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Paulo Ricardo Gherardi Hein, Jean-Marc Bouvet, Eric Mandrou, Philippe Vigneron, Bruno Clair, et al.. Age trends of microfibril angle inheritance and their genetic and environmental correlations with growth, density and chemical properties in *Eucalyptus urophylla* ST Blake wood. *Annals of Forest Science*, 2012, 69 (6), pp.681 - 691. 10.1007/s13595-012-0186-3 . hal-02644062

HAL Id: hal-02644062

<https://hal.inrae.fr/hal-02644062>

Submitted on 28 May 2020

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Age trends of microfibril angle inheritance and their genetic and environmental correlations with growth, density and chemical properties in *Eucalyptus urophylla*

S.T. Blake wood

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Received: 21 April 2011 / Accepted: 20 January 2012 / Published online: 24 February 2012
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Abstract

• **Context** The genetic and environmental control of microfibril angle (MFA) and its genetic correlations with other wood and growth traits are still not well established in *Eucalyptus* sp.

• **Aims** To determine the narrow-sense heritability estimates (h^2) of MFA, wood density (D), Klason lignin (KL) content, syringyl to guaiacyl (S/G) ratio and growth traits, their variation from pith to cambium and their genetic correlations.

Handling Editor: Barry Alan Gardiner

Contribution of the co-authors PRGH was responsible for performing the near infrared (NIR), X-ray diffraction analysis and wood measurements, for generating the phenotypic data set, analysing the results and the writing of the paper. This study is a part of his PhD thesis;

JMB was responsible for planning the experimental design and performing the statistical and all the genetic analysis. He co-writes the paper with PRGH.

EM wrote part of the Material and methods section.

PV was responsible for planning the experimental design and part of the preliminary analysis.

BC was responsible for performing the X-ray diffractometry analysis at Univ. of Montpellier 2. He gave valuable insights to the study and he help to improve part of the discussions.

GC was responsible for planning the measurements and analysing the results. He's co-supervisor of PRGH PhD. He provides comments and corrections to the text.

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• **Methods** Heritability and correlations were assessed in 340 control-pollinated progenies of 14-year-*Eucalyptus urophylla* S.T. Blake using near infrared spectroscopic models.

• **Results** Moderate to high heritability were found for MFA ($h^2=0.43$), D ($h^2=0.61$), S/G ($h^2=0.71$) and LK ($h^2=0.76$). The genetic control of D and MFA and the genetic and residual correlation between chemical and growth traits varied with age. The genetic correlation C×D was always strongly negative ($r<-0.80$) while the correlation D×MFA remained constant and positive in the juvenile wood ($r=0.7$), before disappearing in the mature wood. These results could be explained by gene pleiotropic effect, low microfibril angle compensating for low wood density and fast growth or by linkage disequilibrium induced by sampling. Variations in MFA and KL in the mature wood were also genetically controlled.

• **Conclusions** These findings provide the opportunity for developing breeding strategies for pulpwood, fuelwood and sawntimber production in *Eucalyptus* sp.

Keywords Variance components · MFA · Klason lignin · Syringyl to guaiacyl ratio · Factorial mating design · Wood phenotyping · NIR spectroscopy

1 Introduction

Eucalyptus is one of the most widely cultivated hardwood genera in tropical and subtropical regions of the world. This genus is adapted to a variety of climatic and edaphic conditions and grows at high rates producing raw material adequate for many end uses. Wood quality is determined from the combination of intrinsic factors such as stiffness that depend on microfibril angle (MFA), density and the chemical contents of its main components (Kollmann and Côté 1968). A range of studies have pointed out the importance of microfibril angle for the wood stiffness and wood products quality in *Eucalyptus* sp (Evans and Ilic 2001). In general, wood in which the MFA is low has high rigidity and high economic value. Hence, the determination of the genetic factors contributing to quantitative trait variation of wood properties is essential for

tree breeders (Raymond 2002; Apiolaza 2009) allowing the selection of trees with adequate characteristics for the proposed end use. Furthermore, adverse genetic and environmental correlations between wood quality and growth traits remain as one of the main constraints in tree breeding programmes for pulp and paper, bio-energy or sawnwood industries.

Growth, cellulose content, and wood density (or pilodyn penetration, which is an indirect method for determining wood density; see Raymond and MacDonald (1998) are the main traits considered by tree breeders for *Eucalyptus* pulpwood plantations and many papers have reported heritability estimates and correlations among these properties (Greaves et al. 1997; Raymond 2002; Costa e Silva et al. 2009). However, the genetic parameters of chemical, mechanical and ultrastructural features are rarely reported in *Eucalyptus*. Moreover, the estimation of genetic parameters of these wood traits presents a large variation according to the peculiarities of each study. For instance, the findings reported by Kube et al. (2001) in *Eucalyptus nitens* and Apiolaza et al. (2005) and Poke et al. (2006) in *Eucalyptus globulus* are contrasting, demonstrating that heritability estimates and genetic correlations are specific to the population and site under investigation, but also that they have to be considered with caution due to the low accuracy of their estimation.

With the growing need and interest in establishing plantations for producing stiff and dimensionally stable wood with minimal growth stresses, recent studies have focused on the heritability of traits controlling wood quality (Baltunis et al. 2007). Most of the studies dealing with the genetic control of MFA have been carried out on softwoods because of the generally low stiffness and poor dimensional stability of the juvenile wood. In hardwoods, the degree to which MFA is heritable and its genetic relationship with other traits has not often been reported, especially for *Eucalyptus* wood. To our knowledge, there is only one study reporting narrow-sense heritability estimates (h^2 ; Apiolaza et al. 2005) and another study providing broad-sense heritability estimates (H^2) for MFA in *Eucalyptus* (Lima et al. 2004). References reporting genetic and environmental correlations among MFA, density, lignin, syringyl to guaiacyl ratio and growth traits are rare in *Eucalyptus*. The extent to which microfibril angle is controlled by additive genetic and environmental factors is poorly documented; and the relationship between MFA and the other wood and growth traits, and how these relationships vary with age are still not established in *Eucalyptus*. Knowledge about these issues would be useful to understand how trees adapt their wood traits in order to maintain their erect habit even when they are subjected to bending movements in response to wind and gravity.

Therefore, the main aims of this study were: (1) to assess the level of genetic control of MFA, wood density (D), Klason lignin content (KL), and syringyl to guaiacyl ratio (S/G) and growth traits from a 14-year-old control-pollinated progeny

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test of *Eucalyptus urophylla*; (2) to determine the genetic and residual correlations among MFA, D, KL, S/G and growth traits; and (3) to investigate the age trends of these genetic parameters.

Most studies on genetic parameters of *Eucalyptus* wood traits have been based on averaged values per tree. Because MFA and wood density vary greatly from pith to cambium (Kollmann and Côté 1968) assessing averaged values may not be the best option. In this study, MFA and wood density were assessed in three radial positions namely early juvenile wood, late juvenile wood and early mature wood. Radial variation of genetic parameters and the genetic and residual correlations among these wood traits are presented and the implications for selection are discussed.

2 Material and methods

2.1 Genetic material

Three hundred and forty wood discs were collected at breast height from a 14-year-old *E. urophylla* S.T. Blake progeny trial stand in Pointe Noire, Republic of Congo, in the experimental area of the “Centre de Recherche sur la Durabilité et la Productivité des Plantations Industrielles–CRDPI” (04°45'S, 12°00'E; altitude, 50 m). The climate is tropical humid with a mean annual temperature of 24°C, a mean annual rainfall of 1,200 mm and a dry season from May to October. This progeny trial was established in 1992. It was composed of 33 full-sib families produced by controlled pollination using an incomplete factorial mating design (ratio 33/64) involving 16 parents (eight males and eight females) originally from two provenances (eight from Mont Egon and eight from Mont Lewotobi of the Flores Island in the Sunda Archipelago, 122–127°E, 8–10°S). In 2006, nine to ten trees from each family were selected with the best growth and stem straightness and harvested, resulting in a total of 340 sample trees. Trees were collected from only one block to limit the impact of destructive sampling in the entire trial. This solution did not affect the genetic analysis capacity as the number of trees sampled was high and the harvested block presented very homogenous soil conditions. The circumference at breast height (C) and the height (H) of the trees were measured at 14 years, immediately before harvesting. After harvesting, discs of wood (~30 mm thick) were obtained at breast height of each tree and transported to “Centre de Coopération Internationale en Recherche Agronomique pour le Développement” (CIRAD) in Montpellier, France where the analyses were performed.

2.2 Sample preparation

Defect-free wedges were cut from the discs and milled using a knife mill (Retsch GmbH, Haan, Germany; model SM100)

and an ultracentrifugal mill (Retsch GmbH, Haan, Germany; model ZM 200) in order to produce grounded wood (particle sizes lower than 0.5 mm) for wet-chemistry analysis (Fig. 1, sample a). Pith to bark radial strips (defects free) were removed from the discs using vertical bandsaw machine at random azimuthal directions. Then, small wood samples measuring ~20–30 mm $R \times$ ~20 mm $T \times$ ~30 mm L (Fig. 1, sample b) and tangential sections (Fig. 1, sample c) measuring ~2 mm $R \times$ ~20 mm $T \times$ ~30 mm L were cut, parallel to the growth rings, from each wood strip, for wood density and microfibril angle measurements, respectively. The wood powders of 60 samples were subjected to chemical analysis. In order to select 60 samples from the total sampling, principal component analysis of the NIR spectra was performed using the Unscrambler (CAMO AS, Norway) software version 9.7. Samples well representative in terms of NIR spectra were selected according to their Mahalanobis distances. The samples (milled and solid woods) were kept in a conditioned room at 50% relative humidity and temperature of 20°C before analysis. Under these conditions, the equilibrium moisture content was around 10%.

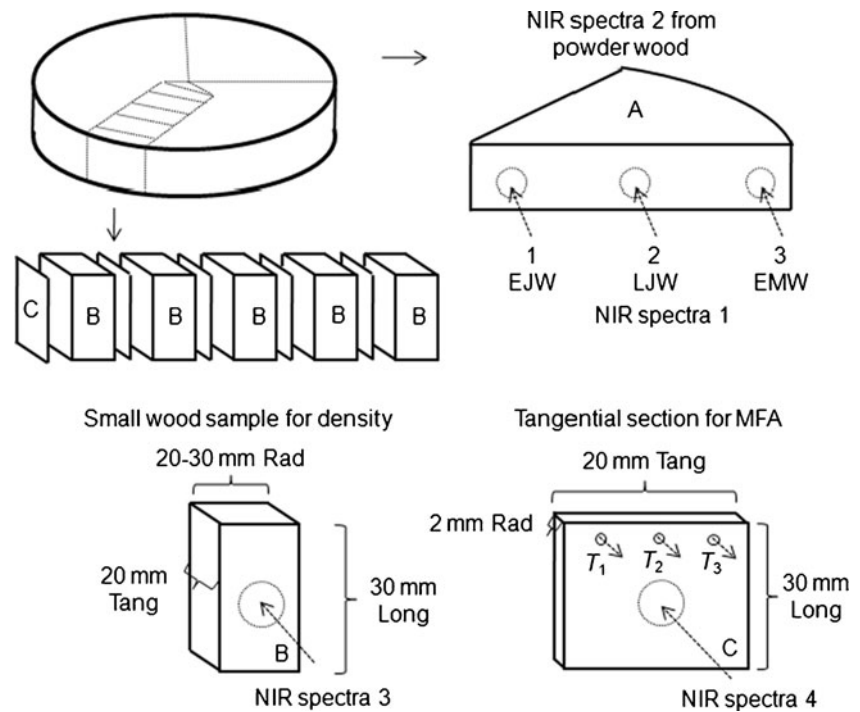
2.3 Wood phenotyping

The KL content and ratio between syringyl (S) and guaiacyl (G) of the wood were determined by the Biological Chemistry Laboratory (INRA-Agro ParisTech, France). The analyses were performed in duplicate for the 60 samples and were previously reported in Hein et al. (2010a). The basic D was determined according to water immersion procedure in 190 small wood samples (~20×25×30 mm) cut from radial strips as previously reported in Hein et al. (2009). All X-ray diffraction data were collected on a diffractometer (Xcalibur-I, Oxford Instruments, USA) with CuK α radiation at “Institut Européen des Membranes” of the University of Montpellier, France. Three X-ray diffraction profiles were recorded and averaged for 175 tangential sections (2×20×30 mm) cut from radial strips. The error of the measure was estimated at 3%. The procedure for microfibril angle measurements was previously reported in Hein et al. (2010b).

2.4 Near-infrared spectroscopy analysis

The wood traits under investigation in this study were based on predictions from NIR spectroscopy calibrations. NIR spectra were recorded using a spectrophotometer (model Vector 22/N, Bruker Optik GmbH, Ettlingen, Germany) on both milled and solid woods. First, NIR spectra were recorded on the radial surface of the wedges (NIR spectra 1) at three radial positions (1, 2, and 3) from pith to bark and on the wood powders (NIR spectra 2) and, for predictions (Fig. 1, sample a).

Fig. 1 Procedure of sample preparation for wood characterization. Wedges (A) and radial strips were cut from discs; small wood samples (B) and tangential sections (C) were cut for density and microfibril angle measurements. Near-infrared (NIR) spectra 1, 3 and 4 were recorded from the early juvenile wood, late juvenile wood and early mature wood. The wedges were ground for wet-chemistry analysis and NIR spectra were measured from the powders (NIR spectra 2)



Point 1 represented the wood formed at ~3–4 years (hereafter called early juvenile wood; point 2, the wood of the ~6–7 years (late juvenile wood); and point 3, the wood of the ~12–14 years (early mature wood). The averaged value per tree for density and microfibril angle are expressed as D and MFA, respectively, while the local D and MFA estimates are expressed as D_1 , D_2 , and D_3 and MFA_1 , MFA_2 , and MFA_3 , accordingly. A single, averaged value per tree was used for the chemical properties. We considered as outliers the samples

Table 1 Mean, minimum and maximum values, coefficient of variation (CV) and number of observations (N) for growth (circumference, C and height, H), chemical properties (Klason lignin, KL and syringyl to guaiacyl ratio, S/G), wood density (D) and microfibril angle (MFA) of the 14-year-old *E. urophylla* population

Trait	Unit	N	Mean	Minimum	Maximum	CV _P (%)
C	cm	340	52.7	24.0	84.0	21.2
H	m	340	21.2	7.8	29.6	17.3
KL	%	321	28.0	25.1	31.7	4.4
S/G	—	321	2.3	1.5	3.5	13.5
D_1	kg m ⁻³	274	443	330	607	10.2
D_2	kg m ⁻³	274	522	391	698	8.4
D_3	kg m ⁻³	274	615	480	761	8.1
D	kg m ⁻³	274	526	423	654	7.2
MFA_1	Degrees	274	18.1	12.7	23.9	10.2
MFA_2	Degrees	274	15.3	10.0	21.1	12.4
MFA_3	Degrees	274	13.3	7.2	19.5	13.9
MFA	Degrees	274	15.6	11.0	20.0	9.8

which presented estimates with high standard errors, and removed them from the original dataset (Table 1).

Partial least square regressions (PLS-R) were developed from the NIR spectra of the wood powders and the chemical analysis of the 60 samples for KL and S/G ratio. These PLS-R calibrations were previously described in Hein et al. (2010a) providing the estimates of single values of KL content and S to G ratio using the NIR spectra 2 taken from the grounded wood of the 340 discs. The PLS-R model for D was based on 190 small wood samples and their NIR spectra 3 (Hein et al. 2009) while the PLS-R model for MFA was based on 175 tangential sections and their NIR spectra 4 (Hein et al. 2010b). These established PLS-R models were applied on the NIR spectra 1 providing the estimates of MFA and wood density in the early and late juvenile wood and in the early mature wood. Table 2 presents fit statistics for the NIR-based models used for phenotyping the woods from this progeny trial.

2.5 Statistical methods for genetic parameter estimation

C, H, KL, S/G, D and MFA were analysed independently to estimate the variance components by using the following mixed linear model:

$$y = Xb + ZQg + Za + e$$

where, y is the vector of observations; b is the vector of fixed effects (in our case the mean value of the trait in the population); g is the vector of fixed effects related to parent groups; a is the vector of genetic effects (individual additive genetic values), e is the vector of residuals; and X , Q and Z

Table 2 Partial least square regression models for estimating the microfibril angle (MFA), wood density (D), Klason lignin (KL) content and syringyl to guaiacyl ratio (S/G) from NIR spectra of *E. urophylla* wood

Trait	R^2_c	RMSEC	R^2_p	RMSECV	RPD
MFA (degrees)	0.72	0.73	0.64	0.84	1.76
D (kg m ⁻³)	0.89	27.0	0.85	30.0	2.70
KL (%)	0.88	0.44	0.85	0.55	2.58
S to G ratio (S/G)	0.92	0.01	0.86	0.13	2.68

R^2_c coefficient of determination of calibration, R^2_p coefficient of determination of validation, $RMSEC$ root mean standard error of calibration, $RMSECV$ root mean standard error of validation, RPD ratio of performance to deviation

are the incidence matrices linking observations to the effects.

This model takes into account the appurtenance of the parent trees to two different groups (provenances) with the incidence matrix Q which links the individuals to the ancestors (see the details of the model in Mrode and Thompson (2005).

The random effect fits a normal distribution whose parameters were:

$$E \begin{bmatrix} a \\ e \end{bmatrix} = \begin{bmatrix} 0 \\ 0 \end{bmatrix} \text{ and } Var \begin{bmatrix} a \\ e \end{bmatrix} = \begin{bmatrix} G & 0 \\ 0 & R \end{bmatrix}$$

The variance-covariance matrices were defined as follows:

$$G = A \cdot \sigma^2_A \text{ and } R = I \cdot \sigma^2_e$$

where, A is the additive genetic relationship matrix computed from a pedigree file that takes into account all the relationships between the individuals, I is the identity matrix, σ^2_A the additive genetic variance and σ^2_e the residual variance. The variances associated with random effects were estimated by restricted maximum likelihood (REML method) using ASReml version 2 (Gilmour et al. 2005).

To compare two (or more) models, we can evaluate the Akaike Information Criteria (AIC; Akaike 1974) or the Bayesian Information Criteria (BIC; Schwarz 1978) for each model. These are given by

$$AIC = -2lR_i + 2t_i \\ BIC = -2lR_i + t_i \log v$$

where, lR_i is the REML log-likelihood of the model, t_i is the number of variance parameters in model i and v is the residual degrees of freedom. AIC and BIC were calculated for each model and the model with the smallest value was chosen as the preferred model.

As the variances are assumed to be independent, the total phenotypic variance σ^2_P was calculated as follows:

$$\sigma^2_P = \sigma^2_A + \sigma^2_E$$

The narrow-sense heritability estimates were calculated as follows:

$$h^2 = \frac{\sigma^2_A}{\sigma^2_P}$$

Variances are not independent of the scale and the mean of the respective traits. Therefore, to compare the genetic and phenotypic variances of the different traits, the genetic (CV_A), residual (CV_E) and phenotypic (CV_P) coefficient of variation were calculated as:

$$CV_{Aj} = \frac{100 \times \sigma_{Aj}}{\bar{x}} \\ CV_{Ej} = \frac{100 \times \sigma_{Ej}}{\bar{x}} \\ CV_{Pj} = \frac{100 \times \sigma_{Pj}}{\bar{x}}$$

where, σ_{Aj} is the square root of the additive genetic variance for the trait, σ_{Ej} is the square root of the residual variance for the trait, σ_{Pj} is the square root of the phenotypic variance for the trait and \bar{x} is the population mean for the trait.

The estimate of phenotypic (r_P), residual (r_E) and genetic additive (r_A) correlations between two traits (X and Y) were performed from a bi-variate analysis using the same individual tree model as for univariate analysis. The variance-covariance matrixes G and R changed and were respectively the Kronecker product of a matrix related to the genetic (residual) effect (A and I) with a genetic (residual) variance-covariance matrix between the two traits (Gt and Rt).

$$G = A \otimes Gt \quad Gt = \begin{bmatrix} \sigma^2_{Ax} & Cov_A(x, y) \\ Cov_A(x, y) & \sigma^2_{Ay} \end{bmatrix} \\ G = I \otimes Rt \quad Rt = \begin{bmatrix} \sigma^2_{Ex} & Cov_E(x, y) \\ Cov_E(x, y) & \sigma^2_{Ey} \end{bmatrix}$$

r_P , r_E and r_A were estimated as follows:

$$r_P = \frac{Cov_P(x, y)}{\sigma_{Px} \cdot \sigma_{Py}} \\ r_E = \frac{Cov_E(x, y)}{\sigma_{Ex} \cdot \sigma_{Ey}} \\ r_A = \frac{Cov_A(x, y)}{\sigma_{Ax} \cdot \sigma_{Ay}}$$

Standard errors of h^2 , σ^2_A , σ^2_P , r_P , r_E and r_A were calculated with ASReml using a standard Taylor series approximation (Gilmour et al. 2005).

3 Results

The descriptive statistics of the wood traits of the 14-year *E. urophylla* are presented in Table 1. As expected for most woods, the MFA were, on average, higher near the pith

decreasing towards the bark and an opposite trend was observed for the basic density of wood (Table 1).

The coefficients of variation were similar to those observed in previous studies on *Eucalyptus* in the Congo (Bouvet et al. 1999). They were higher for growth than for wood property traits in most of the cases (Table 3). Their magnitude showed that the samples used in this study could present an acceptable genetic basis to constitute a breeding population involved in genetic improvement programmes in the Republic of Congo (Bouvet et al. 2009).

3.1 Variance components and heritability estimates

The AIC or the BIC were calculated for each trait and showed that the model including the genetic random effect exhibited the lower AIC and BIC values and was then considered as the best model (details results not shown). This suggests that the genetic variances are significantly different from zero.

The additive genetic and residual variance components and narrow-sense heritability estimates for various traits are given in Table 3. Genetic and residual variations were higher for growth traits than for wood traits as shown by CV_A and CV_E (except for CV_A of S/G and MFA_3). As expected, the narrow-sense h^2 estimates were lower for growth traits: $h^2=0.34$ for height and $h^2=0.14$ for circumference while moderate to high levels of heritability (from 0.33 to 0.76) were obtained for wood traits.

When the mean values of D or MFA per disc were taken into account, higher narrow-sense heritability estimates

were observed ($h^2>0.60$). The high magnitude of these h^2 estimates can be mathematically explained: once the three data values per tree averaged, the phenotypic (and residual) variances are sharply reduced causing an increase in the additive to phenotypic variance ratio.

Patterns in pith to cambium variation of these traits have been known for many years; wood density generally increases while MFA decreases (Kollmann and Coté 1968). The genetic control of the wood density linearly increased from 0.40 to 0.48 towards the bark; however, this linear trend should be considered with caution since the standard errors (SE) were high (~ 0.17). The h_{MFA}^2 was 0.33 in the early juvenile wood (MFA_1), increasing to 0.45 in the late juvenile wood (MFA_2) and decreasing again to 0.41 in the early mature wood (MFA_3). Considering the magnitude of SE (0.105–0.155), no specific trend was observed for the heritability estimates of MFA.

3.2 Genetic and residual correlations

Estimates of genetic and residual correlations among growth, KL, S/G ratio and mean MFA and density are shown in Table 4. As expected, the genetic and residual correlations between growth traits were positive. The genetic correlations between growth and wood traits (MFA, D and KL) were negative. Positive residual correlations were obtained between these traits. The correlations for S/G are low compared to their standard errors.

The strong genetic correlation between C and D was investigated by means of a residual scatter analysis (not

Table 3 Additive genetic and residual variance components, genetic and residual coefficient of variance and narrow-sense heritability estimates for circumference (C), height (H), Klason lignin (KL), syringyl to guaiacyl ratio (S/G), wood density (D), and microfibril angle (MFA)

Trait	Additive genetic variance			Residual variance			Genetic control	
	σ_A^2	SE σ_A^2	CV_A	σ_E^2	SE σ_E^2	CV_E	h^2	SE h^2
C	13.70	9.95	7.0	100.10	9.12	19.0	0.14	0.087
H	3.752	1.896	9.2	11.089	1.180	15.7	0.34	0.136
KL	1.206	0.529	3.9	1.584	0.276	4.5	0.76	0.212
SG	0.067	0.030	11.3	0.095	0.016	13.4	0.71	0.208
D ₁	8.796	4.525	6.7	22.226	2.685	10.6	0.40	0.167
D ₂	8.330	4.153	5.5	20.055	2.454	8.6	0.42	0.168
D ₃	12.384	5.948	5.7	25.657	3.379	8.2	0.48	0.180
D	9.139	3.914	5.7	14.931	2.184	7.3	0.61	0.183
MFA ₁	1.082	0.478	5.8	3.188	0.299	9.9	0.33	0.105
MFA ₂	1.630	0.770	8.3	3.643	0.406	12.4	0.45	0.139
MFA ₃	1.557	0.625	9.4	3.539	0.387	14.1	0.41	0.155
MFA	1.105	0.544	6.8	2.643	0.311	10.4	0.43	0.153

For D and MFA, the index correspond to radial position, 1 is close the pith. The phenotypic values of wood density were divided by 10 for estimating its variance components

σ_A^2 additive genetic variance component, σ_E^2 residual variance component, h^2 narrow-sense heritability estimates, SE standard errors

Table 4 Estimated additive genetic (r_A , below the diagonal) and residual (r_E , above the diagonal) correlations for circumference (C), height (H), microfibril angle (MFA), wood density (D), Klason lignin (KL), and syringyl to guaiacyl ratio (S/G)

	C	H	MFA	D	KL	S/G
C		0.80 (0.05)	0.01 (0.12)	0.45 (0.21)	0.33 (0.21)	-0.17 (0.17)
H	0.47 (0.33)		0.16 (0.19)	0.41 (0.24)	0.40 (0.37)	-0.41 (0.28)
MFA	-0.36 (0.38)	-0.83 (0.17)		-0.68 (0.28)	0.16 (0.32)	0.38 (0.32)
D	-0.94 (0.21)	-0.55 (0.27)	0.54 (0.27)		-0.35 (0.44)	0.03 (0.39)
KL	-0.16 (0.40)	-0.60 (0.25)	0.70 (0.17)	0.28 (0.28)		-0.23 (0.43)
S/G	0.08 (0.41)	0.35 (0.31)	-0.11 (0.31)	-0.61 (0.21)	-0.17 (0.30)	

Standard errors are shown in parentheses

shown). The point dispersions revealed that there was no aberrant point, which could strengthen the correlation. The correlation estimates had large standard errors probably due to the small sample size.

Radial variations of genetic, residual and phenotypic correlations between wood and growth traits are presented in Fig. 2. The genetic correlation between circumference and wood density remained high during all phases of wood formation, while that between circumference and MFA seemed to linearly decrease towards the bark. This trend was also found between MFA and lignin in which there was an increase with age. The genetic correlation between MFA and wood density decreased from juvenile to mature wood.

The residual correlations were low, except for that between MFA and D. The phenotypic and additive genetic correlations between C and MFA, and KL and MFA presented the same trends, with slight differences in their magnitude. The phenotypic correlation between circumference and wood density was null while the genetic correlation was strong. Similarly, the phenotypic correlation between MFA \times D was low while the residual and genetic correlations were high.

In general, additive genetic correlations presented higher magnitudes when compared with the residual correlations. Moreover, the radial variations of the values of the genetic relationships were stronger than those for residual or phenotypic correlations.

4 Discussion

This study compares the genetic control of wood and growth traits, with special focus on microfibril angle which has been poorly reported in hardwood species. Moreover, genetic and environmental correlations among these traits were assessed at different ages, thus improving our knowledge on the functional aspects of wood formation in *E. urophylla*.

4.1 Heritability of MFA and wood traits

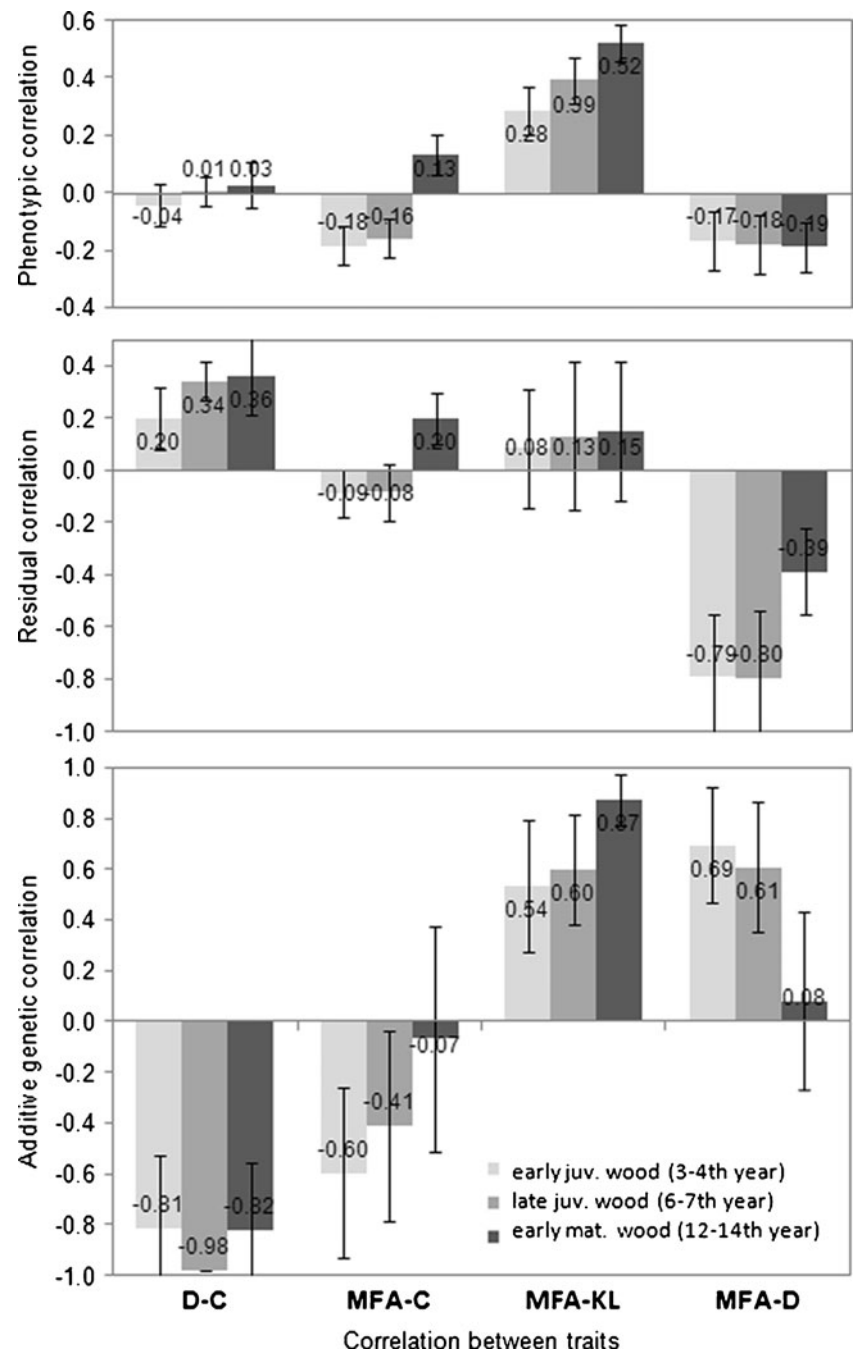
Moderate h^2 estimates were obtained for MFA ($h^2=0.43\pm 0.15$) when calculated from the mean values per disc (Table 3). These narrow-sense heritabilities were higher than those

reported in two previous studies. Apiolaza et al. (2005) used increment cores from 188 open-pollinated progenies of 11-year-old *E. globulus* and obtained $h_{MFA}^2=0.27\pm 0.24$. These relatively lower narrow-sense h^2 estimates of Apiolaza et al. (2005) can be attributed to (1) larger environmental variation since their CV_A (8.36%) is close to the average CV_A of Table 3 (8%); (2) use of a larger number of provenances and (3) fitting provenances as a fixed effect in the model. Lima et al. (2004) reported H^2 of 0.29 in 8-year-old *Eucalyptus* wood. Normally, broad-sense heritability estimates are higher than narrow-sense estimates. The low H^2 estimate of Lima et al. (2004) was expected since they evaluated selected clones of *E. grandis* \times *E. urophylla* for pulpwood so the genetic base was narrow.

Just as for the phenotypic values, the genetic control also varied with age. The heritability estimates of the MFA in the early juvenile wood ($h_{MFA1}^2=0.33\pm 0.105$) increased towards the late juvenile wood ($h_{MFA2}^2=0.45\pm 0.139$), decreasing in the early mature wood ($h_{MFA3}^2=0.41\pm 0.155$). However, due to the high standard error, no specific trend can be demonstrated. This result was similar to the variation pattern reported by Lima et al. (2004) in *Eucalyptus* clones (h_{MFA}^2 increasing from 0.13 to 0.36 and then decreasing to 0.16 towards the cambium). Comparing these findings requires prudence because the experimental designs were different (open- and control-pollinated and clonally propagated tests) and the methods of wood phenotyping were distinct: Apiolaza et al. (2005) estimated the phenotypic MFA values of their study by means of the SilviScan device (based on X-ray diffraction, see Evans 1999); Lima et al. (2004) used polarised light microscopy technique and, here, we associated X-ray-diffraction and NIR spectroscopy to estimate the MFA.

Considerable genetic control was found for mean wood density ($h^2=0.61\pm 0.183$) and for wood density measurements at different radial positions. The heritability estimates linearly increased from the early juvenile wood (0.40 ± 0.167) towards the early mature wood (0.48 ± 0.180). The slight increment in heritability estimates of density with age might have been due a reduction in environmental variation, which may have been associated with canopy closure in the Congolese conditions. Previous studies on *Eucalyptus* in the environmental conditions of the Congo have shown that

Fig. 2 Variation with age of phenotypic, residual and additive genetic correlations of density (D) with circumference (C), and microfibril angle (MFA) with C, Klason lignin (KL) and D and representation of standard error of estimation



before canopy closure, there are large micro-environmental variations between trees while in the following years, tree development depends to a higher magnitude on their genetic potential (Bouvet et al. 2003). Due its relatively easy and simple measurement, the wood density has been extensively investigated in *Eucalyptus* and the heritability of such trait exhibits variable magnitudes: from $H^2=0.51\pm0.13$ (Kube et al. 2001, for all sites) to $h^2=0.73$ (Greaves et al. 1997, for whole-disc density at 1.3 m across sites) in *E. nitens*; from $h^2=0.24\pm0.26$ (Poke et al. 2006) to $h^2=0.44\pm0.22$ (Apiolaza et al. 2005) in *E. globulus* and from $h^2=0.17\pm0.12$ (Raymond et

al. 1998) in *E. regnans*. Most of these studies have shown that the high heritability of this trait is mainly explained by the low environmental influence rather than marked genetic variation (see CV's in Table 3); our study reinforces these previous results.

Lignin content and S/G ratio were shown to be under strong genetic control (Table 3). The high narrow-sense heritability of these chemical traits ($h^2>0.70\pm0.2$) indicates that genetic gain is possible through breeding. The heritability for lignin ($h^2=0.76\pm0.212$) was lower than that reported by Gominho et al. (1997) in *E. globulus* clones

($h^2=0.83$) but higher than that reported by Vigneron et al. (2004) in *E. urophylla* × *grandis* (individual broad-sense $H^2=0.27\pm0.2$), Poke et al. (2006) in *E. globulus* families (narrow-sense $h^2=0.13\pm0.2$ and family means $h^2=0.42\pm0.19$). As lignin content and its composition are key traits of *Eucalyptus*-breeding programmes, especially for pulp, paper and bioenergy production, their high genetic control reported here (Table 3) is important for tree breeders. For instance, high lignin content is desirable for bio-energy because lignin has a high calorific value.

The heritability estimates for growth traits were low ($h^2=0.14\pm0.087$; $h^2=0.34\pm0.136$ for circumference and height, respectively), which are strongly influenced by the environment. These results confirm the trends described in previous studies for other *Eucalyptus*-breeding programmes (Kube et al. 2001; Costa e Silva et al. 2009, 2010) and more specifically in the Congo (Bouvet et al. 2003, 2009). Similar studies on conifers have also reported that growth properties are under less genetic control than wood properties (Zubizarreta Gerendiain et al. 2007).

Although these estimates are population- and site-specific, these findings are coherent with previous studies and useful for tree breeders since they show that MFA, D, KL and S/G present a high heritability and offer possibilities for improvement.

4.2 Correlation at mature age

A lot of literature exists about genetic correlations between growth traits and wood density; however, the findings concerning this issue are inconsistent (see for example Zobel and Van Buijtenen (1989) for a range of wood species). For *Eucalyptus*, several studies also show inconsistent results (Hamilton and Potts 2008; Greaves et al. 1997; Wei and Borralho 1999; Kube et al. 2001) probably due to the small sampling size. The conclusion is that correlation between wood density and growth should be low. In addition, some studies have shown that correlations are influenced by site conditions (Wei and Borralho 1999; Muneri and Raymond 2000; Hamilton et al. 2009) emphasising the environmental impact.

Our results bring new elements to the understanding of the correlation between wood and growth traits. Here, few additive and residual correlations ($C\times H$; $KL\times MFA$) presented the same sign (Table 4 and Fig. 2). This could indicate a pleiotropic gene effect (Falconer 1993), suggesting that these two traits are governed by a given genetic locus (Mode and Robinson 1959).

Most of the additive and residual correlations, and especially those related to MFA (Table 4) had negative sign. This indicates that linkage disequilibrium (non-random associations of genes) between loci may affect the relationship among different wood traits (Falconer 1993). This means that the genes controlling these traits can be statistically associated, but there is no functional relationship among them. Sampling

procedure adopted in this study could have played a role in linkage disequilibrium, since the trees were selected among the best ones with respect to growth and stem straightness. According to Villanueva and Kennedy (1990), selection changes the genetic variance and creates linkage disequilibrium.

4.3 Genetic parameter expression with age

As the tree grows upward, new layers of wood are produced and overlapped in order to withstand the ever-increasing mass of the tree. The variations of the genetic and residual correlations with age may be useful in understanding how the cambial activity and maturation phases are regulated in order to modify the characteristics of the wood making the stem able to withstand gravity and bending movements caused by winds.

The narrow-sense heritability estimates for wood density seems to increase with age (Table 3), but remains more or less constant for MFA. The variances were heterogeneous across ages: the variance of wood density increased with age (trees become more different) while the variance of MFA presented no trends (Table 3). The genetic correlation $C\times D$ is always strongly negative while the correlation $D\times MFA$ remained constant in the juvenile wood, decreasing considerably in the mature wood (Fig. 2). This could mean that trees with a strong potential to grow fast are genetically programmed to produce low-density woods and also to decrease the microfibril angles to help ensure stiffness (the negative correlation $MFA\times C$). However, the sample size (340 trees) is quite small and any genetic correlations were estimated to have large associated errors. Furthermore, the eventual occurrence of linkage disequilibrium within this population means that a functional explanation needs to be treated with caution.

Eucalyptus trees can reach 25–30 m of height in 6–7 years. As the tree grows, enlarging its diameter and increasing its height, the stem becomes more subject to higher bending moments from the wind. It is known that reaction wood is typically formed as the tissues of the periphery are held in “tension” and in many wood species, tension wood is characterised by the presence of a thick and unlignified, inner cell-wall layer that consists of highly crystalline cellulose, in which the MFA is close to zero (Kollmann and Côté 1968). Thus, the low lignin content and MFA of the early mature wood may be associated with tension wood occurrence (Fig. 2). Our results indicate that variations in these wood traits are also genetically controlled since the genetic correlation between MFA and KL increases from 0.54 to 0.87 towards the cambium.

4.4 Implications for selection

These findings bring additional insights for genetic breeding strategies. If the population analysed in this study is to

constitute the first generation of a breeding programme, different elements should be taken into account for selecting candidate trees.

For the purposes of pulpwood production, the objective is to produce trees with high biomass value (volume, density and cellulose content) and reduced lignin content. The negative r_A between growth traits and lignin content ($r_A = -0.60$ for height and -0.16 for circumference) and the positive correlation between MFA and KL ($0.54 < r_A < 0.87$) are favourable for pulp and paper production since the reduction in all of these traits should be simultaneously possible. Lignin is an undesirable compound for pulp and paper production, because the delignification process requires energy and reagent consumption.

For bio-energy production, the positive genetic correlation between lignin and density (0.28) is favourable, despite the low magnitude. However, positive additive correlations between MFA and lignin are unfavourable for bioenergy purposes since decreasing MFA, may result in decreasing lignin content. The problem is that, for energy production, the wood must simultaneously have high lignin content and adequate mechanical properties. Charcoal is used in blast furnaces for iron–steel production and has to support the weight of the iron feedstock during the steps of oxidation–reduction reactions at elevated temperatures (above 1,500°C) without breaking.

For sawn wood production, the objective is to obtain stiff (low MFA), dense and dimensionally stable wood from trees with low growth stresses. Our findings have shown that wood density and MFA were genetically unfavourably correlated with growth. Moreover, selection for increasing density will result in an increase of MFA and decrease of wood stiffness, the major wood trait for structural, furniture and flooring uses.

As pointed out by Baltunis et al. (2007), breeders, forest managers and wood producers will have to strike a balance between overall wood and growth traits, and geneticists should develop breeding strategies to deal with such negative, unfavourable genetic correlations. For *Eucalyptus*, developing breeding objectives may be the first step in dealing with these unfavourable genetic correlations between wood and growth traits.

Acknowledgements The authors would like to thank Arie van der Lee from the Institut Européen des Membranes for assistance with X-ray diffraction measurements; Dr. C Lapierre and her work team from INRA-Agro ParisTech for chemical analysis; N Ognouabi and E Villar from CRDPI (Republic of Congo) and the “Bureau des Ressources Génétiques” (BRG) project for providing material and funding; and especially Dr. JM Gion (coordinator of the BRG project “CCR gene in *Eucalyptus*: a model of functional variability in forest trees”) from CIRAD for providing technical support.

Funding This study is a part of the project “CCR gene in *Eucalyptus*: a model of functional variability in forest trees” funded by the “Bureau des Ressources Génétiques” (BRG). The project was also funded by the Centre de Coopération Internationale en Recherche Agronomique pour le Développement (CIRAD) in Montpellier, France. The first author was supported by the National Council of Technological and Scientific

Development (CNPq, Brazil—process no. 200970/2008-9) during his PhD thesis. He was also supported by the Alfa Gema Project (II-0266-FA) during his masteral studies.

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