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Reinforcement of cellular materials with short fibres: application to a bio-based cork multi-scale foam

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Abstract. A bio-sourced foam, agglomerated cork, was chosen to evaluate the influence of short fibres on the mechanical behaviour of cellular materials. The final material was obtained by mixing cork particles with a thermoset resin. Rigid short fibres were then added before uni-axial compression. Enhancing the foam's mechanical properties without increasing the density is a current problem in transport industries. In this article, we demonstrate how the addition of short fibres strongly modifies the mechanical behaviour of agglomerated foam materials. Dynamic Mechanical Analysis technique revealed that the glass temperature was greater for reinforced foams and more energy loss by heat in visco-elasticity was also noticed for this material. In quasi-static compression, rigidity was strongly enhanced, causing absorbed energy before densification to increase. Maximal force and displacement before fracture were studied by applying Mode I fracture tests, and both were improved by the addition of short fibres. The mesoscopic and microscopic observations revealed it was linked to fracture mechanisms, most of which happen inside cork beads for the reinforced cellular material. The properties of agglomerated foams may then be improved and tailorable by the addition of short fibres and make weight saving possible in several industrial applications.

Keywords: Cellular Material, Microstructure, Cork Agglomerate, Short fibres, Visco-elastic behaviour, Mechanical Properties, Fracture

1 Introduction

Cellular materials are today widely used for a large range of application domains like transport, health, energy and sport. Their main characteristics are their low density, damping properties, thermal insulation and a capability to dissipate energy during impact loadings [1]. This last feature is especially interesting when considering passive safety applications. Depending on the level of stress required, it is possible to modify their composition or structure [2]. However, today this is mainly done by changing the density of the cellular material [3]. As with low-density materials, it is always important to minimise the weight while enhancing the mechanical strength for specific applications [4]. The introduction of reinforcements is one of the most commonly used methods to improve the mechanical properties in usual composite materials [5].

Fibres can be added while the foam is expanding. For rigid closed-cell polyurethane foams reinforced by glass fibres, fibres are located inside foam struts and cause localised changes in the foam morphology [6, 7]. Young's modulus is increased for reinforced foams. When fibres are organised in large bundles, they are not effective as a reinforcing agent because they cause stress concentration, thus weakening the structure. On the other hand, cellular materials in small bundles or lone fibres show longer fracture paths and pull-out fragments because of shear in the

radial struts. A toughening in the mechanical behaviour is then observed with the modification of the mechanism caused by the addition of fibres [8]. However, at extreme temperatures, reinforced foams are more brittle probably because of the difference in the thermal expansion coefficients between fibres and polyurethane [7]. The addition of glass or aramid short fibres to an epoxy foam also improves the mechanical properties globally in shear and to a lesser extent in compression. Increasing fibre length does not have a significant benefit [9].

One of the main drawbacks of adding fibres while expanding foams is the difficulty in controlling fibre dispersion and their layout in the material. Fibres also cause changes in the foam morphology during the foaming process [6, 7, 9].

To overcome these problems, cellular composites reinforced with short fibres can be of interest [4]. They could ensure that the mechanical properties of foams could be tailorable by controlling the compositions and microstructures of the constituent phases, and also the fibre orientation. Multi-scale foams [10] are also conceivable and are investigated here. They consist in agglomerated beads of foam [11]. Before agglomeration, short fibres are added to the preparation [12].

This study focuses on the influence of these short fibres on the foam microstructure and the mechanical behaviour of a multi-scale cellular material, agglomerated cork.

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As such materials are mainly used in compression, the compressive mechanical behaviour of reinforced foam was first studied. The influence of the short fibres on the visco-elastic mechanical behaviour was studied using dynamic mechanical analysis (DMA). The mechanical behaviour for large strains was then studied in quasi-static regime only as it shows the same trends as in dynamic regime when comparing several type of cork agglomerate [13].

The reinforced foam interface, besides containing cork particles and resin, is enriched with short fibres. To know if this involves premature fracture of the material as already observed for composite materials [14], mode I fracture tests were performed. In this kind of multiphase polymer cellular material, it is known that strong non-linear effects can occur during crack initiation and propagation as visco-plasticity effects [15, 16] and sometimes inertia effects [17]. A material parameter such as the critical energy release rate G_{Ic} will therefore not be estimated here as only the tensile strength will be analysed along with post-mortem fractographies.

In both compression and tensile loadings, microstructure was studied after testing in order to determine associated strain and failure mechanisms.

This article demonstrates how the addition of rigid short fibres increases the compressive stiffness of the material but also improves the fracture behaviour of cellular materials without changing the material mean density.

2 Material & Methods

2.1 Materials

Cork is a natural polymeric cellular material with a closed-cell structure [18]. It is formed by small anisotropic prismatic cells arranged in successive layers exhibiting a relatively homogeneous honeycomb structure. The mean length of the cells is around $40\text{ }\mu\text{m}$ and the cell wall thickness around $1\text{ }\mu\text{m}$.

Two agglomerated foam panels were studied, supplied by the Lieges HPK company: this is a classical cork agglomerate and one where short fibres have been added during the material preparation process. This material will be called the reinforced foam and was developed by Lieges HPK and Safran Power Unit for a common project. To obtain agglomerated cork, small beads of cork (with initial diameter $\varnothing_c = 0.5 - 1\text{ mm}$, density $\rho_c = 170\text{ kg m}^{-3}$ and rigidity $E_c = 20\text{ MPa}$ [18]) were mixed together in a tank with polyfurfuryl alcohol (PFA), a bio-based thermoset resin (with density $\rho_r = 1128.5\text{ kg m}^{-3}$ and rigidity $E_r = 2.1\text{ GPa}$ [19]). For the reinforced material, rigid short fibres from milled ex-PAN fibres were added afterwards (with diameter $\varnothing_f = 7 \pm 2\text{ }\mu\text{m}$, median length $l_f = 270\text{ }\mu\text{m}$ and rigidity $E_f = 230\text{ GPa}$) while the mixing step was continued. This addition is the only difference between the two manufacturing methods.

The material was then uni-axially compressed into a $1000 \times 500 \times 150\text{ mm}^3$ block. The compression caused a preferential orientation in the bead shape. Fig. 1(a) shows the two resulting directions: directions in the (Oxy) plane are called in-plane directions (IP) and (Oz) direction is

called out-of-plane direction (OP) [20]. Afterwards the cross-linking was made under compression in an autoclave. Slabs were then machined from the block where samples were collected.

Both materials were made with a cork/resin volume ratio of around 28:1 before compression. The volume fraction of short fibres added to the blending is around 0.02 % of the total volume for the reinforced agglomerate. The relative density is around 0.42, which is a high value in comparison to other cork agglomerates studied previously [21].

A multi-scale cellular material was thus obtained. Fig.1(b) shows the three scales that are considered here. First, the structure scale from which the material parameters will be deduced. After this the bead scale has millimetric lengths. Beads can be defined by orientation, volume and shape descriptors [20]. This scale shows several porosities between beads. Finally each bead is made of a cellular material. Therefore by looking more closely at the beads, cells can be spotted, here with a Scanning Electron Microscope (SEM). This cell scale is characterised by micrometric lengths. Each bead has a specific cellular orientation [20]. Cork cells have variable dimensions [22] but also variable chemical compositions [23].

2.2 SEM observations

In order to observe the structure of cells and interfaces, SEM samples were cut with a razor blade, which was replaced for each cut. This was done in order to cause as little damage as possible to the cell walls [18]. Each cube was cut over-size and then trimmed to the final size (roughly 5 mm sides). Samples were then lightly coated with conductive gold to allow electron conduction at the sample surface. The device used was a Cressington 108 auto. The gold layer was presumed constant on the sample surface.

Samples were then observed with a SEM (Zeiss EVO HD 15). An accelerated tension of 10 keV was used with a current of 200 pA .

2.3 Visco-elastic behaviour

The viscoelastic behaviour was determined by means of Dynamic Mechanical Analysis (DMA) in compression mode with a Mettler Toledo I controlled by STARE software. The sample underwent repeated small-amplitude strains in a cyclic manner. Molecules perturbed in this way store a portion of the imparted energy elastically and dissipate a portion in the form of heat [24]. In this case, the modulus acquires a complex form, $E^* = E' + iE''$. The quantity E' , the storage modulus, is a measure of the energy stored elastically, whereas E'' , the loss modulus, is a measure of the energy lost as heat or molecular rearrangements during the deformation [25]. The loss factor is defined as $\tan \delta = E''/E'$, δ being the angle between the in-phase and out-of-phase components in the cyclic motion.

Temperature scans were performed from $-80\text{ }^\circ\text{C}$ to $150\text{ }^\circ\text{C}$ with $10\text{ }^\circ\text{C}$ steps maintained for 30 min before

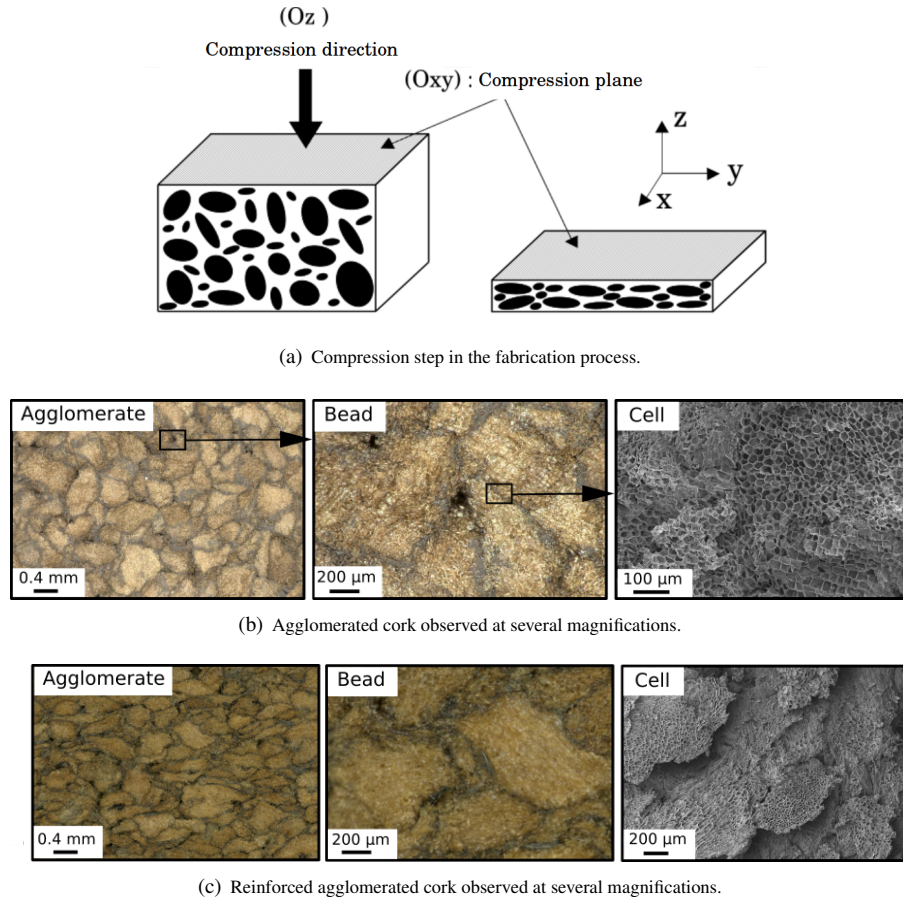


Figure 1. a) Compressive process and principal orientations of the material. b) Different observation scales demonstrating the multi-scale structure of the agglomerated cork obtained. c) Reinforced cork agglomerate observed at different scales. For both b) and c) at the agglomerate and bead scales, pictures are taken with a light microscope in the out-of-plane direction. Cells scale is illustrated with a SEM image of cork cells.

measurement at a 1 Hz frequency and an imposed displacement of $5 \mu\text{m}$. Samples were cubic with a 5 mm side length, the corresponding nominal strain was then 0.1 %. and a pre-strain of 0.01 % was chosen. Such samples are considered as representative of the thermo-mechanical behaviour of such cork agglomerates because of the small grain size (especially after the fabrication process) [20]. As hygrometry deeply influences the mechanical behaviour of polymers and thus cork [26], samples were conditioned beforehand at a relative humidity of 50 %.

2.4 Mechanical behaviour under large strain

2.4.1 Quasi-static compressive tests

An electromechanical traction/compression machine (Zwick Roell 250 from Allround-Line, controlled compliance calibration certified by Cofrac) with a load cell capacity of 250 kN was used. Conditioned cubic samples with a side length of 20 mm were cut. The loading speed was set at 5 mm min^{-1} . Mechanical behaviour was investigated in both directions (in-plane and out-of-plane) at room temperature (around 23°C). For each experiment,

5 to 8 samples were tested. A good repeatability was observed provided that samples were preconditioned.

2.4.2 Post-processing of the compression tests

From the force/machine displacement data, stress/strain curves were deduced by calculating macroscopic nominal stress ($\sigma = F/S_0$) and macroscopic true strain ($\varepsilon = \ln(l/l_0)$). As Poisson's ratio for cork is close to 0, the cross-section did not change during the compression test.

As it can be challenging to clearly distinguish the linear part in polymeric cellular materials [27], the stress/strain curve was fitted with a seven degree polynomial. The end of the linear part of the curve was found with its second derivative minimum, *i.e.* when the tangent starts to decrease. A one degree polynomial was then calculated between $\varepsilon = 0.01$ and this point. Its slope corresponds to Young's modulus E given in this article. The tangent module E_p was defined as the slope of the plateau stage between a 0.1 and a 0.3 strain. The elastic collapse stress σ_{el} marks the end of the linear regime and the start of the plateau stage. It was defined as the stress at which two first degree polynomials with a slope of E and E_p meet.

The densification strain ε_d was identified as the energy efficiency maximum strain [3, 28].

Absorbed energy corresponds to the sum of dissipated and elastic energies. It was obtained by integrating along the displacement using the composite trapezoidal rule. Absorbed energy density at densification was calculated by integrating the force/displacement curve as far as the densification displacement d_d . The energy found was then divided by the sample initial volume.

For each material parameter, uncertainty was calculated as the mean deviation. It gives the average of the absolute deviations of the observations from their arithmetic mean. Variation $v_{a/b}$ of material parameters between two material parameters p_a and p_b was calculated according to $v_{a/b} = (p_b - p_a)/p_a$.

2.4.3 Fracture initiation tests

Mode I fracture loading was studied as it is the least energy-consuming mode for the material. Strip Band Specimen geometry (SBS) was chosen to study the fracture behaviour of both materials (Fig. 2). It consists in a rectangular slab of $300 \times 100 \times 5 \text{ mm}^3$ cut with a band-saw. An initial notch of 100 mm was also machined into the sample with the band-saw to disregard border effects as much as possible and to localise crack initiation. The tip was shaped as a u-notch using a rotary tool. This shape was chosen to obtain a large crack length.

Samples were observed with a Keyence light microscope at $100\times$ magnification. The area near the notch was especially carefully observed. It is depicted by the black dotted line in Fig. 2. Pictures of 5500×3000 pixels were obtained. Samples were conditioned after these observations. The same sample was also observed after testing and was compared to the non-tested picture.

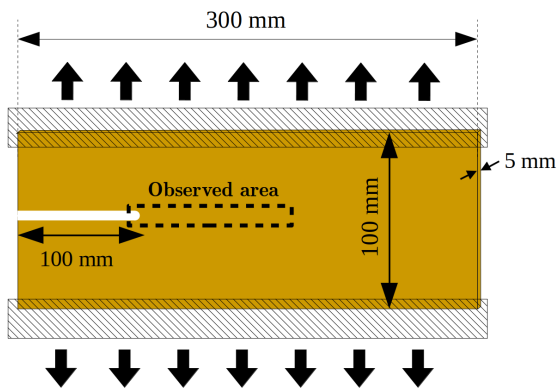


Figure 2. Set-up used to study fracture behaviour with SBS geometry. The black dotted line indicates the area observed with a light microscope before and after testing.

The same electro-mechanical traction/compression machine than previously was used. Displacement was imposed at a specific displacement rate until total degradation of the plate (10 mm/min). This relatively quick displacement rate was chosen in order to limit the activation of the

viscoelastic cork mechanisms (such as stress relaxation) [29].

From the force/displacement curve (Fig.7(a)), the initial slope was measured and assimilated to the initial stiffness, K . The maximal force reached during the test was identified as the fracture force F_{frac} . The corresponding fracture displacement was called d_{frac} . Due to the imposed displacement rate, the displacement of the plate kept increasing while the force was decreasing because of the sample fracture. When the plate was fully broken, the force was null and stable. At this stable force, the final displacement was called d_{max} . The difference between d_{frac} and d_{max} was also calculated and was called Δd .

Crack propagation was observed with a Canon EOS 5D camera at a 2 Hz frequency.

3 Results & Discussion

3.1 Microstructure

Both cellular materials have a very similar structure as can be observed in Fig.1(b) and 1(c). Apart from the interfaces, no major discrepancies were observed at the agglomerate, bead and cell scales. Macroscopic porosities can be seen in both samples (as in Fig. 3(c)). Using data from a previous study [20], the macroscopic porosity total volume was estimated to represent 10 % of the sample total volume for both materials.

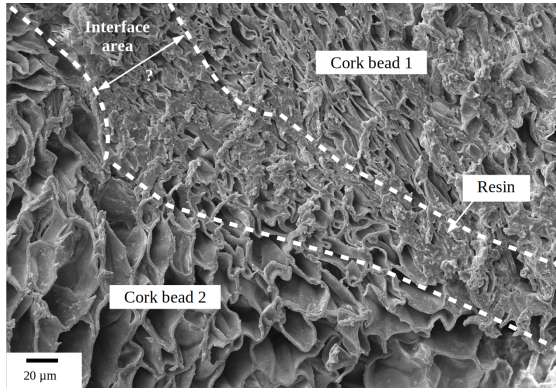
3.1.1 Interphases

Previous work on agglomerated cork revealed that for this kind of materials, interfaces are very thin (around $1 \mu\text{m}$) [20]. The anisotropic nature of cork cells [18] can help to distinguish one bead from another. Fig. 3(a) shows that instead of considering only the sharp resin/particle interface, an interfacial area called the *interphase* should be considered [30]. This is composed of collapsed cells at the bead periphery and resin coating the cork surface [20, 31] and will display a high gradient in its mechanical properties. The thickness of the interphase is variable but is around $30 \mu\text{m}$, as shown in Fig. 3(a).

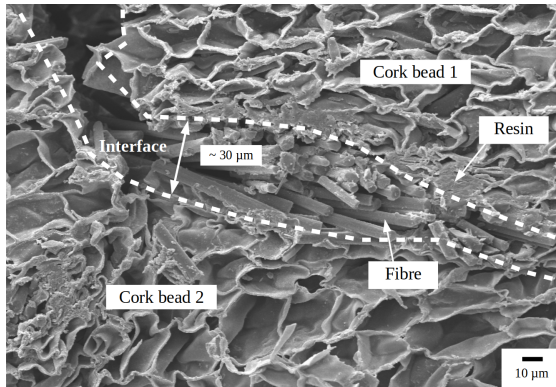
The composite material also presents an interphase with collapsed cells in the vicinity of the interface as shown in Fig.3(b). Resin also forms a thin film at the particle surface. Interphases are thicker, however (around $50 \mu\text{m}$) because of the fibres between the cork particles. The fibres appear to be well dispersed and are present between each bead. Like the beads, most of them seem to be coated with resin. They often organise themselves into bundles of a few coated fibres. Their average diameter (around $20 \mu\text{m}$) is much smaller than that of the foam beads (around $300 \mu\text{m}$ [20]).

3.1.2 Fibre placement and orientation

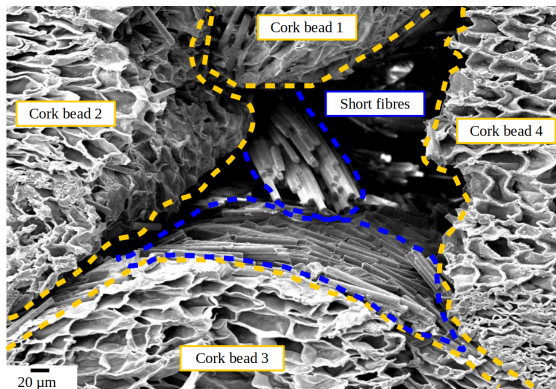
Fig. 3(c) shows that short fibres never damage a foam bead by going across it. For the most part they lie tangentially to the foam beads. As the small mean length of chopped



(a) Intergranular interphases in the unreinforced foam [20].



(b) Intergranular interphases in the reinforced foam.



(c) Short fibres arrangement in the reinforced foam.

Figure 3. SEM pictures of the interphases in cork-based composites.

short fibres is around 230 μm , they fit fairly well onto the bead surface.

As already noted, uniaxial compression manufacturing process causes a transversal anisotropy for high density foam agglomerates [20, 31]. Beads of the cellular material tend to be flattened in the compression plane, i.e. the (Oxy) plane, as shown in the Fig.1(a) and are much more oriented in this plane as proved by X-ray tomography observations [20]. Interphases between beads are then more oriented in this same plane, the in-plane direction. Short fibres are then more oriented in the (Oxy) plane.

3.2 Visco-elastic compressive behaviour

3.2.1 Description of the visco-elastic behaviour of cork agglomerates

Compressive DMA results are shown in Fig. 4 for the cork agglomerate and the reinforced foam samples tested in the out-of-plane and the in-plane directions. For each type of sample the storage modulus (E'), the loss modulus (E'') and the loss factor ($\tan \delta$) are presented in Fig. 4(a), Fig. 4(b) and Fig. 4(c).

Fig. 4(a) shows that for each type of sample, the storage modulus decreases with increasing temperature due to the softening of the cell wall material, as already noted for raw cork and other cork agglomerates [24, 26, 32]. A similar tendency concerning the storage modulus was found for other cork agglomerates with PFA resin [32].

In the early stages of the experiment (*ca* -80° C), a decrease in E' is noticed. This is related to the occurrence of a relaxational process. This relaxation is also observed in the $\tan \delta$ plot in Fig.4(c) and the loss modulus plot in Fig.4(b), where a small peak is observed at this temperature. This dynamic transition can be assigned to a β -relaxation of one of the main cork cell wall components, suberin [26].

From -10° C to 50° C, a stronger decrease in E' is observed. The broad peak on the $\tan \delta$ plot indicates the same relaxational process that was previously identified as an α -transition of some of the components of cork cell wall [26]. A glass transition is then identified at the $\tan \delta$ peak maximum (given in Tab. 1).

The $\tan \delta$ peak observed in Fig.4(c) from 80° C to above 150° C could be due to two processes. The first one is the α -relaxation of the resin and was observed for other cork agglomerates with a similar resin [32]. The glass transition temperature of this resin occurs between 70° C and 130° C depending on the cross-linking conditions [32]. However, on the E' curve, the storage modulus seems to be rather constant in this temperature range as the rubbery plateau of cork components is reached. The α -transition of the resin does not seem to influence the rigidity of the material very much. A relaxation peak was also observed for raw cork at around 80° C but was shifted to a higher temperature and decreased in intensity for an annealed sample. The second process could then be desorption of water molecules in the cork structure [24].

3.2.2 Effect of short fibres on the visco-elastic behaviour of agglomerated foams

When comparing storage moduli E' of cork agglomerate samples and reinforced foam ones in Fig. 4(a), as expected the storage modulus is greater for the reinforced foam samples. Short fibres are indeed much more rigid ($E_f = 230 \text{ GPa}$) than the matrix or cork particles.

A more unexpected result in the modification of the $\tan \delta$ peak corresponds to the α -relaxation in Fig. 4(c). It is less intense but wider and the maximum is shifted to higher temperatures. Tab. 1 shows that from 20° C for cork agglomerates, T_g lies around 40° C for the reinforced

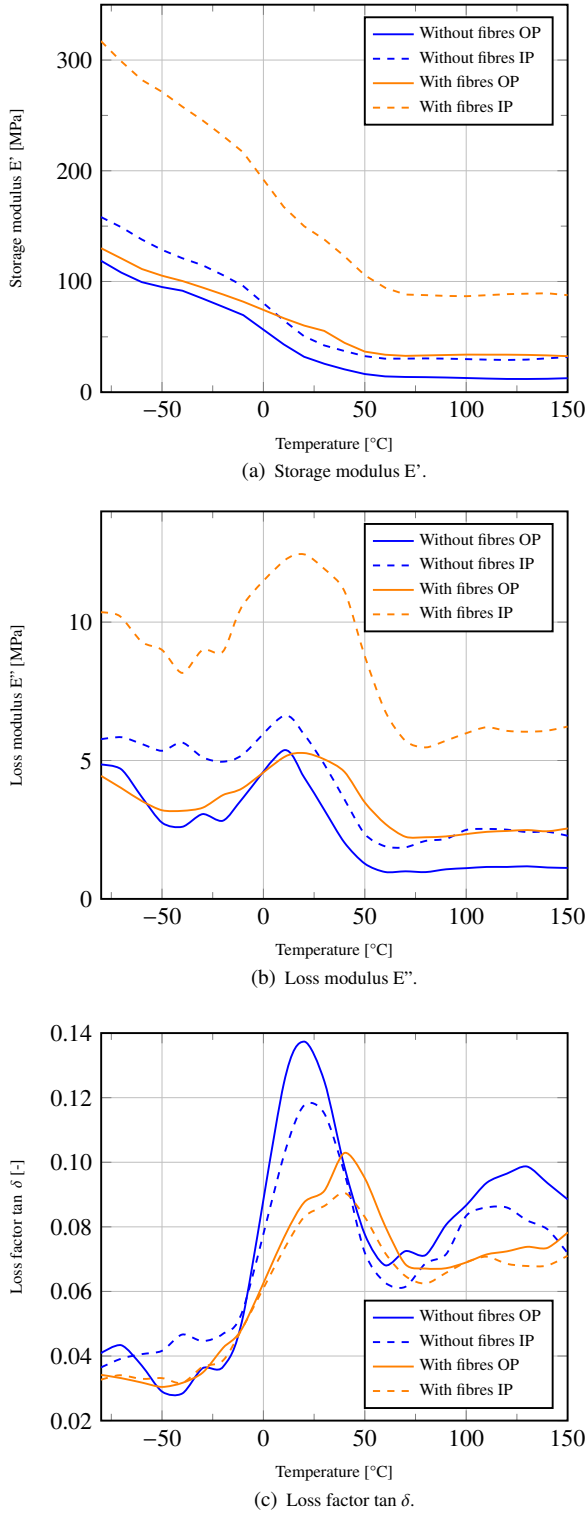


Figure 4. Compressive DMA spectra of a reinforced cork agglomerate and a classic cork agglomerate at a 1 Hz frequency in the in-plane (IP) and out-of-plane (OP) directions.

foam. This would mean that polymer chains in cork cell walls are more constrained in the presence of short fibres compared to a classic cork agglomerate because of the mismatch in thermal expansion coefficient which is relatively low for ex-PAN fibres in comparison to polymers [33].

The loss modulus E'' in Fig. 4(b) indicates the viscous nature of the material. As reinforced samples loaded in the out-of-plane direction show E'' values that are higher but with a similar evolution than the non-reinforced samples, the viscosity of the material does not seem to be changed by the addition of fibres.

The damping $\tan \delta$ of cork agglomerates is decreased by adding fibres. This suggests a good adhesion in the reinforced agglomerate and interfacial bonding plays its role in transferring the load. It should be noted that the damping capacity associated with the surface area of the $\tan \delta$ curve in reinforced agglomerates remains close to that of classical cork agglomerate as shown in Tab. 1. The surface in the out-of-plane (in-plane) direction decreases by 17 % (7 %). The first reason is that the $\tan \delta$ peak is larger for reinforced agglomerates. The other is that even if the rigidity increases with the presence of fibres, E'' also increases, as explained previously.

Material	T_g [°C]	Area [°C]
Agglomerated cork OP	20.2	7.56
Reinforced foam OP	40.3	6.39
Fibre effect OP	-	- 17 %
Agglomerated cork IP	20.2	6.27
Reinforced foam IP	40.3	5.81
Fibre effect IP	-	- 7 %

Table 1. Data extracted from DMA analysis. Glass transition T_g measured at the $\tan \delta$ maximum. Area under $\tan \delta$ between -20° C and 70° C at 1 Hz in compression.

3.2.3 Effect of short fibre orientation on the visco-elastic behaviour of agglomerated foams

The increase in rigidity caused by short fibres is much more obvious for in-plane samples. As the microstructure observations showed, fibres lie along the bead surface. Like beads, fibres will tend to align in the in-plane direction. They will be more solicited when loading happens in this direction. As fibres are more solicited, the is likely to be more friction. This explains why the in-plane reinforced sample in Fig. 4(b) shows a much higher E'' compared with the out-of-plane sample loss modulus, which is similar to non-reinforced samples. As in traditional composites, the storage modulus and the loss modulus depend strongly on the orientation of the fibres with the direction of oscillation [33]. The contribution of the interfacial region to the dynamic properties is more significant when the fibre-dominated properties are measured.

As for the damping properties, it seems that for both materials, samples tested in the out-of-plane direction show a more intense peak in the $\tan \delta$ plot (Fig. 4(c)) and

thus a greater area underneath. However, the higher rigidity in the in-plane direction compensates for the higher energy lost by heat in the in-plane direction and they seem to compensate each other, for both types of material.

3.3 Compressive behaviour for large strains

Cellular materials are partly used for their energy absorption properties in compression. The compressive behaviour of the two materials has thus been studied for quasi-static loading to investigate the influence of short fibres on the mechanical behaviour for large strains.

3.3.1 Macroscopic behaviour in compression

Fig. 5 displays the raw stress/strain curves of each material tested in out-of-plane (OP) or in-plane (IP) direction.

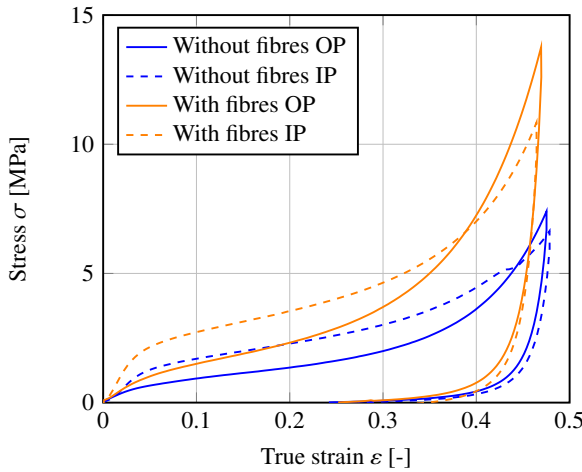


Figure 5. Experimental strain/stress curves of agglomerate cork and of reinforced foam tested in in-plane (IP) and out-of-plane (OP) directions at 5 mm/min at 26°C.

For both materials, a typical foam-like behaviour is noticed [3, 34]. For small strains, they show a linear stress-strain regime. Its slope is characterised by the Young's modulus E . Afterwards stress increases slowly until large strains. This stage is called the plateau stage and is characterised by an elastic collapse stress σ_{el} (where the plateau stage starts). Tangent modulus E_p expresses how the plateau increases due to gas compression [3]. Finally stress increases sharply around ε_d . This is called densification.

All material parameters described above (E , σ_p , E_p , ε_d and $\mathcal{E}_d$) were calculated for the agglomerated cork and the reinforced foam. Their values for each material and direction can be found in Tab.2.

3.3.2 Fibre effect

The comparison of the strain-stress curves indicates no major discrepancies in global mechanical behaviour. Both demonstrate a foam-like behaviour in compression with

the three distinctive stages: linear behaviour, plateau and densification.

In both directions, many material parameters are enhanced by the addition of fibres as indicated by Tab. 2. The Young's modulus increases by 26 and 71 %, in the out-of-plane and in-plane direction, respectively. The elastic collapse stress rises to 30 % (40 %) in the out-of-plane (in-plane) direction. The tangent modulus and the absorbed energy at densification are both also greater for the reinforced foam, by 81 % (53 %) in the out-of-plane (in-plane) direction.

Thus by comparing Young's modulus, the same conclusion is reached as for DMA tests: fibres stiffen foam agglomerates. As compression is pursued at a wider range here, a more complete analysis can be made. When solicited, fibres enhance the rigidity of the interfaces. This leads to a more rigid cellular material, which is characterised (in addition to the Young's modulus) by a higher stress plateau and thus more absorbed energy during compression than observed for other types of reinforced cellular materials [35].

Conversely densification strain ε_d does not vary between agglomerated cork and the reinforced foam. The stability of the densification strain is because the densification happens when many cells are folded. Thus it does not depend on the material rigidity. Nonetheless, densification stress will be higher for the reinforced cellular material because it is stiffer than the material without fibres. To reach such strain, much more stress will be needed.

No such enhancement in the compressive mechanical properties could be observed for graphene-enriched agglomerated cork material [36]. The shape, quantity and size of reinforcements used, while not especially optimised here seem well suited to improving the mechanical behaviour of multi-scale cellular materials like cork agglomerates.

3.3.3 Fibre orientation effect

Discrepancies in material parameters between in-plane and out-of-plane directions were hinted at in the previous section. Tab.3 shows how the anisotropy is modified by the presence of fibres for a large strain. It shows that the reinforced foam displays a modified anisotropy in the material parameters compared to classical high density cork agglomerates.

The enhancement due to the fibres is more present in the plateau stage for the out-of-plane direction (see E_p in Tab.2). The anisotropy difference of E shown in Tab.3 around 88 % between agglomerated cork and the reinforced foam shows that on the other hand mechanical behaviour increases strongly from the linear stage in the in-plane direction (Tab.2).

As reported before, high density agglomerated foam obtained by uni-axial compression displays an anisotropic mechanical behaviour [31] caused by an anisotropic microstructure at the bead scale [20]. X-ray tomography analysis showed that after the fabrication process, beads are flattened in the compression plane, i.e. in the in-plane

Material	E [MPa]	σ_{el} [MPa]	E_p [MPa]	ε_d [-]	$\mathcal{E}_d$ [J/mm ³]
Agglomerated cork OP	22.0 ± 1	0.83 ± 0.06	4.83 ± 0.34	0.38 ± 0.002	0.53 ± 0.06
Reinforced foam OP	27.7 ± 1.9	1.08 ± 0.04	8.74 ± 0.13	0.379 ± 0.001	0.92 ± 0.02
Fibre effect OP	+ 26 %	+ 30 %	+ 81 %	+ 0 %	+ 74 %
Agglomerated cork IP	37.1 ± 1.3	1.46 ± 0.07	4.48 ± 0.48	0.421 ± 0.007	0.94 ± 0.06
Reinforced foam IP	63.6 ± 4.3	2.04 ± 0.03	6.86 ± 0.19	0.407 ± 0.001	1.39 ± 0.02
Fibre effect IP	+ 71 %	+ 40 %	+ 53 %	- 3 %	+ 48 %

Table 2. Material parameters of agglomerated cork and reinforced foam tested in the out-of-plane direction (OP) and in the in-plane direction (IP).

Material	E [MPa]	σ_{el} [MPa]	E_p [MPa]	ε_d [-]	$\mathcal{E}_d$ [J/mm ³]
Anisotropy in agglomerated cork	+ 69 %	+ 76 %	- 7 %	+ 11 %	+ 77 %
Anisotropy in reinforced foam	+ 130 %	+ 89 %	- 22 %	+ 7 %	+ 51 %
Fibre effect	+ 88 %	+ 17 %	+ 214 %	- 36 %	- 34 %

Table 3. Difference in percentage between out-of-plane and in-plane directions for material compression parameters of agglomerated cork and reinforced foam.

direction [20]. As short fibres lie along these beads, according to SEM pictures, fibres will be more oriented in the in-plane direction. Interfaces are then stiffer in this direction. Short fibres constitute a stiff and anisotropic network that is added to the already anisotropic material. This explains why the anisotropy is higher in the composite material.

This preferential orientation also implies that depending on the solicitation direction, fibres will carry more load in a specific direction for the same strain values and thus the for same strain mechanisms. When the reinforced foam is compressed in the out-of-plane direction, fibres are mostly oriented perpendicularly to this direction. They are thus mainly solicited in bending but foam beads will tend to deform before them. This is why their effect is mostly observed after the elastic collapse. On the other hand when the sample is compressed in the in-plane direction, fibres are placed randomly. In order to deform the reinforced foam, fibres will then be solicited either in bending, tension or shear depending on their orientation. This explains why the Young's modulus and the elastic collapse stress are more enhanced in the in-plane direction.

3.3.4 Post-mortem observations at the microscopic scale for compression tests

Both cellular materials were observed after compression. In in-plane and out-of-plane compression directions, observations are very similar at the cell scale.

Shortly after the end of the loading many cells are still collapsed in the loading direction as can be seen in Fig. 6(a). Cells are collapsed mainly in the loading direction. However, according to the local geometry of the bead arrangement, cells can also be collapsed in other directions. So even if the macroscopic loading is uniaxial, because of the complex microstructure, mixed loading modes appear at the microscopic scale.

Observations of the compression surface in Fig. 6(b) show collapsed walls. Some cells were unfolded during

the unloading or, because of the high viscosity of the material [20], will be unfolded later. That is why the height of the sample keeps increasing slowly, even after the end of the test. However, some of the foam cells will stay collapsed, even after a long time.

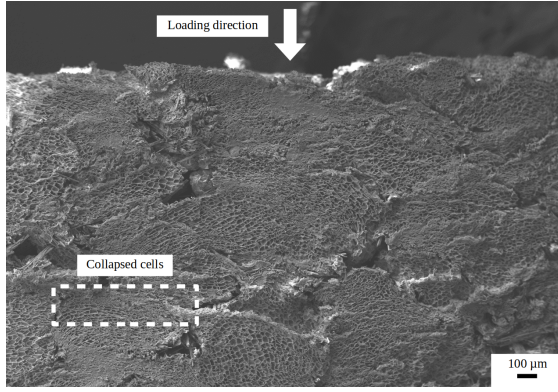
SEM observations are very similar for the two materials. For each type of material, collapsed cells are mostly at the bead periphery after compression, near the interfaces and thus increasing the interphase area observed in Fig. 3(a) and 3(b). Some beads have all their cells collapsed.

Strain mechanisms in compression are therefore not fundamentally different between agglomerated cork and reinforced foam and are close to that presented in a previous work [20]. They are simply a continuation of the damaging process started by the fabrication process and the interphase creation.

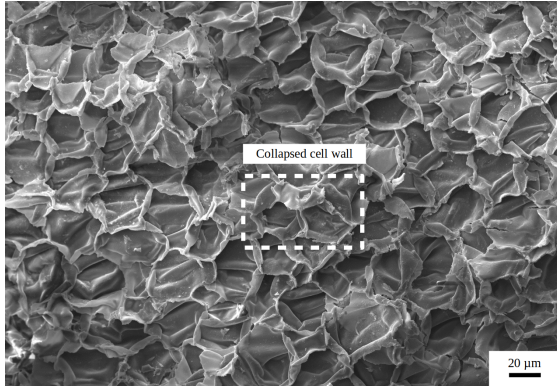
3.4 Fracture Behaviour

The fracture behaviour in mode I was studied in the out-of-plane direction for cork agglomerate and the reinforced foam as it was identified as a critical loading mode (worst-case scenario). In the base cellular material, cork, compression and tension imply very different strain, damage and fracture mechanisms. While compression is mostly about buckling [3], when tension is applied to cork, the cell walls become straighter and aligned in the direction of tension. Further deformation occurs by stretching of the walls [37], before fracture.

The two materials studied here are multi-scale materials. They are constituted by cellular foam beads agglomerated together. Competition between the interfaces and bead toughness is expected in the strain and fracture mechanisms. Besides the macroscopic behaviour of the material, optical and electronic observations of the crack path and of the crack surface are thus of interest.



(a) Profile observation of a reinforced agglomerated foam compressed in the out-of-plane direction.



(b) Surface observation of agglomerated foam compressed in the out-of-plane direction.

Figure 6. Surface and profile of compressed agglomerated foams observed with a SEM.

3.4.1 Macroscopic behaviour in fracture

The fracture behaviour of both multi-scale materials is shown in Fig.7(a). In this figure, the evolution of force as a function of the device displacement is plotted. Both curves display a first linear force/displacement behaviour and then a strain-softening is observed. The same behaviour was observed for cork [37]. The strain-softening could be related to the material viscosity and the material damaging.

As the agglomerated and reinforced cork are both made of randomly oriented cork beads, the three main directions and their combinations can be found. However, the serrations observed in the stress/strain curve for radial tension in raw cork [37] are not observed here in multi-scale foams. The localized fracture events causing these serrations might not then happen in agglomerate foams, contrary to the bulk cellular material.

As summarised in Tab.4, discrepancies can be seen between the reinforced foam and the non-reinforced material. This table shows several parameters measured with the help of the force/displacement curve for the agglomerated cork and the reinforced foam. The initial stiffness (measured as the initial slope) is much higher (+ 84 %) for the reinforced foam. Enhancement of the stiffness by the addition of short fibres is much greater than for cork

agglomerates reinforced by large coconut fibres, which was around + 27 % in tension with a specific coupling agent [38]. This increase in the stiffness is more likely, as for compression, directly linked to the high stiffness of short fibres (230 GPa) even given their small volume ratio. Greater strain softening or non-linear behaviour is also noticed for this sample before fracture. Adding fibres to the agglomerated cork does not enhance viscous effects as previously mentioned. Damage mechanisms are probably modified, as has already been observed in short fibre reinforced polymers [39].

Tab.4 shows a much higher force F_{rupt} (+ 70 %) and displacement d_{rupt} (+ 84 %) before the beginning of the crack propagation for the reinforced cork than for the non-reinforced agglomerate. This higher force is probably also linked to fibre stiffness. The higher displacement before fracture indicates that the reinforced foam is tougher before crack initiation. Damage mechanisms would thus be less critical in this material than in classical agglomerated cork. This may be due to the confinement of these damages in the vicinity of the short fibres, at the interface, as already observed for reinforced polyurethane foams [8]. And despite modifying (without optimising) the interfaces by the addition of short fibres, the chemical bond between cork beads, matrix and fibres seems relatively good as previously noticed for cork agglomerates with PFA resin [32].

Another remarkable measurement is Δd , the distance between d_{rupt} and d_{max} . It quantifies how much the sample still needs to be elongated before complete fracture. For the reinforced cellular material, this resistance to crack propagation Δd is increased by +230 %. Fig.7(d) shows how the crack total length varies as a function of time. Regarding agglomerated cork, crack propagation happens very fast. There are only a few points of measurement. After 3 seconds, the crack has crossed the entire specimen. On the other hand, it takes around 3.5 times longer for the crack to fully propagate in the reinforced foam. For small displacements, the crack seems even not to propagate at all. The addition of short fibres seems thus to also increase resistance against damage extension as it slows down the crack propagation.

At the macroscopic scale, the crack follows a straight path, as observed in Fig.7(b) and Fig.7(c). At the bead scale, however, fibres could act as a field disruptive element. Because of confinement near the fibres, a greater loading and displacement would be needed for the crack to propagate further. A similar influence was reported for soil reinforced by short polypropylene fibres [40]. Fibre addition in such granular media would seem to limit the further development of tension crack thanks to a bridge effect. It also changed the cemented soil's brittle behaviour to a more ductile one as it is observed for cork agglomerate.

3.4.2 Post-mortem observations at the microscopic scale for fissuration tests

After testing, samples were observed in order to identify underlying mechanisms. The area near crack initiation and the crack path on the sample (marked by the black

Material	d_{frac} [mm]	d_{max} [mm]	Δd [mm]	F_{frac} [N]	K [N/mm]
Agglomerated cork	1	1.3	0.3	475	832
Reinforced foam	1.7	2.7	1	875	1531
Fibre effect	+70 %	+108 %	+230 %	+84 %	+84 %

Table 4. Material parameters obtained from mode I fracture tests in the out-of-plane direction for agglomerated cork and for reinforced foam.

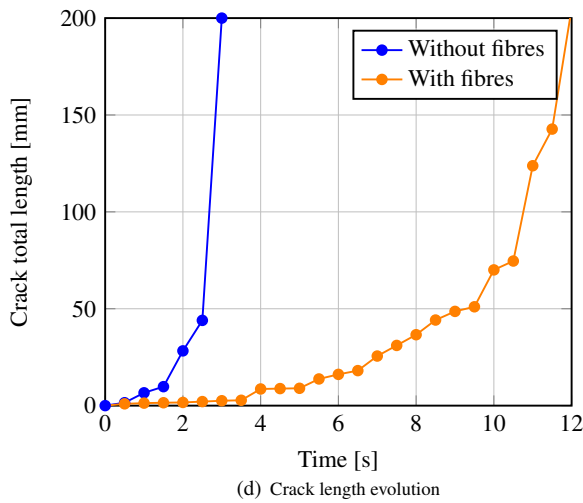
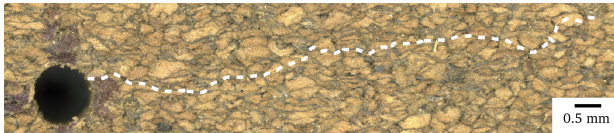
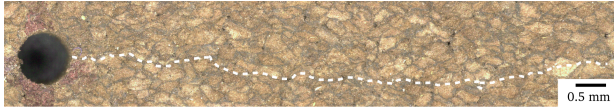
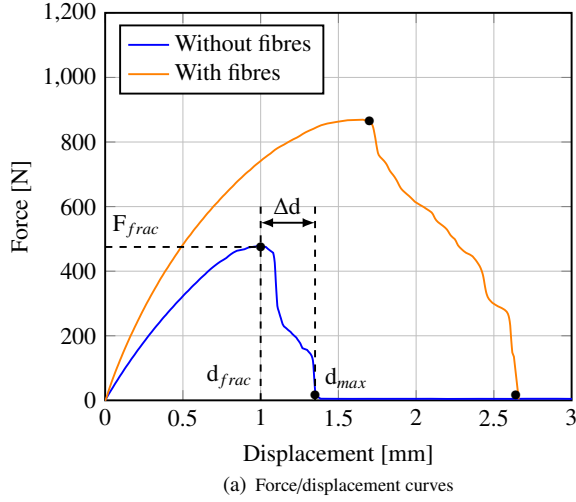


Figure 7. Experimental displacement/force curves obtained from mode I fracture tests. Effect of adding short fibres on the mechanical behaviour of an agglomerated cellular material..

dotted line on Fig. 2) were studied first. By comparing oictures of fractublack samples with the picture taken near the same area before testing, the fracture path was drawn with a white dotted line on the non-tested sample surface (Fig. 7(b) & 7(c)). On both pictures, almost exclusively intraparticle fracture can be observed. This means that between foam and matrix, the cellular material is the one that breaks before the matrix under tension. This confirms the good interfacial adhesion between cork and the PFA matrix [32], even after the addition of fibres.

Nevertheless two types of intraparticle fracture have to be distinguished as schematised in Fig. 8. The first is the intraparticle fracture going straight across the bead, as shown in Fig. 8(a). This will be called the *internal intraparticle fracture*. The other one visible in Fig. 8(b) is an intraparticle fracture occurring near the interface between two foam beads. This will be called a *peripheral intraparticle fracture*. More internal intraparticle fractures could be observed for reinforced agglomerated foam.

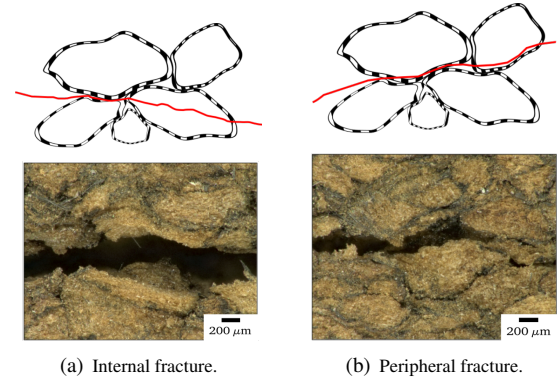


Figure 8. Scheme of the two types of intraparticle fracture observed for multiscale foams after mode I fracture tests.

Fractublack surfaces and profiles are observed with a SEM in order to better understand the numerous fracture mechanisms in both materials. Confirming the optical microscope observation, no fracture is observed in the bulk resin. This means there is no fracture at the interface between two beads. Nor was any bead/matrix decohesion observed for either material.

By observing the fracture surface, the radial direction (hexagonal cells) seems quite present like in Fig. 9(b) and Fig. 9(d). It could come from the different fracture toughnesses of the cork directions. $K_{IC} = 94 \pm 16 \text{ kPa m}^{1/2}$ for non-radial direction and $K_{IC} = 125 \pm 14 \text{ kPa m}^{1/2}$ for the radial direction [37]. To these two toughness values, two fracture mechanisms can be linked. The first for radial tension occurs by transverse breaking of the lateral cell

walls. The other, for non-radial tension, occurs along the walls, which are longitudinally split into two half-walls. However, it is hard to tell whether both these mechanisms happen in agglomerated foam or not. The material microstructure is indeed very intricate. Apart from radial orientation, a quite random orientation of cork cells seems still to be observed. So fracture happens in cork beads, not only in one privileged direction. The anisotropic nature of cork cells could thus matter here in fracture mechanisms of a randomly oriented foam with anisotropic cells but it is not the only parameter to take into account (bead size, orientation, interfaces, cell wall thickness, etc.).

As observed with light microscopy, the two types of intraparticle fracture can be found for both types of foam. Internal intraparticle fracture is noticeable by uncollapsed cell walls. The profile of such fracture across the bead is shown in Fig.9(a). Cells near the crack are fairly undamaged. Microscopic cracks were spotted in the vicinity of the main crack. Fig. 9(b) shows the fracture surface. Cell walls are straight even if they can sometimes be quite damaged.

By looking at the SEM picture of the profile of peripheral intraparticle fracture in Fig.9(c), this type of fracture seems to happen in the already reported pre-collapsed area caused by the fabrication process [20] identified here as the interphase. Indeed, the fracture surface in Fig.9(d) displays cells with plastically deformed cell walls. They are very similar to the collapsed cell walls observed after compression in Fig. 6(b). The presence of resin just behind the damaged cells could also be seen. This indicates that the damaged cells in this picture are just behind the interface. This pre-collapsed zone caused by the fabrication process could imply some pre-damaging of cork cells and thus an important gradient in the mechanical properties leading to the peripheral intraparticle fracture mostly observed for both materials happening in the interphase.

When comparing the two materials, the same observation can be made as for the light microscope observations. The reinforced foam displays a bit more internal intraparticle fracture as some cork cell walls are not collapsed when the surface of the fracture was observed. It is not clear, however, whether this is happening at specific places (like crack cessation) or not because of the high heterogeneity of the material. This kind of fracture could be due to crack confinement caused by fibres at the interface when the crack passes to another bead. Depending on the local mechanical state, it could lead to the crack going through the bead rather than through damaged cells. Undamaged fractured cork cells could be less resilient than pre-damaged cells. This could also explain why Δd is much greater for the reinforced foam.

Resin/fibre debonding is only occasionally noticed for the reinforced foam. Fig.9(e) is a SEM picture of the fracture surface of the reinforced foam. It shows one cork bead with collapsed cells covered with resin. No fibres can be observed at its surface. This indicates that the fracture happened between the matrix and the fibres that can be seen on the other side of the crack. The adhesion between resin and fibres could thus be optimised to improve the reinforced material mechanical behaviour in tension [41].

4 Conclusions

In this article, the reinforcement of cellular materials by rigid short fibres was studied. The composite material consisted of beads of polymeric cellular material (here cork) mixed together with a PFA thermosetting resin and short fibres then uni-axially compressed together before polymerisation in an oven.

- Considering the macroscopic scale, the addition of short fibres improves the compressive and the mode I fracture behaviour much more than long fibres or graphene nanoplates while keeping a foam-like behaviour and a low density.

The glass transition of reinforced agglomerates is shifted to higher temperatures because short fibres seem to constrain polymer chains. Damping properties depend strongly on fibres being oriented with the direction of solicitation.

In compression under large strain, apart from the densification strain, all other material parameters were largely enhanced by the addition of short fibres. A more greater difference between in-plane and out-of-plane direction mechanical behaviour was noticed. This supplementary anisotropy was correlated with the preferential orientation of the short fibres observed in the in-plane direction.

The fracture behaviour of cork agglomerates is improved by the addition of short fibres. Forces reached by the reinforced material before fracture were more than 80 % higher than for the non-reinforced one. Maximal displacements and initial stiffness were also increased by at least 70 % by the addition of fibres. Finally, displacement before complete fracture Δd is much greater for the reinforced foam (by 125 %). This shows that the addition of fibres increases the resistance to damage extension.

- Microscopic observations indicated that, as for mechanisms in compression, short fibres do not seem to change them fundamentally. However, because of the intricate microstructure of both materials, a simple loading like uni-axial compression seems to lead to mixed strain modes at the bead and cell scales.

Conversely, the addition of short fibres strongly influenced the fracture mechanisms. The fracture path was mostly intraparticle for both cellular materials. The non-reinforced material showed fractures mainly inside the beads but in the interphase, in the vicinity of its interfaces where cells are pre-collapsed because of the compression process. As for the reinforced foam, a little more fracture across the beads was also observed. Fibres at the interface might then orientate the crack path by confining the crack.

In order to tailor and enhance cellular material mechanical properties, multi-scale foams and short rigid fibres seem an interesting alternative. Further improvements can be achieved by using fibres with optimised length, amount [9] or surface treatments such as coupling agent specifically tailored to the particular resin [41]. Finally studying fracture mechanisms of multi-scale cellular

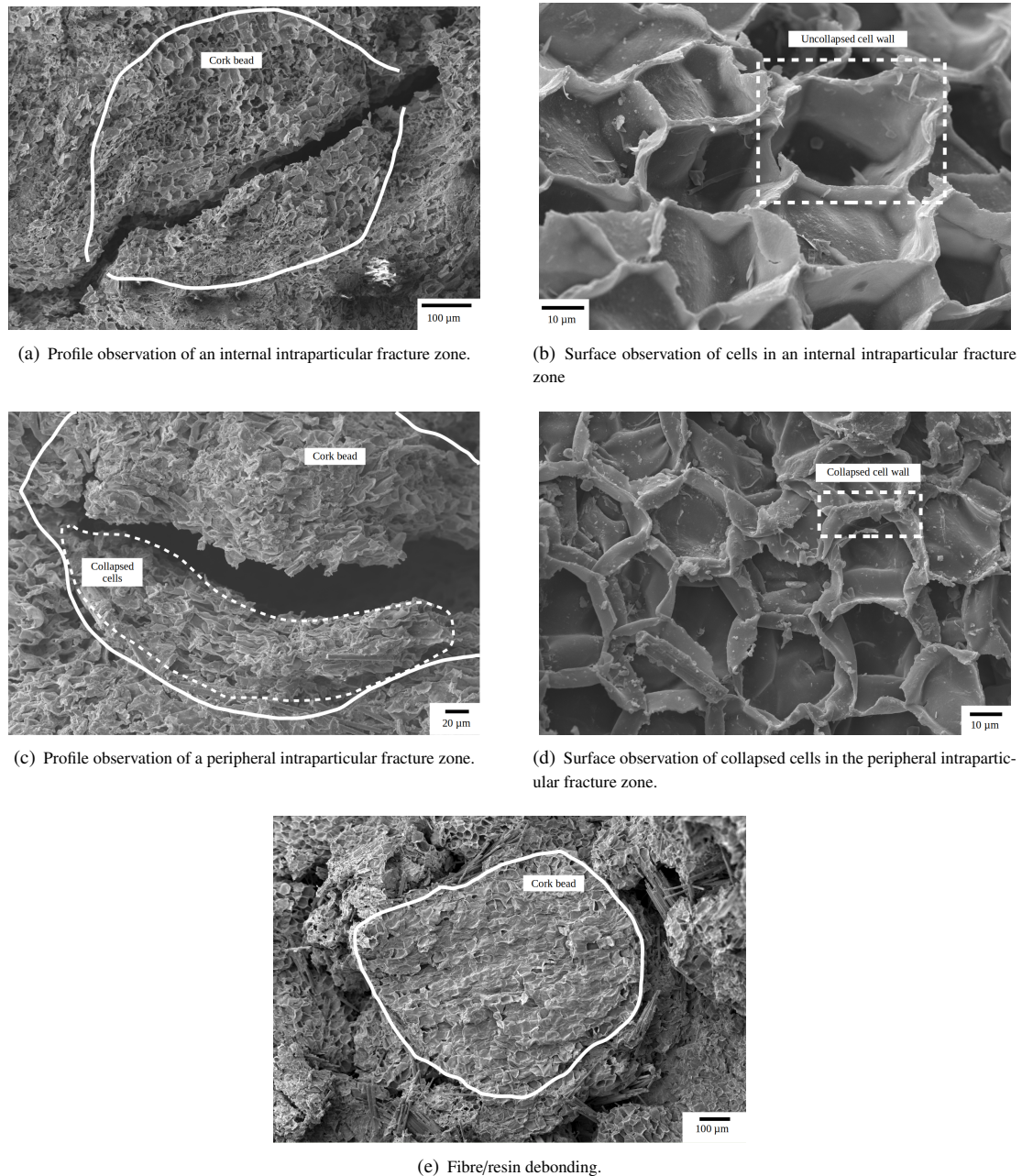


Figure 9. Fracture surface and profile observed with a SEM of multiscale foams after mode I fracture tests.

materials seems consistent for better capturing changes in material interfaces, such as the addition of fibres in this case.

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