GROWTH STRESSES ARE HIGHLY CONTROLLED BY THE AMOUNT OF G-LAYER IN POPLAR TENSION WOOD

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SUMMARY

To determine how gelatinous fibres and gelatinous layers contribute to the magnitude of longitudinal growth stress in tension wood, anatomical measurements of gelatinous fibres were carried out on poplar tension wood (*Populus* I4551). It was found that (a) no gelatinous fibres were observed under a growth strain level of 0.06 to 0.08%; (b) almost 100% of the non-conductive tissues contained gelatinous fibres above a growth strain level of 0.15 to 0.19%; and (c) the area of fibres, the area of fibres with gelatinous layers per unit of tissue area, and the thickness of the gelatinous layers predominantly influenced the magnitude of growth stress.

Key words: Gelatinous fibre, gelatinous layer, growth strain, growth stress, tension wood, poplar.

INTRODUCTION

Trees produce asymmetric growth stresses to maintain the vertical orientation of the main stem or the angle of a branch, in order to receive sufficient light or in response to a strong dominant wind. This is usually achieved by the production of reaction wood, often combined with eccentric growth. While gymnosperms produce compression wood on the lower side of leaning stems, angiosperms produce tension wood generating high tensile stresses on their upper side (Wardrop 1964; Fisher & Stevenson 1981). Both strategies allow strongly heterogeneous growth stress distribution at the periphery of stems, generating the bending moments required to control their shape.

Normal wood fibres are composed of a thin primary wall and a thick secondary wall divided into 3 sub-layers; the S_1 , S_2 and S_3 layers. In many hardwood species such as beech, poplar, oak and chestnut, tension wood contains fibres with a special morphology and chemical composition due to the development of the so-called gelatinous layer (G-layer) (Onaka 1949) that replaces the S_3 layer and a part or the whole of the S_2 layer (Saiki & Ono 1971). The G-layer is known to have a high cellulose content with a high degree of crystallinity (Norberg & Meier 1966; Côté *et al.* 1969) and to contain microfibrils oriented along the axis of the cell (Fujita *et al.* 1974).

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There is some disagreement about the origin of growth stresses in wood (Boyd 1985; Bamber 1987; Yamamoto & Okuyama 1988; Okuyama *et al.* 1994; Yamamoto 1998; Bamber 2001). While it is known that some species do not need to produce a G-layer to induce high growth stresses (Okuyama *et al.* 1994; Yoshida *et al.* 2000; Clair *et al.* 2006b), tension wood with a G-layer is a good model for trying to understand growth stress generation. In this paper we will concentrate on the contribution of the G-layer to the magnitude of growth stresses in tension wood. Is it the percentage of fibres, the percentage of fibres with a G-layer (G-fibres) or the thickness of the G-layer in the G-fibres?

Previous studies (Okuyama *et al.* 1994; Yamamoto *et al.* 2005) have examined similar questions, but G-layer quantification was biased by its swollen appearance always observed on sliding microtome sections (Clair *et al.* 2005a). Moreover, this artefact was possibly influenced by the growth stress level, so that the bias introduced in the previous findings could have been even greater. In this study measurements were done on embedded sections to avoid this artefact and thus allow a correct quantification of the G-layer.

MATERIALS AND METHODS

The experiments were performed on poplar tension wood (*Populus* I4551). Poplar tension wood has fibres with a gelatinous (G-)layer and exibits high longitudinal tensile stress. The tension wood samples were obtained from a 15-year-old leaning tree growing in a plantation near Montpellier in the south of France. The tree was chosen because it was leaning and there was evidence that there was an active process to restore the stem to a vertical orientation.

Growth strain (GS)

The presence of tension wood was confirmed by the measurement of residual growth strains using the strain gauge method described in Yoshida & Okuyama (2002). Measurement of longitudinal growth strain (GS) was done at 25 positions around the surface of an inclined poplar trunk at four different heights. GS is directly correlated to the growth stress level within trees of a same species (Archer 1986; Fournier *et al.* 1994). As all the GS values were negative, absolute GS values were used to simplify representation and analysis. GS values ranging from 0.01 to 0.23% were obtained with the highest values from the upper side of the stem and lowest values from the lateral and lower sides of the stem. As the study focused on the role of the G-layer and none or very few G-layers were found microscopically in the samples with GS values up to 0.06%, 5 samples with GS values regularly spread from 0.08 to 0.23% (0.23, 0.19, 0.15, 0.12 and 0.08%) were chosen for anatomical studies.

Sample preparation

Samples were taken from the respective GS measurement positions and placed in water as soon as they were taken from the tree. As normal sectioning methods with a sliding microtome results in an uncontrolled transverse swelling and detachment of the G-layer in poplar (Clair *et al.* 2005a, 2005b), embedded wood samples were used and serial-sectioning was performed with a glass knife.

Wood samples (2 mm in longitudinal direction, 1×1 mm in cross section) were longitudinally cut by splitting. They were then cut mid-length, perpendicular to the fibre direction, with a new razor blade to obtain two matched samples (one was used for this study, and the other used to examine the drying shrinkage of the G-layer (Fang *et al.* 2007)).

The samples were dehydrated with ethanol and embedded in LR White resin (two exchanges of resin/ethanol mixture for 1 hour, followed by two exchanges in pure resin for 1 hour and kept overnight at room temperature, then polymerised at 65 °C overnight). After polymerisation of the resin, tissue deformation is prevented, and further sectioning will not alter the shape and the size of the cell wall layers.

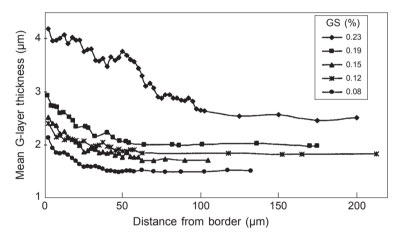


Figure 1. Mean G-layer thickness (MGLT, μm) variation with the distance from the border for 5 samples.

Sectioning

Serial transverse sections ($2.5 \mu m$ thickness) were performed with a glass knife and distance from the upper surface (border) was recorded for each section. For each sample more than 100 sections were obtained, mounted on glass slides and observed under an optical microscope.

To avoid measurement of the G-layer in a swollen state (Clair *et al.* 2005a), we plotted, for each of the 5 samples, the variation of the mean G-layer thickness (MGLT, measured as explained below) with the distance from the border (Fig. 1). MGLT became almost stable when the distance from the border reached 70 to 120 μ m, depending on the sample. In this paper we will focus on these stabilised values, as they provide a good indication of the undisturbed morphology of the cell wall of tension wood cells, and in our opinion, as the cell wall was in the living tree. For each sample the values of the last 5 or 6 measured sections were used for measurements.

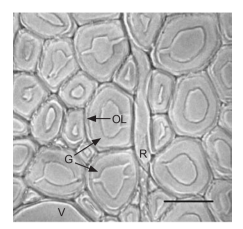
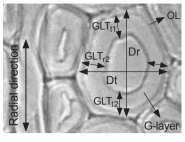


Figure 2. Transverse section of sample 1. G = G-layer; $OL = other cell wall layers including compound lamella, <math>S_1$ and S_2 ; V = vessel; R = ray. — Scale bar = 20 μm .

Measurement

Images (Fig. 2) were obtained with a digital camera and measurements obtained with ImageJ 1.34s and Optimas v6.5 image analysis software.



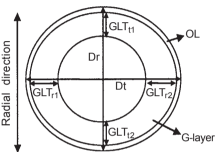


Figure 3. Detail of Fig. 2, presenting the G-layer thickness and cell diameter measurements. G-layer thickness was always measured in the same 4 positions: 2 radial (GLT_{r1} , GLT_{r2}) and 2 tangential (GLT_{t1} , GLT_{t2}). Cell diameter was measured in two directions (radial, Dr, and tangential, Dt).

At the tissue scale, for each sample, measurements of the vessel area were performed on images (magnification \times 100) covering the whole section and measurements of the G-fibres area were performed on 5 images (magnification \times 500) ordered in the radial direction. The following parameters were measured:

Total area: A_T Vessels area: A_V G-fibres area: A_{GFs}

This allows the calculation of the following parameters:

Fibre area: $A_F = A_T - A_V$ (assuming the ray area is negligible)

Fibre area ratio: $FR = A_F / A_T = 1 - (A_V / A_T)$

Area ratio of G-fibres among fibres: $GFR_F = A_{GFs} / A_F$

Area ratio of G-fibres among the total area: $GFR_T = GFR_F \times FR$

At the fibre scale, for each sample, on each section, the same 10 to 12 G-fibres were followed from the sample border to 100–200 µm deep in the sample (Fig. 1). The following parameters (Fig. 3) were measured for each of the 10 to 12 fibres (radial and tangential directions determined as parallel and perpendicular to the rays respectively):

Fibre diameters: D_r , D_t (respectively in radial and tangential directions);

G-layer thickness (measured on both sides of the fibre): GLT_{rl} , GLT_{r2} in radial direction and GLT_{tl} , GLT_{t2} in tangential direction.

In order to estimate the surface area of the G-fibre and G-layer, two simplifying assumptions were made: (1) the shape of the cell is circular; and (2) the thickness of other cell wall layers is ignored (since they are usually very thin in the observed G-fibre). Based on these assumptions, the following parameters were calculated:

Mean fibre diameter: FD = (Dr + Dt) / 2

Mean G-layer thickness in a fibre: $GLT = (GLT_{r1} + GLT_{r2} + GLT_{t1} + GLT_{t2}) / 4$

Mean G-layer thickness in a section: $MGLT = \sum GLT / n$ (n = 10 to 12)

G-fibre area: $A_{GF} = (\pi/4) \times FD^2$

G-layer area: $A_{GL} = (\pi/4) \times [FD^2 - (FD - 2 \times GLT)^2]$

Area ratio of G-layer in G-fibre:

$$GLR_{GF} = A_{GL}/A_{GF} = 4 \times (GLT/FD) \times (1 - GLT/FD)$$

This allows the area ratio of the G-layer in the whole section to be estimated:

Area ratio of the G-layer among the total area: $GLR_T = GLR_{GF} \times GFR_T$

RESULTS AND DISCUSSION

Relationship between GS and tissue surface ratios

Table 1 shows the average values of GFR_F , FR and GFR_T for the different samples. In the sample with the GS value of 0.06% or less, none or very few G-layers were observed. The possible existence of a threshold of G-fibre occurrence between 0.06 and 0.08% can be hypothesised. A similar result was obtained by Washusen $et\ al.\ (2003)$ in $Eucalyptus\ globulus$. Another threshold was also observed above 0.15 and 0.19% where almost all fibres were G-fibre. Both thresholds, however, are hypothetical as they would need to be confirmed by other observations.

Jourez et al. (2001) found a lower vessel lumen ratio in tension wood than in opposite wood for poplar and Ruelle et al. (2006) confirmed this observation in 21 tropical species. The present study confirms that this tendency holds within tension wood samples with a different GS since the total fibre ratio (FR) was significantly correlated to GS (at the 0.05 level with a 2-tailed test, r = 0.909). However, this ratio varies in a very narrow range (Table 1) and it appears doubtful that fibre percentage could explain the change in GS. On the other hand, the GFR_T has a significant positive correlation with GS (at the 0.05 level with a 2-tailed test, r = 0.884), as previously observed (Clair et al. 2003; Washusen et al. 2003). In combination with the results of this study, we can presume that fibre ratio does play some role in growth stress generation. However, GFR_T has the most important effect.

Table 1. Average values of GFR_F , FR and GFR_T for different GS values.								
	Sample	GS (%)	$GFR_{r_{r}}(\%)$	FR (%)	GF			

Sample	GS (%)	$GFR_F\left(\%\right)$	FR (%)	$GFR_{T}(\%)$	
1	0.23	≈100.0	75.0	75.0	
2	0.19	≈100.0	74.9	74.9	
3	0.15	69.0	74.1	51.1	
4	0.12	73.8	72.8	53.7	
5	0.08	68.0	73.3	49.9	

Relationships between GS and microscopic features

Table 1 allows us to separate our G-fibre samples into two groups. For samples 1 and 2, where almost 100% of the fibre tissues were identified as G-fibres, GFR_T was approximately 75%, while samples 3 to 5 had a GFR_T close to 50%. However, both groups correspond to large ranges of GS values. Clearly an analysis at a finer scale is required to understand the origins of these GS variations: were more G-layers produced, or different G-layers?

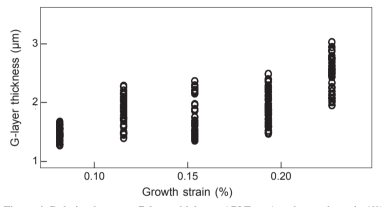


Figure 4. Relation between G-layer thickness (GLT, μm) and growth strain (%).

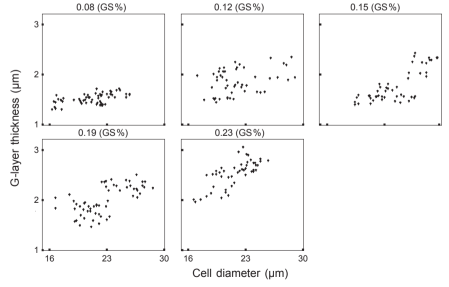


Figure 5. Relation between G-layer thickness (GLT, μm) and cell diameter (FD, μm) for different GS values (%).

The thickness of the G-layers, given here by GLT, is a first approach to quantify the amount of G-layers. Figure 4 shows a considerable scatter of GLT measurements for each GS level, although a positive trend can be observed in the relationship between GLT and GS. Within each sample corresponding to a given GS, GLT was positively correlated with FD, with similar slopes (Fig. 5). This can be explained by GLT variation along a G-fibre. Okumura $et\ al.\ (1977)$ reported that the G-layer is thickest in the mid-region of the fibre and apparently gets thinner toward the tips. Hence it is necessary to control the fibre diameter when comparing GLT. When a partial correlation analysis method controlling cell diameter FD was used, a highly significant positive correlation (r = 0.734, p < 0.001) was found between GS and GLT which indicates that at the same level of cell diameter, thicker G-layer accompanies higher GS. This can be explained by the accumulation effect of each unit of microfibrils. It also confirms that in G-fibres it is the G-layer that plays the major role in the growth stress generation process.

Sample	GS (%)	$GLR_T(\%)$	N	Standard deviation
1	0.23	30.6	48	2.1
2	0.19	23.9	58	2.7
3	0.15	13.3	50	1.4
4	0.12	16.4	50	2.3
5	0.08	13.2	60	1.3

Table 2. Average value of GLR_T for different GS values.

Relationships between GS and G-layer proportion (GLR_T)

Table 2 shows that the GLR_T is significantly correlated to GS (Pearson r = 0.846, p < 0.001) (Fig. 6) and indicates that a higher proportion of G-layer in tension wood produces higher growth stress. The relationship is highly significant and suggests that the amount of G-layer is largely controlling the stress level.

Some inconsistency was observed between samples 3 and 4. As shown in Table 1, Figure 4 and 6, sample 3 has lower GFR_T , thinner mean GLT and lower GLR_T than

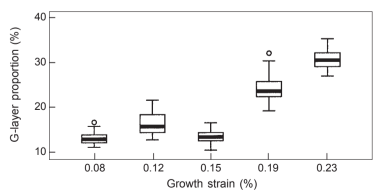


Figure 6. Relation between G-layer area ratio (GLR_T , %) and growth strain (GS, %).

sample 4, but a higher GS. Similarly, Washusen *et al.* (2003) reported that some tissue exhibited high GS with few G-fibres. They explained that it could be attributed to a local heterogeneity in the amount of G-layer.

Some authors have shown differences in cellulose organisation or crystallite size between normal and G-fibre secondary wall (Washusen & Evans 2001; Donaldson 2007; Ruelle *et al.* 2007b); however, these studies did not check if these changes occur in the G-layer of samples having low to high tension wood. Our results show that a change of structure or composition of G-layer is not needed to explain the increase of GS: the amount of G-layer could be sufficient to control the tensile stress level. Recently, Ruelle *et al.* (2007a) showed that crystal size increases with growth stress, even in species not producing tension wood with a G-layer.

CONCLUSIONS

No G-fibres were observed for a GS up to 0.06% while their surface ratio amounted to 50% or more for GS greater than 0.08%, suggesting a hypothetical threshold for G-fibres occurrence between these two GS values. Almost 100% of the fibres contained G-fibres above another hypothetical GS threshold between 0.15 and 0.19%.

In the samples examined, more G-fibres per unit of tissue area and thicker G-layer accompany higher longitudinal growth stress (proportional to GS) in tension wood with G-fibres and suggests that these factors contribute to growth stress generation and therefore the G-layer plays the most important role in high growth stress generation. This may be explained by the hypothesis that the tensile stress of microfibrils governs the longitudinal tensile stress in tension wood (Bamber 1978; Okuyama *et al.* 1986; Bamber 1987; Clair *et al.* 2006a).

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