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Wood versus Plant Fibers: Similarities and Differences in Composite Applications

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The work on cellulose fiber composites is typically strictly divided into two separated research fields depending on the fiber origin, that is, from wood and from annual plants, representing the two different industries of forest and agriculture, respectively. The present paper evaluates in parallel wood fibers and plant fibers to highlight their similarities and differences regarding their use as reinforcement in composites and to enable mutual transfer of knowledge and technology between the two research fields. The paper gives an introduction to the morphology, chemistry, and ultrastructure of the fibers, the modeling of the mechanical properties of the fibers, the fiber preforms available for manufacturing of composites, the typical mechanical properties of the composites, the modeling of the mechanical properties with focus on composites having a random fiber orientation and a non-negligible porosity content, and finally, the moisture sensitivity of the composites. The performance of wood and plant fiber composites is compared to the synthetic glass and carbon fibers conventionally used for composites, and advantages and disadvantages of the different fibers are discussed.

1. Introduction

Composites based on cellulose fibers from wood and plants constitute a relatively new and promising class of composite materials [1–4]. They are environmentally friendly, and they offer good technical performance. For several load-carrying applications, where glass or carbon fiber composites are conventionally used, cellulose fiber composites can be a worthwhile alternative. This is particularly the case for applications where the green advantages (renewability, biodegradability) play an important role, and top-end mechanical properties are not the primary motivation. A vast amount of scientific literature on cellulose fibers for composite applications has been compiled during the last decade (e.g., see recent reviews [5–8]), although the publications tend to be divided into two separate fields depending on the origin of the fibers, that is, from wood or annual plants. The reason for this division is perhaps that the raw materials producers are looking for new markets for their fibers (technology pull), and that the end-users (market pull) have yet to exploit the potential of cellulose fibers, independent of the origin of the fibers. The raw materials producers in this case, that is, forestry for wood fibers and agriculture for plant fibers, have developed their specific technologies along the value chain to produce fibers depending on the traditional usage of the fibers. For wood fibers, pulp mills have been built to produce raw materials for making paper and board. For plant fibers, textile technologies are refined to produce yarns and fabrics. In view of the maturing research field of cellulose fibers shifting towards achieving the technical performance demands of the end-users, and the rather independent existence of research communities of wood and plant fibers, respectively, this paper has been written to shed some further light on the similarities and differences of these two types of cellulose fibers (wood and plant origin), with regard to industrial usage to produce cellulose fiber composites for structural applications. The
performance of the fibers will also be compared to the main current competitors, that is, composites made from synthetic fibers, such as glass and carbon fibers. Similarities and differences of the different fibers for composites will be discussed, in light of future potentials in engineering applications. By bringing the disciplines of wood and plant fiber science closer together, one could hope for a mutual transfer of knowledge, as the two research fields have evolved rather independently and have thus reached different levels of understanding with regard to various aspects, such as characterization methods, fiber treatment, fiber preform processing, and composite manufacturing.

The paper presents an introduction to (i) the morphology, chemistry and ultrastructure of the fibers, (ii) the modeling of the mechanical properties of the fibers, (iii) the fiber preforms available for manufacturing of composites, (iv) the typical mechanical properties of the composites, (v) the modeling of the mechanical properties of the composites with special focus on composites having a random fiber orientation and a non-negligible porosity content, and (vi) the moisture sensitivity of the composites. Furthermore, examples of new composite applications are given, followed by considerations of the future perspective of using wood and plant fibers to produce cellulose nanofiber composites. Finally, an outline of the differences and resulting advantages of the two types of cellulose fibers, namely, wood and plant fibers, is given, as well as the differences and resulting advantages of cellulose fibers versus synthetic fibers.

2. Morphology, Chemistry, and Ultrastructure of Fibers

Plantae is the one of the five kingdoms of living organisms that includes green plants, that is, mosses, ferns, gymnosperms (e.g., softwood), and angiosperms (e.g., hardwood and annual plants). The cells of green plants are surrounded by a rigid cell wall, and this is the main characteristic distinguishing them from cells in animals. In some types of cells, the cell walls are enlarged to have superior mechanical properties, which provide the required structural performance of the plants. The dimensions of these so-called fibers vary between different plants but their overall shape is most often elongated with lengths in the range 1–50 mm, and diameters in the range 15–30 μm. In the perspective of composite reinforcement, it is convenient to group the fibers by their lengths.

(i) Short fibers (1–5 mm), originating typically from wood species (e.g., spruce, pine, birch, eucalyptus), and typically used for making composites with in-plane isotropic properties, that is, composites with a non-specific (random) fiber orientation.

(ii) Long fibers (5–50 mm), originating typically from annual plant species (e.g., flax, hemp, jute), and typically used for making composites with anisotropic properties, that is, composites with a specific fiber orientation.

In the living green plants, when the fibers are fully developed, their intracellular organelles start to degenerate resulting in fibers having an empty central cavity, the so-called lumen. In wood fibers, the luminal area is in the range 20–70% of the fiber cross-sectional area [9]. In contrast, annual plant fibers, such as hemp and flax, have a relatively smaller luminal area in the range 0–5% [3, 10].

The main chemical constituent of the cell wall is cellulose, which is a non-branched polysaccharide polymer made up of glucose units. For wood fibers, the cellulose chain is having an average length of 5 μm corresponding to a degree of polymerization (i.e., glucose units) of 10,000 [9]. This molecular linearity makes cellulose highly anisotropic with a theoretical stiffness and strength of about 130 and 15 GPa, respectively, in the chain direction [11]. The cellulose chains are arranged in parallel to form bundles, which are denoted microfibrils. In some regions of the microfibrils, the glucose molecules of the cellulose chains are arranged in a highly ordered crystalline structure. The two other principal chemical constituents of the cell wall are hemicellulose and lignin. Hemicellulose is a heterogeneous group of polysaccharides characterized by being short and branched. Lignin is a highly branched polymer composed of phenylpropane units organized in a complex three-dimensional structure. In addition to the organization of the three chemical constituents, the structural complexity of the cell wall is increased by being organized into a number of layers differing by the angle of the cellulose microfibrils to the longitudinal fiber axis. The angle of the cellulose microfibrils in the various layers, in addition to the relative layer thicknesses, dictates the overall mechanical performance of the fibers. Thus, altogether, the cell wall of wood and plant fibers is essentially organized like a composite laminate with a number of laminae with differently oriented, stiff and strong semicrystalline cellulose microfibrils embedded in a matrix of hemicellulose and lignin.

In contrast to cellulose fibers, the synthetic fibers that traditionally are used for reinforcement in composites, such as glass and carbon fibers, are monolithic and with a much more simple ultrastructure. Glass fibers are primarily composed of silicon oxide molecules organized in an amorphous configuration. Carbon fibers are composed of carbon atoms in graphite layers that are organized in a stackwise turbostratic configuration.

Table 1 shows key numbers of chemical composition and ultrastructure of cellulose fibers. The cellulose content of unprocessed fibers is in the range of 40–50% w/w for wood fibers, and in the range of 60–70% w/w for plant fibers. Accordingly, the content of hemicellulose and lignin is higher in wood fibers, and this is particularly true for lignin which shows a content of about 30% w/w in wood fibers, in comparison to only about 5% w/w in plant fibers. The chemical composition of wood and plant fibers is clearly different from each other. In addition, wood fibers show lower cellulose crystallinity than plant fibers, with typical values in the ranges of 55–70 and 90–95% w/w, respectively. The microfibril angle in wood fibers vary in the range 3–50° depending on the type and location of the fibers in the wood (e.g., late and early wood) [12], whereas the microfibril angle in plant fibers is more constant in the range 6–10° [13].
Table 1: Chemical composition and ultrastructure of wood and plant fibers.

<table>
<thead>
<tr>
<th>Wood fibers</th>
<th>Chemical composition</th>
<th>Ultrastructure</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Cellulose (% w/w)</td>
<td>Hemicellulose (% w/w)</td>
<td>Lignin (% w/w)</td>
</tr>
<tr>
<td>Spruce</td>
<td>49</td>
<td>20</td>
<td>29</td>
</tr>
<tr>
<td>Pine</td>
<td>42</td>
<td>29</td>
<td>28</td>
</tr>
<tr>
<td>Pine (kraft pulp)</td>
<td>76</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cedar</td>
<td>44</td>
<td>21</td>
<td>30</td>
</tr>
<tr>
<td>Balsa</td>
<td>48</td>
<td>28</td>
<td>22</td>
</tr>
<tr>
<td>Birch</td>
<td>41</td>
<td>32</td>
<td>22</td>
</tr>
<tr>
<td>Poplar</td>
<td>39</td>
<td>28</td>
<td>30</td>
</tr>
<tr>
<td>Soft wood</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

| Plant fibers         | Hemp (textile)       | 91                      | 7          | 2                      | 6–10 | [13] |

The effect of processing treatments on the chemical composition of the fibers is shown in Table 1. In general, for both wood and plant fibers, the cellulose content is increased after processing treatments, due to removal of non-cellulose residues of the fibers (e.g., pectins and waxes). The effect of processing treatment is most clearly seen for the highly processed textile hemp fibers in the study by Madsen et al. [14] where the cellulose content was measured to be as high as 91% w/w.

The influence of growth conditions and processing treatments on the chemistry and ultrastructure of cellulose fibers lead typically to fibers with more variable properties than seen for synthetic fibers. This is frequently considered to be one of the major disadvantages of using cellulose fibers for reinforcement in composites. It is however believed that this concern is caused by a general uncertainty about the cause for the variability in properties, and the lack of a system for classification of the quality of cellulose fibers, for example, similar to the system that exist for classification of solid wood. It should also be mentioned that variability in mechanical properties of fibers can have a positive effect on the notch sensitivity and the fracture toughness of composites [15, 16].

3. Modeling of Mechanical Properties of Fibers

Micromechanical models can be useful in understanding how the chemical composition and ultrastructure of cellulose fibers affect their mechanical properties. From a geometrical point of view, the cell wall in cellulose fibers can be approximated by layers of concentric cylindrical shells.

Figure 1 shows such an idealized fiber geometry. Summaries of how the ultrastructural features of the cell wall affect the mechanical properties of the fibers have been compiled by Neagu et al. [17], Salmén and Burgert [18], and Salmén [19].

In principle, the most important mechanical properties of fibers when used in composites are the stiffness and strength in the axial direction, that is, in the fiber length direction. It is in this direction that the fibers are supposed to carry load when used in composites. In the case of cellulose fibers, the key ultrastructural features that affect the axial mechanical properties of the fibers are as follows.

(i) Lumen Size. Only the cell wall carries load, that is, the fiber mechanical properties are proportional to the cell wall cross-sectional area. The larger the relative
lumen size, the lower the stiffness and strength of the fibers.

(ii) **Cellulose Content.** In a study by Thygesen et al. [20], an increase in the cellulose content of the fibers was found to be well correlated with an increase of their stiffness and strength. In addition, the cellulose crystallinity and the crystallite aspect ratio are known to affect the stiffness of the cell wall in the microfibril direction (e.g., [21]).

(iii) **Microfibril Angle.** It can be demonstrated with classic laminate theory (in-plane rotation of an orthotropic plate) that the effective elastic properties of the fibers in the axial direction scales with the local stiffness in the microfibril direction multiplied with $\cos^4 \theta$, where $\theta$ is the microfibril angle. The fiber stiffness (and strength) is thus very sensitive to the microfibril angle, even if the mechanical properties in the microfibril direction are constant. This trend is also captured by more accurate and detailed micromechanical models (e.g., in the study by Hofstetter et al. [22]). The low microfibril angle of plant fibers makes them highly anisotropic (which also is the case for the synthetic carbon fibers, but not for glass fibers), and this leads to relatively low transverse mechanical properties.

Of the three above-mentioned ultrastructural features, the most important factor to be addressed by the modeling of the mechanical properties of the fibers is probably the microfibril angle, since fiber lumens can either be collapsed (as for earlywood in chemically pulped fibers), or filled with low-viscosity resin during manufacturing of composites, and the cellulose content is an intrinsic property, which is roughly constant for plant fibers and constant, albeit lower, for wood fibers (cf. Table 1). The dependence of the fiber stiffness on the microfibril angle is a well-known effect (e.g., [23]), and can be described by classic laminate theory [24].

### 4. Preforms of Fibers

The types of preforms of cellulose fibers, to be used for manufacturing of composites, are in principle identical to the ones for synthetic fibers, although concerns must be addressed to some particular characteristics. Here follows details of the preforms of wood and plant fibers.

#### 4.1. Wood Fiber Preforms.

Wood fibers are available at a low cost as pulp fibers (Figure 2(a)). These are used to make paper sheets or board materials for packaging. One way to make composites based on wood fibers is to use such fiber mats (Figure 2(b)), which can be impregnated by using for example, a resin transfer molding technique (e.g., [46]). A viscous thermoset resin is impregnating the enclosed wood fiber mat by the aid of a pressure vessel attached to the mold inlet and sometimes also assisted by vacuum suction at the outlet. This manufacturing technique is only adequate for low-viscosity resins, typically thermosets. Thermoplastics usually have a high viscosity in the molten state, and resin transfer molding is not suitable since the impregnation times would be too high, or the required high pressure would induce severe deformation of the fiber mat. Instead, a commingling technology can be advocated. By using papermaking machines, for example, so-called French or Finnish sheet formers in the laboratory scale, one can produce mats composed of commingled wood pulp fibers and thermoplastic fibers (e.g., [46]). The thermoplastic fibers should preferably have similar dimensions as the pulp fibers, in order to have approximately the same hydrodynamic properties during the formation process, which facilitates efficient mixing. Thermoplastic fibers can be spun to have diameters around 30 $\mu$m and be chopped to roughly 3 mm lengths (similar to the dimensions of the pulp fibers). When the commingled fiber mat has been dried, it can be placed in a hot press and composite components can be molded. This method is not only limited to flat plates for materials testing, but complex parts with double curvatures can also be made [47].

The papermaking industry encompasses a huge infrastructure to produce wood fiber mats. Anticipated volumes of such fiber preforms for composite applications are extremely small compared with produced volumes of conventional paper and board. Nevertheless, there is an opportunity to build upon the experiences and use small-scale paper mills to produce composite preforms. In the laboratory scale, two main techniques are used to mimic the paper manufacturing process. The most common are sheets produced by dynamic sheet forming and regular handsheets. In dynamic sheet forming, a fiber suspension jet is directed towards a rotating wire drum [48]. The fibers will deposit onto the wire whereas
4.2. Plant Fiber Preforms. The types of plant fiber preforms available for composites are shown in Figure 3. Here follows descriptions of their processing and characteristics.

After the fibers have been extracted from the plants by a retting process, followed by a series of mechanical processes, the fibers can be converted into non-woven mats by air-laid and needle-punching techniques [49]. The fiber orientation in non-woven mats is nominally in-plane random, but they can show a preferred fiber orientation in the machine direction [50]. Alternatively, the fibers can be converted into a continuous yarn by using various spinning techniques, such as ring spinning, rotor spinning, wrap spinning, and air-jet spinning [51]. Ring spinning is the most widely used method. During spinning, the continuous bundle of almost parallelized fibers (a so-called sliver) is twisted so that the fibers take up a helical configuration. The effect of the fiber twisting angle on the mechanical properties of composites has been addressed in a few studies [52–54]. Furthermore, the cross-sectional area of the yarn (which is specified indirectly by its linear density given in units of g/1000 m), and the degree of yarn compaction are other important yarn characteristics, which however have received limited attention in the perspective of composite reinforcement [14]. It can be speculated that the degree of yarn compaction is correlated with the permeability of the yarn for matrix impregnation during manufacturing of composites. More studies are needed to improve the understanding of the correlation between the various structural characteristics of plant fiber yarns, and the mechanical performance of the yarns in composites.

Plant fiber yarn preforms can be used directly to produce composites by commingled filament-winding together with a thermoplastic filament yarn, followed by compression molding [41], or the yarn can be used to make preforms of woven fabrics and non-crimp fabrics. Woven fabrics are fabricated with a range of weaving patterns, such as plain, twill and satin weave, in which the yarns are differently interlaced in the two main, orthogonal, planar directions. The yarns in the two directions can have different linear densities, and they can be placed with different distances to each other. The woven fabrics offer the possibility of having a planar yarn configuration in two dimensions designed to meet the loading profile of a given composite application. Woven fabrics of flax, jute and cotton fibers are widely available, but they are most often tailored for textile applications, and not for composite applications. Non-crimp fabrics consist of yarns that are not held together by being woven into each other, but instead they are stitched together by thin and flexible threads (typically thermoplastic polyester). This means that the yarns are fully stretched; that is, they have no crimp, since they do not have to go over and under each other. Single layers of parallel yarns held together by transversely directed stitching threads are denoted uniaxial non-crimp fabrics. Such uniaxial layers are stacked and stitched together to form biaxial or multi-axial non-crimp fabrics with specific planar yarn orientations, for example, ±45°, 0°/90°, and 0°/+45°−45°/90°. Recently, a number of European companies have started production of non-crimp fabrics of flax fibers. Thus, for the first time, fabrics of plant fibers that are specifically tailored for composites are commercially available.

5. Mechanical Properties of Composites

The mechanical properties of wood and plant fiber composites have been extensively characterized and analyzed. However, mostly tensile properties, as well as bending and to some extent also impact properties have been characterized, since they are relatively straightforward to measure, and they are commonly used to benchmark different materials in the process of materials development. Other more complex mechanical properties, such as fatigue [55–58] and creep [59] have been studied to a lesser extent.

Table 2 presents typically reported tensile properties (stiffness and strength) of wood and plant fiber composites, together with values for glass and carbon fiber composites. The remarkably high stiffness and strength on 26 GPa and 247 MPa, respectively, for Kraft paper impregnated phenol formaldehyde composites [34] have hitherto not been reached for wood fiber composites. These materials were developed during World War II for use in skins of aircraft wings. Apart from these extreme results by Cox and Pepper [34], it can be observed that cellulose fiber composites (both wood and plant fibers) with a nominal in-plane random fiber orientation, made by using the preforms of loose fibers, paper, and non-woven mats, possess moderate tensile properties with stiffnesses in the range 4–8 GPa and strengths in the range 30–60 MPa. With respect to glass fiber composites, with a similar in-plane random fiber orientation, showing stiffnesses in the range 5–7 GPa and strengths in the range 80–100 MPa, cellulose fiber composites show in general comparable stiffnesses, and slightly lower strengths. It is well known that various chemical approaches can be used to control the interface bonding in order to improve the strength of cellulose fiber composites. Acetylation is one type of surface treatment that can be used to reduce the polarity of the fibers making them more compatible with the (typically) non-polar matrix [60]. Also, coupling agents, such as maleic anhydride, can be used to form covalent bonds between the fibers and the matrix [61]. In the study by Clemons [30] (Table 2),
the strength of wood fiber/PP composites were shown to increase from 28 to 52 MPa by using maleic anhydride as a coupling agent, whereas the stiffness did not show any changes. A similar large increase in strength from 40 to 60 MPa has been found in the study of jute fiber/PP composites by Andersen and Plackett [37].

When preforms with nominally unidirectional fibers, such as yarns and non-crimp fabrics are used, the tensile properties of the composites are markedly increased with stiffnesses in the range 20–32 GPa and strengths in the range 130–340 MPa (Table 2). In addition, in comparison to the composites with an in-plane random fiber orientation, the fiber volume content of the unidirectional composites is in general higher (up to 50–55%) [62]. This is related to the better fiber packing ability of aligned fibers as compared to randomly oriented fibers. However, in comparison to synthetic fibers, assemblies of cellulose fibers generally have a lower packing ability [63], which means that the maximum fiber volume content is typically lower in cellulose fiber composites. This is part of the explanation for the lower stiffness of unidirectional cellulose fiber composites, as compared to glass fiber composites, with values of about 30 and 45 GPa, respectively. The contributing stiffness of cellulose fibers in composites has been estimated to be in the range 20–90 GPa [41], which for the best quality cellulose fibers is comparable to glass fibers with stiffnesses in the range 70–87 GPa [45]. In terms of strength, unidirectional cellulose fiber composites show radically lower values of about 300 MPa compared to about 1000 MPa for glass fiber composites. The explanation for the low strength of unidirectional cellulose fiber composites is currently not known, however, it is expected that fiber defects, which are introduced to the fibers during their processing, play a large role [64].

The tensile properties in Table 2 well illustrate the current status of cellulose fiber composites where stiffness is acceptable, and comparable to glass fiber composites, but strength needs to be improved. Due to the low density of cellulose fibers, the specific mechanical properties of cellulose fiber composites is particularly competitive compared with glass fiber composites. Furthermore, if these specific properties are normalized with respect to cost, cellulose fiber composites compare well also with carbon fiber composites. In other words, for large volume applications where weight is an issue, for example, in packaging and transport, cellulose fiber composites are likely to be the main contending materials.

6. Modeling of Mechanical Properties of Composites

For composite materials, the quantitative relation between microstructure and mechanical properties is generally
Table 2: Tensile properties of wood and plant fiber composites. The type of fiber preforms used for the composites is given, in addition to their nominal fiber orientation; in-plane random (RD) and unidirectional (UD). For means of comparison, tensile properties of glass and carbon fiber composites are shown.

<table>
<thead>
<tr>
<th>Fiber content (% v/v)</th>
<th>Stiffness (GPa)</th>
<th>Strength (MPa)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Wood fiber composites</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wood pulp/PP¹; RD</td>
<td>27</td>
<td>4.2</td>
<td>28</td>
</tr>
<tr>
<td>Eucalyptus sawdust/UP¹; RD</td>
<td>46</td>
<td>6.2</td>
<td>60</td>
</tr>
<tr>
<td>Kraft + TMP/PP; RD</td>
<td>40</td>
<td>4.5</td>
<td>43</td>
</tr>
<tr>
<td>Sulphite pulp/PP¹; RD</td>
<td>50</td>
<td>3.9</td>
<td>51</td>
</tr>
<tr>
<td>Kraft/PF—paper; RD</td>
<td>72</td>
<td>²6.2</td>
<td>247</td>
</tr>
<tr>
<td>Kraft/PF—paper; RD</td>
<td>72</td>
<td>11.7</td>
<td>156</td>
</tr>
<tr>
<td><strong>Plant fiber composites</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Flax/starch—loose fibers¹; RD</td>
<td>37</td>
<td>8.3</td>
<td>51</td>
</tr>
<tr>
<td>Jute/PP—non-woven mat; RD</td>
<td>32</td>
<td>8.4</td>
<td>39</td>
</tr>
<tr>
<td>Jute/PP—non-woven mat; RD</td>
<td>30</td>
<td>5.2</td>
<td>40</td>
</tr>
<tr>
<td>Flax/PLA—non-crimp fabric; UD</td>
<td>39</td>
<td>19.5</td>
<td>150</td>
</tr>
<tr>
<td>Flax/epoxy—non-crimp fabric; UD</td>
<td>35</td>
<td>19.8</td>
<td>234</td>
</tr>
<tr>
<td>Flax/epoxy—yarn²; UD</td>
<td>40</td>
<td>28.0</td>
<td>133</td>
</tr>
<tr>
<td>Flax/PET—yarn²; UD</td>
<td>48</td>
<td>32.0</td>
<td>344</td>
</tr>
<tr>
<td><strong>Glass fiber composites</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Glass/PP—loose fibers¹; RD</td>
<td>30</td>
<td>7.3</td>
<td>100</td>
</tr>
<tr>
<td>Glass/PP—chopped strand mat; RD</td>
<td>20</td>
<td>5.4</td>
<td>77</td>
</tr>
<tr>
<td>Glass/epoxy—roving; UD</td>
<td>55</td>
<td>39.0</td>
<td>1080</td>
</tr>
<tr>
<td>Glass/PP—roving²; UD</td>
<td>60</td>
<td>45.0</td>
<td>1020</td>
</tr>
<tr>
<td><strong>Carbon fiber composites</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Carbon/epoxy—roving; UD</td>
<td>60</td>
<td>313.0</td>
<td>1140</td>
</tr>
<tr>
<td>Carbon²/epoxy—roving; UD</td>
<td>60</td>
<td>142.0</td>
<td>2140</td>
</tr>
</tbody>
</table>

¹ Injection molding; ² filament-winding.
² Machine direction; ³ cross direction; ⁴ high modulus fibers; ⁵ high strength fibers.

6.1. Composites with Random Fiber Orientation. Composites with an in-plane random fiber orientation distribution, which is usually the case for wood fiber composites, can be regarded as a stack of unidirectional plies, where the relative thickness of each ply is determined from the fiber orientation distribution. This is known as a laminate analogy, where classic laminate mechanics can be used to relate the elastic properties of the hypothetical unidirectional ply to those of the composite plate. The laminate analogy is schematically illustrated in Figure 4. It is assumed that the fiber orientation distribution is symmetric, which is typically the case for wood fiber mats produced with conventional wet-forming techniques [65]. This means that the materials are globally orthotropic, and the global stiffness matrix can be described by five elastic constants, namely the longitudinal and transverse Young’s moduli, the major and minor Poisson ratios, and the shear modulus. The components in the global stiffness matrix can be determined from standardized macroscopic testing, and the fiber orientation distribution can be found by image analysis of scanned sections of the fiber mat [66].

Softwood pulp fibers have an aspect ratio of about 100 [67]. From a mechanical point of view, these fibers can be regarded as continuous, that is, of infinite length, since the ineffective lengths close to the fiber ends are relatively small, as can be calculated by shear-lag theories [68, 69]. The stiffness contribution of wood fibers to the unidirectional plies in the laminate analogy can then be described by simple mechanical models, such as the rule of mixtures model for the longitudinal elastic properties, and the Halpin-Tsai model for the transverse and shear elastic properties (e.g., [70]). For the off-axis properties, Hashin’s concentric cylinder model is more accurate [71]. The latter model has been used by Neagu et al. [72] to back-calculate the contributing stiffness of wood fibers from the measured stiffness of composites, and thereby...
Figure 4: Schematic illustration of a laminate analogy for randomly oriented cellulose fiber composites.

Figure 5: The effect of bleaching level, as characterized by the kappa number, on the contributing fiber stiffness in softwood fiber composites. Shown are results from laboratory softwood kraft fibers and prehydrolyzed kraft fibers [72].

6.2. Composites with Non-Negligible Porosity Content. In cellulose fiber composites, the porosity typically makes a noteworthy contribution to the overall composite volume with porosity contents up to 10% [62]. In contrast, in glass and carbon fiber composites, considerable knowledge has been accumulated to diminish the porosity contents below 1% [73]. Altogether, porosity can typically not be neglected in cellulose fiber composites, and it should be integrated in the evaluation of composite performance.

Figure 6 shows examples of the three types of porosity that typically can be found in cellulose fiber composites: fiber lumen porosity, interface porosity, and impregnation porosity. In a study by Madsen et al. [62], the porosity content is correlated with the fiber and matrix contents, and a model for the numerical correlation between weight and volume contents of the composite constituents is presented. Input parameters are (i) the density of fibers and matrix, which can be measured by pycnometry and buoyancy methods, (ii) a number of empirical porosity constants, which can be measured from images of composite microstructures, and (iii) the maximum obtainable fiber volume fraction, which can be determined from the compaction behavior of the fiber assembly. The model predicts the volume fractions of fibers, matrix and porosity as a function of the fiber weight fraction. The model applies to composites in general, but it is particularly relevant to composites with a relatively high porosity content, which is typically the case for cellulose fiber composites.

Figure 7(a) shows experimental data and model predictions of the volumetric composition of a series of unidirectional flax fiber/thermoplastic matrix composites with variable fiber weight fractions. The volume fractions of fibers and porosity are increased as a function of the fiber weight fraction, until a certain value where after the fiber volume fraction is constant, and the porosity starts to increase more dramatically. The transition fiber weight fraction is determined to be 0.61. Thus, the given composites should be manufactured with a fiber weight fraction of 0.61 to have the best possible combination of high fiber volume fraction, and low porosity, and as will be shown next, this leads to composites with a maximum obtainable tensile stiffness.

The predictions of the volumetric composition in composites can be integrated with micromechanical models. This has been done in the study by Madsen et al. [74] by applying the rule of mixtures model for stiffness of composites. A modified version of the model was used in which the effect
Figure 6: Types of porosity in cellulose fiber composites. Shown are cross-sectional images of unidirectional hemp fiber/polyethylene terephthalate composites. The optical microscope image in (a) shows a hemp fiber yarn, and the scanning electron microscope image in (b) shows a close-up of the fibers in the yarn [62].

The types of porosity present in the composites are illustrated in the figure. These include impregnation porosity, interface porosity, and fiber porosity. Impregnation porosity refers to the porosity that results from the infiltration of the matrix into the fiber surfaces. Interface porosity is the porosity at the fiber/matrix interface, and fiber porosity refers to the porosity within the fibers themselves.

7. Moisture Sensitivity of Composites

Compared to composites with conventional fibers, the Achilles’ heel of cellulose fiber composites is their propensity to take up moisture, which leads to swelling, dimensional instability, and potential degradation of mechanical properties. The hydrophilicity of the fibers is due to the abundance of available hydroxyl groups in hemicellulose, in amorphous cellulose and at the surface of cellulose crystallites. For structural materials, moisture sensitivity is generally considered to be a disadvantage, and should be reduced, if possible. For cellulose fiber composites, this can be done by cross-linking of the cellulose polymer within the fibers [76], use of a stiff and hydrophobic matrix [77], and use of a moisture barrier coating [78].

How the swelling of the fibers affects the dimensional stability of the composites is complicated due to the irregular microstructure of the fiber assembly. One way to isolate the hygroexpansion of the fibers, and to quantify its contribution to the hygroexpansion of the composites, is to use micromechanical models. These are similar to models primarily developed for thermal expansion and residual stresses in ceramic-matrix composites. Thermal expansion and hygroexpansion are governed by the same physical equations, where thermal and hygral strains are governed by temperature and moisture, respectively. The micromechanical models for hygroexpansion of composites include also parameters for the elastic properties of the fiber and matrix constituents. In a study by Neagu et al. [48], curl measurements of strips of wood fiber composites and wood fiber mats were used to determine the
transverse hygroexpansion coefficient of wood fibers to be approximately 0.10 strain per relative moisture content. This is in accordance with a few scarce data found in the literature on the hygroexpansion properties of cellulose fibers [79].

In a study by Madsen et al. [80] of the hygroexpansion of unidirectional hemp fiber/thermoplastic matrix composites, the dimensional changes were found to be well correlated with the fiber volume fraction. Selected results are shown in Figure 8. By using a reference humidity of 65% RH, the transverse hygroexpansion was found to be 0.9% and 3.3% at humidities of 85 and 100% RH, respectively, for composites with the highest fiber volume fraction of 0.43. The hygroexpansion in the longitudinal direction was found to be low, and slightly negative, which presumably is due to moisture-induced relaxation of residual tensile stresses in the matrix. It can be observed in the figure that the experimental data points are well simulated by the micromechanical model lines.

Glass and carbon fibers do not take up any moisture, although glass fibers are sensitive to environmental stress corrosion in the presence of moisture and tensile stress [81]. The moisture sensitivity is larger for wood fibers than for plant fibers, since the former contains a larger relative amount of hemicellulose which is the most hydrophilic polymer in the cell wall.

**Figure 8: Hygroexpansion of unidirectional hemp fiber/polyethylene terephthalate composites as function of the fiber volume fraction. Hygroexpansion was measured in the longitudinal and transverse directions, and at the humidities of 85 and 100% RH, with respect to a reference humidity of 65% RH. Lines are calculated by micromechanical models. Modified from [80].**

**8. Applications of Composites**

In Europe, cellulose fiber composites are mainly used by the automotive industry. The applied fiber preforms are loose fibers used for injection molding techniques, and non-woven mats used for compression molding techniques. Due to the nominal random fiber orientation in these composites, they possess only moderate mechanical properties (see Table 2), but this makes them nevertheless well qualified to be used in non-structural components such as doorliners, bootliners, and parcel shelves. The low prices of loose fibers and non-woven mats of cellulose fibers, compared to their synthetic counterparts, form a strong motivation for the use of these two preforms in the automotive industry. Outside Europe, the use of non-structural components based on cellulose fibers is more widespread, and wood fibers are by far the preferred fiber type. In North America, the main applications are building components, such as decks, windows profiles and floorings.

Recently, in the context of research and development projects, a number of demonstrators have been made to reveal the good potential of cellulose fiber composites in new kinds of applications (see Figure 9).

(i) Sculpture shown at the Louisiana Museum of Modern Arts, Denmark, special exhibition “Green architecture for the future”, winner of the JEC innovation award 2010, joint venture of 20 companies coordinated by 3XN architects, Denmark.


(iii) Chair for children, developed by the research institute Innventia AB together with pulp industry and architects, exhibited at the Milan furniture fair 2009.


(v) Small-scale rotor blade to be used for a wind turbine car, EU 7th Framework Programme project, WOODY (2009–2012).


**9. Future Perspectives: Nanofiber Composites**

In recent years, considerable attention has been directed towards composites made from cellulose nanofibers (e.g., [82]). As already described, the cell wall of wood and plant fibers is structured like composites with cellulose microfibrils embedded in a matrix of hemicellulose and lignin. The cellulose microfibrils are having lateral dimensions in the 10–100 nm range and axial dimensions in the micrometer range, and they are therefore suitable as reinforcement in nanofiber composites. The idea is to achieve considerable improvements in engineering properties with the addition of nanofibers, beyond those obtained with fibers in the micrometer range. This can be attributed to the high specific surface area of the nanoscale fibers, which will affect the properties of the surrounding matrix. The success of nanofiber composites is particularly obvious if only a minute addition of fibers is considered, and the dispersion of nanofibers is preserved.

Carbon nanotube composites have shown great promise for a relatively long time, but have yet to deliver in large volume applications [83]. A difference between cellulose nanofibers and carbon nanotubes is the ability of the cellulose nanofibers to bond to each other, by hydrogen bonding, whereas the carbon nanotube surface is chemically inert.
Figure 9: Examples of new applications of cellulose fiber composites: (top, left) sculpture, (top, right) wheel rim, (middle, left) chair for children, (middle, right) double-curvature panels, (bottom, left) small-scale rotor blade, and (bottom, right) exhibition stand. See text for more details.

The cellulose nanofibers can form a very strong network, and furthermore bond well to polymer matrix materials with polar groups. This leads however also to processing difficulties, since the cellulose nanofibers tend to aggregate and take a long time to dry after wet processing. Processability and performance are thus complementary and mutually opposing behaviors. The functional hydroxyl groups of the fibers can be modified to improve dispersion and processability, although this is typically accompanied with increased costs. The main challenges for cellulose nanofiber composites are probably to learn how to manufacture bulk composite components with retained nanofiber slenderness and dispersion. The raw materials are the cellulose fibers themselves, from wood pulp or plant fibers, making the raw materials costs negligible compared with manufacturing costs.

10. Overall Comparison between Fibers

As shown in the sections above, wood and plant fibers are similar in some respects and differ in others. Thus, depending on the intended application, one particular fiber type is more suitable than the other. In the following, an overall comparison is given to highlight some advantages of wood versus plant fibers, and vice versa. Similarly, cellulose fibers are compared to their synthetic counterparts, glass and carbon fibers.

Advantages of wood fibers, as compared with plant fibers, are as follows.

(i) Low cost, readily available from pulp mills.
(ii) Relatively short fibers mean better processability.
(iii) Mature infrastructure available in pulp and paper mills to produce large quantities at low cost.
(iv) Preforms can be made using paper-making technologies.
(v) Rather uniform batches of pulp qualities can be achieved.
(vi) Does not compete with cultivation of food crops.
Advantages of plant fibers, as compared with wood fibers, are as follows.

(i) High productivity and yield.

(ii) High cellulose content, high degree of cellulose crystallinity, low microfibril angle, small lumen mean excellent mechanical properties of fibers.

(iii) Relatively long fibers means possibility to control fiber orientation and lay-up.

(iv) Textile technologies can be used to produce yarns, woven fabrics and non-crimp fabrics.

Despite the differences, wood and plant fibers have more in common than in what differ them from one another. Some advantages of cellulose fibers as compared with glass and carbon fibers can be mentioned.

(i) Renewable.

(ii) Biodegradable.

(iii) Light, that is, the composites have good specific properties which are important in automotive and packaging.

(iv) Low cost raw materials.

The main disadvantages of cellulose fibers as compared with glass and carbon fibers are as follows.

(i) Moderate mechanical properties.

(ii) Sensitivity to moisture, leading to dimensional instability, and potential degradation of mechanical properties.

(iii) Not fully developed composite manufacturing techniques.

The above lists are by no means comprehensive, but only serve to show some of the traits of wood and plant fibers in an applied composite context. In the further development of cellulose fiber composites, both advantages and disadvantages play an important role. The specific advantages guide which application areas that are relevant. For instance, the combination of low cost, renewability and biodegradability make cellulose fiber composites suitable materials for packaging applications. The disadvantages limit their applications. Research on how to alleviate these shortcomings can expand the proliferation of cellulose fibers as an eco-friendly alternative to synthetic fibers. If cellulose fibers can be processed to retain better their innate high stiffness and strength, and can be modified to become less hydrophilic they are also potential reinforcement fibers in advanced structural outdoor applications, for example, in rotor blades for wind turbines and in load-carrying components in transport applications.

References


