginate and X-chitosan) biopolymer aerogel beads

"The number (%) in parentheses refers to the initial concentration (w/w) of the sodium alginate or the chitosan solution. ^b Porosity calculated according to the formula: $(\rho_s - \rho_0)/\rho_s$, where ρ_s : skeletal density and ρ_0 : bulk density. C Micropore surface area via t-plot analysis, according to the Harkins and Jura model. ^d Total pore volume calculated according to formula: 1/ ρ_0 - 1/ ρ_s e Cumulative volume of pores between 1.7 and 300 nm from N₂-sorption data and the BJH desorption method. f Calculated by the 4 V/ σ method; \geq was set equal to the maximum volume of N2 adsorbed along the isotherm as $P/P_0 \rightarrow 1.0$. For the number in parentheses, \geq was set equal to V_{Total} from the previous column. ^g Particle radius calculated by the formula: $r = 3/(\rho_s \times \sigma)$, where σ : BET surface area. For the number in parentheses, σ was set equal to the external surface area, $\sigma_{\rm ext}$ calculated from the BET surface area minus the micropore surface area.

from Desmodur N3300.9,10,37 More specifically, they show the characteristic stretching vibrations of the carbonyls on the isocyanurate ring and on the urea group at 1686 cm^{-1} and 1630 cm^{-1} , the bending vibration of C-H at 1463 cm^{-1} , and the scissoring vibration of the N-H group at 1535 cm^{-1} . In the spectrum of X-Ca-alginate aerogels, a shoulder at 1733 cm^{-1} can be assigned to the urethane carbonyl group.^{6,11,12,61,62} No peak is shown at 2266 cm^{-1} , assigned to the vibration of the isocyanate groups,⁹ showing that all isocyanate groups of Desmodur N3300 have reacted.

The ¹³C CPMAS spectra of X-Ca-alginate aerogel beads (Fig. 6) show the characteristic broad peaks at 175, 100 and 71 ppm, corresponding to the acetate carbonyls, to the acetal carbons –C–O–C–O– on the alginate rings and to the alginate ring carbons attached to oxygen (–OH or ether), respectively. They also show the characteristic peaks of the corresponding isocyanate-derived PUA, and more specifically the peaks at 160 ppm (urea carbonyl), 150 ppm (isocyanurate carbonyl), 43 ppm (– CH_2 – next to –NH–) and 27 ppm (– CH_2 –).^{9,10,37} Peaks corresponding to the urethane carbonyl are observed as a small shoulder at 157 ppm, reflecting the fact that the amount of urethane groups is small, as those are only the links of PUA on the alginate framework. By comparison between X-Ca-alginate aerogels form G35 or G75 alginate, the alginate/PUA ratio seems to be higher for G75-derived materials. This is in agreement with the PUA content of those materials shown in Table 2, and our previous findings for monolithic X-alginate aerogels, that G75-derived X-alginate aerogels have a higher PUA content compared to their G35-derived analogues (97% vs. 84%, respectively, for monolithic aerogels prepared with the same initial sodium alginate concentration (2% w/w) and Desmodur N3300 concentration (0.6 M) as the beads).³⁷

The 13 C CPMAS spectra of X-chitosan aerogel beads (Fig. 7) show the characteristic peaks of chitosan, e.g. the ones at 173 and 105 ppm, corresponding to the amide carbonyls and to the acetal carbons $-C-O-C-O-$ on the sugar rings, respectively,⁶³ and the characteristic peaks of the corresponding PUA at 159 ppm (urea carbonyl), 149 ppm (isocyanurate carbonyl), 42 ppm (-CH₂- next to -NH-) and 27 ppm (-CH₂-).^{9,10,37} The ¹⁵N CPMAS spectrum of X-chitosan (Fig. 8) clearly shows the two characteristic peaks of PUA at 140 and 83 ppm, corresponding to the nitrogens of the isocyanurate ring and the urea groups, respectively.³⁷ The peak at 126 ppm corresponds to the nitrogen of the amide groups of chitosan,⁶⁴ and is present in the spectra of both native chitosan and X-chitosan aerogels, while the peak at 23 ppm, corresponding to the free amine groups of chitosan,⁶⁴ has almost disappeared in the spectrum of X-chitosan aerogels, indicating that almost all free amine groups of chitosan had reacted with Desmodur N3300.

Thermogravimetric analysis (Fig. 9 left and S4⁺ left) revealed that the thermal decomposition of native alginate aerogels (G35 and G75) was very similar, especially at temperatures below 500 °C, showing three weight loss steps: one at temperatures below 100 \degree C, which can be assigned to water loss^{20,65} (weight loss between 10–20%), another one at temperatures between 220 and 300 °C (weight loss between 20-40%), and a third one at temperatures between 300 and 500 C (weight loss between 15–

Table 2 Polyurea (PUA) content of crosslinked (X-Ca-alginate and Xchitosan) biopolymer aerogel beads

Sample	PUA^{a} (% w/w)
X-Ca-alginate (G35 2%)	43
X-Ca-alginate (G75 2%)	53
X-Ca-alginate (G35 1%)	49
X-Ca-alginate (G75 1%)	66
X-chitosan (1%)	51
X-chitosan (0.5%)	31

^a PUA content calculated according to formula: %PUA (w/w) = $[(\rho_{s} \text{PUA}/\rho_{s})$ $_{\text{X-alginate}}$) \times (ρ_{s} native alginate ρ_{s} $_{\text{X-alginate}}$)/(ρ_{s} native alginate ρ_{s} $_{\text{PUA}}$)] \times 100.³ Values for ρ_s native alginate and ρ_s x-alginate were taken from Table 1 and the value for ρ_s _{PUA} was taken equal to 1.20 g cm⁻³ for Desmodur N3300.⁹

20%). The residue at 800 $^{\circ}$ C was between 10-28%. The decomposition was slower and smaller for G75-derived materials, compared to G35-derived materials from the same initial concentration of sodium alginate. Lower initial concentration of sodium alginate led to materials that lost more mass during decomposition, which is more pronounced in the case of G35 derived materials. X-alginate aerogels behaved similarly up to 220 °C, but weight loss was different. For the first step, <100 °C, the weight loss was between 3–8%, in agreement with the hydrophobic character of X-alginates.³⁷ At temperatures between 220 and 300 °C the weight loss was between 8-20%, and at temperatures between 300 and 500 $^{\circ}$ C the weight loss was between 38–42% (in two steps) for X-Ca aerogels from G35 and between 60–68% for X-Ca aerogels from G75. The latter can be assigned to the decomposition of urethane and urea groups, as well as the isocyanurate rings.^{10,12,37,66} The residue at 800 °C was between 10–20%. Thermogravimetric analysis of chitosanbased materials (Fig. 9 right and S4† right) showed that the decomposition of native chitosan aerogels occurred in two steps: the first one up to 100 $^{\circ}$ C with weight loss 11% and the second one at temperatures between 240 and 360 \degree C with weight **Paper**

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loss 48%. The residue at 800 $^{\circ}$ C was 24%. The initial concentration of chitosan had no effect on the thermal decomposition of the aerogels. The decomposition of X-chitosan aerogels occurred in three steps: the first one up to 100 \degree C with weight loss 6%, the second one at temperatures between 250 and 350 °C with weight loss 38% and the third one at temperatures between 350 and 500 $^{\circ}$ C with weight loss 30%. The residue at 800 \degree C was 18%. As in the case of X-Ca-alginate aerogels, the last step can be assigned to the decomposition of urea groups, as well as the isocyanurate rings.^{10,12,37,66}

3.3 Material properties of native and crosslinked alginate aerogel beads

Selected material properties of all aerogels are summarized in Table 1. Bulk densities of native Ca-alginate aerogels were very low (<0.1 g cm⁻³ for G35 and <0.15 g cm⁻³ for G75) and followed the trend observed previously for the corresponding monolithic aerogels,³⁷ although the densities of the beads are a bit higher. Skeletal densities were also a bit higher compared to monoliths (around 2 g cm^{-3} for both G35 and G75). As expected from the above values, porosities were high, up to 99% v/v (G35) or 98% v/ v (G75). BET surface areas were around 300 m^2 g⁻¹ for G35 Caalginates (similar to those observed for monolithic aerogels) and above 500 m^2 g⁻¹ for G75 alginates (higher compared to those observed for monolithic aerogels); both G35- and G75 derived aerogels also showed a small amount of microporosity (15-23% of the BET surface area). By comparison of V_{Total} and $V_{1.7-300\text{ nm}}$, as well as from the N₂ sorption plots (Fig. S5 and S6†), it is evident that the materials are mostly macroporous (especially the G35-derived aerogels). Primary particle radii, r, were calculated from skeletal densities, ρ _s, and the external surface areas, σ_{ext} , via $r = 1/(3 \times \rho_s \times \sigma_{ext})$ and were found around 6–7 nm for G35 and around 3 nm for G75.

The bulk densities of X-Ca-alginate aerogels were slightly higher compared to native alginate aerogels, because of the

500 nm

Fig. 10 Representative SEM images of native biopolymer and X-biopolymer aerogel beads, as indicated. Scale bar in all cases at 500 nm (magnification: 50 000 \times).

polymer (PUA) covering the alginate skeleton. The bulk density of X-Ca-alginate aerogels was still very low, as low as 0.047 g cm⁻³ for G35- and 0.10 g cm⁻³ for G75-derived materials. Skeletal densities were lower; they depend on the relative ratio of Ca-alginate and PUA (Table 2). The PUA content was in the range of 43–66%, and it was higher for the lower initial concentration of sodium alginate. X-Ca-alginate aerogels were highly porous, with porosities up to 94% v/v. X-Ca-alginate aerogels were still mostly macroporous materials ($V_{\text{Total}} > V_{1.7-}$ _{300 nm}), but there was no microporosity. In agreement with the above, the form of N_2 -sorption plots (Fig. S5 and S6†) indicates macroporous materials with some mesoporosity. The Barrett– Joyner–Halenda (BJH) curves (Fig. S5 and S6†), for pores in the range of 1.7–300 nm, showed maxima in the range of 22–43 nm for all native and crosslinked materials, and broad distributions, as expected for networks formed after particle aggregation. BET surface areas were still high, although lower compared to native aerogels. That trend was considered as an indication that PUA entered secondary particles, coated primary particles and filled the space within secondary particles.^{37,40} As expected, particle radii were bigger compared to native alginate aerogels; the more PUA accumulates on the network, the larger the radius of the particles. RSC Advances Wextare on 100 November 2020. The bulk density gradient, which caused a forey results are considered on 0.5 AM. This area of 0.5 AM and the set of the Data Creative Commonstration of the data Creative Commons

Very similar trends were observed for chitosan-derived materials. Chitosan also yielded aerogel beads with low bulk densities (as low as 0.029 g cm⁻³ for native and 0.031 g cm⁻³ for X-chitosan aerogels) and high porosity (up to 99% v/v for native and 98% v/v for X-chitosan aerogels), that are mostly macroporous (Table 1 and Fig. S7†). The BJH curves, for pores in the range of 1.7–300 nm, showed maxima at 23–27 nm for all native and crosslinked materials (Fig. S7†), and broad distributions. BET surface areas for X-chitosan aerogel beads were about half (or less than half) of those of native chitosan aerogel beads. The maximum BET surface area was 217 $\mathrm{m}^2\ \mathrm{g}^{-1}.$ Like in alginate monoliths and many other oxide aerogels,⁶⁷ particle radii increased after crosslinking in proportion to the PUA uptake (31–51%; Table 2).

All indications above show very small particles. SEM shows that all aerogels are fibrous, leading to the conclusion that bers consist of particle aggregates, in analogy to what has been reported previously, most notably with certain PUA^{9,68} and polyimide⁶⁹ aerogels. SEM (Fig. 10 and S8-S11[†]) showed that all samples were fibrous. The micromorphology of both native and crosslinked aerogels is the same, showing that PUA is formed on the biopolymer skeletal network, coating the network conformally. SEM images of the surface of the beads revealed the presence of a thin skin with a different, denser, morphology. That is more pronounced in the case of Ca-alginate materials. Although much denser, the skin is porous because it allowed gasses to penetrate the interior, *i.e.*, during N_2 -sorption porosimetry or He-pycnometry. An analogous observation has been made in PUA beads prepared from Desmodur N3300 and ethylenediamine,¹⁰ and it was attributed to the higher concentration of ethylenediamine on the surface of the droplet containing Desmodur N3300, which caused a fast reaction and the formation of a crust on the surface, while the diffusion of ethylenediamine to the interior established a concentration

gradient, which caused a slower reaction. In this case, the concentration of Ca^{2+} or H_3O^+ ions is higher on the surface of the droplet containing sodium alginate or chitosan, respectively, and causes gelation of the surface, and afterwards the diffusion of Ca²⁺ or H_3O^+ ions to the interior reacts slower and forms the fibrous structure of Ca-alginate or chitosan. That fibrous structure of the hydrogel is retained in the native biopolymer aerogel,⁷⁰ and later on in the X-biopolymer aerogel. The different skin effect between alginate-based and chitosanbased aerogels probably points at different gelation mechanisms.

4. Conclusions

Polyurea-crosslinked calcium alginate (X-Ca-alginate) monoliths have been reported recently and are very strong materials, in contrast to the native Ca-alginate aerogels that, like most biopolymer aerogels, are mechanically weak. In this work we have prepared analogous materials in the form of beads, which have several advantages compared to monoliths, for example easier processing via accelerated solvent exchange and supercritical drying, fewer defects, and homogeneity even at relatively high concentrations. In addition, this new class of materials was expanded to include crosslinked chitosan (X-chitosan) aerogel beads, suggesting a general applicability of this method to other biopolymer aerogels as well.

X-Ca-alginate and X-chitosan aerogel beads have been prepared from the reaction of an aliphatic triisocyanate (Desmodur N3300) with the –OH groups of calcium alginate or the $-NH₂$ groups of chitosan preformed wet gels, forming anchoring urethane and urea groups, respectively, on the surface of the biopolymer network; subsequent formation of a polyurea conformal coating of the skeletal networks took place via reaction of the surface-bound triisocyanate with adsorbed water retained from gelation on the inner surfaces of the wet gels. Despite an up to 66% increase in mass by the crosslinking process, SEM showed that all aerogels, native and crosslinked, were fibrous and their micromorphology did not change after crosslinking. Applications under consideration for these materials include environmental remediation and conversion to carbon aerogels. Results show that both types of X-biopolymer aerogels are carbonizable and can be used as precursors to isomorphic (fibrous) carbon aerogels.

Conflicts of interest

There are no conflicts to declare.

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