Mesoporosity as a new parameter for understanding tension stress generation in trees

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Abstract

The mechanism for tree orientation in angiosperms is based on the production of high tensile stress on the upper side of the inclined axis. In many species, the stress level is strongly related to the presence of a peculiar layer, called the G-layer, in the fibre cell wall. The structure of the G-layer has recently been described as a hydrogel thanks to N₂ adsorption–desorption isotherms of supercritically dried samples showing a high mesoporosity (pores size from 2–50 nm). This led us to revisit the concept of the G-layer that had been, until now, only described from anatomical observation. Adsorption isotherms of both normal wood and tension wood have been measured on six tropical species. Measurements show that mesoporosity is high in tension wood with a typical thick G-layer while it is much less with a thinner G-layer, sometimes no more than normal wood. The mesoporosity of tension wood species without a G-layer is as low as in normal wood. Not depending on the amount of pores, the pore size distribution is always centred around 6–12 nm. These results suggest that, among species producing fibres with a G-layer, large structural differences of the G-layer exist between species.

Key words: Growth stress, hydrogel, mesoporosity, tension wood.

Introduction

Tension wood (TW) is a peculiar wood tissue that is often formed in the upper side of leaning trunks and branches in hardwood species (Isebrands and Bensend, 1972) to control the orientation of the growth axis by generating high tensile stress (Wardrop, 1964; Fisher and Stevenson, 1981). For many commonly studied species such as beech, poplar, oak, or chestnut, TW is characterized by the occurrence of fibres with a particular morphology and chemical composition due to the development of the so-called gelatinous layer (G-layer). This layer is composed of cellulosic microfibrils that are nearly parallel to the fibre axis (Dadswell and Wardrop, 1955; Wardrop, 1964; Côté et al., 1969) embedded in a highly hydrated polysaccharide matrix (Nishikubo et al., 2007; Bowling and Vaughn, 2008; Mellerowicz et al., 2008).

Although it has been well established that the G-layer is the driving force of the high tensile stress generated in TW (Trénard and Guéneau, 1975; Yamamoto et al., 2005; Fang et al., 2008), the underlying mechanism is still a subject of debate. In previous research, the structure of the G-layer has been described as possessing gel-like characteristics: large shrinkage (Clair and Thibaut, 2001; Fang et al., 2007) and high rigidification during drying (Clair et al., 2003). Recently, the hydrogel structure of the chestnut G-layer has been characterized thanks to nitrogen adsorption. The...
chestnut G-layer contains mesopores (pore size between 2 nm and 50 nm) and the pore surface areas is more than 30 times higher than that in normal wood (NW) (Clair et al., 2008). The swelling of the G-layer matrix has been suggested recently by several authors as the possible driving force of the growth stress generation in TW (Nishikubo et al., 2007; Goswami et al., 2008). However, it is known that many species (Onaka, 1949; Fisher and Stevenson, 1981; Clair et al., 2006) are able to produce tensile stress without forming a typical G-layer. Different anatomical patterns of TW exist, from fibres with a typical G-layer to fibres exhibiting no difference at the fibre level (Clair et al., 2006; Ruelle et al., 2006, 2007).

These results led us to revisit the concept of the G-layer and to propose an objective description of the TW cell wall structure, until now only defined from visual assessments (stain, detachment or swollen aspect) which are the subject of much debate (Clair et al., 2005a, b, 2006).

Let us briefly introduce the principles of mesoporosity measurements by the nitrogen adsorption method. This technique, based on the measurement of the adsorption isotherm of nitrogen at its boiling temperature (77 K) on an outgassed sample, allows the pore size and surface area of materials with cavities smaller than 50 nm to be estimated (Gregg and Sing, 1982; Rouquerol et al., 1999) and has recently been applied to the study of the texture of polysaccharide aerogels (Valentin et al., 2005; Quignard et al., 2008). The majority of isotherms have been grouped into six types by IUPAC classification (Sing et al., 1985), but only three are commonly found in the adsorption on polar materials (type I, type II, and type IV in Fig. 1).

Type I is obtained with microporous (pore size <2 nm) solids. The adsorption takes place at very low relative pressure regions (the ratio between pressure and saturation pressure $P/P^0 <0.3$) because of multidirectional interactions between the pore walls and the adsorbate. The reversible type II isotherm is characteristic of the non-porous or macroporous (pore size >50 nm) solids. If the knee of the isotherm is sharp, the uptake at point B—the beginning of the middle linear section—provides a measure of the monolayer capacity, from which the surface area of the adsorbent can be calculated. The type IV isotherm is obtained with mesoporous (2 nm <pore size <50 nm) solids. The hysteresis loop is associated with the secondary process of capillary condensation, which results in the complete filling of the mesopores at $P/P^0 <1$.

Pores can have a regular or, more commonly, an irregular shape, either an ink-bottle shape (pore body larger than pore mouth) or a funnel shape (the opposite). Pores can be closed (not accessible from the outside), blind (open only at one end), or through (open at both ends). Each pore can be isolated or, more frequently, connected to other pores to form a porous network (Rouquerol et al., 1999) (Fig. 2). The surface and structural properties of the pores control the interactions of material with gases, fluids, and other solids.

The adsorbate desorption is the opposite of adsorption, but evaporation from mesopores usually takes place at a pressure lower than that of capillary condensation giving a hysteresis loop. The reason for the hysteresis is that the formation of the meniscus in capillary condensation is an activated phenomenon, while the retreat of the meniscus in evaporation is usually an equilibrium phenomenon. Pore shape affects the mechanisms of condensation and evaporation and four types of hysteresis have been recognized according to IUPAC classification (Sing et al., 1985) (Fig. 3).

Type H1 hysteresis is characteristic of solids crossed by channels with uniform sizes and shapes. Type H2 corresponds to channels with a pore mouth smaller than the pore body (this is the case of ink-bottle-shaped pores). Type H3
hysteresis is usually found on solids with a very wide distribution of pore size and type H4 corresponds to limited amounts of mesopores limited by micropores.

The aim of this study was to adapt experimental techniques used in material chemistry to the specificity of biological material such as wood. It was applied to tension wood of several angiosperm species to answer the following questions: is the occurrence of a G-layer always associated with high porosity? Does non-G-layer TW contain mesopores? Do all TW types have the same pore shape and pore size distribution? Can we check the presence/absence of the G-layer by an N₂ adsorption–desorption isotherm?

Materials and methods

Experiments were performed on six tropical rainforest species (one tree per species) distributed in three families (Table 1). Sampling was carried out near the Paracou experimental field in French Guyana. Care was taken to maintain the samples in wet conditions in storage from the living tree to the experiments. TW occurrence was demonstrated by the mechanical measurement of released strains at the circumference of the leaning trees.

Growth stress measurements

The strain gauge method (Yoshida and Okuyama, 2002; Jullien and Gril, 2008) was used to estimate the residual growth stress close to the trunk surface on the living tree. Measurements of longitudinal growth strains (GS expressed in μm m⁻¹ or microstrain) were performed at eight positions every 45° around the circumference of the inclined trunks at breast height.

Tension stress gives a negative strain value while compression stress gives a positive strain value. A high local level of growth stress is always related to the presence at TW (Trenard and Gueneau, 1975). For each species, a TW sample was taken near the highest absolute value of growth stress, while a NW sample was taken near the lowest absolute value.

Anatomical observations

Cross-sections (12 μm in thickness) were cut with a sliding microtome equipped with disposable razor blades. Double staining with safranin/fast green was used to demonstrate the presence of the G-layer. Safranin stains lignified tissues red, and fast green stains both lignified and unlignified cell walls green. Consequently, lignified tissues were red mixed with varying degrees of green while essentially cellulosic cell wall layers, like the G-layer, were green. Sections were observed with an optical microscope at ×400 magnification.

N₂ adsorption–desorption measurement

As direct drying usually results in the collapse of mesoporosity (Clair et al., 2008), samples were supercritically dried as follows.

**Table 1.** Characteristics of the six species studied

<table>
<thead>
<tr>
<th>Family</th>
<th>Species</th>
<th>Tree diameter (cm)</th>
<th>GS (μstrain)</th>
<th>Fibre pattern</th>
<th>Vₘₚ (cm³ g⁻¹)</th>
<th>Sₐ (m² g⁻¹)</th>
<th>D₃₄ (nm)</th>
<th>D₅₃ (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lauraceae</td>
<td>Ocotea guyanensis</td>
<td>19</td>
<td>−2097</td>
<td>−399</td>
<td>Gthin</td>
<td>0.003</td>
<td>&lt;0.001</td>
<td>2.8</td>
</tr>
<tr>
<td>Lauraceae</td>
<td>Sextonia rubra</td>
<td>25</td>
<td>−2362</td>
<td>−328</td>
<td>Gthick</td>
<td>0.133</td>
<td>0.002</td>
<td>64.6</td>
</tr>
<tr>
<td>Fabaceae</td>
<td>Inga alba</td>
<td>29</td>
<td>−2408</td>
<td>−310</td>
<td>Gthin</td>
<td>0.014</td>
<td>0.016</td>
<td>8.8</td>
</tr>
<tr>
<td>Fabaceae</td>
<td>Tachigali melinonii</td>
<td>18</td>
<td>−1488</td>
<td>−480</td>
<td>Gthick</td>
<td>0.004</td>
<td>0.001</td>
<td>4.8</td>
</tr>
<tr>
<td>Myristicaceae</td>
<td>Virola micheli</td>
<td>36</td>
<td>−1708</td>
<td>+45</td>
<td>no G</td>
<td>0.003</td>
<td>0.003</td>
<td>2.6</td>
</tr>
<tr>
<td>Myristicaceae</td>
<td>Iryanthera sagotiana</td>
<td>26</td>
<td>−1485</td>
<td>+12</td>
<td>no G</td>
<td>0.005</td>
<td>0.002</td>
<td>4.7</td>
</tr>
</tbody>
</table>
Wood sticks (5 mm in a longitudinal direction, 1×1 mm² in cross-section) were dehydrated by immersion in increasing concentrations (10, 30, 50, 70, 90, 100%, and anhydrous) of ethanol solutions for 3 d each. The dehydrated samples were then introduced into a Polaron 3100 apparatus which was filled with liquid CO₂. Samples were left for 2 h in liquid CO₂ before ethanol evacuation. The temperature was raised to 32 °C so that CO₂ reached its critical point (74 bar, 31.5 °C). Depressurization took 30 min. During this procedure the liquid phase has been transformed in a supercritical fluid with a null surface tension. In this way, the shrinkage due to capillary pressure was prevented and the solid obtained, an aerogel, was expected to reproduce the texture of the original hydrogel in the dry state (Pierre and Pajonk, 2002; Cansell et al., 2003).

Nitrogen adsorption–desorption was performed on a micromeritics ASAP 2010 volumetric apparatus at 77 K. Before adsorption measurements, samples (1.0–1.5 g) were outgassed at 373 K under vacuum until a stable 3×10⁻⁵ Torr pressure was obtained without pumping. This is done to remove physically absorbed gases from the sample surface, in particular, water vapour. The adsorption at low relative pressure allows the specific surface area of the samples to be evaluated by the BET method (Brunauer et al., 1938), assuming an adsorbed N₂ molecule covers 0.162 nm². Pore size distributions calculated from adsorption and desorption data using the method of Broekhoff and de Boer, proved more accurate than the commonly used BJH method (Barrett et al., 1951; Galarneau et al., 1999). The pore volume was evaluated by the a₋₀ method (Gregg and Sing, 1982) and is given in Table 1 by assuming for condensed N₂ the density of liquid N₂. The adsorbed volume in the figures is given as the volume of gas at standard temperature and pressure, with a density 647 times lower than the liquid density.

Results

Anatomical observations

Anatomical sections of TW and NW in the six species are illustrated in Fig. 4. Octeoa guyanensis, Sextonia rubra, Tachigalia melinonii, and Inga alba TW have fibres with a well-differentiated G-layer that is obviously green after double staining with safranin/fast green (Fig. 4A, C, E, G). However, there are notable differences in morphology, thickness, and distribution. In TW fibres of S. rubra, a thick G-layer almost fills up the entire lumen, whereas in O. guyanensis, I. alba or T. melinonii TW, the G-layer is thinner, sometimes delaminated from the adjacent S₂ and folded into the lumen towards the same direction. This detachment phenomenon results from sample preparation with classical sectioning (Clair et al., 2005b). In T. melinonii, a G-layer was only observed in a limited number of fibres whereas, in the other three species, most fibres contained a G-layer.

For the other two species (Virola michellii and Iriartea sagotiana) there is no G-layer and no obvious difference between TW and NW fibres can be seen simply by anatomical observation. Irrespective of the fibre pattern, all species can produce a high level of mechanical tensile stress (ranging from −2408 to −1485 μstrain). As already shown earlier (Clair et al., 2006; Ruelle et al., 2006), the general relations between tensile stress level in tension wood and macroscopic anatomical variations are not visible, while observation at an ultrastructural level allows some common features in cellulose organization to be seen (Ruelle et al., 2006, 2007).

Textural characterization

The textural parameters of TW and NW in six species are reported in Table 1. S. rubra, the only specimen with a thick G-layer, is the only one to present a TW with a very high mesopore volume. The other samples, with a thin G-layer or no observable G-layer, present a much lower mesopore volume difference between TW and NW.

Pore surface area (S\textsubscript{BET})

Examples of nitrogen adsorption–desorption isotherms (77 K) of S. rubra and I. sagotiana are shown in Fig. 5 (the other four species were studied and the observed isotherms were similar to I. sagotiana, data not shown). Among the six species studied only O. guyanensis, S. rubra, T. melinonii, and I. sagotiana TW aerogels clearly show higher specific surface areas than the corresponding NW aerogels. The surface area data are strongly correlated with the mesopore volume.

S. rubra TW adsorbed a 60 times larger amount of nitrogen than NW, yielding a surface area of 64.6 m² g⁻¹ for TW and 2.6 m² g⁻¹ for NW (Table 1). A similar difference has also been observed in species with a thick G-layer such as chestnut (Clair et al., 2008) or poplar (B Clair, unpublished data). However, a large range of specific surface areas (from 3 m² g⁻¹ to 65 m² g⁻¹) was observed in TW aerogels among the six species, proving that there are large difference in mesopore amounts between the thick and thin G-layers.

Pore shape and size distribution

According to the IUPAC classification (Sing et al., 1985), the isotherm of S. rubra TW is type IV (Fig. 1) with a H₃ type hysteresis loop (Fig. 2), indicating the presence of mesopores between the cellulose microfibrils or in the matrix forming slit-shaped pores with a non-uniform size. The isotherm of I. sagotiana TW (as well as the other four species) is intermediate between type IV and type II, indicating the presence of large mesopores with a broad size distribution that continues into the macropore domain. The hysteresis loop is very narrow, the adsorption and desorption branches being almost vertical and nearly parallel above 0.8 relative pressure, confirming the presence of a significant outer surface.

The peak pore sizes estimated from the adsorption and desorption are listed in Table 1. Examples of pore volume...
The pore size distributions against pore diameter are plotted in Fig. 6 for S. rubra and I. sagotiana TW aerogels. All the samples present broad pore size distributions, measured both on the adsorption and desorption branches of the isotherms. The sharp peak between 4 nm and 5 nm on the desorption branch is a typical artefact due to the tensional instability of the N₂ meniscus (Trens et al., 2005). Pore size distribution determined from the adsorption branch of the isotherm corresponds to the cavity size, while that of the desorption branch corresponds to the throat size of a pore (Groen and Pérez-Ramirez, 2004). Thus, comparing the adsorption and desorption pore size distribution leads to information about the pore shapes. For S. rubra and O. guyanensis aerogels, the pores have an ink-bottle shape with pore cavities that are less than twice the diameter of the throats. The pore size distribution of I. alba (as well as I. sagotiana and T. melinonii) aerogels, the pore size measured on the adsorption branch is smaller than the pore size measured on the desorption branch. This behaviour indicates that the usual correlations between condensation pressure and pore size do not apply to the geometry of the measured pores. This effect, albeit infrequent, is far from being unprecedented and has been observed in adsorbents with highly connected...
tridimensional pore systems (Fan et al., 2001) and can be attributed to funnel-like pores with a wide opening, which decreases the activation energy of condensation.

All porosity parameters observed in *S. rubra* correspond to what was described for the chestnut G-layer (Clair et al., 2008) and also observed for the poplar G-layer (B Clair, unpublished data) which also present thick G-layers; i.e. an isotherm of type IV, presenting a typical hysteresis loop of mesoporous adsorbents with a pore size diameter between 2 nm and 50 nm (Clair et al., 2008) with a maximum around 6 nm. *S. rubra* was the only sample of the present study with a thick G-layer. Other samples with a thin G-layer, like *O. guyanensis* and *T. melinonii*, present an extremely low mesopore volume, suggesting that not all G-layers present a porosity which can be easily stabilized by supercritical drying.

In the case of *I. alba*, another species with a thin G-layer in TW, the mesoporosity of NW has been measured at a much higher level than that of any other NW, and was even slightly higher than that of its TW. Moreover, the shape of the pores seems to be different from the ink-bottle pores of chestnut and *S. rubra* TW.

In species without a G-layer, mesoporosity was low and at the same level in NW and TW (a similar observation was made on *Simarouba amara*; B Clair, unpublished data). Moreover, for *V. michellii*, *I. sagotiana*, and *T. melinonii* the shape of the pores seems to correspond to a pore system more accessible than the interfibrillar mesoporosity of chestnut TW (Clair et al., 2008).

**Discussion**

This study shows the ability of the nitrogen adsorption–desorption isotherm method to characterize mesoporosity in wood. It appears that mesoporosity can always be detected in never-dried wood samples. The number of pores can be very different between the samples and the structures of the pore networks are also different. In wood with a thick G-layer, the large amount of mesopores can easily be attributed to the G-layer itself and provides indications about the nature of pores. Pores have ink-bottle shapes with a pore body size distributed around 6–11 nm and a pore mouth around 6 nm and can be attributed to cavities between macrofibrils. In wood without a G-layer, the mesopore volume is very small and the narrow hysteresis could correspond to a network of large-mouth funnel pores. *I. alba*, a species with a thin G-layer and an intermediate mesopore volume, presents an intermediate width of the hysteresis loop, which could be attributed to the simultaneous presence of both pore systems, the one related to the G-layer and the one typical of woods without a G-layer.

The G-layer is known to be composed of pectin and hemicelluloses such as xyloglucan and (1,4)-β-galactan (Nishikubo et al., 2007; Arend, 2008; Bowling and Vaughn, 2008; Mellerowicz et al., 2008). These components would also be present in other cell wall layers (in NW cell wall) in a highly hydrated state allowing it to keep its mesoporosity. The pit membrane, which is known to be composed of pectin gel (Zwieniecki et al., 2001; Van Ieperen, 2007), would also be a good candidate for the place where mesoporosity can be concentrated. In *I. alba*, where the mesoporosity has been measured higher in NW than in TW, axial parenchyma could be susceptible to contain such
mesoporosity since its abundance was measured around 40% in NW and 20% in TW.

This study provides new insights into the understanding of the G-layer since it confirms that major differences between species can be observed in TW fibre secondary walls. All these species are able to produce high tensile stress but their anatomy and nanostructure can differ widely. The following conclusions can be drawn. (i) TW that does not develop a G-layer does not contain a mesoporosity significantly higher than the NW. (ii) TW developing a G-layer can be separated into two classes, a thin G-layer presenting little mesoporosity and a thick G-layer showing a high mesoporosity. The fact that mesoporosity is observable only when the G-layer is thick leads to a new open question. Is the G-layer thick merely because of its high mesoporosity allowing it to remain highly hydrated in a swollen stage? In that case, the total amount of cellulose would be not so different in a swollen G-layer compared to a thin G-layer. This would explain why both TW types can produce similar high tensile growth stresses. It would then be interesting to follow the maturation process of these woods and detect the swelling or shrinkage in the gel, which is currently suspected as being responsible for the growth stress generation in TW.

It is quite likely that several mechanisms contribute to the generation of high tensile stress to maintain the vertical orientation of the main stem. The characterization of microstructural features of different woods and their correlation with mechanical parameters are a first step towards an assessment of the possible mechanisms. Then we can hypothesize that mechanisms differ from species to species and, in some cases, are not directly linked to mesoporosity. An alternative hypothesis would be a common mechanism, but apparent mesoporosity would be a residual state of the maturation process that was later hidden in some species.

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