# Wood properties and chemical composition of the eccentric growth branch of Viburnum odoratissimum var. awabuki 

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#### Abstract

To clarify the wood properties and chemical composition of branches of Viburnum odoratissimum produced by unusual eccentric growth, we investigated growth strain (GS), basic density ( $D_{\mathrm{b}}$ ), microfibril angle (MFA), elastic moduli ( $E_{\mathrm{L}}$ and $E_{\mathrm{L}} / D_{\mathrm{b}}$ ), creep deformation, cellulose crystalline features, and lignin structure in upper and lower sides of the branches, and considered the correlations among these factors. In most measuring positions, the distribution of GS showed that higher tensile GS was in the upper side and compressive GS was in the lower side of the branch, which combines GS features of reaction wood. However, the generation of GS in the lower side was different from that in compression wood, because $E_{\mathrm{L}} / D_{\mathrm{b}}$ and MFA had a negative correlation. The creep compliance curves show that the upper-side wood had low rigidity and high viscosity, whereas the lower-side wood had large rigidity and low viscosity. Relative creep had a negative relation with MFA in the upper side, which is unusual. The cellulose crystalline features showed no obvious difference between both sides of the branch; however, the lignin with


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less $\beta-O-4$ proportion and less $S$ units but more $G$ units seemed to exist in the lower side because of a decreased syringyl/guaiacyl $(S / G)$ molar ratio. This suggests that cell wall could be reinforced by lignin resulting in lower viscosity in the lower side of the branch. Additionally, the $S / G$ ratio showed a relatively high correlation with GS in the lower side. These results suggest that lignin structure plays an important role in adapting to environmental changes during eccentric growth for V. odoratissimum.


Keywords Growth strain • Syringyl/guaiacyl ratio • Creep • Elastic modulus • Cellulose crystallinity

## Introduction

In angiosperms, tension wood is generally induced on the upper side of inclined stems with eccentric growth, and has higher cellulose content than normal wood; in gymnosperms, compression wood is induced on the lower side of inclined stem with eccentric growth, and is characterized by higher lignin content (Onaka 1949; Timell 1986a, b, c). However, some special cases in angiosperms have occasionally been reported. For instance, Buxus microphylla Siebold \& Zucc. shows pronounced growth on the lower side of inclined stems with large compressive stress. Its xylem has some anatomical features similar to those of compression wood (Yoshizawa et al. 1993), and the chemical structure of lignin shows a progressive grading from normal wood to reaction wood (Baillères et al. 1997). Similarly, growth eccentricity of Pseudowintera colorata (Raoul) Dandy occurs on the lower side of the inclined branches and the lower-side tracheids have a large microfibril angle (MFA) (Kucera and Philipson 1977, 1978; Meylan 1981). Regardless of reaction wood or special
cases, wood properties are closely related to chemical composition of wood.

We recently found that inclined branches of Viburnum odoratissimum var. awabuki (K. Koch) Zabel also have pronounced growth on the lower side (Fig. 1). Compared with B. microphylla and $P$. colorata, the anatomy of the branches of $V$. odoratissimum does not obviously differ between the upper and lower sides: the cell wall lacks gelatinous fiber and MFAs show no great differences. Uniquely, a relatively larger growth strain (GS) is commonly observed on the upper side opposite to the eccentric growth side. These results show that the branch of V. odoratissimum forms neither tension wood nor compression wood (Wang et al. 2009). Meanwhile, these results take some questions. For example, how does the branch keep a stably biomechanical situation, and why does eccentric growth occur in the lower side of branch even though $V$. odoratissimum is angiosperm.

In this study, we focused on understanding the abnormal biomechanics of branches exhibiting unusual eccentric growth. The wood properties, such as basic density $\left(D_{\mathrm{b}}\right)$, the elastic modulus in the longitudinal direction $\left(E_{\mathrm{L}}\right)$, creep deformation, and chemical composition of the branch were investigated. First, we measured the released surface GS in seven branches from two $V$. odoratissimum trees, and examined some physical and mechanical properties
( $D_{\mathrm{b}}, \mathrm{MFA}, E_{\mathrm{L}}, E_{\mathrm{L}} / D_{\mathrm{b}}$, creep compliance) of the branch wood. Second, we examined the cellulose crystalline features using attenuated total reflection infrared spectroscopy and X-ray diffraction (XRD) methods, and then analyzed the composition of guaiacyl unit (vanillin) and syringyl unit products comprising lignin by gas chromatography (GC). Finally, the relationship between wood properties and chemical composition in the branches was considered. Correlations among these factors would be advantageous to understand the biomechanics of branches.

## Materials and methods

## Plant material

GS of a V. odoratissimum tree (I) was measured in a previous study (Wang et al. 2009). Materials from this tree were used for chemical analysis. Two more 12-year-old $V$. odoratissimum trees (II and III) growing in Uji campus $\left(130^{\circ} 82^{\prime} \mathrm{E}, 34^{\circ} 81^{\prime} \mathrm{N}-130^{\circ} 80^{\prime} \mathrm{E}, 34^{\circ} 81^{\prime} \mathrm{N}\right)$, Kyoto University, Japan were studied. The seven branches were 210250 cm in length, and were inclined at angles between $50^{\circ}$ and $60^{\circ}$ from the vertical (the angle of branch no. 1 of tree III was $80^{\circ}$ ). After GS measurement, mechanical and chemical experiments were performed.

Fig. 1 Macro photo of the branch no. 3B of tree I and the cross sections stained with the mixed solution of zinc chloride and iodine. There is no G-layer in either side


Growth strain measurement

For trees II and III, GSs in two measuring positions (A and B) of seven branches were measured using a direct method in December 2008. The selection of measuring points was as described in a previous study (Wang et al. 2009). The direct method means that no twigs and branches were cut from the trees; after removing the bark at the measuring position, GS ( $\varepsilon_{\mathrm{r}}$ ) of the branch was directly measured using electrical resistance strain gauges by cutting grooves in the standing trees. The value of $\varepsilon_{\mathrm{r}}\left(\varepsilon_{\mathrm{r}}=\delta \varepsilon_{\mathrm{r}}\right)$ including the influence of self-loading is equal to the strain increment at stress release. Theoretically, values of $\varepsilon_{\mathrm{r}}$ obtained from both the direct and cumulative methods (Yoshida and Okuyama 2002; Wang et al. 2009) approach the real growth stress of the standing tree (Yamamoto et al. 1989; Jullien and Gril 2008).

Creep experiment, microfibril angle, and density
The branches of trees II and III were cut, and samples $\left[25(L) \times 2(T) \times 1(R) \mathrm{mm}^{3}\right]$ at the mid-point of measuring positions A and B were cut from relatively new xylem of the upper and lower sides for corresponding to the results of released surface GS (Fig. 2). These samples were kept in water as green wood. The creep deformation of each green wood was tested using the cantilever condition for 5.7 h . The load was 0.45 N , well within the proportional limits, and the span was $20 \mathrm{~mm} . E_{\mathrm{L}}$ and creep compliance were calculated. MFAs of the air-dried samples [ $20(L) \times 2(T) \times$ $\left.1(R) \mathrm{mm}^{3}\right]$ were measured using an XRD method. Experiments were performed on a 4-circle diffractometer (Oxford Diffraction Gemini S) equipped with a $1,024 \times$ $1,024 \mathrm{CCD}$ camera. $\mathrm{CuK} \alpha$ radiation was generated by an X-ray generator operating at $50 \mathrm{kV}, 25 \mathrm{~mA}$. Diffraction patterns were integrated between $2 \theta=21.5$ and 23.5 along the whole 360 azimuthal interval to plot the intensity diagram of the (200) plane. The average MFA of each sample was estimated using the method proposed by Cave (1966). The $T$ parameter was measured as the half distance between intersections of tangents at inflexion points with the baseline. Finally, the $D_{\mathrm{b}}$ of the sample was expressed on the basis of the oven-dried weight and the swollen volume, and specific Young's modulus ( $E_{\mathrm{L}} / D_{\mathrm{b}}$ ) was calculated.

## Wood powder for chemical analysis

The wood chips of tree I were cut from two measuring positions of four branches, and then wood powder was prepared for chemical component analysis. The samples of trees II and III were cut from the same sample blocks which provided the samples for wood property experiments


Fig. 2 Sample preparation. The samples for wood property experiments and chemical analysis were cut from relatively new xylem (US upper side, $L S$ lower side)
(Fig. 2). The wood powder was extracted with hot water at $60^{\circ} \mathrm{C}$ for 4 h , and successively extracted with benzeneethanol $(1 / 1, v / v)$ for $6 h$.

Attenuated total reflection infrared spectroscopy and X-ray diffraction

Smooth radial-section samples were analyzed using a Spectrum One (PerkinElmer Co.) with ATR model. The angle of incidence was $45^{\circ}$. Attenuated total reflection infrared spectroscopy (ATR-FTIR) spectra were recorded between 4,000 and $400 \mathrm{~cm}^{-1}$ at a $4 \mathrm{~cm}^{-1}$ resolution.

The crystallinity index and crystallite width of cellulose in the wood samples $\left[5(T) \times 1(R) \times 15(L) \mathrm{mm}^{3}\right]$ were measured using the XRD method. XRD was performed using a Rigaku X-ray diffractometer (UltraX 18HF), with $\mathrm{CuK} \alpha$ radiation $(\lambda=0.15418 \mathrm{~nm})$, operated at 40 kV , 300 mA and $2 \theta=5^{\circ}-40^{\circ}$. The crystallinity index was determined from the ratio of the separated crystalline peak area to the total reflection area including the background. The apparent crystallite width in the direction perpendicular to the 200 crystal plane was calculated using the Scherrer's equation (Cutter and Murphey 1972) as follows: $L=0.9 \lambda / H \cos \theta$,
where $\lambda$ is X-ray wavelength, $H$ is the full-width at halfmaximum at the radian, and $\theta$ is the Bragg angle. The Scherrer's equation theoretically applies to a perfect crystal. In reality, the peak broadening, quantified by parameter $H$, is affected by other factors than just the crystal size. Several studies (Burghammer et al. 2003; Kennedy et al. 2007) show that the Scherrer's equation is affected by the disorder within the crystal so it will be preferred to talk about apparent crystal size.

Klason lignin

The wood powder $(0.5 \mathrm{~g})$ was hydrolyzed at $20^{\circ} \mathrm{C}$ with $72 \%$ sulfuric acid ( 10 ml ) for 4 h . The solute was diluted with distilled water until the sulfuric acid concentration was $3 \%$ and then boiled for 4 h . After rinsing and filtration, the oven-dried weight of Klason lignin was measured.

Nitrobenzene oxidation and acetylation
The extractives-free wood powder ( 100 mg ) was mixed with 2 M sodium hydroxide ( 4 ml ) and nitrobenzene $(0.24 \mathrm{ml})$, and heated at $170^{\circ} \mathrm{C}$ for 2 h . After cooling, 2 ml of 5-iodovanillin-dioxane $(4.19 \mathrm{mg} / \mathrm{ml}$ dioxane solution, internal standard) was added to the mixture. The mixture reaction solution was filtered and cleaned with 0.1 M sodium hydroxide, and pH of the rinsing solution was adjusted to $2-3$ with 2 M hydrochloric acid. The acid aqueous solution was extracted with ethyl acetate and washed with saturated sodium chloride solution. The extract solution was successively dried with anhydrous sodium sulfate and the solvent was evaporated off under vacuum.

The dried 16 nitrobenzene oxidation samples from three trees were dissolved in acetic anhydride ( 0.5 ml ) and pyridine $(0.5 \mathrm{ml})$, and placed at $25^{\circ} \mathrm{C}$ for 24 h . Then the solvent of the mixture solution was evaporated off with ethanol under vacuum. The evaporated sample was dissolved in acetone ( 1 ml ), and the solution ( $1 \mu \mathrm{l}$ ) was used to analyze lignin structure by GC measurement.

## Gas chromatography

The product mixture was analyzed by GC-FID (GC-2014, Shimadzu) equipped with a SP-2330 capillary column ( $25 \mathrm{~m} \times 0.25 \mathrm{~mm}, 0.25 \mu \mathrm{~m}$ film thickness). The GC conditions were as follows: injector $250^{\circ} \mathrm{C}$, detector $270^{\circ} \mathrm{C}$, column $230^{\circ} \mathrm{C}$, and temperature program $5^{\circ} \mathrm{C} / \mathrm{min}$. The carrier gas was helium delivered at $30 \mathrm{ml} / \mathrm{min}$. The correlation coefficients of the standard curves for $p$-hydroxybenzaldehyde, vanillin, and syringaldehyde were 0.9871 , 0.9939 , and 0.9964 , respectively. The ratio of syringyl unit to guaiacyl unit ( $S / G$ ) was calculated from the peak areas assigned to syringyl and vanillin type phenols.

## Results and discussion

## Growth strain of branches

Table 1 shows growth eccentricity and GSs of the branches of trees II and III. Growth eccentricity was found on the lower side of all branches except for 1B of tree II. The

Table 1 Growth strains in the branches of trees II and III

| Branch no. | $R_{1}(\mathrm{~mm})$ | $R_{2}(\mathrm{~mm})$ |  | Growth strain $(\mu \mathrm{m} / \mathrm{m})$ |  |
| :---: | :--- | :--- | :--- | :--- | ---: |
|  | Upper radius | Lower radius |  | Upper | Lower |
| Tree I |  |  |  |  |  |
| 1A | 17 | 22 |  | -205 | 16 |
| 1B | 15 | 18 |  | -706 | 433 |
| 2A | 14 | 50 |  | -191 | -326 |
| 2B | 10 | 20 |  | -494 | 23 |
| 3A | 22 | 55 |  | -669 | -53 |
| 3B | 24 | 45 |  | -553 | -57 |
| 4A | 25 | 55 |  | -545 | -212 |
| 4B | 25 | 51 |  | -520 | -152 |
| Tree II |  |  |  |  |  |
| 1A | 11 | 31 |  | -445 | 181 |
| 1B | 15 | 15 |  | -304 | 345 |
| 2A | 12 | 21 |  | -236 | 474 |
| 2B | 12 | 16 |  | -329 | 100 |
| 3A | 11 | 30 |  | -41 | 220 |
| 3B | 18 | 24 |  | -848 | 35 |
| Tree III |  |  |  |  |  |
| 1A | 12 | 36 |  | $-1,395$ | -91 |
| 1B | 18 | 24 |  | -357 | -141 |
| 2A | 8 | 15 |  | -471 | 373 |
| 2B | 9 | 12 |  | -466 | 323 |
| 3A | 14 | 28 |  | -602 | 200 |
| 3B | 12 | 19 |  | -648 | 329 |
| 4A | 12 | 18 |  | -287 | 279 |
| 4B | 12 | 14 |  | -430 | 392 |

The data of tree I is from the previous study (Wang et al. 2009)
distribution of GS showed two patterns. One is tensile stress in the upper side and compressive stress in the lower side (the main pattern in trees II and III); the other is relatively higher tensile stress in the upper side and slight tensile stress in the lower side, like in the case of tension wood occurrence but with smaller GS levels (the main pattern in tree I). Although the $E_{\mathrm{L}}$ (see Table 2) of the lower side was slightly larger than that of the upper side ( $P<0.1$ ) except for tree II-3 and tree III-3, we could calculate that the growth stress in the upper side was greater than that in the lower side. This confirms that in contrast to the most usual pattern in angiosperm species, there was not a larger growth stress in the eccentric growth side of the stem.

According to the above results, we found that the eccentric growth in the lower side of the branch was a normal growth pattern for $V$. odoratissimum tree. The distribution of GS in the branch depended on the individual tree, but was in all cases characterized by a positive difference between the upper and lower sides to fulfill the

Table 2 Basic density, microfibril angle (MFA), elastic modulus in longitudinal direction ( $E_{\mathrm{L}}$ ), and specific modulus ( $E_{\mathrm{L}} / D_{\mathrm{b}}$ ) of the branches

| Branch no. | Basic density ( $\mathrm{g} / \mathrm{mm}^{3}$ ) |  | MFA by XRD ( ${ }^{\circ}$ ) |  | $E_{\mathrm{L}}(\mathrm{GPa})$ |  | $E_{\mathrm{L}} / D_{\mathrm{b}}\left(\mathrm{GPa} \mathrm{mm}{ }^{3} / \mathrm{g}\right)$ |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Upper | Lower | Upper | Lower | Upper | Lower | Upper | Lower |
| Tree II |  |  |  |  |  |  |  |  |
| 1 | 0.51 | 0.57 | 21.5 | 17.8 | 4.69 | 9.94 | 9.20 | 17.4 |
| 2 | 0.48 | 0.51 | 19.6 | 17.9 | 6.25 | 8.57 | 13.0 | 16.8 |
| 3 | 0.52 | 0.59 | 23.4 | 22.3 | 6.06 | 5.80 | 11.7 | 9.83 |
| Tree III |  |  |  |  |  |  |  |  |
| 1 | 0.52 | 0.58 | 19.2 | 20.3 | 5.91 | 7.51 | 11.4 | 13.0 |
| 2 | 0.47 | 0.51 | 16.5 | 17.9 | 6.18 | 7.50 | 13.1 | 14.7 |
| 3 | 0.48 | 0.53 | 15.5 | 18.8 | 6.28 | 6.16 | 13.1 | 11.6 |
| 4 | 0.51 | 0.48 | 17.7 | 20.4 | 4.63 | 5.05 | 9.08 | 10.5 |

biomechanical requirements. This suggests that the mechanism responsible for eccentric growth of branch in $V$. odoratissimum trees is different from that for eccentric growth of branch in reaction wood of usual angiosperms species.

Physical and mechanical properties of branch wood
The $D_{\mathrm{b}}$, MFA, $E_{\mathrm{L}}$, and $E_{\mathrm{L}} / D_{\mathrm{b}}$ of the branch wood in trees II and III are shown in Table 2. Because the lower-side wood had a large basic density, except for tree III-4, that can induce an increased elastic modulus of wood, the effect of density on $E_{\mathrm{L}}$ is shown in Fig. 3. We found that there was not a distinct relationship between both factors. Moreover, Fig. 4 shows the relationships between $E_{\mathrm{L}} / D_{\mathrm{b}}$ (relating to the mean modulus), $E_{\mathrm{L}}$ (including the effect of cell-wall thickness) and MFA. In this case, there was no great difference in two relationships because the range of density was small. This also proves that the mechanical property of branch wood is not influenced by density. Meanwhile, the relationship between MFA and $E_{\mathrm{L}} / D_{\mathrm{b}}$ was different in trees II and III: a clearly negative correlation in tree II ( $R^{2}=0.75$ ), whereas a weak correlation in tree III ( $R^{2}=0.09$ ), which may be due to branch architecture of individual tree. However, for trees II and III, the $E_{\mathrm{L}} / D_{\mathrm{b}}$ of the lower side of the branch has a negative relation with MFA ( $R^{2}=0.71$ ). This means that the generation of compressive GS in lower side is different from compression wood, because compression wood had a low $E_{\mathrm{L}} / D_{\mathrm{b}}$ that obliges the tree to be denser in order to compensate for the lower cell-wall modulus resulting from high MFA.

Then, we investigated the viscoelastic property of the branch wood at green condition. Figure 5 shows the creep curves of green wood cut from the upper and lower sides of the branches. The creep compliance $\left(J_{\mathrm{t}}\right)$ of most upper-side samples was larger than that of the lower-side samples, suggesting that the upper-side sample had lower rigidity and higher viscosity. Higher creep is usually related to


Fig. 3 Relationship between the basic density and $E_{\mathrm{L}}$ in the green branch wood
higher MFA. However, in Fig. 6, effect of MFA on relative creep $\left(J_{10,000} / J_{10}-1\right)$ showed a unique result, that is, there is a negative relationship between both factors in the upper side, which is in contradiction with expectation.

After considering GS and the wood properties of the branch, we found that the branch of V. odoratissimum has some features similar to reaction wood. The branch showed a distribution of GS similar either to tension wood (larger tensile growth stress in the upper side), or to compression wood (compressive growth stress in the lower side). Eccentric growth did not occur in the upper side, where no anatomical feature (for example, G-layer) of tension wood was visible. Although no particular anatomical feature was observed on the lower side either, from a biomechanical viewpoint of the generation of compression stress, in the case of the GS pattern (dominant in trees II and III), is consistent with the direction of eccentric growth and both are typical features of compression wood of gymnosperms, but a much lower specific modulus $\left(E_{\mathrm{L}} / D_{\mathrm{b}}\right)$ associated with a higher MFA. The higher rigidity and lower fluidity of the lower side acts also consistently with the biomechanical requirement of the branch, which would not be necessarily

Fig. 4 Relationships between $E_{\mathrm{L}}, E_{\mathrm{L}} / D_{\mathrm{b}}$ and MFA in trees II (hollow mark) and III (solid mark)




Fig. 5 The creep compliance curves of green wood cut from the upper and lower sides of branches (solid mark the upper side; hollow mark the lower side)


Fig. 6 Relationship between the relative creep $\left(J_{10,000} / J_{10}-1\right)$ and MFA in trees
characterized by a higher density like in the case of compression wood. Although negative GSs were measured, especially in tree I, their level was not sufficient to indicate compression wood. Moreover, the MFA values and their negative correlation with creep show that this branch wood does not correspond to usual trend of normal wood versus reaction wood. Therefore, it is necessary to consider the influences of chemical composition and matrix structure of cell wall on viscoelastic property of the branch.

Cellulose crystalline features in branches

The ATR-FTIR spectra of branch (tree I-3B) are shown in Fig. 7. Peaks of the spectra showed the typical characteristic of hardwood. Peak heights of $1,734 \mathrm{~cm}^{-1}$ for unconjugated $\mathrm{C}=\mathrm{O}$ in xylans and $1,505 \mathrm{~cm}^{-1}$ for aromatic skeletons in lignin (Pandey 1999; Pandey and Pitman 2003) were not obviously different between both sides of the branch. The same result was also obtained from other branches.

Table 3 shows the crystallinity index and the apparent crystallite width in the branches exhibiting unusual eccentric growth. The development of crystallinity in the lower side due to eccentric growth was expected because crystallinity influences the mechanical properties of the cellulose fiber (Lee 1961). In fact, although the crystallinity index in the lower side seems to be less than that in the upper side for most of the samples, there was no significant


Fig. 7 ATR-FTIR spectra of the branch (tree I-3B) for $V$. odoratissimum
difference, while the apparent crystallite width had no change in both sides of the branches. Therefore, although the lower side of the branch had a larger $E_{\mathrm{L}}$ in $V$. odoratissimum, this property is not explained by cellulose crystallinity.

In addition, Washusen and Evans (2001) reported that tension wood has wider crystallites than normal wood in Eucalyptus globulus. The apparent crystallite width was less in compression wood than in the opposite wood of Japanese red pine (Tanaka et al. 1981). Thus, it is generally believed that crystallinity changes with eccentric growth. Compared with these cases, in V. odoratissimum, the lower side of the branch, which exhibits eccentric growth, did not obviously have different crystalline features, confirming that growth eccentricity in V. odoratissimum is not associated with reaction wood production.

## Lignin structure in branches

Lignin is an important chemical composition of wood. Its content is $25-35 \%$ in softwood and $17-25 \%$ in hardwood (Tsoumis 1991). The lignin content in the branches of $V$. odoratissimum is shown in Table 3. The Klason lignin content was about $20-30 \%$ in both sides, depending on the individual tree, exhibiting a good lignification. Moreover, Klason lignin had a slight but significant distribution pattern in the branch. That is, the Klason lignin content in the lower side was higher than that in the upper side of the branch except for the branch no. 1 of tree III ( $P<0.05$ ). This distribution pattern also suggests that the upper side should have higher content of cellulose. Although this
trend is consistent with what is observed in the case of reaction wood occurrence, either tension wood in the upper side or compression wood in the lower side, when the relationship between the lignin content and GS was considered (Fig. 8), a relatively weak correlation between both factors in the lower side was observed. In fact, the branches did not display a larger compressive stress in the lower side or a remarkable tensile stress in the upper side as in reaction wood.

Furthermore, lignin is described as a three-dimensional structure formed by phenylpropane units. In general, softwood is rich in guaiacyl $(G)$ units while guaiacyl and syringyl $(S)$ units are present in hardwood. The p-hydroxyphenyl $(H)$ unit is typical of compression wood lignin (Timell 1986a). Yoshida et al. (2002a) reported that the $S / G$ ratio increases with tensile released strain in yellow poplar that does not form G-layer. Similar results were obtained in black locust (Yoshida et al. 2002b) and eucalyptus (Baillères et al. 1995; Aoyama et al. 2001). To investigate the effect of $S / G$ ratio on GS in $V$. odoratissimum, the lignin structure was analyzed using GC measurement.

The relative molar distributions of $S / G$ in the upper and lower sides of the branches are summarized in Table 3. Although eccentric growth occurs on the lower side of the branch, the H unit was not detected in either side, implying that growth eccentricity in V. odoratissimum is different from that in compression wood and Buxus wood (this result should be also checked using thioacidolysis). $S$ and $G$ units were widely present in the branches, indicating that relatively reactive ether bond $(\beta-O-4)$ linked $G$ and $S$ units is a

Table 3 The factors of chemical structures in the branches of V. odoratissimum

| Branch no. | Crystallinity <br> index (\%) |  | Crystallite width ( nm ) |  | Klason lignin content (\%) |  | Total yield of products ( $\mathrm{S}+\mathrm{G}, \mu \mathrm{mol} / \mathrm{g}$ of Klason lignin) |  | $S / G$ ratio |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Upper | Lower | Upper | Lower | Upper | Lower | Upper | Lower | Upper | Lower |
| Tree I |  |  |  |  |  |  |  |  |  |  |
| 1A | 43.0 | 36.1 | 4.0 | 4.0 | 26.5 | 27.8 | 3,487 | 3,366 | 2.24 | 1.41 |
| 1B | 35.7 | 35.3 | 4.0 | 4.0 | 25.3 | 26.9 | 3,956 | 2,340 | 1.86 | 1.63 |
| 2 A | 36.7 | 36.1 | 4.0 | 4.0 | 27.1 | 28.8 | 2,291 | 1,701 | 1.78 | 1.27 |
| 2B | 35.0 | 38.1 | 4.0 | 4.0 | 26.7 | 28.3 | 2,035 | 2,230 | 1.94 | 1.56 |
| 3A | 32.9 | 33.9 | 4.0 | 4.0 | 27.2 | 28.2 | 2,610 | 2,062 | 1.94 | 1.27 |
| 3B | 34.4 | 33.1 | 4.0 | 4.0 | 26.5 | 28.9 | 3,593 | 2,353 | 2.03 | 1.44 |
| 4A | 32.1 | 36.6 | 4.0 | 4.0 | 27.7 | 29.1 | 3,517 | 2,298 | 1.50 | 1.06 |
| 4B | 36.1 | 34.7 | 4.0 | 4.0 | 27.8 | 29.9 | 3,226 | 3,191 | 1.94 | 1.13 |
| Tree II |  |  |  |  |  |  |  |  |  |  |
| 1 | 35.2 | 32.4 | 4.0 | 4.0 | 21.0 | 23.8 | 2,566 | 1,963 | 2.33 | 1.30 |
| 3 | 35.3 | 31.9 | 4.0 | 4.0 | 19.9 | 21.4 | 2,758 | 1,988 | 2.57 | 1.49 |
| Tree III |  |  |  |  |  |  |  |  |  |  |
| 1 | 34.7 | 31.9 | 4.0 | 4.0 | 25.3 | 25.1 | 1,674 | 1,407 | 2.57 | 1.33 |
| 3 | 33.1 | 31.0 | 4.0 | 4.0 | 25.4 | 26.0 | 1,654 | 1,438 | 2.03 | 1.56 |



Fig. 8 Relationship between lignin content and growth strain. GS values of tree I are from the previous study (Wang et al. 2009). The GSs of the branch of trees II and III are average values of two measuring positions
structural feature in $V$. odoratissimum common to angiosperms. By comparing the total yield in both sides, we observed that the yield of products in the lower side was less than that in the upper side except for tree I-2B ( $P<0.05$ ). This result suggests that the $\beta-O-4$ structure, which is the noncondensed-type, is less common in the lower side of the branch. The varying distribution of $S / G$ shows the following trend: $S$ units decreased and $G$ units increased in the lower side of the branch, that is, the $S / G$ ratio in the eccentric growth side (lower side) was lower than that in the normal growth side (upper side).

These results suggest that the quality of lignin in the branch is different due to the proportion of $\beta-O-4$ and $S / G$ ratio. Less $\beta-O-4$ proportion, and less $S$ units but more $G$ units seem to exist in the lignin of the lower side, whereas more $\beta-O-4$ proportion, and more $S$ units but less $G$ units seem to be present in the lignin of the upper side of the branch, suggesting that cell wall is reinforced by different type of lignin from that of the upper side during lignification in the lower side because lignin serves as glue to stiffen the matrix structure of cell wall. Hence, the mechanical properties of the branch seem to be controlled by the lignin structure. Particularly, regard as a compressive GS in the lower side, the lower $S / G$ ratio can replace a low $E_{\mathrm{L}} / D_{\mathrm{b}}$ of compression wood, and is more efficient. Therefore, the lower side of the branch had higher rigidity and lower viscosity to sustain downward bending due to the gravity.

Then, the effect of $S / G$ ratio on the GS of the branches was considered. A general trend is found in Fig. 9: higher $S / G$ ratio with tensile GS and lower $S / G$ ratio with mostly compressive GS. This is consistent with usual trend between tension wood and normal wood (Baillères et al. 1995; Aoyama et al. 2001; Yoshida et al. 2002a). Furthermore, we found that GS decreased with a decreased $S / G$ ratio in the lower side of the branch. There was a


Fig. 9 Relationship between $S / G$ ratio and growth strain. GS values of tree I are from the previous study (Wang et al. 2009). The GSs of trees II and III are average values of two measuring positions of each branch
relatively high correlation $\left(R^{2}=0.52\right)$ between GS and $S / G$ ratio in the lower side. It seems that the matrix of cell wall reinforced by the lignin with less $\beta-O-4$ proportion and smaller $S / G$ ratio resists growth stress in the lower side of the branch. In the upper side, the lignin with more $\beta-O-4$ proportion and larger $S / G$ ratio was more common so that the amount of GS was not restricted by lignin structure. Thus, for most of the branches in V. odoratissimum, a larger GS occurs in the upper side opposite to the eccentric growth side. These results indicate that the rigid lower side would restrain the downward deformation of the upper side of the branch.

## Conclusion

Although eccentric growth is found in the lower side of the branches of $V$. odoratissimum, the distribution of GS combined the characters of tension wood (tensile GS in the upper side) and compression wood (compressive GS in the lower side) without any anatomical features of reaction wood. However, the lower-side wood has not a lower $E_{\mathrm{L}} / D_{\mathrm{b}}$ to compensate for the low cell-wall modulus resulting from high MFA, like in the case of compression wood. The upper-side wood had low rigidity and high viscosity, whereas the lower-side wood had large rigidity and low viscosity. Relative creep of the upper-side wood had a negative relation with MFA, which is in contradiction with usual belief. The cellulose crystalline features in the lower side were not significantly different from those in the upper side. On the other hand, the lignin structure showed a clear distinction between both sides of the branch, that is, $S / G$ ratio decreased in the lower side, suggesting that the lignin with less $\beta-O-4$ proportion, and less $S$ units but more $G$ units exists in the lower side, and greatly influenced the GS of the branch. It seems that the matrix structure of cell wall of the
lower side become stiffer due to an increased $S / G$ ratio. Therefore, the chemical properties of lignin may control the amount of GS and the mechanical properties of the branches. These results also indicate that $V$. odoratissimum has different growth biomechanics to other woody plants in order to adapt to changes in the environment.

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