BREEDING ZONES BASED ON GENOTYPE-ENVIRONMENT INTERACTION IN SEEDLING SEED ORCHARDS OF Eucalyptus pellita IN INDONESIA

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ABSTRACT

The magnitude of the genotype-environment interaction (GEI) in seven six years old seedling seed orchards of Eucalyptus pellita established at three locations in Indonesia was examined by analyzing height and diameter. The seed orchard at each location consists of 121 families from three provenances of Papua New Guinea. The pooled sum of squares derived from analysis of variance of growth of the three provenances was used to estimate genetic parameters. These parameters were then used to calculate genetic gain as a result of both direct and indirect selections. The GEI across the three locations was very strong, while the results of pair-wise site analysis varied between the pairs of locations. The genetic correlations for pairs of locations were moderately high between South Sumatra and Riau; 0.68 for dbh and 0.85 for height, but were lower between South Kalimantan and the other locations (0.32 to 0.56). The gains resulting from direct selection were apparently greater than those resulting from indirect selection. The genetic gains achieved in South Sumatra and Riau would decrease by 24% as a result of transfer to other sites, while transfer from South Kalimantan to Sumatra Island (and vice versa) would reduce in the genetic gain by 60%. Hence, the breeding population in South Kalimantan should be developed separately from those in South Sumatra and Riau.

Keywords: Breeding population, direct and indirect selection, *Eucalyptus pellita*, genetic correlation, genetic gain

I. INTRODUCTION

The concept of genotype-environment interaction, or GEI, has been defined as the variation in relative performance of genotypes with environment (Burdon, 1977). The effects of GEI complicate tree breeding programs and may reduce the rate of genetic progress by selection. Heterogeneity among forest sites, due to variations in soil and/or climatic conditions, may cause GEI (Gwaze *et al.*, 2001). Therefore, the magnitude of the GEI will affect the decisions on testing, selection and deployment. GEI has been studied for various pine species, including: *Pinus radiata* (Pederick, 1990; Johnson, 1992), *P. caribaea* (Woolaston *et al.*, 1991), *P. elliotii* (Pswarayi *et al.*, 1997) and *P. taeda* (Gwaze *et al.*, 2001). The results of genetic correlation analysis suggest that the greatest gains are almost always result from selecting and replanting at the same site. However, the genetic loss as a result of selecting and planting

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at different sites is generally small enough to suggest that selecting for general adaptations (using selections adapted to a wide range of sites) is a better strategy than regionalizing breeding operations (separate breeding populations or seed orchards, each focusing on "locally" adapted selections). However, if GEI is found to be important, two approaches can be applied: the breeding can focus on either the environments or the genotypes. The first alternative aims to determine groups of environments within which GEI approaches zero; it thus aims to maximize the accuracy of selection and the precision of the estimated genetic parameters. In the second approach, genotypes may be classified as stable or unstable and the latter removed from the breeding population, thus greatly reducing GEI (Raymond and Lindgren, 1990). Therefore, the breeder has to decide between a number of approaches for choosing source populations and selection zones for planting.

Eucalyptus pellita is one of the most promising Eucalyptus species for plantation in tropical low lands. It has been planted in a wide range of sites in Indonesia, especially in Sumatra and Kalimantan, as part of the development of industrial plantation program. In the first rotation *E. pellita* is considered to be less productive than the tropical acacias (Harwood et al., 1997). The tree improvement program for E. pellita, currently under-way with seedling seed orchards established at several locations in Indonesia, is expected to increase the level of productivity by supplying genetically improved seed. However, no data are available describing the change in productivity at each location resulting from the first generation tree improvement and the use of optimum seed transfer zone to reduce loss of genetic gains due to genotype-environment interactions. A number of different methods have been used by breeders of pine species to investigate the magnitude and predict the effect of GEI on genetic gain. In the present study, several established analytical methods were applied to height and diameter growth data from six year-old trees, to determine the magnitude of the GEI in seven seedling seed orchards of E. pellita established at three locations in Indonesia. These methods included analysis of variance (ANOVA) between pairs of locations and between all three locations, to obtain variance components for family and GEI. The variance components from the analysis were used to calculate a ratio between GEI and family variance components (Shelbourne, 1972), and genetic correlations between pairs of locations (Yamada, 1962). These were then used to predict genetic gains from indirect selection between pairs of locations. We also examined the ratios of gain by selection between pairs of locations, and compared the results with gain when applied to other locations. The main objective of this study was to examine the magnitude of GEI by predicting expected genetic gains for single site selection compared to multi-site selection in seedling seed orchards of *E. pellita* to obtain breeding zones of *E. pellita* in Indonesia.

II. MATERIALS AND METHODS

A. Seedling Seed Orchard

Data used in this study were obtained from seedling seed orchards of *E. pellita* in Indonesia: one location in South Kalimantan containing a single seed orchard and two locations in South Sumatra and Riau established in January 1995 and March 1996, respectively (Figure 1). At each of the sites in Sumatra there were three seed orchards, each with genetic materials derived from three different provenances in Papua New Guinea: South Kiriwo, North Kiriwo and Serissa Village. The seed orchard in South Kalimantan was established in January 1994, and the three provenances were established as a mixed population.

The layout of the seed orchards at the three locations was randomized complete blocks. The number of families of South Kiriwo, North Kiriwo and Serissa Village provenances were 48, 39 and 34, respectively. Families were usually represented by initially planting five trees in a row plot, with 10 replicates in South Kalimantan and Riau, and 12 replicates in South Sumatra. In all cases spacing was 4 x 1.5 m. Site characteristics for the three locations are presented in Table 1.

Height and diameter were measured when the seed orchards were six years old. Prior to the measurements selections within family-plot based on growth and stem form criteria were conducted three times until the best tree in each plot remained. The data from the single retained tree in each plot were used in the analysis.



Figure 1. Approximate location of the study area in Kalimantan and Sumatra, Indonesia

Information	South Kalimantan	South Sumatra	Riau		
Site	Pelaihari	Pendopo	Lipat Kain		
Latitude (South)	3°58'	4°	0°		
Longitude (East)	114°38'	104°	100°32'		
Altitude (m asl)	30	80	50		
Rainfall (mm/year)	2,730	2,781	2,783		
Dry season	May-September	May-September	May-September		
Soil type	Ferralsols	Acrisols	Podzolic		
Soil orders (USDA)	Oxisols	Ultisols	Ultisols		
Temperature (min- max)	23°-33° C	24°-33° C	22°-32°		
Wind	Middle	Low	Low		
Slope	0%	3%	2%		
Vegetation cover	Imperata cylindrica	Imperata cylindrica	Secondary forest		
Ground cultivation	Plowing 2 X	Plowing 2 X	Felling & burning		
Fertilizer	NPK 2 X/year	NPK 1 X/year	NPK 2 X/year		

Table 1. Site characteristics of the seedling seed orchards of *E. pellita* at the three locations

B. Data Analysis

Analysis of variance was undertaken using the individual tree data for height and diameter (dbh) in each seed orchard, using the linear model:

 $Y_{ii} = \mu + R_i + F_i + \varepsilon_{iik}$

where, Y_{ij} is the individual tree data in the *i*-th replicate of the *j*-th family; μ is the overall mean; R_p , F_j and ε_{ijk} are, respectively, the *i*-th replicate effect, the *j*-th family effect, and the error for Y_{ij} . The variance components at each location were estimated by pooling the sums of squares and the degrees of freedom for the three populations of seed orchard; and then comparing the respective pooled mean squares with their expected values. The pooled estimates were used to estimate genetic parameters (Woolaston *et al.*, 1990). The individual tree heritability (h_i^2) was calculated using the following formula (Williams and Matheson, 1994):

 $h_i^2 = 2.5 \, \sigma_f^2 / \left[\sigma_f^2 + \sigma_e^2 \right] \dots (2)$

where, σ_f^2 and σ_e^2 are estimates of the variance component for family and error, respectively.

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The cross-location data set was analyzed to estimate variance components for families and for the family-location interaction (GEI) and then to calculate genetic correlations between pairs of locations, using the following linear model:

 $Y_{iik} = \mu + S_i + R_i(S_i) + F_k + FS_{ik} + \varepsilon_{ijk}.$ (3)

where, Y_{ijk} is the *j*-th replication within the *i*-th site and the *k*-th family; μ and S_i are the overall mean and the *i*-th site effect; respectively; $R_j(S_i)$, F_k , FS_{ik} and \mathcal{E}_{ijk} , respectively, are the *j*-th replicate effect in the *i*-th site, the *k*-th family effect, the interaction of the *i*-th site and *k*-th family, and the error for Y_{ijk} . The variance components across-locations were also calculated by pooling the estimates for the three populations of the seed orchards.

A ratio of the variance components was calculated between the family-site interactions (GEI) and the family variance components $(\sigma_{s}^2/\sigma_{p}^2)$. Shelbourne (1972) suggested that the effects of GEI are "likely to be serious on gains from selection and testing" when the GEI component reaches 50% or more of the family variance component.

The genetic correlation between locations x and y for the same trait $[r_{g(xy)}]$ was calculated using the following formula (Yamada, 1962; Burdon, 1977):

 $r_{g(xy)} = \sigma_f^2 / [\sigma_f^2 + \sigma_{fs}^2]$ (4) where, σ_f^2 and σ_{fs}^2 are estimates of the variance component for family and family-site interaction across pairs locations, respectively.

The predicted genetic gain by indirect selection between two locations (Δg_i) and by direct selection at a single location (Δg_d) were calculated using the following formula (Falconer and Mackay, 1996; Burdon, 1977):

$$\Delta g_i = i_x \cdot h_x \cdot h_y \cdot r_{g \ xy} \cdot \sigma_{Py} \tag{5}$$

where, i_x and i_y are selection intensity, *i* was assumed to be 1.0 for ease of calculation; h_x and h_y are the square root of heritability at locations *x* and *y*, respectively; and σ_{py} is the phenotypic standard deviation of family means at location *y*.

III. RESULTS AND DISCUSSION

A. Genetic Parameter Estimation

The results of the ANOVA across the three locations are presented in Table 2. Family variations were highly significant for both height and diameter. Significant differences were also observed for the family-site interaction (GEI) for both traits, indicating that the GEI was very strong across the three locations. Therefore, genetic gain for the three locations cannot be optimal when selection is based on data from a single environment. This is because GEI reduces heritability and hence genetic gain. It is, therefore, necessary to create and maintain separate breeding populations (Barnes *et.al.*, 1984).

Source of variation	d.f.	Mean square	Prob.	Mean square	Prob.	
		Height	> F	Diameter	> F	
Three locations:						
Family	118	16.266	0.0001	23.548	0.0001	
Family x Site	235	8.670	0.0011	11.558	0.0004	
Error	2508	6.546		8.487		
Two locations:						
South Kalimantan and South Sumatra						
Family	118	11.674	0.0150	15.014	0.0387	
Family x Site	118	7.813	0.0014	10.831	0.0002	
Error	1836	5.373	6.928			
South Kalimantan and Riau						
Family	118	15.119	0.0310	22.512	0.0015	
Family x Site	117	10.696	0.0027	12.979	0.0081	
Error	1635	7.513	9.553			
South Sumatra and Riau						
Family	118	14.422	0.0002	20.683	0.0006	
Family x Site	117	7.494	0.2621	11.317	0.0548	
Error	1545	6.916		9.214		

Table 2. Means squares for height and diameter from the ANOVA across locations

In order to examine the differences in GEI between the three locations, pairwise site analysis were conducted. These show very clear differences in GEI between the pairs of locations. The GEI is highly significant between South Kalimantan and the other locations (South Sumatra and Riau), but is not significant between South Sumatra and Riau (Table 2). These findings are consistent with ratios of GEI to family variance components ($\sigma_{ff}^2/\sigma_{ff}^2$), which, for both traits, is high between South Kalimantan and the other locations (>0.5) and lower between South Sumatra and Riau (Table 3). The differences between ratios suggest that the interactions in these seed orchards are likely to be of practical importance in selection and testing (Shelbourne, 1972).

	Va	Genetic		
Across locations/Trait	Family (σ_f^2)	Interaction (σ_{sf}^{2})	Ratio $(\sigma_{sf}^2 / \sigma_{f}^2)$	correlation $[r_{g(xy)}]$
Three locations				
Height	0.306	0.269	0.880	
Diameter	0.484	0.389	0.804	
Two locations:				
South Kalimantan and South Sumatra				
Height	0.211	0.287	1.362	0.423
Diameter	0.224	0.459	2.051	0.328
South Kalimantan and Riau				
Height	0.260	0.420	1.614	0.383
Diameter	0.581	0.452	0.777	0.563
South Sumatra and Riau				
Height	0.456	0.080	0.175	0.851
Diameter	0.608	0.291	0.478	0.677

Table 3.	Variance components	across locations	and genetic	correlation	between
	pairs of locations for 1	height and diam	eter $[r_{g(xy)}]$		

 $r_{g(xy)}$ was calculated using Eq. 4

Genetic correlations between pairs of locations for the two traits, calculated using Eq. (4), are presented in Table 3, together with the variance components across locations. The correlations were moderately high between South Sumatra and Riau; 0.68 for dbh and 0.85 for height. They were lower between South Kalimantan and the other locations (0.32 to 0.56). The correlation results support the ratios of GEI to family variance components for the three locations (Table 3) and suggest that family ranking in Riau is similar to that in South Sumatra, but that they are different from the ranking in South Kalimantan. This is probably due to the close geographic distances between South Sumatra and Riau as both are on the island of Sumatra and associated with differences in environmental conditions. However, the climate of the three locations (Table 1) is similar, with a pronounced dry season from May to September and a wet season from October to April, so the differences may be associated with soil type or edaphic factors. According to the USDA soil classification, the Acrisols in South Sumatra and Podzolic soils in Riau are in the same order as the Ultisols, they are red to yellow in color and quite acidic, often with a pH of less than 5. In contrast, the Ferralsols in South Kalimantan are classified as Oxisols, best known for their occurrence in tropical rain forests, and they have a red or yellowish color due to the high concentration of iron (III) and aluminum oxides, and hydroxides (Buol *et al.*, 2003). This observation concurs with Shelbourne (1972) and Zobel and Talbert (1984), who suggested that, although there are many causes of GEI, in general they are generally more closely related to edaphic factors than to climatic factors.

B. Predicted Genetic Gains and Practical Implication

The genetic gains from indirect selection predicted by Eq. (5) and direct selection, Eq. (6), comparing pairs of locations, are shown in Table 4 and Figure 2. The predicted genetic gain at each location is the result of direct selection, and those of the correlated response when it is applied at other locations are indirect selections. The genetic gains by direct selection appeared to be greater than those associated with indirect selection. With respect to direct selection, the genetic gains for diameter and height in South Kalimantan were 13-32% greater than those in Riau, and 64-137% greater than those in South Sumatra. The differences in the genetic gains for both traits may be due to the higher heritability in South Kalimantan than that at the other locations (Table 4). Except in South Sumatra, the genetic gains for diameter were greater than for height.

The relative ratios of genetic gains, comparing indirect selection with direct selection, were 76% between South Sumatra and Riau. In contrast, the average ratio between South Kalimantan and the other locations was 40% (Table 4). Thus, the genetic gains obtained by direct selection in South Kalimantan would be reduced by 60% if the selected entries were transferred to the other locations, and vice versa, while the reductions in genetic gain resulting from such transfers between South Sumatra and Riau would be only 24% (Figure 3).



Figure 2. Expected gain by selection between pairs of locations. Charts were drawn using the data presented in Table 4. Black bars and light bars represent absolute gain by direct selection and indirect selection, respectively

	Phenotypic	Herita	Selection					
Correlated response	$SD(\sigma_p)$	$\begin{array}{ccc} { m SD} & { m bility} & { m Sou} \ (\sigma_{ ho}) & (b_i^{\ 2}) & { m Kalima} \end{array}$		South Sumatra	Riau			
Height:								
South Kalimantan	2.297	0.22	<u>0.510 (100.0)</u>	0.169 (33.21)	0.152 (29.87)			
South Sumatra	2.267	0.14	0.131 (42.35)	0.310 (100.0)	0.262 (84.71)			
Riau	2.851	0.14	0.189 (49.01)	0.330 (85.50)	0.386 (100.0)			
Diameter:								
South Kalimantan	2.739	0.24	<u>0.658 (100.0)</u>	0.162 (24.59)	0.310 (47.18)			
South Sumatra	2.048	0.14	0.091 (33.76)	0.277 (100.0)	0.209 (75.58)			
Riau	3.437	0.17	0.389 (67.13)	0.351 (60.59)	0.580 (100.0)			

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Table 4.	Expected	l gain and	ratio of	t gain l	ov se.	lection	between	pairs o)t I	locations
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Note: b_i^2 (individual heritability) was calculated using Eq. 2, Absolute gains were calculated using Eqs. 5 and 6, Gain predictions by direct selection at the location (Δg_a) are underlined and ratios of gains by indirect selection (%) are given in parentheses.



Figure 3. Seed transfer and reduction of genetic gain by selection between pairs of locations

Most breeders of pine species suggest that the GEIs in their trials are not of practical importance in selection for growth traits, because the ratio of GEI to family variance components is low (Pederick, 1990; Woolaston *et al.*, 1991; Johnson, 1992; Pswarayi et al., 1997; Gwaze et al., 2001). Such results suggest that selection for generally adapted populations is possible for these species, without incurring any significant losses in genetic gain. However, the results from the current study identified very strong GEI, both when comparing the three sites together and as pairs (Table 2). Moreover, the effects of GEI from the pair-wise site analysis predicted a serious loss of genetic gain if the seeds from Kalimantan orchard were transferred to Sumatra and vice versa (Table 4). These findings suggested that the genetic gains could be maximized by breeding within regions or by grouping locations with similar environments; a ratio between GEI and family components lower than 0.5 would be appropriate within such group. This would provide a better approach to choosing source populations and selecting planting zones.

Although, several private companies concerns are undertaking tree improvement programs for *E. pellita* in Indonesia, they do not collaborate and each has a limited range of testing sites. Thus, the expected gains at other locations have not yet been confirmed. However, the plantation establishment program in Indonesia is being conducted at national scale, focusing particularly on Sumatra and Kalimantan, so the results reported herein could be applied in the form of guidelines to control seed transfer for *E. pellita*, thus minimizing the loss of genetic gain due to GEI.

IV. CONCLUSION

The magnitude of GEI in the seedling seed orchards of *E. pellita*, indicated by the ratio of variance components between GEI and family, and genetic correlations between pairs of locations suggests that the effect of GEI is of practical importance when the results of selections are used on other islands. Calculated losses in genetic gain as a result of inter-island transfer (from Kalimantan to Sumatra and vice versa) were generally high; reductions of approximately 60%. In contrast, the loss of genetic gains resulting from transfer on a single island, Sumatra, was only 24%. Therefore, to maximize the efficiency of selection between different locations and to minimize losses in genetic gain, the breeding population in South Kalimantan should be developed separately from those in South Sumatra and Riau.

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