

Merging flexibility with superinsulation: machinable, 1 nanofibrous pullulan-silica aerogel composites 2 3 Shanyu Zhao^{1*}, Olivier Emery¹, Anja Wohlhauser², Matthias M. Koebel¹, Christian Adlhart 4 ², Wim J. Malfait ^{1*} 5 6 ¹ Laboratory for Building Energy Materials and Components, Empa, Überlandstrasse 129, 8600 Dübendorf, Switzerland 7 ² Institute of Chemistry and Biotechnology, Zurich University of Applied Sciences ZHAW, 8 9 Einsiedlerstrasse 31, CH-8820 Wädenswil, Switzerland 10

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Abstract: Freeze-dried nanofibrous scaffolds are flexible, but typically have high thermal 13 conductivities. Conversely, silica aerogel has an ultra-low thermal conductivity, but is brittle. 14 Here, the impregnation of pullulan/PVA nanofiber scaffolds with hydrophobic silica aerogel 15 decreased the thermal conductivity from 31.4 to 17.7 mW/($m \cdot K$). The compatibility between 16 the silvlated nanofibers and the silica aerogel promotes the overgrowth of silica particles onto 17 the fiber surfaces and the fiber incorporation. The composites display improved compressive 18 and tensile properties compared to the neat pullulan scaffold and silica aerogel. The 19 20 composite's E-modulus is 234 kPa compared to 4 kPa for the pullulan scaffold and 102 kPa for the silica aerogel. The composite's tensile strength is five times higher than that of the 21 22 silica aerogel. Because of its reduced brittleness, the pullulan-silica aerogel composites can be 23 shaped using a sharp blade. The composites can sustain uniaxial compression up to 80%

24	strain, but the decompressed composites display two times higher densities because the strain
25	is partially irreversible. This densification reduces thermal conductivity to 16.3 mW/(m·K)
26	and increases final compressive strength by a factor of seven. Both the as prepared and
27	densified composites demonstrate unique material properties in terms of thermal conductivity,
28	mechanical strength and machinability.
29	
30	Keywords: nanocomposite; polysaccharide; nanofibers; hybrid aerogels; densification

32 **1. Introduction**

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Biopolymer aerogels were among the first aerogels ever synthesized [1] and have seen a 34 strong revival in the last decade, motivated by the potential for new technical applications of 35 aerogels, the search for more sustainable aerogel precursors, and a strong motivation from the 36 academic community to add value to biopolymer materials [2]. Among biopolymers, 37 polysaccharides have received particular attention, because they are available in large 38 quantities from agricultural production, often as a by-product (pectin) or waste material 39 (cellulose, chitin), and their use as a feedstock for advanced materials could be a great value 40 41 proposition. Many biopolymer aerogel studies target applications in thermal insulation [3-7], the most prominent industrial application of silica aerogel, but the vast majority of materials 42 produced do not reach the ultra-low thermal conductivities associated with silica aerogels [2, 43 8], despite some notable exceptions with thermal conductivities below 20 mW/(m·K), for 44 example from pectin [3, 9], chitosan [10, 11] and cellulose [8, 12-14]. In addition, concerns 45 about long-term stability have not yet been fully addressed. 46

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48 Cellulose, alginate, pectin, starch carrageenan and chitin/chitosan account for the vast majority of polysaccharide aerogel studies [2, 15-17], but alternative polysaccharides have 49 received far less attention in aerogel research. Pullulan is a starch-derived polysaccharide 50 51 based on maltotriose units connected by α -1,6 glycosidic bonds. Recently, low-density, freeze-dried scaffolds have been prepared from electrospun pullulan-polyvinyl alcohol 52 53 nanofibers [18-21]. These novel materials have a strong application potential in wound healing, gas filtration and oil-water separation, but do not possess a sufficiently high fraction 54 of mesopores to qualify as thermal superinsulators, which is confirmed by the thermal 55 conductivity measurements presented in this work (see below). 56

Here, we impregnated nanofibrous, silvlated pullulan scaffolds [18] with silica aerogel 58 59 through a sol impregnation based process. The rationale is that the small pore sizes in silica aerogels, which are below the mean free path length of air (~70 nm at 1 bar and 25°C), will 60 61 reduce the gas phase thermal conduction. At the same time, the tortuosity of the silica and pullulan nanofiber networks should limit the solid phase conduction. Recent publications on 62 aerogel materials and composites have combined silica aerogel with (bio)polymers and fibers 63 over a wide range of length scales. The industrially most relevant approach is the 64 impregnation of silica aerogel into macroscopic fiber blankets [22-25]. The use of nanofibers 65 represents an intermediate length scale, where either the nanofibers are added to the silica sol 66 67 in a cogelation approach [26-34]or a preformed nanofibrous scaffold is impregnated with a silica sol [35-37]. The latter approach is the one pursued in this study. At the smallest length 68 scale, the (bio)polymers or their monomeric precursors have been incorporated as individual 69 70 molecules, first through post-modification [38-40] and more recently through co-gelation approaches [41-46]. Finally, biopolymer-free, silicone-like rather than silica-like aerogels 71 72 with excellent thermal and mechanical properties have been prepared from functionalized 73 silanes [47, 48].

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Here, we aim to synthesize a material that combines the excellent thermal conductivity of silica aerogel with the flexibility of nanofibrous pullulan scaffolds. Therefore, previously fabricated, silylated nanofibrous pullulan scaffolds were impregnated with a silica sol, followed by gelation, aging, hydrophobization and supercritical drying. The resulting nanocomposites were then analysed in terms of their thermal and mechanical properties.

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81 **2. Experimental**

- 82
- 83 2.1 Synthesis

85 The preparation and characterization of the pullulan/PVA nanofibrous scaffolds is detailed in a previous publication [18] and will only be summarized here. Pullulan/PVA nanofibers were 86 prepared by free liquid surface electrospinning an aqueous solution of 4 wt% pullulan and 6 87 wt% polyvinylalcohol (PVA) at 80 kV. The recovered nanofibrous membrane was cut into ca. 88 1 cm² pieces and dispersed in 1,4-dioxane for 20 minutes at 13000 rpm. The homogenous 89 90 dispersion was cast into a mold, frozen, freeze-dried for 48 h and thermally cross-linked at 180°C for 40 minutes. Finally, the cross-linked pullulan/PVA scaffolds were hydrophobized 91 for 24 h by chemical vapour deposition in a desiccator saturated in trichloro(octyl)silane, 92 93 followed by the elimination of excess silane and HCl under vacuum (10 mbar, 2h).

The pullulan-silica aerogel composites were prepared according to Figure 1. A 94 polyethoxydisiloxane (PEDS) sol stock solution, i.e. pre-polymerized tetraethoxysilane 95 96 (TEOS) with a water-to-TEOS molar ratio of 1.5 and a SiO₂ equivalent concentration of 20 wt% in ethanol, was used as silica precursor. 30 ml of PEDS solution was diluted with 120 ml 97 ethanol (F25-A-MEK, 94 wt% ethanol denatured with 2% methyl ethyl ketone, Alcosuisse, 98 Switzerland) to a silica concentration of 4 wt%. Then, 5 ml of water and 1.2 ml of 5.5 M 99 NH₄OH aqueous solation were added to trigger gelation. The activated sol was portioned into 100 101 6 aliquots of 20 ml and cast into a prefabricated pullulan/PVA nanofibrous scaffolds, followed by 5 minutes degassing in a vacuum chamber (~10 mbar) to remove air bubbles 102 trapped in the fiber scaffold. Gelation typically occurred 10 to 15 minutes after the base 103 104 addition at room temperature. The gelled composites were covered with ethanol and aged 105 overnight at 55°C. The aged gels (120 ml in total) were hydrophobized by immersing them in a mixture of hexamethyldisiloxane (240 ml), ethanol (8.82 ml) and 37% HCl (0.96 ml). After 106 washing with ethanol once, the gels were dried supercritically in a SCF extractor (Autoclave 107 4334/A21-1, Separex, France). A set of reference silica aerogels was also prepared according 108 to the same procedure, but without the nanofibrous pullulan scaffold. The surfaces of the 109

dried samples were polished prior to performing the mechanical tests and thermal
conductivity measurements, with 400 grit sandpaper to remove the edges (menisci), and 2000
grit paper for finishing.

113



115 Figure 1. Synthesis scheme for the pullulan/PVA-silica aerogel nanocomposites.

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117 2.2 Characterization

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Density and Brunauer-Emmett-Teller (BET). The bulk density was calculated from the 119 mass and volume of the regularly shaped, cylindrical samples and the skeletal density was 120 approximated by the linear combination of the known skeletal densities of the constituents: 121 silica aerogel 2.0 g cm⁻³, pullulan 1.5 g cm⁻³, and PVA 1.2 g cm⁻³. The BET specific surface 122 of the composites was determined by nitrogen sorption after degassing at 100°C and 133 mbar 123 for 20 h. Nitrogen adsorption and desorption isotherms were obtained at liquid nitrogen 124 temperature on a Micromeritics TriFlex instrument with 15 seconds equilibration time. The 125 specific surface area of the samples was determined by the Brunauer–Emmett–Teller (BET) 126 method. The pore size distributions and pore diameter D'_{pore} were obtained from the 127

desorption branch of the isotherm using the Barrett–Joyner–Halender (BJH) model, despite
the known limitations of nitrogen sorption analysis in determining pore volume and pore size
[49].

SEM characterization and Energy Dispersive X-ray (EDX). Composite aerogels were 131 analyzed after coating with a platinum layer of nominally 10 nm, which refers to the thickness 132 of the Pt coating on the quartz sensor, whereas the actual thickness on the sample may be 133 quite a bit thinner because of the high surface area of the samples. SEM analysis of all 134 materials was performed on a FEI Nova NanoSEM 230 instrument (FEI, Hillsboro, Oregon, 135 USA) at an accelerating voltage of 10 kV and a working distance of 5 mm. The elemental 136 composition was obtained by using energy dispersive X-ray spectrometry (EDX, INCA X-137 Act, Oxford Instruments, UK) using 20 kV acceleration voltage and a 6mm working distance. 138

Solid-state NMR spectra were collected on a Bruker Avance III spectrometer equipped with a wide-bore 9.4 T magnet with ¹H, ¹³C and ²⁹Si Larmor frequencies of 400.2, 100.6 and 79.5 MHz, respectively. ¹H-¹³C cross polarization (CP) and ¹H-¹³C CP spectra were collected with respective contact times of 2 and 5 ms, 7 mm zirconia rotors, a magic angle spinning (MAS) rate of 4 kHz, a recycle delay of 2 s and between 180 and 27000 scans, depending on the sample.

145 Mechanical properties

Uniaxial compression tests of the composites were performed on monolithic cylindrical 146 samples (~22 mm diameter, ~ 30 mm high) using a universal materials testing machine 147 (Zwick/Z010, Zwick/Roell, Germany), equipped with a 10 kN force transducer (KAP-S, AST 148 Gruppe GmbH, Germany) in a controlled environment (23°C, 50% relative humidity). Elastic 149 moduli were measured in compression mode and were calculated from the linear region of the 150 151 stress-strain curves which typically occurred at 3 ± 2 % strain. A constant deformation rate of 1 mm/min was used and compressive strength values were taken at the first noticeable sign of 152 cracking. Tensile strength was estimated by the Brazilian split test, which is typically used in 153

154 cements and ceramic materials and entails the compression of a cylindrical sample lying on its 155 side [50]. The tensile strength σ_T can be calculated from the geometry and the compressive 156 force F as shown in Eq. 1, where D and L are the diameter and length of the samples. The test 157 setup was the same apparatus as used for the uniaxial compression test, but the loading cell 158 was adapted with a customized holder to fit to the cylindrical samples.

159
$$\sigma = \frac{F}{\pi \left(\frac{D}{2}\right)L}$$
(1)

160 **Thermal conductivity**

Thermal conductivity measurements of flat cylindrical tiles of approximately 45 mm in 161 162 diameter and 7 mm thickness were carried out on a custom-built guarded hot plate device designed for small samples / low thermal conductivity materials (guarded zone: 50×50 mm², 163 measuring zone: 25×25 mm²) at a temperature of 25°C on the hot side and 10°C on the cold 164 165 side (the apparatus is shown in Figure S1, SI).[51] In order to be consistent with measurements according to the European standards [52], calibration measurements were 166 carried out using conventional expanded polystyrene samples measured first in the standard 167 168 test equipment (conventional guarded hot-plate device) and then cut into smaller pieces to be measured in a second run in the smaller apparatus. The accuracy of the custom-built device 169 was determined to be $\pm 1.0 \text{ mW/(m \cdot K)}$ by measuring over 20 samples with known thermal 170 conductivities in the range of 15 to 25 mW/($m \cdot K$). 171

172 Water contact angle

The surface wettability of samples was evaluated by water contact angle measurement using a Contact Angle System OCA (Dataphysics TBU 90E, Germany), combined with a high-speed camera. Water droplets were deposited directly on the top or bottom surfaces of the samples. Two measurements were performed per sample and averaged. The volume of the water droplet was 5 µL, and the tip used was a precision stainless steel tip (Gauge 32, EFD).

179 **3. Results and Discussion**

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181 3.1 Chemistry, microstructure and the pullulan-silica aerogel interface

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The ¹H-²⁹Si CP MAS NMR spectrum of the pullulan/PVA scaffolds (Figure 2a) displays no 183 observable signals, indicating that the concentration of grafted octylsilyl groups is too low to 184 be detected by our current solid-state NMR experiment. This is not surprising, given the 185 relatively low surface area (~10 m²/g predicted from a nanofiber diameter of 240 \pm 55 nm) and 186 resulting low concentration of octylsilyl groups from the chemical vapour deposition of 187 trichloro(octyl)silane, with <5 wt% weight gain or <1 wt% of Si [18]. The grafted silanes 188 however can be detected as a (weak) Si signal in the EDX spectra of the silvlated 189 pullulan/PVA sponges (Figure S1). The ¹H-²⁹Si CP MAS NMR spectrum of both the 190 191 reference silica aerogel and the pullulan-silica aerogel nanocomposites displays the typical peaks from the trimethylsilyl (TMS) groups and the Q^n species of the silica aerogel, where Q^n 192 193 is a Si atom coordinated by n bridging oxygens (=Si-O-Si=) and 4-n non-bridging oxygens 194 $(\equiv$ Si-OH or \equiv Si-OCH₂CH₃).

195

The ¹H-¹³C CP MAS NMR spectrum of the pullulan/PVA scaffold displays the bands 196 197 expected for a mixture of pullulan [53] and PVA [54]. In addition, bands related to the caramelization products that originate from the thermal cross-linking step are present [55]. 198 Note that, as for the ¹H-²⁹Si CP spectrum, the signal from the grafted octylsilyl groups is not 199 intense enough to be detected in the ¹H-¹³C CP spectrum. The ¹H-¹³C CP MAS NMR 200 spectrum of the reference silica aerogel displays three main bands, related to trimethylsilyl 201 202 groups and the methyl and methylene groups of the ethoxy groups (\equiv Si-OCH₂CH₃), as expected for a PEDS derived silica aerogel prepared in an ethanol-based solvent [56, 57]. The 203 ¹H-¹³C CP MAS NMR spectrum of the pullulan-silica aerogel nanocomposite closely 204

resembles a linear combination of the spectra of the neat pullulan/PVA sponge (~25% of signal) and that of the reference silica aerogel (~75% of signal), in proportions that are consistent with the densities of the pullulan/PVA scaffold (0.030 g/cm³), reference silica aerogel (0.071 g/cm³) and pullulan-silica aerogel nanocomposite (0.099 g/cm³).

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Although CP NMR spectra are not quantitative in nature, a comparison of the relative peak 210 intensities for different samples does provide a qualitative measure of species concentrations 211 212 because the CP conditions were kept constant between the different measurements. The TMS/Q^n peak intensity ratio is higher in the ¹H-²⁹Si spectrum of the reference silica aerogel 213 than for the pullulan-silica aerogel composite (Figure 2a), indicating that the former is more 214 completely hydrophobized. A higher degree of hydrophobization should be associated with a 215 higher degree of polymerization because oxygen bridges (\equiv Si-O-Si(CH₃)₃) are formed during 216 the grafting reaction of TMS, and this is confirmed by the higher Q^4/Q^3 peak ratio for the 217 218 reference silica aerogel. Finally, the higher hydrophobization degree for the reference silica aerogel is also confirmed by the higher TMS/ethoxy peak ratios in the ¹H-¹³C spectrum 219 220 (Figure 2b). In previous studies on silica-pectin and silica-chitosan hybrid aerogels, prepared from molecular rather than nanofibrous polysaccharides, we have observed a similar effect, 221 i.e. a less complete hydrophobization for the hybrid aerogels compared to neat silica aerogel 222 223 [42, 43]. Note that this difference in TMS concentration does not affect the water contact angle, which is the same for both the reference silica aerogel and the composite (~140°, Table 224 225 1).



Figure 2. Solid-state MAS NMR spectra of the reference silica aerogel, silylated pullulan/PVA scaffold, and pullulan-silica aerogel composite; a) ¹H-¹³C CP spectra; b) ¹H-²⁹Si CP spectra.

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232 The pullulan/PVA scaffolds display a hierarchical pore structure with major cellular pores from the freeze-drying process, separated by pore walls with high local concentrations of 233 entangled nanofibers with minor pores in between. The pullulan/PVA nanofibers themselves 234 235 have fiber diameters on the order of 250 nm, consistent with those prepared in an earlier study [18], and have smooth surfaces. Locally, nanoparticle aggregates formed on the pullulan/PVA 236 nanofibers (Figure 3b) and EDX analysis (Figure S1) confirms these to be rich in Si. These 237 238 aggregates are clearly originating from the trichloro(octyl)silane chemical vapour deposition process. The Si content of the smooth fiber surfaces, further away from the nanoparticle 239 aggregates, is close to the EDX detection limit, i.e. either just below or just above the 240 detection limit, depending on the location of the analysis (Figure S1). 241



243

Figure 3. SEM images of a silylated pullulan/PVA scaffold (a,b) and the pullulan-silica aerogel composites (c,d).

247 The SEM images of the pullulan-silica aerogel nanocomposites confirm that the silica aerogel 248 phase completely fills the interstitial pores between the pullulan/PVA nanofibers and no macroscopic voids are present in the composites (Figure 3c,d). The silica aerogel phase itself 249 displays the particle-network mesostructure typical for silica aerogels, with secondary 250 particles of around 20 to 50 nm in diameter (Figures 4, 3d). The mesoporosity of the silica 251 aerogel inside the pullulan-silica aerogel nanocomposites is confirmed by nitrogen sorption 252 analysis (Figure S2), with a strong signal in the capillary condensation regime, a specific 253 surface area (S_{BET}) of 617 m²/g and a BJH mesopore volume of 2.4 cm³/g. Note that the 254 specific surface area and specific BJH mesopore volume of the silica aerogel phase within the 255 composite is most likely higher than the values reported above, because approximately 30% 256

of the mass of the composite consists of low surface area pullulan/PVA fibers. The average
pore diameter, as approximated from the density and surface area, assuming cylindrical pores,
shown in Equation 1.

$$D'_{pore} = 4 V_{pore} / S_{BE}$$
(1)

261 where $V_{pore} = 1/\rho_{envelope} - 1/\rho_{skeletal}$, is around 60 nm.

262

In a recent study, we observed a strong beneficial effect of silvlation on the mechanical 263 properties of cellulose foam-silica aerogel nanocomposites [35] and the same mechanism is 264 probably acting in the pullulan-silica aerogel system. The pullulan/PVA nanofiber surfaces in 265 266 the composites are overgrown with colloidal silica secondary aerogel particles (Figure 4), suggesting a good interfacial compatibility between the silvlated pullulan/PVA surface and 267 268 the silicon alkoxide based silica sol. The high compatibility is mediated by the nanofiber 269 surface modification with a trichloro(octyl)silane derived polysiloxane CVD layer that benefits from the high reactivity of the chlorosilanes to form \equiv C-O-Si \equiv bonds on the 270 biopolymer surface. The silica growth on the silvlated nanofibers progresses through ≡Si-O-271 Si≡ bond formation, which offsets the somewhat lower reactivity of the alkoxysilanes during 272 the silica sol-gel impregnation step compared to the higher reactivity of the chlorosilanes 273 274 during the chemical vapour deposition. CVD silane mediated silica overgrowth on the pullulan/PVA nanofibers (see schematic sketch in Figure 4) ensures that the nanofibers are 275 fully incorporated into the silica aerogel structure, also mechanically, and improves the 276 277 reinforcement effect that the nanofibrous scaffold has on the composite.



Figure 4. SEM image of a pullulan/PVA-silica aerogel composite and a schematic overview of the chemical interactions at the interface between the silylated nanofiber and the silica aerogel.

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283 **3.2 Thermal and mechanical properties**

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With a thermal conductivity of 31.4 mW/(m·K) (Table 1, Figure 5b), the hydrophobic, freeze-285 dried pullulan/PVA scaffolds are good thermal insulators in their own right, with a 286 performance that is similar to high performance mineral wool or polystyrene insulation. 287 However, they are not superinsulators because the pores between the pullulan/PVA 288 nanofibers are large (between 1 to 1000 µm) compared to the mean free path length of the air 289 290 molecules (~70 nm STP) and hence, gas phase conduction is not reduced by the Knudsen effect. The reference silica aerogel in this study has a thermal conductivity of 17.5 291 $mW/(m \cdot K)$, i.e. well below the ~26 mW/(m \cdot K) typical for standing air, because the gas phase 292 293 conduction within the aerogel mesopores is limited by the Knudsen effect. However, the

thermal conductivity of the reference silica aerogel is somewhat higher than that of an 294 optimized silica aerogel because of its specifically designed density (0.071 g/cm³) below the 295 optimum for minimal thermal conductivity (typically near 0.120 g/cm³ [50, 58]). With a value 296 of 17.7 mW/($m \cdot K$), the pullulan-silica aerogel composite has a near-identical thermal 297 conductivity to the reference silica aerogel, which indicates that the solid conduction through 298 the pullulan-fiber network is not significant. This is not unexpected as solid conduction 299 through low-density fiber networks is generally low [59, 60]. The ultra-low thermal 300 301 conductivity of the pullulan-silica aerogel composite is comparable to that of high quality silica aerogel and places this hybrid material among the very best silica-(bio)polymer hybrid 302 aerogels in terms of thermal conductivity [41-43, 61-63], in contrast to classical polymer-303 reinforced aerogels (X-aerogels) for which the, admittedly much stronger, increase in 304 mechanical strength is accompanied by a large penalty in density and thermal conductivity 305 306 [38, 64, 65].

307

308 The compression properties of the freeze-dried pullulan/PVA scaffolds have been described in 309 detail before [18]: the materials can sustain compression stress without rupture up to at least 80%, but display a low E modulus and final compressive strength commensurate with their 310 low density (Table 1, Figure 5a,b). The reference silica aerogel displays compressive 311 properties expected for low density silica aerogels (0.071 g/cm³): unlike the more elastic and 312 brittle intermediate and high density silica aerogels [50], the reference aerogels studied 313 plastically deform and can sustain uniaxial compression up to at least 80% strain. However, 314 several minor stress release events - linked to multiple crack initiations - are evident from the 315 stress-strain curves between 50 and 80% strain for some samples, leading to a relatively wide 316 scatter in final compressive strength, i.e. σ_{80} (Figure 5a). The pullulan-silica aerogel 317 composites also sustain uniaxial compression up to at least 80%, but have smoother, more 318 reproducible stress-strain curves. The composites display an increase in E modulus by a factor 319

of 56 and 2.3 compared to the neat pullulan/PVA scaffolds and the reference silica aerogel, 320 respectively, with similar increases in σ_{80} . The tensile strength of the silica aerogel and 321 pullulan-silica aerogel composite was estimated from the Brazilian split test [50] and is nearly 322 five times higher for the composite than for the reference silica aerogel (Table 1). As for the 323 324 compression tests, the stress-strain curves of the pullulan-silica aerogel composite are smooth, whereas those of the reference silica aerogel display minor episodes of step-wise stress 325 releases that decrease the tensile strength and lead to a large sample-to-sample variation 326 327 (Figure S3). Thanks to their improved compression and tensile properties, the pullulan-silica aerogel composites can be shaped simply by cutting with a sharp blade (Figure 5c), a 328 remarkable feature for silica aerogel composites. 329



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Figure 5. a) Stress-strain curves during uniaxial compression upon first compression; each curve corresponds to a different sample. b) Thermal and mechanical properties. c) Demonstration of the ability to shape the pullulan-silica aerogel composites by cutting.

Table 1. Density, thermal conductivity and mechanical properties.

						Compression (cylinders)			Compression (plates)				Brazilian test	
	ρ		λ		WCA	E		٤ _{max}	σ ₅₀		σ ₈₀		σ_T	
	[g/cm ³]	±	[mW/(m·K)]	±	[°]	[kPa]	±	[%]	[kPa]	±	[kPa]	±	[kPa]	±
Pullulan/PVA scaffolds	0.030	0.004	31.4	1.7	132	4	-	>80	8	-	48	-	n.a.	
Silica aerogel	0.071	0.002	17.5	0.6	140	102	27	>80	57	15	470	84	7	8
Pullulan-silica composite	0.099	0.005	17.7	0.9	141	234	23	>80	164	30	818	105	34	7
Compressed composite	0.182	0.015	16.3	1.1	n.a.	n.a.		n.a.	1057	-	6121	-	n.a.	
n.a.: not analyzed														

WCA: water contact angle

338 **3.3** Improving thermal and mechanical properties through uniaxial irreversible 339 compression

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A relatively low SiO₂ equivalent concentration in the silica sol was selected for the synthesis 341 of the reference and composite aerogels and this leads to a low density of the resulting 342 aerogels (Table 1). This low density in turn leads to thermal conductivities that are somewhat 343 344 higher than usual for silica aerogel, presumably because the low densities leads to larger pore sizes and a sub-optimal suppression of the gas phase thermal conductivity [50, 58]. Recently, 345 Plappert et al. [12] irreversibly compressed pre-formed low-density cellulose aerogels to 346 decrease the thermal conductivity and increase the mechanical strength. The same approach 347 was applied here. The pullulan-silica aerogel nanocomposites can sustain uniaxial 348 349 compression up to 80% strain, but display double the density after decompression, compared to the original composite (0.183 versus 0.101 g/cm^3). In other words, part of the strain is 350 irreversible. The compressed pullulan-silica aerogel composite displays a 1.4 mW/(m·K) 351 352 lower thermal conductivity of 16.3 mW/($m \cdot K$). The reduction in thermal conductivity is consistent with a decrease in gas phase conduction due to a decrease in average pore size: the 353 compression treatment reduces the average pore size from ~60 to 35 nm, as calculated from 354 the density, pore volume and BET surface area (560 m^2/g), to well below the mean free path 355 of air (70 nm at STP). More strikingly, the mechanical properties are improved by nearly an 356

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order of magnitude, for example the maximum compressive strength (σ_{80}) increased from 357 0.82 to 6.12 MPa (Table 1, Figure 6). The mechanical properties of the pullulan foam, silica 358 aerogel, composite aerogel, and compressed composite aerogel follow a single power-law 359 behaviour as a function of density, for example both σ_{80} and σ_{50} correlate with $\rho^{2.7}$ (Figure S4). 360 Power law behaviour is common for (biopolymer) aerogels [2, 26], but it is somewhat 361 surprising that the very different materials studied here define a single trend. This indicates 362 that, at least for the investigated systems, density is the primary factor that determines the 363 compressive properties, with less prominent effects from the composition of the solid phase 364 (pullulan/PVA versus silica) or the morphology of the microstructure (nanofibrous foam 365 versus particle based network). In summary, permanent densification through uniaxial 366 compression provides a substantial improvement of the thermal and the mechanical properties 367 of the composites, without compromising machinability (Figure S5). 368



Figure 6. a) Stress-strain curves under uniaxial compression of sample plates. b) Ashby plot of thermal conductivity versus σ_{80} .

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374 **3.4** Comparison with constituent reference compounds and other hybrid aerogels

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The thermal and mechanical properties illustrate that the pullulan-silica aerogel composite is more than just the sum of its parts and has superior properties compared to both the neat pullulan/PVA scaffold and the neat silica aerogel. This is true for the mechanical properties collected under lab scale conditions, e.g. compression testing on polished cylinders, but even more so for the real world behaviour, e.g. in terms of machinability, fragility and dust release. The spider plot in Figure 7a compares the four materials studied here in terms of their thermal

insulation performance, machinability, non-brittleness, dust release, hydrophobicity and E 382 383 modulus. The silica aerogel phase imparts the as-prepared and densified composites with an ultra-low thermal conductivity. The pullulan/PVA scaffolds impart the composites with a low 384 dust release and machinability. As discussed above, the E modulus of the composites benefits 385 from the higher density, particularly for the densified composite, but also for the as-prepared 386 samples (Table 1, Figures 5, 6, S4) and because of the power law dependence of the 387 388 compression properties on density, these effects exceed the benefits expected for a simple additive behaviour. Importantly, no application relevant properties are worse in the composite 389 390 than in the neat counterparts.

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Some application relevant properties of silica aerogel hybrids from the literature are plotted in 392 Figure 7b for comparison. Standard silica aerogel, with a density of 0.100 to 0.120 g/cm³, 393 394 displays elastic compression behaviour, but is very brittle and not machinable [50]. The closest materials in terms of microstructure and synthesis procedure are the silica aerogel -395 396 nanofibrillated cellulose foam composites we prepared in a previous study, which were also a 397 major inspiration for the current work. In that study, silvlation of the biopolymer foam was found to improve the interfacial compatibility with silica aerogels [18, 66], but because the 398 399 scaffold was impregnated with a standard density silica aerogel, the mechanical reinforcement effects were rather limited: because of its relatively low density, the silica aerogel employed 400 in the current study is non-brittle [50], and combined with the flexible, highly machinable 401 pullulan-PVA scaffolds, the composites inherit the merits from each component and 402 403 synergistically enhance the overall properties, particularly in terms of dust release and machinability. Macroscopic fiber blankets filled with silica aerogel [22, 23] are also easy to 404 405 cut into shape, but typically suffer from a high dust release, in contrast to the composites presented in this study, which display a very low dust release, presumably because of the high 406 interfacial compatibility (Figure 4) and because the nanofibrous scaffolds are more effective 407

at locking fractured aerogel particles in place (Figure 3). Classical polymer reinforced 408 aerogels (X-aerogels) have excellent mechanical properties, with particularly high E-moduli 409 [65], but strongly increased thermal conductivities. In contrast, recent (bio)polymer aerogel 410 411 reinforced silica aerogels prepared by co-gelation approaches have good thermal properties, a low dust release, but poor machinability [41-43]. Finally, single phase organic-silica aerogels 412 based on functional organosilanes have been developed with excellent thermal and 413 mechanical properties [47, 48]. Most of the cited materials above, and also the pullulan-silica 414 aerogel composites developed here, have not yet been evaluated for a range of other 415 application relevant properties, including fire behaviour and long term stability, but these are 416 beyond the scope of a scientific study. 417





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Figure 7. Comparison of thermal, mechanical and hydrophobic properties of a) the compressible silica aerogel (0.070 g/cm³), freeze dried pullulan scaffolds, and their asprepared and compressed hybrids developed in this study; b) PU-silica X-aerogels [65], cogelation polymer-silica aerogel hybrids [41-43], macroscopic fiber aerogel blankets [22, 23], NFC-silica aerogel nanocomposites [35], elastic silica aerogel (0.120 g/cm³) [50].

425

426 Conclusions

Recently, a wide variety of sol-gel derived, freeze-dried scaffolds have been prepared from 428 (bio)polymer nanofibers. Although these materials often have a low thermal conductivity, in 429 our case equivalent to that of high quality conventional thermal insulation, this is not low 430 enough to offset the more complex synthesis procedure. In addition, the materials are not 431 competitive with classical aerogels in terms of thermal conductivity. In this study, we have 432 demonstrated that the impregnation of silica aerogel decreases the thermal conductivity of the 433 pullulan/PVA scaffold from 31.4 to 17.7 mW/(m·K). At the same time, the pullulan-silica 434 aerogel nanocomposites display strongly improved compression and tensile properties by a 435 factor of 2 and 5 as compared with pure silica aerogel, which makes them easy to shape and 436 machine. Densification of the as-prepared, low-density nanocomposites further reduces 437 thermal conductivity to 16.3 mW/($m \cdot K$) and strongly increases the mechanical strength about 438 an order of magnitude and densification through uniaxial compression provides a simple way 439 440 to tune the thermal and mechanical properties towards specific applications.

441

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443

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447

448 Data availability

449 The raw/processed data required to reproduce these findings cannot be shared at this time due450 to technical or time limitations.

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