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Structure of the specific combining ability between two species of *Eucalyptus*.

I. RAPD data

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Abstract Within the context of the reciprocal recurrent selection scheme developed in 1989 by CIRAD-Forêt on *Eucalyptus*, RAPD essays were performed to assess the genetic diversity in the two species *E. urophylla* and *E. grandis*. The molecular markers were split into two parts: the specific markers (present with different frequencies in the two species) and the common markers (present with similar frequencies in the two species). The study analyses the structure of genetic diversity within and between the two species of *Eucalyptus*. Different genetic distances are worked out for use in prediction equations of the individual tree trunk volume of hybrids at 38 months. Each distance is expressed as the sum of the general genetic distance and the specific genetic distance. The general genetic distance based on the double presence plus the double absence of bands seems to be an interesting co-variate to use in a factor regression model. Through this model the distance calculated between species explains the general combining ability (GCA) and the specific combining ability (SCA) of the phenotypic character with a global coefficient of determination of 81.6%.

Key words RAPD · Genetic distances · Heterosis breeding · Prediction · *Eucalyptus* · Interaction

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Introduction

Eucalyptus which originated in Australia, is a highly polymorphic genus (around 600 species) belonging to the family *Myrtaceae* which is primarily grown for firewood and wood pulp (mainly for paper). It was introduced in the Congo during the fifties and today represents the second major resource of this country. The fortuitous observation of a natural hybrid between two species from the sub-genus *Symphomyrtus*, namely *Eucalyptus urophylla* and *E. grandis*, revealing an important heterosis, led breeders to use a reciprocal recurrent selection scheme (Vigneron 1991). This strategy is especially adapted to interspecific crosses (Gallais 1990) between two highly divergent and complementary populations that have evolved in different environments, with no genetic exchanges between them. *E. urophylla*, for which 85 samples were collected, is adapted to local conditions and is high-yielding, while *E. grandis*, for which 25 samples were collected, is less well adapted but has high growth potential.

Since the selection process in tree improvement is very time consuming, any means of predicting tree performance has to be considered with interest. The information required for the effective breeding of hybrids felled at 7 years, can be obtained after 2 or 3 years (Bouvet and Vigneron 1995), but one question remains: which crosses have to be made among all the possible ones? The aim of this paper is to answer this question using genetic distances obtained through the use of RAPDs.

Materials and methods

In 1990, 13 maternal trees of *E. urophylla* were crossed with 13 paternal trees of *E. grandis* in a factorial mating design (Bouvet and Vigneron 1995). While the maternal trees came from two highly differentiated provenances in the island of Flores, namely Monte Lewotobi and Monte Egon, all the pollen came from trees uniformly

distributed over the northern part of the natural area near Atherton, Queensland, Australia. Unfortunately, technical problems in controlled pollination prevented the mating design from being complete and balanced, with only 87 families among the 169 possible ones (i.e. a ratio equal to 51%). Moreover, the number of replicates varied from one to four according to the cross. These constraints led to a reduced factorial design involving nine *E. urophylla* and nine *E. grandis* in which 49 families were present among the 81 possible ones (i.e. a ratio equal to 60%) with three or four replications per family. In this reduced design, each female and male parent is represented by at least three or four families, respectively. In each experimental unit (square plot of $4 \times 4 = 16$ trees), height and circumference at 1.3 m were measured at 38 months. In the Congo, this corresponds to the half-rotation age in commercial plantations. Volume was calculated by considering the tree trunk as a cone. All of the following analyses were performed using the means of the three or four family replications.

RAPD assays

The details of total genomic DNA isolation from dry leaves and DNA amplification is presented in Verhaegen et al. (1995). Oligonucleotide primers (10-mers) were used as single primers for the amplification of RAPD sequences according to Williams et al. (1990). The occurrences of a specific band of amplified DNA is scored as 1 and absence as 0, leading to the characterization of each individual by a binary variable. Pairwise comparisons of individuals were employed to calculate two similarity coefficients: Jaccard's coefficient (Jaccard 1908; Jain et al. 1994) and Sokal and Michener's coefficient (Sokal and Michener 1958), also called a simple matching coefficient (Skroch et al. 1992; Lamboy 1994). From these similarity coefficients, two indexes of dissimilarity between individuals A and B were calculated:

$$D_2 = 1 - Sim_1,$$

where $Sim_1 = N_{ab}/N_a + N_b - N_{ab}$, N_{ab} is the number of fragments shared by A and B (double presence), while N_a and N_b are the number of fragments present in individual A and in individual B, respectively.

$$D_2 = 1 - Sim_2,$$

where $Sim_2 = N_{AB}/N$, N_{AB} is the number of fragments present and absent for individuals A and B (double presence plus double absence), and N is the total number of fragments.

Note that $N \times D_2$ is the euclidian distance between two individuals.

Let $d^2(i, i')$ be the Euclidian distance between individuals i and i' , let $j = 1, \dots, N$. Then:

$$d^2(i, i') = \sum_j (x_{ij} - x_{i'j})^2,$$

where $x_{ij} = 0$ if the band corresponding to the j^{th} marker is absent and $x_{ij} = 1$ if the band is present. Consequently $x_{ij} - x_{i'j} = 0$ either if $x_{ij} = x_{i'j} = 0$ (double absence) or if $x_{ij} = x_{i'j} = 1$ (double presence). Hence $d^2(i, i')$ is the number of non-coincidences in the two individuals of 0 or 1.

From the sample of 26 parents (13 *E. urophylla* and 13 *E. grandis*), 415 reliable RAPD bands were obtained using 15 primers.

Structure of the genetic diversity

Firstly, factorial analysis of distance tables seems to be a good tool for visualizing the structure of the genetic diversity. Secondly, in each population each random amplified product can present a number of bands varying from zero (if no individual exhibits the frag-

Table 1 Breakdown table of the number of markers presenting specific combinations of band frequencies in the two *Eucalyptus* populations (*E. urophylla* in rows and *E. grandis* in columns). Italic numbers symbolize the six different groups of markers

	0	1	2	3	4	5	6	7	8	9	10	11	12	13
0
1	.	<i>1</i>	<i>4</i>	<i>6</i>
2
3
4
5
6
7	.	<i>4</i>	<i>2</i>	<i>5</i>
8
9
10
11
12	.	<i>6</i>	<i>5</i>	<i>3</i>	.
13

ment) to 13 (if all the individuals show the fragment). A two-way table is constructed with 14×14 cells corresponding to the coincidence of a specific frequency of bands in the female population with a specific frequency of bands in the male population. The contingency table shows the number of variables concerned with each combination of band frequencies. An exact Fisher test allows one to discriminate markers shared with equal frequency by the two species (dotted area in Table 1) from those showing significant frequency differences between the two species. The contingency table was further subdivided into nine zones corresponding to different values of the exact Fisher test (dashed lines in Table 1). These zones correspond to the combination of three areas in each population, namely the bands whose frequency does not differ significantly from zero [zone (1) if the two species show this type of frequency], the bands whose frequency does not significantly differ from one [zone (3) if the two species show this type of frequency], and the bands with intermediate frequencies [zone (2) if the two species show this type of frequency]. Reciprocal areas have been merged in order to create six different zones (given in italic letters).

Prediction of specific crossing values

The two genetic distances D_1 and D_2 have been calculated between and within species using all the RAPD variables, and also using separately either the common variables or the specific variables. Eighteen different distances [six classes of variables \times (two intraspecific distances + one interspecific distance)] have been defined and calculated on the 9×9 contingency table corresponding to the reduced design. Each distance table has been adjusted to the additive model (1) and the estimates of the additive parameters have been retained as co-variables to be introduced in a factor regression model

in order to explain GCA (general combining ability) and SCA (specific combining ability). The additive model with two factors applied to distance variables is:

$$X_{ij} = \mu + Y_i + Z_j + R_{ij} \quad (1)$$

where X_{ij} is the genetic distance between individual i and individual j , μ is the general mean distance, Y_i is the mean distance of individual i minus the general mean, Z_j is the mean distance of individual j minus the general mean and R_{ij} is the residual term.

A large positive or negative Y_i indicates that the individual possesses many bands that occur at low or high frequencies, respectively, in the other individuals. If individuals i and j possess the same bands with low frequencies their R_{ij} will be highly negative, whereas if two individuals possess different bands with low frequencies their R_{ij} will be highly positive. This partition has earlier been proposed on Rogers' distance with RFLP data by Melchinger et al. (1990a, b). These general genetic distances can be defined between and within species. In the case of distance tables within species, the null diagonal has been removed before adjustment to the additive model. In this model the interaction term is merged into the residual term. The estimates of additive parameters Y_i and Z_j are then used as co-variables in the factor regression model (Denis 1988; Baril 1992) applied to the phenotypic variable:

$$W_{ij} = \mu + \gamma \cdot Y_i + \alpha'_i + \delta \cdot Z_j + \beta'_j + \rho \cdot Y_i \cdot Z_j + v_j \cdot Y_i + \lambda_i \cdot Z_j + \varepsilon_{ij} \quad (2)$$

where W_{ij} is the individual tree trunk volume at 38 months of the family descending from the cross between mother i and father j , μ is the general mean, $\alpha_i = \gamma \cdot Y_i + \alpha'_i$ is the GCA of mother i , $\beta_j = \delta \cdot Z_j + \beta'_j$ is the GCA of father j , and $\rho \cdot Y_i \cdot Z_j + v_j \cdot Y_i + \lambda_i \cdot Z_j$ is that part of the SCA explained by the model.

Unlike co-variance analysis in which the co-variables depend on the two factors, factor regression allows the partitioning of SCA into functions of only one factor, each multiplied by a regression coefficient depending on the other factor. In order to keep the logical symmetry between the two parents, the co-variables have always been introduced in the model by pairs, i.e. one for *E. urophylla* and the other for *E. grandis*. This model provides both an explanation of the additive part of the variability (i.e. GCA) and an explanation of the interactive part (i.e. SCA). Each parental additive contribution to the whole variability is split into two terms. The first represents that part of GCA explained by the regression over the female (or male) co-variate and the second is the rest of GCA (not explained by the general distance). The interaction between the two parents is split into four terms. The first term is the combined regression over the product of the two co-variables (male and female); the two following terms are the remaining regressions over the female co-variate on the one hand and the male co-variate on the other hand. Finally, the rest, ε_{ij} , is the residual term.

The application of models with many parameters when there are numerous missing values in the data set can give fallacious estimates (Denis and Baril 1992). In order to avoid aberrant estimates of phenotypic data, the missing values were estimated by the bi-joint regression model. This model is named by analogy with the joint regression model, well known in studies of genotype \times environment interactions (Finlay and Wilkinson 1963), which allows one to regress the interaction term on one of the two additive terms (the environment effect). In the bi-joint regression model, the interaction between the two factors, namely father and mother, is split into four parts as in a factor regression model where the co-variables depending on each factor are the estimates of the two additive effects (GCA). The first part of the interaction is the regression on the product of the two GCAs, which corresponds to the model proposed by Tukey (1949).

The bi-joint regression model applied to the phenotypic variable gives:

$$W_{ij} = \mu + \alpha_i + \beta_j + \rho \cdot \hat{\alpha}_i \cdot \hat{\beta}_j + v_j \cdot \hat{\alpha}_i + \lambda_i \cdot \hat{\beta}_j + \varepsilon_{ij} \quad (3)$$

As for the simple joint regression, the data are first adjusted to the additive model in order to estimate the additive parameters (namely, $\hat{\alpha}_i$ and $\hat{\beta}_j$), and then adjusted to the complete model using these estimations as co-variables. The bi-joint regression model not only fits the data well (coefficient of determination = 90.8%) but also provides estimates which retain part of the interaction. Once the missing values were estimated by this model, the factor regression model has been used on the new completed data set. The estimates of missing data have also been obtained through the additive model, whose results are not presented in this paper, and have given almost similar results in the following analyses. All these analyses were performed using the computer package INTERA (Decoux and Denis 1991), which provides least-squares estimates of parameters. Finally, factorial analyses of the two distance tables D_1 and D_2 were performed with a Statistical Analysis System (SAS Institute 1988).

Results

Structure of the genetic diversity

Factorial analyses of the two distance tables D_1 and D_2 show a clear aggregation of individuals for each species. The first, second and third principal factors, based on D_1 distance, account for 24.9%, 8.6% and 8.0%, respectively (that is to say a sum equal to 41.5%), while the three principal factors, based on D_2 distance, account for 33.9%, 8.2% and 7.9%, respectively (that is to say a sum equal to 50%). The plots of principal factors worked out from the two genetic distances show a tendency for the D_2 distance (Fig. 1) to provide the

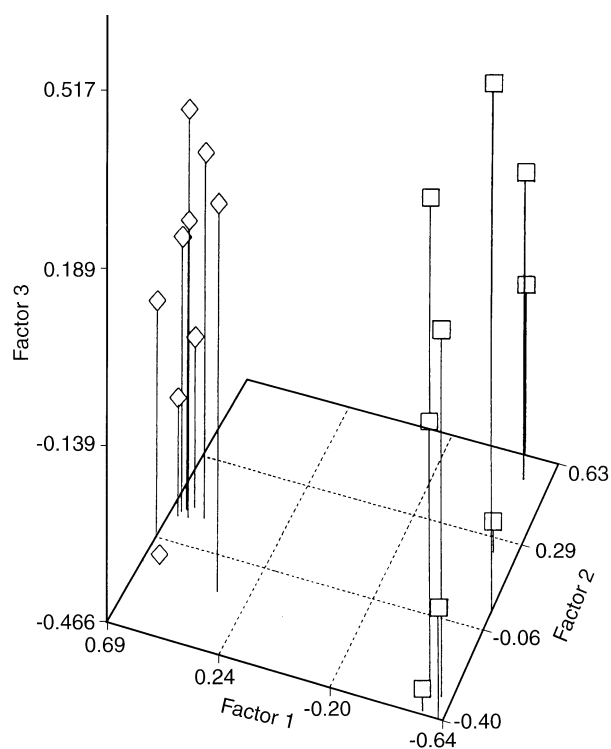


Fig. 1 Plot of the three first factors of FADT computed from D_2 . Species: \square *E. urophylla*, \diamond *E. grandis*

Table 2 Contingency table of the number of markers presenting specific combinations of band frequencies in the two *Eucalyptus* populations (*urophylla* in rows and *grandis* in columns). Σ_u and Σ_g are the sum of RAPD markers with a particular frequency of bands in *E. urophylla* and *E. grandis*, respectively

	0	1	2	3	4	5	6	7	8	9	10	11	12	13	Σ_u
0	-	15	16	17	6	5	7	5	2	1	1	1	1	2	79
1	39	13	8	6	4	2	2	0	0	1	2	0	1	0	78
2	15	8	4	3	3	1	0	2	0	1	1	0	0	0	38
3	11	10	5	1	1	2	4	0	1	2	1	0	0	1	39
4	9	4	4	2	2	0	4	0	1	3	1	2	3	0	35
5	11	4	2	3	2	3	2	0	0	2	0	1	2	1	33
6	6	5	3	1	0	0	2	0	0	1	0	2	0	1	21
7	5	1	2	1	1	2	2	0	1	2	0	2	1	0	20
8	4	1	2	0	2	1	0	1	0	2	2	1	3	0	19
9	1	1	1	2	1	4	0	0	3	1	0	0	0	0	14
10	1	1	0	1	0	0	0	0	1	1	0	0	0	0	5
11	2	1	1	0	0	1	0	1	0	0	1	2	1	4	14
12	4	0	0	0	1	1	1	0	2	1	1	0	1	1	13
13	0	0	1	0	0	0	0	0	0	1	2	1	0	2	7
Σ_g	108	64	49	37	23	22	24	9	10	19	13	12	13	12	415

best differentiation index between the two species of *Eucalyptus*. Each cell of the contingency table presented in Table 1 is filled in Table 2 with the number of bands with a specific combination of frequencies in the *E. urophylla* population and the *E. grandis* population.

The contingency table is split in two parts, one along the second bisecting line which contains the common markers (dotted area with 226 RAPD variables, i.e. 54.5% of the total number of markers) and the remaining parts of the table which contain the specific markers (189 RAPD variables, i.e. 45.5% of the marker sample). Among these markers, 99 are significantly more frequent in the *E. urophylla* population, and 90 are more frequent in the *E. grandis* population. The threshold values of the exact Fisher tests were calculated at the 25% probability level in order to split the markers into two balanced parts, namely the specific ones (with different frequencies for the two species, DIF) and the common ones (with common frequencies for the two species, COM). The sum of markers present in each row (Σ_u for a particular proportion of bands in *E. urophylla*) and column (Σ_g for a particular proportion of bands in *E. grandis*) is shown in the margin of Table 2. For instance, 108 markers were present in at least one *E. urophylla* tree and never in *E. grandis* trees. The reciprocal situation concerns only 79 markers, never present in *E. urophylla*. These results are consistent with Fig. 1 which shows a larger variability of the *urophylla* population when the distance table is based on D_2 , i.e. when the calculation of the distance between two individuals depends on the whole sample of individuals. The lower

Table 3 Frequency of RAPD variables in the nine areas of the contingency table according to the exact Fisher tests calculated at the P-level = 25% (a) and 5% (b). p_u (or p_g) are the band frequencies among *E. urophylla* (or among *E. grandis*)

a

	$p_g \approx 0$	$0 < p_g < 1$	$p_g \approx 1$
<i>p_u ≈ 0</i>	(1) 28%	(4) 17%	(6) 1%
<i>0 < p_u < 1</i>	(4) 23%	(2) 17%	(5) 5%
<i>p_u ≈ 1</i>	(6) 2%	(5) 3%	(3) 3%

b

	$p_g \approx 0$	$0 < p_g < 1$	$p_g \approx 1$
<i>p_u ≈ 0</i>	(1) 50%	(4) 9%	(6) 6%
<i>0 < p_u < 1</i>	(4) 13%	(2) 3%	(5) 6%
<i>p_u ≈ 1</i>	(6) 5%	(5) 3%	(3) 5%

distances between *grandis* individuals reflect the higher number of coincidences of zero in the 415 RAPD variables relatively to the *urophylla* individuals. The contingency table was further subdivided in nine parts (dashed lines in Table 1) according to whether or not the frequency of bands is significantly different from 0 or 1. The proportion of markers present in the each zone is shown in Table 3 (a). The six different groups (in italic letters in Table 1), defined the merging of the symmetric area, from number one to number six: 118 (28%), 72 (17%), 12 (3%), 166 (40%), 33 (8%) and 14 (3%) RAPD variables.

Common variables with low band frequencies are represented in high frequency (group 1) and comprise those present once either in *E. urophylla* (39 of them) or in *E. grandis* (15 of them). In order to prevent the intervention of artefacts (a source of “noise”) in the calculation of genetic distances we worked out the different distances excluding these two cells [namely, (0, 1) and (1, 0)]. In fact, the results obtained with these genetic distances were of no interest and hence will not be mentioned further in this paper. Finally, one may note that the cell (0, 0) is structurally empty. The contingency table was then subdivided into nine parts according to the exact Fisher test calculated at the 5% probability level. The proportion of markers present in each new group is shown in Table 3 (b). The six different groups corresponding to this new structure contain from number one to number six: 206 (50%), 14 (3%), 21 (5%), 94 (23%), 36 (9%) and 44 (11%) RAPD variables.

Prediction of specific crossing values

The specific genetic distance R_{ij} (first model) is analogous to the specific Roger’s distance proposed by Melchinger et al. (1990). The correlation coefficients between R_{ij} of distances D_1 and D_2 and the SCA of tree

trunk volume at 38 months are not significant (0.07 and 0.05, respectively). These results justify the use of Y_i and Z_j , the general genetic distances, as co-variates in the factor regression model. Two distance tables within species (suffix URO and suffix GRA) and one distance table between species (suffix U × G) have been created for three groups of RAPD variables: the total number of bands (prefix ALL), the subset of common bands (prefix COM) and the subset of different bands (prefix DIF). The sum of the common bands and the different bands forms the total number of RAPD variables. If all

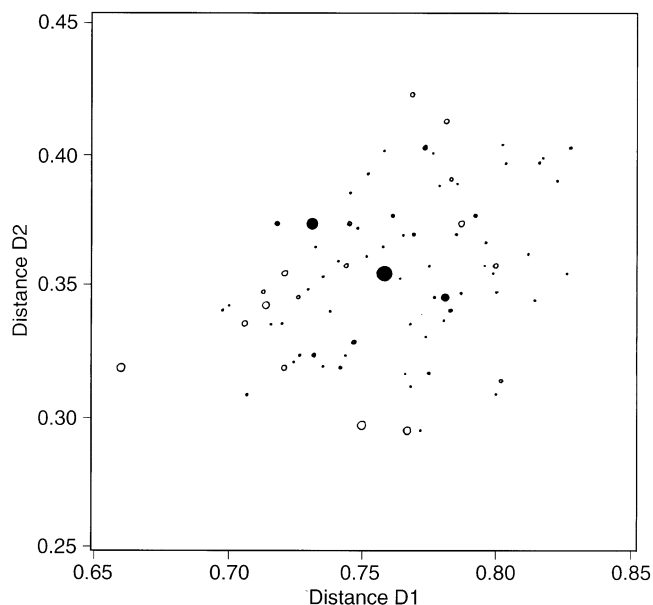


Fig. 2 Plot of genetic distance D_2 versus D_1 , Circle size stands for the value of SCA; solid circles show a positive SCA and empty bubbles show a negative SCA

the bands are considered, the correlation coefficients between distances D_1 and D_2 calculated within species (without the null diagonal values) are $\rho = 0.77$ and $\rho = 0.76$ for *E. urophylla* and *E. grandis*, respectively. The correlation coefficient between the two types (D_1 and D_2) of between-species distance is 0.36 (significant at the 0.1% level). This rather small value reflects an actual difference between the two genetic distances. Figure 2 shows the plot of between species D_2 in terms of between species D_1 where circle-size stands for the SCA value of tree trunk volume at 38 months. Positive SCAs are solid and negative ones are empty. It seems that highly positive SCA values (heterosis) are confined to medium values of genetic distance.

The additive model was performed on distances D_1 and D_2 defined within and between species and worked out from different groups of RAPD variables. The distributions of different sources of variability for each distance are presented in Table 4 with the mean value of each distance table.

The distance tables within species are naturally symmetric and therefore the two additive effects are equal. The amount of variability corresponding to the specific genetic distance R_{ij} is, as expected, greater within species than between species. The greater the amount of the specific genetic distance, the smaller the meaning of the use of general genetic distances. The mean distances within *E. urophylla* are systematically greater than the mean distances within *E. grandis*. This remark is especially true for the whole group and *a fortiori* for the specific group. For these two groups the mean distances between species are clearly greater than the mean distances within species. Finally, the mean distances D_2 , involving double presences plus double absences, are smaller than the mean distances D_1 which only involve double presences. Correlation coefficients between co-variates associated to the *E. urophylla*

Table 4 General mean and percentages of the variability of genetic distances explained by the main effects and interaction term. % Y_i , % Z_j and % R_{ij} are the percentages of variability of the distance between individuals i and j explained by the two additive effects and the interaction effect, respectively. When additive effects are not

significant (at the 5% level in the ANOVA) the corresponding percentage of the whole sum of squares is replaced by the NS. μ_{D_1} and μ_{D_2} are the general means of D_1 and D_2 , respectively, for each distance table

Item		D_1				D_2			
		% Y_i	% Z_j	% R_{ij}	μ_{D_1}	% Y_i	% Z_j	% R_{ij}	μ_{D_2}
ALL	URO	NS	–	98.3	0.65	25.8	–	48.3	0.28
	GRA	–	NS	97.1	0.61	–	27.2	45.6	0.25
	U × G	44.5	27.3	28.3	0.76	43.4	35.5	21.1	0.35
COM	URO	30.7	–	38.6	0.69	36.2	–	27.6	0.25
	GRA	–	25.8	48.9	0.67	–	31.8	36.3	0.23
	U × G	36.9	14.2	48.9	0.68	48.3	23.6	28.4	0.24
DIF	URO	NS	–	65.4	0.61	NS	–	75.9	0.31
	GRA	–	27.5	45	0.55	–	NS	65.5	0.26
	U × G	55.9	31.5	12.6	0.82	44.2	42.9	12.9	0.49

population (Y_i) and the *E. grandis* population (Z_j) have been calculated (data not shown).

The low correlation coefficients between D_1 and D_2 , COM and DIF, and W and B, justify the calculation of these parameters. The estimates of additive parameters based on different distance tables are used in the factor regression model to explain the tree trunk volume of hybrids at 38 months (Table 5).

The superiority of the explicative power of the D_2 distance versus the D_1 distance is clear. Considering the D_2 distances, those calculated between species are more explicative than those calculated within species. Moreover, distances calculated within *E. grandis* never explain any part of the variability of the phenotypic character. The co-variates used in the factor regression model are extracted from the between-species D_2 distances calculated over all the markers (ALL-U \times G). The corresponding ANOVA table is shown in Table 6.

The estimates of the parameters γ , δ and ρ of this model are all positive. Hence the GCA of tree trunk volume at 38 months is positively correlated with the

genetic distance D_2 . If the additive model is applied to the volume at 38 months, the proportion of variability taken into account by the additive part is 60.5% for the *urophylla* parent plus 14.1% for the *grandis* parent, and the interaction part (SCA) represents 25.4%. In the factor regression model, these three components are split into regressions on the co-variates and residual terms from the regressions. The explicative power of the two co-variates Y_i and Z_j , reflected by the coefficient of determination, is 81.6%.

Finally, the factor regression model based on the D_2 between-species distance table calculated with the common bands (COM-U \times G) presents a determination coefficient equal to 84.2% while the determination coefficient corresponding to the D_2 between-species distance table calculated with the different bands (DIF-U \times G) is 77.8%.

This results shows that the effective RAPD markers for predicting the performance of *Eucalyptus* hybrids are those presenting common frequencies in the two species.

Table 5 Results of the factor regression model on the tree trunk volume of hybrids at 38 months using the estimates of general genetic distances as co-variates, giving the percentages of variability explained by the different terms of model. When the effects are not significant (at the 5% level in the ANOVA) the corresponding

percentage of the whole sum of squares is replaced by NS. The interaction term consisting of the regression on the product of the male and female co-variates ($\rho \cdot Y_i \cdot Z_j$) is never significant and hence has not been presented in this table

		D_1				D_2			
		% $\gamma \cdot Y_i$	% $\delta \cdot Z_j$	% $v_j \cdot Y_i$	$\lambda_i \cdot Z_j$	% $\gamma \cdot Y_i$	% $\delta \cdot Z_j$	% $v_j \cdot Y_i$	$\lambda_i \cdot Z_j$
ALL	URO	NS	–	NS	–	18.8	–	6.6	–
	GRA	–	NS	–	NS	–	NS	–	NS
	U \times G	NS	NS	NS	NS	37.2	NS	6.8	NS
COM	URO	NS	–	NS	–	15.2	–	NS	–
	GRA	–	NS	–	NS	–	NS	–	NS
	U \times G	6.1	3.5	NS	NS	26.9	1.8	8.2	NS
DIF	URO	NS	–	NS	–	18.2	–	NS	–
	GRA	–	NS	–	NS	–	NS	–	NS
	U \times G	NS	NS	NS	NS	13.7	NS	NS	NS

Table 6 ANOVA of the factor regression model with one covariate associated to each parent (derived from the ALL-U \times G D_2 distance). The last column of the ANOVA table shows the amount of the variability of the phenotypic character explained by each term of the factor regression model

Source of variation	Sum of squares	df	Mean squares	F-test	P-level	% Sum of squares
<i>E. urophylla</i>	4597.82	8	574.73	–	–	60.5%
$\gamma \cdot Y_i$	2827.41	1	2827.41	99.1	< 0.001	37.2%
α'_i	1770.41	7	252.91	8.9	< 0.001	23.3%
<i>E. grandis</i>	1072.03	8	134.00	–	–	14.1%
$\delta \cdot Z_j$	1.03	1	1.03	0.0	0.844	\approx 0.0%
β'_j	1070.99	7	153.00	5.4	< 0.001	14.1%
<i>E. uro</i> \times <i>E. gra</i>	1934.26	81	23.88	–	–	25.4%
$\rho \cdot Y_i \cdot Z_j$	0.68	1	0.68	0.0	0.873	\approx 0.0%
$v_j \cdot Y_i$	517.35	7	73.91	2.6	0.023	6.8%
$\lambda_i \cdot Z_j$	17.56	7	2.51	0.1	0.998	0.2%
ε_{ij}	1398.67	49	28.54			18.4%

Discussion

Without any available references concerning the use of RAPDs in *Eucalyptus* breeding, our approach is mainly prospective.

Firstly, this study shows that genetic distance based on RAPDs provides a useful tool to differentiate the two species of *Eucalyptus*. If the RAPD variables are clustered in nine different groups according to their frequency in both species tested at the 5% probability level, bands with low frequencies in both species (group 1) represented 50% of the total number. This structure of the genetic diversity provides results characterized by a large number of bands with low frequencies and a small number of bands with high frequencies. Surprisingly, the bands predicting hybrid performance are those with similar frequencies common to the two species.

Secondly, the factor regression model gives an interesting partitioning of the GCA and the SCA on tree trunk volume data at 38 months into linear functions of relevant male and female co-variables. The global coefficient of determination, 81.6%, is satisfactory, knowing that the general genetic distance of *E. urophylla* individuals explains 27% of SCA of tree trunk volume at 38 months. The prediction of heterosis through genetic markers is an old dream of plant breeders. The usefulness of RAPD-based genetic distance measures in predicting the performance of between-species hybrids in *Eucalyptus* is thus demonstrated.

Thirdly, this study examines the problem of the choice of genetic distance for predicting crossing values. The best co-variables are defined from genetic distances calculated between the species of *Eucalyptus* and based on the simple matching coefficient. The greater efficiency of co-variables derived from between-species distance versus co-variables derived from within-species distance is intuitively obvious. However, the greater efficiency of co-variables corresponding to general genetic distances defined by the double presence plus the double absence (D_2) of bands is not evident. The two genetic distances D_1 and D_2 have been defined at the two extremities of the spectrum of the possible distances with no *a priori* preference. There are convincing arguments in favour of each definition of genetic distance. Jaccard's coefficient does not depend on the individual samples studied but its use implicitly presupposes a positive effect of the presence of bands. Peltier et al. (1994) compared these two genetic distances using RAPD markers in the polygenetic reconstruction of seven species of *Petunia*. The authors showed that the best fit between the *a priori* taxonomy and the *a posteriori* grouping is obtained with D_1 for *a priori* distant species, while the best fit is obtained with D_2 for *a priori* near species. Two phenomena can generate the absence of a RAPD band: a modification of a flanking sequence and the entire absence of the amplified se-

quence. In the first case, D_2 is the more accurate genetic distance, while in the second case D_1 is more accurate. One could argue that it would be interesting to test other distances. Let us consider the weighted Euclidian distance $d'^2(i, i')$ between the two individuals i and i' : $d'^2(i, i') = \sum_j w_j (x_{ij} - x_{i'j})^2$ where $w_j = 1/\text{var}(V_j)$ with $\text{var}(V_j)$ the variance of the j^{th} marker. This distance would emphasize the loci presenting a great imbalance between a proportion of 0 (absence of the band) and a proportion of 1 (presence of the band). The inverse weighting would emphasize the loci with a similar proportion of presence and absence of bands and might be more interesting. Other kinds of weights could be imagined employing different criteria. For example, the number of bases constituting the primers used in the polymerase chain reaction (PCR) could be one of them.

To conclude, this study shows the necessity of using a genetic distance based on common markers for two species in order to predict the value of crosses. Instead of using a large number of RAPD bands, it would be more efficient to find specific bands for the genomic regions actually contributing to heterosis for the agronomic traits of interest. A thorough reflection on the definition of genetic distances and localization in the between-species contingency table of bands linked to quantitative traits will be the subject of a further paper.

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