Novel non-cytotoxic alginate–lignin hybrid aerogels as scaffolds for tissue engineering

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Abstract

This paper presents a novel approach toward the production of hybrid alginate–lignin aerogels. The key idea of the approach is to employ pressurized carbon dioxide for gelation. Exposure of alginate and lignin aqueous alkali solution containing calcium carbonate to CO2 at 4.5 MPa resulted in a hydrogel formation. Various lignin and CaCO3 concentrations were studied. Stable hydrogels could be formed up to 2:1 (w/w) alginate-to-lignin ratio (1.5 wt% overall biopolymer concentration). Upon substitution of water with ethanol, gels were dried in supercritical CO2 to produce aerogels. Aerogels with bulk density in the range 0.03–0.07 g/cm3, surface area up to 564 m2/g and pore volume up to 7.2 cm3/g were obtained. To introduce macroporosity, the CO2 induced gelation was supplemented with rapid depressurization (foaming process). Macroporosity up to 31.3 ± 1.9% with interconnectedness up to 33.2 ± 8.3% could be achieved at depressurization rate of 3 MPa/min as assessed by micro-CT. Young's modulus of alginate–lignin aerogels was measured in both dry and wet states. Cell studies revealed that alginate–lignin aerogels are non-cytotoxic and feature good cell adhesion making them attractive candidates for a wide range of applications including tissue engineering and regenerative medicine.

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between hydrophilicity and hydrophobicity of the surface is an important factor of cell adhesion. Lignin is expected to reduce hydrophilicity of alginates and hence provide more suitable environment for cells to adhere, grow, and differentiate. Bearing in mind ultimate stability of lignin, it was also expected that the presence of lignin may abate the scaffold degradation rate and help to match it with the rate of new bone tissue regeneration.

Due to its abundance and low price, it is of definite interest to usher lignin into high-value products, i.e., biomaterials, adsorbents, thermal insulators. Several attempts have been reported in the literature on lignin as a part of biomaterials exemplified by composites with hydroxypatite [27,28]; as a carrier in laxative formulations [29]; allergenicity reducer for latex rubber [30]. Potential applications in food industry are also reported [31]. For comprehensive overview on other application of lignin and lignin-based products readers are referred to recently published reviews [32–34].

One objection against lignin as a material for biomedical and pharmaceutical applications is its phenolic nature. Organosolv lignin has been reported to be slightly cytotoxic for peripheral blood mononuclear cells [28]. One lignin derivative, sulphonated lignin, when blended with fish gelatin, showed cytotoxicity only at very high concentrations (IC50 in the range 1500–1750 μg/ml) [31]. IC50 values in the range 400–1200 μg/ml were found for lignins from different sources by Ugartondo et al. [35]. Microalgae (Chlamydomonas reinhardtii) and Backer’s yeast (Saccharomyces cerevisiae) show indistinguishable loss of viability after incubation with lignin nanoparticles compare with a control sample [36]. From this data, it can be surmised that generally lignin is not cytotoxic up to moderate concentration. One aim of this work is to prove whether Ca-crosslinked alginic–lignin aerogels are non-cytotoxic and to evaluate them as potential biomaterials.

Apart from lower hydrophilicity and higher stability another potential advantage of lignin is its antimicrobial activity. Although antimicrobial properties of the phenolic units of lignin are well documented [32], there has been some controversy in the literature whether lignin and lignin-containing materials have antimicrobial activity. Erakovic et al. [28] have found no significant antimicrobial activity of films obtained by electrophoretic deposition from 1 wt% suspension of organosolv lignin in the presence of hydroxypatite. Some antimicrobial activity was detected for sulphonated lignin [31]. However, no direct comparison of water insoluble lignin with sulphonated lignin is possible. Antimicrobial action of the latter may be ascribed to its surface active properties. Study of Dizhbite et al. [37] revealed antibacterial effect of Kraft lignin and related it to the high activity as radical scavenger. Lignin-related compounds from pine cone are found to induce varieties of antiviral activity [38].

Composites and blends of lignin with cellulose [39], cellulose acetate [40], xanthan gum [41], PEG [42], PVA [43], PLA [44], PVP [45,46] are known from the literature. Even though there may be only weak interaction between lignin and principal constituent, addition of lignin may offer advantages such as more control over water uptake [41] and improved mechanical properties [31,45].

Importance of conjugating lignin with polysaccharides for in vivo expression of various kinds of immunopotentiating activity is also reported [38]. These features may also have a beneficial effect with respect to biomedical applications.

Gelation by a reaction with crosslinkers is a common technique to obtain lignin aerogels. Gelation with resorcinol formaldehyde [47], phenol formaldehyde [48], tannin formaldehyde systems [49] and α,ω-diglycidyl ethers [50] are reported. To the best of our knowledge, ionic crosslinking of pure lignin or polymer blends containing lignin has not been reported. In this work a goal was set to use alginic as a “glue” for lignin. Presence of alginic allows the use of ionotropic gelation instead of chemical crosslinking.

Gelation of alginates induced by pressurized carbon dioxide was recently developed [51] and is used in this work to gel alginic–lignin mixtures. In processing of biomedical materials, CO2 induced gelation have certain advantages over internal and diffusion gelation methods: (i) carbon dioxide, being volatile acid in water media, can be recovered at post-processing stages; (ii) fast depressurization leads to macroporous foam-like hydrogels; (iii) bactericidal activity of pressurized CO2 simplifies preparation of food and medical materials [52]; and (iv) the process potentially allows to avoid ambient pressure solvent exchange and can be directly combined with subsequent supercritical drying [51,53].

2. Materials and methods

2.1. Chemicals

Alginic acid sodium salt (suitable for immobilization of microorganisms grade, catalogue no. 71238) was obtained from Sigma Life science, Germany. Lignin was produced as described below (Section 2.2). Calcium carbonate (light, precipitated powder, particle size ca. 1 μm) was purchased from Magnesia GmbH, Germany. Sodium hydroxide (>99%) and anhydrous ethanol (99.9%) for the solvent exchange were purchased from Carl Roth GmbH and H. Möller GmbH & Co. KG, respectively. Carbon dioxide used for drying (99.9 mol% purity) was procured from AGA Gas GmbH (Hamburg, Germany). In case of in vitro cell culture studies, the chemicals used were of analytical reagent or tissue culture grade. Deionized water was used throughout the study.

2.2. Starting solutions

Lignin was obtained from wheat straw as described elsewhere [50–54]. This process was carried out by the biorefinery research group at the Institute of Thermal Separation Processes, Hamburg University of Technology (Germany). Briefly, wheat straw was fractioned by a hydrothermal pretreatment with liquid hot water at 473 K and 5 MPa followed by an enzymatic hydrolysis step (50 °C, pH 5, Novozymes CTeC2, 72 h). Water insoluble lignin was collected after the enzymatic cleavage. Lignin was washed with water and dried at 70 °C for 50 h. 3 wt% solution of lignin was prepared by mixing a certain amount of dried lignin with 1 M NaOH and overnight stirring.

3 wt% sodium alginate solution was prepared by gentle overnight stirring of Na-Alg powder with water. After the preparation both solutions were bottled and stored at 5 °C.

Calcium carbonate powder was dispersed in Na-Alg solution with a high speed homogenizer Ultra-turrax (IKA, Staufen, Germany). Then lignin solution was added to obtain desired alginic-to-lignin ratio: 2:1, 3:1, 4:1 or 5:1 (w/w). Mixture was diluted with water to keep 1.5 wt% overall biopolymer concentration (alginic + lignin) and once again homogenized (Ultra-turrax) for 1 min. Two crosslinking degrees (q) were used: alginic-to-CaCO3 of 1:0.1825 (w/w) is referred as q = 1, q = 2 corresponds to the doubled amount of CaCO3. Resulting suspension was filled into a standard 48 multiwell plate (BD Biosciences, USA) and subjected to CO2 induced gelation.

2.3. CO2 induced gelation and hydrogel foaming

Multiwell plates with Na-Alg/lignin/CO2 mixture were placed into an autoclave and exposed to gaseous carbon dioxide at 4.5 ± 0.5 MPa and room temperature for 24 h. The autoclave described elsewhere [55] was used for both gelation and supercritical drying. To study effect of the depressurization rate on macroporosity of the gels, pressure release was employed at 0.8 MPa/min and 3 MPa/min. The gels were left in the air till...
formation of bubbles ceased, then washed with water and finally transferred into ethanol–water mixture to perform solvent exchange as described below.

2.4. Solvent exchange and supercritical drying

Hydrogels were immersed in grades of aqueous ethanol (30, 60, 90 and 99.9 vol.%) for 3 h at each ethanol concentration. The final solvent exchange was done twice or thrice before the hydrogels were supercritically dried. A density meter DMA 4500 (Anton Paar Company, Austria) was used to control completeness of the solvent exchange. Gels were wrapped in filter paper and placed into pre-heated autoclave (318 K). Supercritical drying was performed using the same autoclave as for gelation. The autoclave was sealed and CO2 was filled in by a compressor. Once 12 ± 1 MPa was reached, outlet was opened and constant flow (0.2 kg/h) was set for 5 h such that 6–7 residence volumes of CO2 were used. Then system was depressurized in 30 min followed by cooling down to room temperature.

2.5. Textural and morphological properties

Bulk density of the samples was calculated as ratio of mass to volume. The length and diameter of the aerogels were measured with Vernier calipers. SEM pictures were taken by a Leo 1530 microscope (Carl Zeiss, Germany). Samples were sputtered with gold (7 nm). Pictures were taken at an accelerating voltage of 5 kV and working distances in the range of 4.0–6.0 mm. Surface area, pore volume and pore diameter were analyzed by nitrogen adsorption desorption techniques using Nova 3000e (Quantachrome Instruments, USA). Surface area was obtained from multipoint BET. Pore size distribution and volume of mesopores were calculated from desorption branch using BJH method. Porosity, interconnectivity and mean pore size in the macroporous range were evaluated by micro-CT using Scanco 20 equipment (SkyScan 1702, Belgium) with penetrative X-rays of 30 kV and 167 μA, in high resolution mode with a pixel size of 14.71 μm and 1.5 s of exposure time. A CT analyzer (v1.5.1.5, SkyScan) was used to visualize the samples and calculate the parameters from 2D aerogel structures. The analysis was done thrice within different regions of interest. Results are given as mean ± standard deviation.

2.6. Mechanical properties

Compressive properties of the aerogels were measured using an INSTRON 5540 universal testing machine (Instron Int. Ltd, High Wycombe, UK) with a load cell of 1 kN. Compression tests were carried out at a crosshead of 2 mm/min, until a maximum deformation of 60%. Young’s modulus was calculated as the initial linear modulus on the stress–strain curves. The results are presented as the average of three experiments ± standard deviation. In wet state, the samples were immersed for 10 min in PBS solution before compression tests.

2.7. Water uptake

Aerogel were placed into test tubes, filled with adequate amount of Tris–HCl buffer solution (pH 7.4) and placed in a water bath (37 °C, 60 rpm). Weight of the swollen sample at was measured after removing excess of the buffer with filter paper after 1, 3, 7 and 14 days. For each time point three parallel samples were measured and the water uptake WU was calculated relative to the initial weight W0 as follows:

\[ WU\% = \frac{W_t - W_i}{W_0} \times 100. \]

2.8. In vitro biological performance

2.8.1. Cell culture

A mouse fibroblast-like cell line (L929 cell line, European Collection of Cell Cultures, UK) was maintained in DMEM (Sigma–Aldrich, Germany) supplemented with 10% heat-inactivated fetal bovine serum (Biochrom AG, Germany) and 1% antibiotic–antimycotic solution (Gibco, UK). Cells were cultured in a humidified incubator at 37 °C in a 5% CO2 atmosphere.

2.8.2. Indirect contact assay

Aerogels extracts were prepared according ISO/EN 10993 in DMEM culture medium, L929 cells at a concentration 1.5 × 10^4 cell/mL were cultured in a 48–well plate for 24 h at 37 °C. At this time, medium was replaced by aerogels extract. Cell viability was evaluated by the MTS assay after 72 h of culture time.

2.8.3. Direct contact assay

Confluent L929 cells were harvested and seeded in the aerogel samples as follows. Samples were distributed in a 48-well cell culture plate. Samples were initially immersed in sterile PBS to swell the matrix. Later, PBS was removed and a drop (20 μl) of a cell suspension with a concentration of 1.5 × 10^3 cells/mL was added to each aerogel. These constructs were statically cultured for 1, 3 and 7 days under the culture conditions of 37 °C at 5% CO2 in an incubator. Triplicates were used for each time point.

2.8.4. MTS assay

Cell viability of the aerogels was determined after the predetermined culture times by the MTS assay using the Cell Titer 96 AQueous One Solution Cell Proliferation Assay (Promega, USA) according to the manufacturer instructions. This assay is based on bioreduction of tetrazolium compound into water-soluble formazan derivative. The formazan absorbance which is directly proportional to the number of living cells was measured at 490 nm in a microplate reader (Synergie HT, Bio-Tek, USA).

In case of indirect contact, effect of the leachable released from the aerogels on cellular metabolism was evaluated by culturing L929 cells in the extracts obtained from aerogels. Latex was used as a negative control and TCP (tissue culture polystyrene) was used as a positive control. In direct contact assay the cell-scaffolds were transferred to a new culture plate in order to evaluate the presence of viable cells only on the surface of the aerogel. In this case, TCP was used as a positive control. All cytotoxicity screening tests were performed in three replicates and the results are presented as mean ± standard deviation.

2.9. Statistical analysis

Statistical analysis of the data was conducted using IBM SPSS Statistics version 20 software. Shapiro–Wilks test was employed to evaluate the normality of the data sets. Once the results obtained did not follow a normal distribution, non-parametric tests, in particular, Kruskal–Wallis test was used to infer statistical significant differences. Differences between the groups with p < 0.05 were considered to be statistically significant.

3. Results and discussion

Reports on alginate-based aerogels for biomedical application are limited. To the best of our knowledge, alginate aerogels were evaluated to date as drug delivery systems by Mehling et al. [56], García-González et al. [57]; Veronovski et al. [58,59]; Ulker and Erkey [60] and as bio-superabsorbents by Mallepally et al. [61].

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Production of aerogels with controlled pore size and dual pore size distribution still remains a challenge and restrains aerogels from filling a niche in regenerative medicine where macroporosity of the scaffold is of concern. As pointed out by Reverchon et al. [3], it is very difficult to obtain the coexistence of the macro and microstructural characteristics within one scaffold. Various techniques have been proposed to address this issue: addition of solid/liquid porogen and subsequent leaching [3]; emulsion templating [62,63] including supercritical carbon dioxide as a dispersed phase [64,65]; in situ generation of gas bubbles confined in the gel [63] or rapid expansion of a gas dissolved in the gel (see below). In this report gelation induced by pressurized CO2 with subsequent foaming was performed to create macroporous aerogels.

3.1. CO2 induced gelation

Solubility of carbon dioxide in water increases with rising pressure along with lowering of pH down to 3 [66]. The drop in pH causes in turn an increase in solubility of calcium carbonate along with the release of calcium ions. At conditions used in this study for gelation (298 K and 4.5 MPa), CaCO3 solubility is much larger (ca. 2.8 g/L [67]) than at ambient conditions (0.006–0.01 g/L [68]) so considerable amount of Ca2+ ions is available for the reaction with alginate. To support that Ca2+ ions act as crosslinker a blank experiment was performed. It showed that alginate does not form a gel in the absence of CaCO3. Experiments in a tilting viewing cell showed noticeable increase in viscosity neither for Na-Alg solution alone nor for Na-Alg/alginate mixture. These findings can be attributed to moderate pH change; in pure CO2/water system at 25°C pH approaches value of around 3 and remains constant above 3 MPa [69]. Apparently, this pH is not low enough to form a stable acid alginate gel (pKₐ of M and G units in the range of 3.4–3.7 [70]). Moreover, sodium hydroxide introduced with lignin solution reacts with CO2 yielding bicarbonate, which possesses buffer properties: bicarbonate buffer at 5 MPa CO2 pressure is able to maintain pH around 6–7 (only drop 0.5–1.0 pH units compared to ambient conditions, [71]). In this regard, this gelation method can be classified as the internal setting method exploiting acidic properties of CO2–water mixture.

In this study the CO2 induced gelation method was extended over polymer compositions. SEM analysis of aerogels showed the net-like structure, which is typical to alginate aerogels (Fig. 1). Visual inspection of hydrogels and SEM revealed no sign of lignin inclusions. Some authors have found that lignin has limited compatibility with other biopolymers, e.g. with cellulose [39] and xanthan gum [41]. These findings support rather interpenetrating than co-crosslinking structure of the hybrid network. Rudaz [40] have prepared hybrid cellulose–lignin hydrogels and noticed that lignin can be washed out from the hydrogels during the solvent exchange due to weak cellulose–lignin interaction. This in turn led to the increase in porosity of cellulose aerogels since lignin acted as a porogen. In this study an opposite trend was found. As lignin concentration increases the BJH pore volume decreases (see Fig. 3). However, it was not possible to obtain stable hydrogels with lower alginate-to-lignin ratio than 2:1 (w/w). Additional experiments with pure lignin with and without Ca2+ resulted in lignin precipitation demonstrating that lignin of itself is unable to form a gel at this condition. Taking into account the high affinity of Ca2+ to lignin [72], we suppose that OH-groups of lignin may participate in the formation of egg-box junctions, but only to certain extent. Partial substitution of alginate COO− groups with phenolic OH-groups of lignin in the egg-box junctions may explain the absence of lignin inclusions in the aerogels.

3.2. Foaming of hydrogels

Foaming of hydrogels is a well-known process exemplified by cellulose [73], chitin [74] and gelatin [75]. However, to the best of our knowledge, combination of both gelation and foaming into a one-pot approach has not been reported. Moreover, such a combination opens up an inviting prospect to realize all steps of aerogel processing (gelation, foaming, solvent exchange and supercritical drying and loading) under carbon dioxide pressure as an integrated process [51,53].

For the purposes of tissue engineering scaffolds the important conclusion is that CO2 induced gelation should be coupled with fast pressure release to obtain macroporosity. Indeed, our results indicate great impact of the depressurization rate: 3 MPa/min favors formation of numerous pores of approximately 200 µm in size, whereas slow pressure release (0.8 MPa/min) led to significantly low porosity with two-fold larger pores (Fig. 2). Very slow depressurization at 0.02 MPa/min gave no detectable macroporosity (data not shown).

Table 1 summarizes results of micro-CT assessment for the aerogels produced through preceding foaming. Foaming allowed to introduce macropores in the range of 200–450 µm. Aerogels foamed at higher depressurization rate demonstrate two-fold increase in overall macroporosity along with almost two-fold decrease in mean pore size. This decrease in pore size is however well above a minimal size (38–63 µm), which allows cell to grow and proliferate [76]. These results indicate that CO2 induced gelation followed by hydrogel foaming seems to be an efficient method to introduce macroporosity into hydrogels and aerogels, which are intrinsically micro- and mesoporous.

In the context of this study it is interesting to adduce results from Floren et al. [77] for silk protein hydrogels prepared under high pressure CO2 (0.5–15 MPa). In this work acidification of silk fibroin aqueous solution by pressurized CO2 led to the formation of stable hydrogel through the development of extensive β-sheet structures. The results of Floren et al. indicate that protein hydrogels prepared under CO2 pressure followed by slow depressurization (0.02–0.5 MPa/min) display distinctly more homogeneous pore structure compare to fibrin hydrogels acidified by citric acid at ambient conditions [77]. This clearly shows that carbon dioxide induced gelation, not followed by fast depressurization, leads to more compact hydrogels compared to ambient conditions. This conclusion is in agreement with observations made by Annabi et al. [78]. Elastin-based hydrogels produced in pressurized CO2 were found to be stiffer (in terms of compression modulus) than those

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produced at atmospheric conditions. In addition, another study revealed that gelation at high pressure reduces the pore size of the hydrogels [79]. One possible explanation for these findings is that high pressure CO₂ facilitates coacervation of the polymer leading to densification of the polymer junctions. We can speculate that a similar phenomenon allowed us to prepare pure alginate hydrogels from Na-Alg with concentration as low as 0.25 wt%, whereas conventional methods led to unsatisfactory results [51].

3.3. Textural properties

To study the effect of lignin concentration on the textural properties of the aerogels, alginate-to-lignin ratios of 2:1, 3:1, 4:1 or 5:1 (w/w) were studied keeping overall biopolymer concentration at 1.5 wt%. The effect of the crosslinking degree, q, on textural properties was also studied at two different levels (Fig. 3a and b). All alginate–lignin aerogels showed bulk densities in the range 0.03–0.07 g/cm³. No clear trend was observed with the crosslinking degree or the lignin concentration. Conversion of hydrogels into aerogels implies shrinkage of certain extent [57]. Overall linear shrinkage caused by solvent exchange and supercritical drying was in the range of 20–35% across all samples. Despite the pronounced shrinkage all aerogels remained cylindrical shape and showed quite high surface area compared to the state of the art (150–600 m²/g and up to 450 m²/g for alginate and lignin aerogels, respectively, [49,57]). Doubled crosslinker amount (q = 2) leads to moderate reduction in surface area (Fig. 3a), whereas reduction in pore volume is more pronounced (Fig. 3b). At q = 2, lignin concentration does not exert much influence on the surface area. In other words, higher crosslinking degree results in more compact aerogel structures, whereas q = 1 and lower crosslinking degree led to soft and difficult-to-handle hydrogels. Moreover, foaming of a less crosslinked gel often resulted in its disruption. In search of a compromise between possibly high lignin concentration, good textural properties (high surface area, pore volume) and ability to perform foaming the crosslinking degree was kept constant at 2 and alginate-to-lignin ratio at 4:1 (w/w). All further in vitro studies were performed with this formulation, which exhibited the density of 0.07 ± 0.01 g/cm³ and surface area of 382 m²/g.

Table 1

<table>
<thead>
<tr>
<th>Depressurization rate, MPa/min</th>
<th>Porosity, %</th>
<th>Mean pore size, µm</th>
<th>Interconnectivity, %</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.8</td>
<td>14.28 ± 0.96</td>
<td>423 ± 7</td>
<td>27.6 ± 4.6</td>
</tr>
<tr>
<td>3</td>
<td>31.3 ± 1.9</td>
<td>220 ± 18</td>
<td>33.2 ± 8.3</td>
</tr>
</tbody>
</table>

Fig. 2. Micro-CT image of alginate–lignin aerogels produced depressurization rate of 0.8 MPa/min (a) and 3 MPa/min (b).

Fig. 3. BET surface area (a) and BJH pore volume (b) of alginate–lignin aerogels with two crosslinking degree: q = 1 (white bars) and q = 2 (shaded bars). Depressurization rate is 0.8 MPa/min.

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3.4. Water uptake

Water uptake study was done with alginate–lignin aerogels in Tris–HCl buffer. The latter was chosen instead of commonly used PBS due to its lower affinity to calcium ions. Phosphate ions presented in PBS leads to fast dissolution of the alginate materials [80] so that water uptake may be distorted due to fast calcium leakage [81]. The water uptake gradually increased from day 1 to day 14 and reached a plateau after about 1 week (Fig. 4). Compared to pure alginate aerogels and starch–alginate hybrids [82], it was found that lignin slows down the water uptake kinetics, consistent with its hydrophobic nature. Equilibrium water uptake of alginate-based materials presented in the literature varies in the wide range from 30 to 35,000% [83–85]. On account of vast variety of production methods a direct comparison is difficult. Kulkarni et al. [84] have found similar water uptake for chemically crosslinked alginate, but with much faster kinetics (equilibrium reached in 2–4 h). It was noticed [81,85] that almost no swelling happened upon contact with Tris–HCl buffer due to the lack of specific interaction between buffer and Ca-crosslinked alginate. Our results however show that alginate–lignin aerogels are able to uptake up to 1613% of Tris–HCl buffer. Swelling of the material was also noticed during the study. Although detailed mechanism of water uptake needs to be elucidated it is clear that not only pore filling contributes into the equilibrium uptake but also the swelling of the matrix.

3.5. Mechanical properties

In the context of tissue engineering applications, mechanical properties are an important characteristic. The mechanical response of the alginate–lignin aerogels prepared at two different depressurization rates were evaluated in the compression mode. Table 2 compares Young’s modulus of dry and wet aerogels. As can be seen from this data alginate–lignin aerogels can be classified as materials with low stiffness both in dry and wet states. Their Young moduli are in the range of granulation and fibrous tissues [86]. It was also found that Young’s modulus is affected by the depressurization rate: the value was three times lower for the aerogel foamed at 3 MPa/min than at 0.8 MPa/min, whereas wetting makes aerogels almost insensitive to the rate of depressurization.

Due to various compression conditions reported in the literature (compression rate, range of strain for Young’s modulus) and variation in aerogel densities a comprehensive comparison is infeasible. Native silica aerogels are brittle and break at small tensile strains [2]. Viggiano and Schiraldi [87] have reported the compressive modulus of 1.78 MPa for a cryogel composed of alginate and lignin (1:1 w/w, ratio with 5% overall solid content). This result is close to our results for dry aerogels. Alginate–lignin aerogels reported here demonstrate compressibility and become flexible when compressed, similar to pure alginate aerogels produced by CO2 induced gelation [51]. This behavior is rather unusual for biopolymer aerogels and has mainly been observed for polymer crosslinked silica aerogels, e.g. isocyanate-coated silica aerogels [2].

3.6. In vitro biological performance

In a first approach the cytotoxicity of the samples prepared was evaluated. Indirect studies were conducted to check the effect of the leachables of the matrices on cells cultured in a tissue plate.

**Table 2**

<table>
<thead>
<tr>
<th>Sample</th>
<th>Rate of depressurization, MPa/min</th>
<th>Young’s modulus, MPa</th>
</tr>
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<tbody>
<tr>
<td>Alginate–lignin dry: 0.8</td>
<td>1.36 ± 0.24</td>
<td></td>
</tr>
<tr>
<td>Alginate–lignin dry: 3</td>
<td>0.38 ± 0.05</td>
<td></td>
</tr>
<tr>
<td>Alginate–lignin wet: 0.8</td>
<td>0.05 ± 0.02</td>
<td></td>
</tr>
<tr>
<td>Alginate–lignin wet: 3</td>
<td>0.02 ± 0.01</td>
<td></td>
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**Fig. 5.** In vitro biological studies: indirect cytotoxicity MTS assay after 72 h (a); and direct contact MTS assay with cells cultured on the surface of lignin aerogels for 3 and 7 days (b).

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As negative control we used latex rubber and as positive control cells cultured in DMEM culture media. Cytotoxicity screening by indirect contact assay studies revealed that alginate–lignin aerogels did not show any evidence of toxic effects of the leachables over the fibroblast like L929 cells (Fig. 5a). Cell viability after 72 h for alginate–lignin was comparable to TCP, whereas latex showed clear cytotoxic effect. These results demonstrate that despite the phenolic nature of lignin, it can be used as a material for biomedical and pharmaceutical applications, at least in the concentration range used for aerogel preparation.

The main result of this indirect study is that alginate–lignin aerogel does not hinder cell growth and thus can be recognized as non-cytotoxic material. Cell adhesion tests revealed that cells are able to adhere on the surface of the materials and the metabolic activity has increased from day 3 to day 7, comparable to TCP results (Fig. 5b).

These results give an account that the alginate–lignin aerogel demonstrate no cytotoxicity and good cell adhesion properties, at least in the range of lignin concentration studied. This clearly indicates that lignin-containing aerogels can be viewed as candidates for further in vitro and in vivo testing.

4. Conclusions

The present work deals with the production of alginate–lignin aerogels using CO2 induced gelation followed by solvent exchange and supercritical drying. Pressurized carbon dioxide acts as an acidifier to liberate Ca2+ ions for the crosslinking of alginate–lignin mixture. Foaming by rapid expansion of carbon dioxide can be readily implemented to introduce macroporosity in the aerogels. Foaming procedure is free of templating agents and shown to be an effective way to introduce macropores of few hundred microns into hydrogels and subsequently aerogels. Despite the pronounced shrinkage, aerogels produced by CO2 induced gelation followed by foaming demonstrate low density and good textural properties both at meso and macroscale. Apart from readily available foaming there are several additional advantages in using pressurized CO2 to induce gelation. First, carbon dioxide strengthened the hydrogel, whereas hydrogels formed from the same formulation at ambient conditions are more soft and often do not preserve the shape. Second, wide range of polymers can be mixed with alginate leading to hybrid hydrogels with modified properties. Third, the use of carbon dioxide as a volatile acidifier, allows for efficient recovery of CO2 at post-processing stages. Finally, the process can be directly combined with subsequent supercritical drying into a one-pot approach. In this work we have proven the feasibility of alginate–lignin aerogels to be used in a tissue engineering perspective.

The alginate–lignin aerogels present textural and morphological properties suitable for tissue engineering applications. Furthermore they have high equilibrium water uptake. In terms of Young’s modulus studied aerogels can be classified as material with low stiffness both in dry and wet states. In vitro cytotoxicity screening has demonstrated that lignin does not compromise cell viability and it has been shown that alginate–lignin aerogels possess good cell adhesion properties prompting possible further in vitro and in vivo assessment.

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