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DEFORMATION RESPONSE OF GELLAN GUM BASED BONE SCAFFOLD SUBJECTED TO UNIAXIAL QUASI-STATIC LOADING

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ABSTRACT. This study is focused on an investigation of the reinforcement effect of the bioactive glass nano-particles in the gellan gum (GG) scaffolds used in bone tissue engineering. The investigated material was synthesized as the porous spongy-like structure improved by the bioactive glass (BAG) nano-particles. Cylindrical samples were subjected to a uniaxial quasi-static loading in tension and compression. Very soft nature of the material, which makes the sample susceptible to damage, required employment of a custom designed experimental device for the mechanical testing. Moreover, as the mechanical properties are significantly influenced by testing conditions the experiment was performed using dry samples and also using samples immersed in the simulated body fluid. Material properties of the pure GG scaffold and the GG-BAG reinforced scaffold were derived from a set of tensile and compression tests under dry and simulated physiological conditions. The results are represented in the form of stress-strain curves calculated from the acquired force and displacement data.

Keywords: gellan gum scaffold; reinforcement; uni-axial loading.

1. Introduction

The worldwide incidence of bone disorders and conditions is growing by steeply increasing trend. Particularly in the high income regions, a twofold increase between 2010 and 2020 is expected [1]. This is the tribute for populations aging coupled with improper nutrient consumption and poor physical activity. Globally, more than $40\,\%$ of women and $30\,\%$ of men are under increased risk of occurence of bone disorders [2]. Annually, only in the USA more than half a million bone defects are reported. The treatment expenditures reach more than \$2.5 billion per year.

Treatment of bone disorders using engineered bone tissue has been considered as a promising alternative to traditional medical treatment methods including use of autografts and allografts. Currently, the field of artificial tissue engineering represents overcoming problems, such as donor site morbidity, loss of bone inductive factors, and scaffold resorption during healing [3]. Generally, several fundamental requirements have to be simultaneously fullfilled by the artificial tissue: i) normal cellular activity without toxicity effects, ii) minimizing of the stress shielding effect, iii) successful diffusion of nutrients and oxygen, and iv) controlled degradation coupled with the resorption of the artificial material [4].

The presented paper deals with a uni-axial quasistatic testing of the artificial spongy-like structure [5] proposed for bone tissue engineering purposes as the bone scaffold. The investigated GG-BAG material combines organic (polysacchariditic) components with inorganic (Silicon-Calcium based) nanoparticles. This approach effectively enables the adaptation of physical and mechanical properties of the synthesized material according to the desired application [6].

Mechanical properties of gellan-gums can vary significantly depending on the procedure of synthetization. Young's modulus in range $0.15-150\,\mathrm{kPa}$ was reported in [7]. Therefore, the newly synthesized material was subjected to the quasi-static loading in both tension and compression to evaluate primarily its stiffness and yield behaviour. Firstly, the testing was carried out under dry conditions to evaluated the material properties of the synthesized material itself. Then, the experiment was repeated under simulated physiological conditions by employment of the bioreactor with circulating synthetic plasma.

2. Gellan-gum based scaffolds

The GG-BAG material investigated in this study was synthesized at the Jozef Stefan Institute (Slovenia) as a porous spongy-like structure improved by the BAG nano-particles with size of $\approx 200\,\mathrm{nm}$ [8]. The GG is a microbial extracted polysaccharide used in the food and pharmaceutical industry [9, 10]. It is composed of repeating units consisting of two D-glucose and one of each L-rhamnose and D-glucoronic acids [11]. The main

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Sample	h [mm]	D_{\min} [mm]	D_{\max} [mm]	m [mg]
GG00 1	8.77	4.97	5.09	10.9
GG00 2	8.73	5.03	5.04	10.9
GG00 3	8.93	4.95	5.04	10.4
GG00 4	9.18	5.01	5.02	10.5
GG00 5	9.01	4.93	4.96	10.0

Table 1. Dimensions of pure GG samples for the compression test under dry conditions.

Sample	h [mm]	D_{\min} [mm]	D_{\max} [mm]	m [mg]
GG00 1 GG00 2	8.35 8.28	4.94 5.05	4.97 5.06	12.0 10.6
GG003	8.23	5.04	5.11	10.1
GG00 4 GG00 5	$8.22 \\ 8.17$	$4.99 \\ 5.08$	$5.05 \\ 5.10$	$11.3 \\ 11.4$

Table 3. Dimensions of pure GG samples for the compression test under wet conditions.

Sample	h [mm]	D_{\min} [mm]	D_{\max} [mm]	m [mg]
GG50 1	8.87	4.99	5.10	14.8
GG50 2	9.03	5.09	5.09	14.5
GG50 3	9.10	4.94	5.10	15.0
GG50 4	8.80	5.04	5.06	14.0
GG50 5	8.97	4.92	4.97	14.2

Table 5. Dimensions of GG samples with $50\,\mathrm{wt}\%$ BAG for the compression test under dry conditions.

advantage is an ability to form highly porous 3D structures, when properly cross-linked and fabricated [12]. In the terms of bone regeneration characteristics, the GG is biocompatible and biodegradable, but its mechanical properties are inappropriate to bear stresses during normal movement of a patient and to enable successful bone regeneration. The material also does not promote natural bone formation. Therefore, the GG needs to be reinforced by the bioactive glass particles (BAG).

The BAG is a nano-particulate amorphous material with the chemical composition of $70\,\mathrm{n}\%$ SiO₂ and $30\,\mathrm{n}\%$ CaO, which is prepared by the modified sol-gel method. The BAG is unique since, during degradation, it can induce precipitation of hydroxyapatite formation (even *in vitro*) and consequently bonds toward soft and hard tissues. Therefore, the BAG particles are highly interesting for bone regeneration applications.

During the production process of the investigated GG-BAG samples, the gellan gum was dissolved in ultrapure water by heating the solution for 30 minutes at 90 °C. To the hot GG solution, a dispersion of the BAG was admixed and 0.18 wt% $CaCl_2$ was added. Kept under high temperatures, this mixture was subsequently

Sample	h [mm]	D_{\min} [mm]	D_{\max} [mm]	m [mg]
GG00 1	9.12	5.04	5.05	10.4
GG00 2	8.92	4.91	5.00	10.6
GG00 3	8.91	4.92	4.94	10.6
GG00 4	8.90	4.94	5.00	10.7
GG00 5	9.04	4.91	5.02	11.0

TABLE 2. Dimensions of pure GG samples for the tensile test under dry conditions.

Sample	h [mm]	D_{\min} [mm]	D_{\max} [mm]	m [mg]
GG00 1	8.25	4.96	5.02	10.6
GG00 2	8.24	4.93	4.98	10.7
GG00 3	8.21	4.97	5.01	10.8
GG00 4	8.13	4.96	5.02	11.5
GG00 5	8.37	4.99	5.02	10.1

Table 4. Dimensions of pure GG samples for the tensile test under wet conditions.

Sample	h [mm]	D_{\min} [mm]	D_{\max} [mm]	m [mg]
GG50 1	8.98	4.93	5.07	15.1
GG50 2	8.62	5.06	5.08	14.4
GG50 3	8.74	4.97	5.05	14.7
GG50 4	9.09	4.94	4.95	14.6
GG50 5	9.05	4.90	5.01	14.4

Table 6. Dimensions of GG samples with $50\,\mathrm{wt}\%$ BAG for the tensile test under dry conditions.

poured into the required mould and left there to be spontaneously jellified. The weight ratio of the GG and the BAG was 1:1 and the final concentration of $CaCl_2$ was 0.03 wt% in all samples. Such samples were frozen for 12 hours at -80 °C and freeze-dried for three days in a freeze dryer.

Cylindrical samples with height $h \approx 9 \, \mathrm{mm}$, diameter $h \approx 5 \, \mathrm{mm}$, and weight $m \approx 11 \, \mathrm{mg}$, $\approx 16 \, \mathrm{mg}$ and $\approx 24 \, \mathrm{mg}$ for the pure GG scaffold, GG-BAG reinforced scaffold with 50 wt% and 70 wt% BAG respectively were used for the testing. Dimensions are listed in detail in Tabs. 1–12.

X-ray microtomography imaging was performed on all three types of material. Based on reconstructed volumetric data porosity and pore size analysis was performed. Region of interest with face of inscribed square to the face of the sample and height of 75% of the sample was binarized. Using this data the porosity 67.46, 61.56, and 35.79% for pure GG, 50 wt% BAG and 70 wt% BAG content samples respectively was obtained. Microstructure of the samples were irregular with closed pores (but with some broken cell-walls). Pore size was assessed from medial, lateral and frontal section of each imaged sample. The pores in shape of

Sample	h [mm]	D_{\min} [mm]	D_{\max} [mm]	m [mg]
GG50 1	8.40	4.93	4.98	18.0
GG50 2	8.56	5.07	5.10	17.7
GG50 3	8.38	4.91	4.96	18.0
GG50 4	8.24	4.93	5.01	16.5
GG50 5	8.34	5.06	5.11	17.1

Table 7. Dimensions of GG samples with $50\,\mathrm{wt}\%$ BAG for the compression test under wet conditions.

Sample	h [mm]	D_{\min} [mm]	D_{\max} [mm]	m [mg]
GG70 1	9.16	5.01	5.08	23.6
GG702	9.12	5.00	5.09	24.4
GG703	8.85	4.93	4.94	24.1
GG704	8.88	5.04	5.05	24.5
GG705	8.66	5.05	5.09	24.5

Table 9. Dimensions of GG samples with $70 \,\mathrm{wt}\%$ BAG for the compression test under dry conditions.

Sample	h [mm]	D_{\min} [mm]	D_{\max} [mm]	m [mg]
GG70 1	8.64	5.06	5.09	24.1
GG70 2	9.10	4.95	5.02	24.4
GG70 3	8.62	4.90	5.04	24.3
GG70 4	9.05	4.95	4.96	24.4
GG70 5	8.66	4.94	5.07	23.3

Table 11. Dimensions of GG samples with $70\,\mathrm{wt}\%$ BAG for the compression test under wet conditions.

prolate ellipsoids were dominantly oriented in loading direction. Length of pores were $796\pm334\,,\,565\pm127$ and $429\pm72\,\mu\mathrm{m}$ for pure GG, $50\,\mathrm{wt}\%$ BAG and $70\,\mathrm{wt}\%$ BAG content samples respectively.

3. Methods

To obtain information about deformation characteristics of the synthesized material, sets of quasi-static experiments were performed. The first goal was to demonstrate possibilities of the in-house developed experimental infrastructure for such measurement. Expected collapse force ranged within single Newton orders and precise loading plate positioning was required as well. To obtain more relevant results, modifications of the experimental devices presented in detail in Section 3.2 were carried out. Then, material properties in terms of stress-strain response were obtained.

3.1. Experimental procedure

Cylindrical samples with the dimensions and weights listed in Tabs. 1–12 were subjected to tensile and compressive loading under dry and wet conditions. Here, the dry conditions represented the final state of the

Sample	h [mm]	D_{\min} [mm]	D_{\max} [mm]	m [mg]
GG50 1	8.27	5.05	5.07	17.2
GG502	8.30	5.05	5.08	16.5
GG503	8.20	5.07	5.08	17.0
GG504	8.18	5.03	5.05	17.3
GG505	8.31	5.10	5.14	17.0

TABLE 8. Dimensions of GG samples with 50 wt% BAG for the tensile test under wet conditions.

Sample	h [mm]	D_{\min} [mm]	D_{\max} [mm]	m [mg]
GG70 1	8.61	4.93	4.96	23.0
GG70 2	8.86	4.91	4.94	23.6
GG70 3	8.63	4.93	5.00	24.2
GG70 4	8.91	4.97	5.10	24.4
GG70 5	9.12	4.94	5.00	24.5

Table 10. Dimensions of GG samples with 70 wt% BAG for the tensile test under dry conditions.

Sample	h [mm]	D_{\min} [mm]	D_{\max} [mm]	m [mg]
GG70 1	8.20	5.09	5.15	22.6
GG702	8.34	5.10	5.15	23.2
GG70 3	8.26	4.93	4.99	22.2
GG70 4	8.24	4.97	5.01	23.3
GG705	8.25	5.05	5.07	24.7

Table 12. Dimensions of GG samples with 70 wt% BAG for the tensile test under wet conditions.

scaffold synthetization, whereas the wet conditions simulated physiological environment of human body using artificial plasma with the content listed in Tab. 13. Dry samples were loaded at room temperature (22 °C). The samples placed in the bioreactor chamber were fully immersed in the solution with temperature 37 ± 2 °C. Because of permeability (required for transport of nutrients) of the material, full saturation by simulated body fluid was used during the test. Loading platen displacement was set to approximately 1000 µm which corresponds to $\approx 11-12\%$ deformation sufficient to induce some observable damage to the samples' microstructure [13, 14]. Loading rate was set to $2 \,\mu m \, s^{-1}$ yielding quasi-static strain rate $\approx 2 \cdot 10^{-4} \, \mathrm{s}^{-1}$. The force and position was read-out with the sampling frequency 50 sps.

Natrii chloridum	$5.26{\rm g/l}$
Kalii chloridum	$0.37{\rm g/l}$
Magnesii chloridum hexahydricum	$0.30{\rm g/l}$
Natrii acetas trihydricus	$3.68{\rm g/l}$
Natrii gluconas	$5.02\mathrm{g/l}$

Table 13. Content of the artificial plasma.

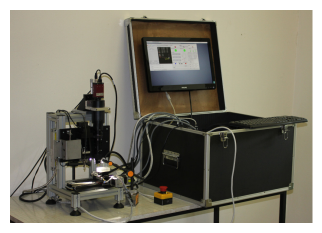


FIGURE 1. Custom designed micro-loading device with control unit.

3.2. Instrumentation

The in-house developed loading device (depicted in Fig. 1) for low-force indentation was adapted for both tensile and compression testing. For the indentation, the device was designed using the stiff modular aluminium ($30 \times 30 \,\mathrm{mm}$ hollow profiles) frame bearing the following components: (i) two motorized axes KK40 (HIWIN, Japan) for sample positioning accuracy $10 \,\mathrm{\mu m}$, (ii) loading axis based on linear stage MGW12 (HIWIN, Japan), and (iii) linear actuator 43-series (Haydon Kerk, USA) with positioning full-step resolution $3 \,\mathrm{\mu m}$ with mounting for U9B/C series (HBM, Germany) load-cell. This axis was upgraded using the same series linear actuator with positioning full-step resolution $1.5 \,\mathrm{\mu m}$, encoder with resolution $0.5 \,\mathrm{\mu m}$, and U9B/C load cell with nominal force $50 \,\mathrm{N}$.

For the testing under the wet conditions, the device was equipped with a custom designed bioreactor (see Fig. 2). The bioreactor enables full control over flow-rate and temperature of the fluid used for the experiments. The fluid is pumped from a heated reservoir to a basin surrounding the samples and loading platens; circulation of the fluid is provided by a peristaltic pump with adjustable stream velocity.

The control unit of the experimental setup was designed specifically for controlling stepper- or servomotor based positioning devices. The control unit is also equipped with drivers, an I/O board for controlling peripheral devices (i.e. lights, etc.), and a unit for readout signals from load-cells. The used drivers are capable of microstepping up to 64 microsteps per steps to achieve a maximum possible smoothness of motion, when stepper motors are used for the experiments.

3.3. STRAIN CALCULATION

The investigated material was expected to exhibit very low stiffness, which, coupled with high porosity and suboptimal geometry of the samples, induces high potential for significant boundary effects (i.e. localized plastic deformation of the sample near the contact with the loading platens) yielding yielding low reliability

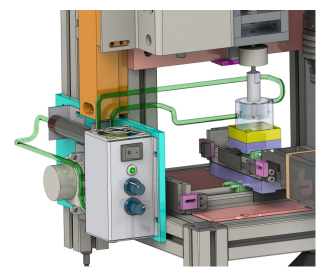


FIGURE 2. Visualization of bioreactor mounted on loading device.

of strain evaluation based on displacement of loading platens.

The accuracy of the presented measurements was reduced due to geometrical properties of the samples, the type of boundary conditions during experiments, and also by very low stiffness of the material. Currently, the production process of the GG-BAG samples does not allow to produce cylindrical samples too reliably. The diameter of the specimens varied in average by $\pm 100\,\mu\text{m}$, the loaded faces were rough and not planparallel. Furthermore, the character of the material, particularly its brittleness that is making it prone to severe damage during any type of preparation procedures, made machining of the samples impossible.

Thus, the non-contact optical displacement measurement was employed instead of calculations based on the known position of the loading platens. The strains were evaluated from optically acquired displacements using the digital image correlation (DIC) method. This method is based on the comparison of differences within the sequence of the deforming object images. When the image with identifiable texture is divided to subsets, the center coordinates (x', y') of arbitrary subsets in a subsequent image are given by:

$$x' = x + u + \frac{\partial u}{\partial x} \Delta x + \frac{\partial u}{\partial y} \Delta y, \tag{1}$$

$$y' = y + v + \frac{\partial v}{\partial x} \Delta x + \frac{\partial v}{\partial y} \Delta y, \tag{2}$$

where u and v are the displacements of the subset centroid in the X and Y directions respectively. Δx and Δy represent the distances from the centroid of the sub-image to the point x and y. This enables to create any arbitrary correlation pattern on the investigated surface, including a matrix for full-field displacement tracking.

Here, the matrix of correlation points for full field measurements was constructed on every measured sample and paths of the correlation points were observed for

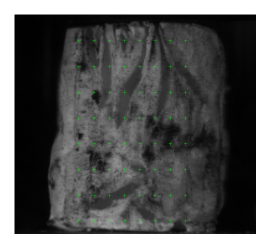


FIGURE 3. Surface of well manufactured sample with artificial patters and grid of correlation points for optical strain measurement.

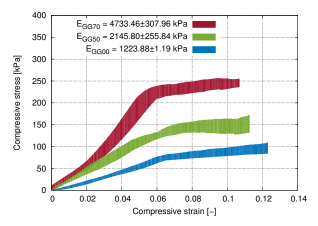


FIGURE 5. Stress-strain curves envelope dry scaffolds under the compressive loading.

calculating local deformations over the whole image sequence using the Lucas-Kanade tracking algorithm [15]. Using the obtained data, strain fields were derived and used for the calculation of stress-strain relations.

The full-field optical strain measurement of the wet samples placed in the fluid basin was not possible with the existing setup. Therefore, only comparative measurements on the dry pure GG samples and GG with 50 wt% BAG were performed. Here the average displacement of the bottom and top line of the correlation points (see Fig. 3) was used for strain calculation together with the encoder output. The resulting stress-strain curves of two compressive experiments are nearly identical in the elastic region. The strain measurement using the encoder signal can be considered acceptable only for such a measurement that can be seen in the comparative stress-strain diagrams depicted in Fig. 4.

3.4. Stress calculation

The stress σ in all experimental analyses was considered as engineering stress obtained using:

$$\sigma = \frac{F}{A_c},\tag{3}$$

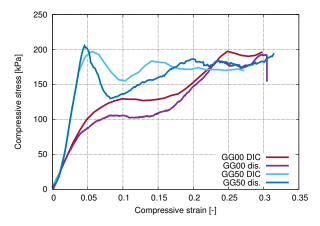


FIGURE 4. Stress-strain curves of two samples (pure GG in blue and 50 wt% BAG content in red) subjected to compressive loading under dry condition obtained from optical measurement and encoder signal.

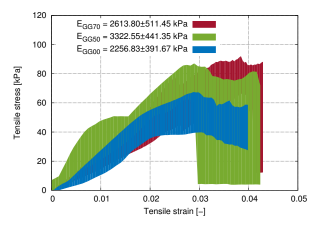


FIGURE 6. Stress-strain curves envelope of dry scaffolds under the tensile loading.

where A_c is the cross-sectional area of the specimen calculated from the minimal sample diameter measured before deformation. The force F was acquired by the load-cell. For the purpose of stress calculations, there were considered the ideally cylindrical samples neglecting all geometrical irregularities.

4. Results

Material properties and the deformation behaviour of the pure GG scaffold and the GG-BAG reinforced scaffold with $50\,\mathrm{wt}\%$ and $70\,\mathrm{wt}\%$ the BAG content were studied during the tensile and compression tests under the dry and wet conditions. Five experiments for each type of material, ambient conditions, and loading mode were performed. The stress–strain curves for each experimental batch are presented in Figs. 5–8. Young's modulus was calculated using the linear regression applied on the elastic part of the stress–strain diagrams. Regression error R was calculated using:

$$R = \sqrt{\frac{\sum_{i=1}^{n} (\sigma_i - \hat{\sigma}_i)}{n}},$$
 (4)

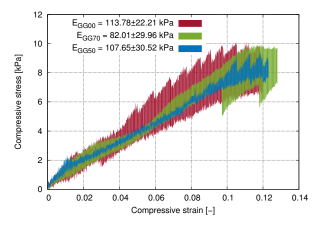


FIGURE 7. Stress-strain curves envelope wet scaffolds under the compressive loading.

Sample	E [kPa]	σ_Y [kPa]
GG00 1	1194.32 ± 4.16	61.52
GG00 2	1479.81 ± 6.27	76.65
GG00 3 GG00 4	1146.19 ± 4.77 1119.64 ± 5.26	$88.51 \\ 56.71$
GG00 5	1179.43 ± 3.20 1179.43 ± 3.07	62.28
Average	1223.88 ± 1.19	69.14 ± 13.14

Table 14. Elastic properties and yield stresses of pure GG samples for the compression test under dry conditions.

Sample	E [kPa]
GG00 1	88.58 ± 0.14
GG002	119.50 ± 0.15
GG003	126.15 ± 0.14
GG004	93.57 ± 0.14
GG00 5	141.08 ± 0.14
Average	113.78 ± 22.21

Table 16. Elastic properties of pure GG samples for the compression test under wet conditions.

where $\hat{\sigma}$ represent the calculated values of σ . For the yield stress $\sigma_{\rm Y}$ evaluation, the Christensen [16] second derivation criteria was used:

$$\sigma_{\rm Y} = \sigma \quad \text{at} \quad \left| \frac{\mathrm{d}^2 \sigma}{\mathrm{d}\epsilon^2} \right| = \text{max}.$$
 (5)

For noise reduction, significantly distorting resulting derived function, a rolling average filter was computed from five values as:

$$\sigma_{n_a}' = \frac{\sigma_{n-2}' + \sigma_{n-1}' + \sigma_n' + \sigma_{n+1}' + \sigma_{n+2}'}{5}.$$
 (6)

The calculated Young's modulus, regression error, presented as $E \pm R$ and the yield stresses σ_Y are listed in Tabs. 14–25 together with mean values of each set of measurements and standard deviation of such calculated values. The yield stresses were presented only

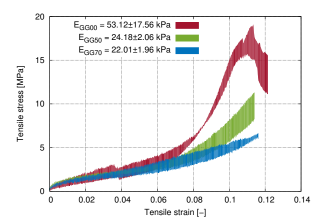


FIGURE 8. Stress-strain curves envelope of wet scaffolds under the tensile loading.

Sample	E [kPa]	σ_Y [kPa]
GG00 1	2164.31 ± 0.92	36.18
GG002	2259.77 ± 0.62	45.83
GG00 3	2915.55 ± 1.10	67.12
GG00 4	2036.73 ± 0.40	48.78
GG00 5	1906.54 ± 0.42	45.18
Average	2256.58 ± 391.67	48.62 ± 11.36

Table 15. Elastic properties and yield stresses of pure GG samples for the tensile test under dry conditions.

Sample	E [kPa]
GG00 1	56.34 ± 0.84
GG002	33.40 ± 0.20
GG003	36.38 ± 0.41
GG004	69.07 ± 0.94
GG00 5	70.43 ± 0.95
Average	53.12 ± 17.56

Table 17. Elastic properties of pure GG samples for the tensile test under wet conditions.

for the dry samples. In case of testing under the wet conditions, the loadcell output signal-to-noise was too low for the yield stress evaluation. All obtained results in the form of the enveloped strain–stress curves are plotted in Figs. 5–8.

5. Conclusion and discussion

The GG-BAG samples with different fraction of the reinforcing BAG particles were subjected to optically evaluated uni-axial measurements under both dry and wet conditions. It was found out that the ambient environment has significant influence on the mechanical response of the material and measured properties of the dry and wet scaffolds were apparently different.

The scaffolds wetted by a synthetic plasma solution exhibited radical loss of stiffness while the elastic modulus decreased more than ten times. In addition,

Sample	E [kPa]	σ_Y [kPa]
GG50 1	2100.57 ± 0.46	132.52
GG502	2066.95 ± 3.01	127.33
GG503	1818.36 ± 1.57	93.46
GG504	2520.59 ± 1.01	132.68
GG50 5	2222.56 ± 0.42	140.42
Average	2145.81 ± 255.84	125.28 ± 18.39

Table 18. Elastic properties and yield stresses of GG samples with $50\,\mathrm{wt}\%$ BAG for the compression test under dry conditions.

Sample	E [kPa]
GG50 1	81.45 ± 0.14
GG502	131.88 ± 0.13
GG503	70.84 ± 0.14
GG504	113.89 ± 0.28
GG50 5	140.22 ± 0.20
Average	107.65 ± 30.52

Table 20. Elastic properties of GG samples with 50 wt% BAG for the compression test under wet conditions.

the values of the peak force, which was measured during the tests and averaged at $\approx 0.5\,\mathrm{N},$ may be influenced by the load-cell nonlinearity and low signal-to-noise ratios.

Therefore, it wasn't possible to evaluate the yield stress properly using the acquired load-cell signal.

All compressed wet samples showed very similar deformation behaviour. The standard deviation for the samples from one group was higher than the effects of the BAG reinforcement. Unexpectedly, the results obtained for the wet samples subjected to tensile loading showed higher elastic modulus and ultimate stresses of the pure GG samples. This can be attributed to the fact that, on the microstructural level, the BAG particles disrupt integrity of the wet gellan gum. In case of the dry samples subjected to compression loading, a significant reinforcement effect of the BAG was observed. The values of elastic modulus and yield stresses are two and three times higher for 50 wt% and 70 wt% BAG improved samples respectively.

During the tensile loading, the elastic properties of the samples were very similar independently on the content of the BAG reinforcing particles, but higher reinforcement increased the measured ultimate stress. The yield stresses of both the $50\,\mathrm{wt}\%$ and $70\,\mathrm{wt}\%$ BAG were $20\,\%$ compared to the pure GG samples. The samples loaded in tension were also able to resist the deformation up to $4\,\%$.

Possibilities of experimental infrastructure for the testing of the newly synthetized GG based scaffolds was successfully demonstrated. The most limiting part of the experimental setup is the loadcell signal-to-noise ratio at the desired loading level in case of the wet

Sample	E [kPa]	σ_Y [kPa]
$GG50\ 1$	5805.60 ± 0.30	40.15
GG50 2 GG50 3	3137.46 ± 0.23 2520.69 ± 0.27	85.71 67.99
GG50 4	2520.09 ± 0.27 2157.34 ± 0.31	63.40
GG50 5	2991.67 ± 1.02	85.25
Average	3322.55 ± 1441.35	68.50 ± 18.75

Table 19. Elastic properties and yield stresses of GG samples with 50 wt% BAG for the tensile test under dry conditions.

Sample	E [kPa]
GG50 1	24.40 ± 0.09
GG502	27.15 ± 0.08
GG503	24.28 ± 0.08
GG504	23.71 ± 0.08
GG505	21.37 ± 0.08
Average	24.18 ± 2.06

TABLE 21. Elastic properties of GG samples with 50 wt% BAG for the tensile test under wet conditions.

samples and generally suboptimal geometrical characteristics of the samples inducing shear stresses during loading. For more complex investigation including the detailed analysis of the deforming microstructure, the time-lapse radiographical methods will be used [17, 18].

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Sample	E [kPa]	σ_Y [kPa]
GG70 1	4857.36 ± 0.34	208.05
GG70 2 GG70 3	4441.11 ± 0.38 5058.45 ± 0.32	204.68 202.55
GG70 4	4371.74 ± 4.38	224.20
GG70 5	4938.64 ± 0.23	213.25
Average	4733.46 ± 307.96	210.55 ± 8.63

TABLE 22. Elastic properties and yield stresses of GG samples with 70 wt% BAG for the compression test under dry conditions.

Sample	E [kPa]
GG70 1	134.42 ± 0.12
GG702	63.56 ± 0.07
GG703	79.59 ± 0.07
GG704	67.37 ± 0.08
GG705	65.13 ± 0.07
Average	82.01 ± 29.96

Table 24. Elastic properties of GG samples with $70\,\mathrm{wt}\%$ BAG for the compression test under wet conditions.

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Sample	E [kPa]	σ_Y [kPa]
GG70 1	2565.45 ± 0.68	83.81
GG70 2 GG70 3	3087.56 ± 0.22 2847.83 ± 1.07	56.13 68.55
GG70 4	1761.72 ± 0.67	65.98
GG70 5	2806.60 ± 1.06	88.45
Average	2613.83 ± 511.06	70.78 ± 13.30

TABLE 23. Elastic properties and yield stresses of GG samples with 70 wt% BAG for the tensile test under dry conditions.

Sample	E [kPa]
GG70 1	22.76 ± 0.08
GG702	23.44 ± 0.08
GG703	20.90 ± 0.08
GG704	21.14 ± 0.10
GG70 5	22.70 ± 0.07
Average	22.19 ± 1.10

TABLE 25. Elastic properties of GG samples with 70 wt% BAG for the tensile test under wet conditions.

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