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Genotypic variation of *Cunninghamia lanceolata* revealed by phenotypic traits and SRAP markers

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Abstract: The success of a tree breeding program largely depended on the available genetic variability of the germplasms. Our present study aimed to assess the phenotypic variation and DNA variability using sequence-related amplified polymorphism (SRAP) markers among 50 *Cunninghamia lanceolata* (Chinese fir) genotypes. Extensive phenotypic variations ($p < 0.05$ or 0.01) were found for all the growth and wood property traits (height, diameter at breast height, stem volume, and wood basic density, hygrosopicity, heart-wood ratio, tracheid length, tracheid diameter and tracheid length-diameter ratio) with coefficients of variation spanning from 6.8 to 31.3%. At the DNA level, thirty-five SRAP primer combinations produced 498 bands with 89.4% polymorphism across genotypes; moreover, the Nei's gene diversity was detected to be ranged between 0.204 and 0.373 (mean = 0.279), while the Shannon's Information Index stretched from 0.324 to 0.555 with an average value of 0.427. Significance ($p < 0.01$) of the variability of SRAP polymorphism among genotypes was further demonstrated by AMOVA. These results indicated a relatively high level of genetic diversity in genotypes. The SRAP' dendrogram additionally revealed that these genotypes could be split into 7 clusters with higher discriminating capacity over that of phenotype. Notably, a total of 99 statistically significant ($p < 0.05$) marker-trait associations related to the growth and wood property traits were identified. These marker-trait associations corresponded to 77 different SRAP markers with R^2 (percentage of the phenotypic variation explained by marker) ranging from 8.3 to 26.4%.

Additional key words: Chinese fir, wood property, molecular marker, dendrogram and marker-trait association.

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Introduction

Cunninghamia lanceolata (Lamb.) Hook, commonly known as Chinese fir, is one of the most important

conifer tree species in China due to its great afforestation values in both timbers and ecological contributions. It has been cultivated for over 3,000 years and currently occupies ~25% of man-made planta-

Table 1. Descriptive statistics of the growth and wood property traits for fifty 24-year-old Chinese fir genotypes. H, height; DBH, diameter at breast height; V, stem volume; WBD, wood basic density; Hy, hygroscopicity; HR, heart-wood ratio; TL, tracheid length; TD, tracheid diameter; TLDR, tracheid length-diameter ratio; SD, standard deviation; F, the ANOVA (analysis of variance) parameter F (*p < 0.05, **p < 0.01); CV, coefficient of variation; R, repeatability

No.	Genotype	H (m)	DBH (cm)	V (m ³)	WBD (g/cm ³)	Hy (%)	HR (%)	TL (μm)	TD (μm)	TLDR
1	c17	13.0	26.0	0.3792	0.2713	304.3	53.8	3030.3	46.0	73.1
2	c18	13.5	28.7	0.4565	0.3170	250.9	55.4	3065.7	47.4	78.3
3	c22	12.3	27.2	0.3924	0.3093	264.6	47.3	3097.4	42.5	71.0
4	c49	13.7	30.3	0.5157	0.2965	272.7	52.3	3423.7	46.5	75.1
5	c51	15.0	35.7	0.7749	0.3038	264.0	59.0	3244.7	40.3	72.1
6	c53	12.8	28.4	0.4322	0.3134	254.8	50.0	3174.0	40.8	81.6
7	c54	14.8	34.0	0.6939	0.2673	309.0	60.9	3318.4	46.3	72.0
8	c58	11.7	20.9	0.2163	0.2800	292.6	42.2	3094.1	41.3	75.7
9	c59	13.5	30.5	0.5126	0.3027	267.4	51.1	3283.6	41.1	68.5
10	c62	12.5	27.7	0.3959	0.3153	251.9	51.1	3304.5	40.8	73.5
11	c70	13.0	28.7	0.4419	0.3215	246.9	44.2	3120.5	46.0	74.3
12	c73	12.3	29.3	0.4319	0.2720	304.4	49.6	3508.9	43.7	78.5
13	c78	12.7	26.9	0.3877	0.3662	207.9	52.9	3334.4	41.4	80.2
14	c80	14.3	34.7	0.6933	0.3054	262.4	54.6	3078.8	42.7	75.8
15	c87	11.0	21.8	0.2193	0.3319	237.3	44.7	3173.5	44.7	72.5
16	c91	12.0	24.0	0.2844	0.2959	274.3	46.4	2827.4	45.1	67.7
17	c97	12.8	27.7	0.4384	0.3119	259.4	57.7	3571.3	48.6	77.9
18	c98	13.0	30.0	0.4805	0.3252	242.9	57.8	3398.1	54.5	90.4
19	c99	13.3	29.6	0.4956	0.3389	230.3	55.8	2729.5	42.5	63.8
20	c100	12.0	25.2	0.3146	0.3246	243.1	51.1	3398.8	45.1	67.9
21	c101	12.7	27.8	0.4138	0.3292	241.5	48.5	3162.2	44.3	66.6
22	c102	15.5	35.8	0.7891	0.3225	244.7	67.4	3224.4	42.1	71.5
23	c103	15.7	35.7	0.7952	0.2790	293.2	58.5	3555.6	49.0	77.2
24	c104	14.7	32.4	0.6453	0.2790	293.2	58.5	3555.6	49.0	77.2
25	c106	14.0	33.9	0.6507	0.3204	247.3	60.8	3363.3	44.1	72.9
26	c107	11.5	27.4	0.3586	0.2995	268.5	48.3	3286.9	50.8	67.9
27	c108	13.5	31.6	0.5465	0.3479	224.8	66.0	3461.5	42.0	71.1
28	c109	14.0	33.1	0.6248	0.3012	266.7	68.3	3311.7	44.6	72.2
29	c125	11.5	26.1	0.3280	0.2667	310.0	59.9	3285.8	53.1	75.2
30	c129	12.0	27.6	0.3867	0.3013	266.6	61.9	3748.5	43.7	63.0
31	c130	13.5	38.9	0.8666	0.2970	272.8	61.5	3282.3	41.5	62.2
32	c134	13.7	30.7	0.5247	0.2563	325.8	50.0	3125.9	52.8	73.8
33	c135	15.7	39.7	0.9800	0.2774	298.8	56.0	3071.5	49.9	70.3
34	c139	12.3	27.0	0.3893	0.2903	284.5	58.1	3223.8	45.9	75.2
35	c142	11.8	23.5	0.2748	0.3467	229.7	57.1	3298.5	48.0	71.8
36	c39	11.3	28.8	0.3903	0.3226	254.4	60.9	3122.8	44.9	67.6
37	c43	15.8	42.8	1.1422	0.2584	321.7	50.8	3242.6	47.0	75.8
38	c44	13.0	27.9	0.4223	0.2975	270.8	58.9	3688.3	48.0	82.7
39	c28	15.5	37.9	0.8874	0.3042	264.0	57.9	3191.6	42.8	68.1
40	c69	15.0	34.3	0.7216	0.2951	285.1	52.9	3168.2	48.1	84.7
41	c148	14.3	30.9	0.5560	0.3250	243.2	57.7	3370.9	48.3	79.7
42	c36	12.5	30.9	0.4878	0.2900	279.5	60.0	3647.6	44.7	61.9
43	c37	12.0	29.6	0.4355	0.3282	239.5	63.8	3761.0	50.7	70.7
44	c149	12.7	27.1	0.4280	0.2767	297.9	51.0	3490.5	47.8	59.2
45	c6	12.7	28.8	0.4467	0.3461	223.9	60.5	3620.7	44.5	69.5
46	c16	12.3	25.6	0.3374	0.2931	277.0	55.2	2989.7	41.0	70.4
47	c63	12.3	24.0	0.2964	0.3115	256.1	44.1	2834.1	42.9	71.3
48	c10	14.5	34.7	0.7087	0.2933	279.1	46.6	3093.4	47.6	61.9
49	c23	12.8	28.1	0.4155	0.3249	244.5	56.9	3213.1	43.4	76.0
50	c29	13.3	29.7	0.4825	0.2898	283.8	59.6	3195.6	44.5	65.9
	Mean	13.2	30.0	0.5138	0.3048	266.6	55.1	3275.3	45.5	72.5
	SD	1.3	4.6	0.1992	0.0246	26.7	6.2	227.7	3.5	6.2
	F	4.64**	4.63**	4.87**	2.50**	2.50**	1.71*	1.56*	3.65**	1.75**
	CV (%)	7.7	12.7	31.3	9.2	11.5	10.0	9.7	6.8	11.8
	R	0.78	0.78	0.79	0.60	0.60	0.42	0.36	0.73	0.43

tions in southern China (Shi et al. 2010). Based on the knowledge and availability of genetic variability for selection (Bian et al. 2014), remarkable successes in breeding have already been achieved for this species, e.g. large-scale collection of elite germplasms, establishment of first and second and third generation seed orchards, and significant gains of superior clones. However, in most of the case, only a small proportion of the variability is actually used in the breeding programs, which finally results in a narrowing genetic base for further breeding. While a considerable number of potential elite germplasms (genotypes) conserved in *ex situ* banks were underutilized and frequently redundant because of the lack of adequate passport data for most of the lines. Thus, proper characterization of these genetic resources appeared to be highly required.

Diversity in trees could be assessed by measuring variation in phenotypic/morphological traits such as flowers, fruits, growth habit, and quantitative agronomic/economical traits like yield potential and wood property traits, etc., which are of direct interest to breeders, but this approach has certain limitations: (1) insufficient genetic information obtained, and (2) strong influence of genotype \times environment (G \times E) interactions (Rao 2004; Nybom et al. 2014). Molecular technology offers an avenue for the determination of informative DNA variation complemented to phenotype regardless of growth, differentiation, development, or environmental effects (Agarwal et al. 2008; Zhao et al. 2009). Polymorphism uncovered by DNA markers enables the breeders to discriminate the germplasm at a very precise level and to elucidate the genetic structure, diversity and relationship, as well as the marker-trait associations. Among the present available marker systems, the PCR-based markers have become popular because their application does not need any prior sequence information, and is easy to performed (Tatikonda et al. 2009). SRAP (sequence-related amplified polymorphism) technique is one such PCR-base marker system widely used for the plant genetic/genomic studies (Li and Quiros 2001; Budak et al. 2004; Baloch et al. 2010). It appeared to be simple, efficient and cost-effective. Typically, it specifically targeted to the genome open reading frame (ORF) sequences providing more genetic information associated with phenotype (Ferriol et al. 2003; Uzun et al. 2009; Castonguay et al. 2010; Rana et al. 2013).

The objectives of this study were to survey the variability of growth and wood property traits of 50 *ex situ* conserved Chinese fir genotypes and their SRAP polymorphisms in terms of genetic diversity and relationship and significant marker-trait associations, aiming to profile these germplasms properly for further breeding use.

Materials and Methods

Plant materials

Fifty Chinese fir genotypes were analyzed in this study (Table 1). The top 35 genotypes (No. 1 to 35) belonged to the Lechang provenance (Guangdong, China), and the next every three genotypes were derived from the provenance of Guangxi (No. 36 to 38), Guizhou (No. 39 to 41), Hunan (No. 42 to 44), Jiangxi (No. 45 to 47) and Fujian (No. 48 to 50) respectively (China). These genotypes were conserved in the Longshan State Forest Farm (Guangdong, China) by grafting with a Latin square design of 5 \times 5 m spacing since 1985. Each genotype has four ramets that presented to be similar in size and vigor. Trees were maintained through standard commercial practices. 24-year-old plants were then measured for the traits of height (H) and diameter (diameter at breast height, DBH), and the randomly selected three ramets for each genotype were subjected to the wood quality analysis assay.

Measurement of growth and wood property traits

DBH was measured at 1.3 m by a measuring tape and H by poles for each tree. Stem volume (V) was calculated according to the formula of $V = 5.8777042 \times 10^{-5} \times \text{DBH}^{1.9699831} \times \text{H}^{0.8964616}$ (Zheng et al. 2012).

A 5.02 mm increment core was taken at breast height (1.3 m) from every sampled tree by using a tree growth cone, and immediately stored in a plastic tube with two ends sealed, and then subjected to the wood quality analysis assays. Wood basic density (WBD) was evaluated by using a water displacement method: weight in water (W_1) and oven dry weight (W_2) in grams were taken for every sample, and the WBD (g/cm^3) was then estimated following the formula of $\text{WBD} = 1 / ((W_1 / W_2) - 0.346)$ (Zheng et al. 2012). While the hygroscopicity (Hy) can be evaluated according to the formula of $\text{Hy} = (W_1 - W_2) / W_2$. The heart-wood ratio (HR) was determined theoretically based the value of $(r^2 \times \Pi) / (R^2 \times \Pi)$ (r and R stand for the radius of heartwood and whole wood respectively). The tracheid length (TL) and diameter (TD) were assessed by the method as described by Huang et al. (2004)

DNA extraction and SRAP procedure

Total genomic DNA was extracted from the Chinese fir mature leaves with a DNAsure Plant Kit (TIANGEN, Beijing, China). SRAP PCR amplifications were carried out using 35 optimal primer combinations including 16 forward primers (Me1, Me2, Me3, Me4, Me9, Me10, Me11, Me12, Me13,

Table 2. Phenotypic correlations among growth and wood property traits in Chinese fir. H, height; DBH, diameter at breast height; V, stem volume; WBD, wood basic density; Hy, hygroscopicity; HR, heart-wood ratio; TL, tracheid length; TD, tracheid diameter; TLDR, tracheid length-diameter ratio. *p < 0.05, **p < 0.01

	H	DBH	V	WBD	Hy	HR	TL	TD	TLDR
H	1.0000								
DBH	0.8678**	1.0000							
V	0.9024**	0.9798**	1.0000						
WBD	0.1577*	-0.2083**	-0.2141**	1.0000					
Hy	0.1493	0.1983*	0.2067**	-0.9912**	1.0000				
HR	0.2117**	0.3181**	0.2724**	0.0121	-0.0200	1.0000			
TL	-0.0216	0.0446	-0.0036	-0.0369	0.0354	0.1987*	1.0000		
TD	0.0550	0.0350	0.0490	-0.1107	0.1253	0.0892	0.1135	1.0000	
TLDR	0.0492	-0.0641	-0.0402	0.0490	-0.0308	0.0318	0.0634	0.1632*	1.0000

Table 3. The employed SRAP primer combinations and their performance in Chinese fir (n=50). TNB, total number of bands; NPB, number of polymorphic bands; PPB, percentage of polymorphic bands; PIC, polymorphic information content; MI, marker index; RP, resolving power; h, Nei's gene diversity; I, Shannon's Information Index; SD, standard deviation

SRAP Primer combination (forward/reverse, sequence 5' to 3')	TNB	NPB	PPB (%)	PIC	MI	RP	h	I
Me1(TGAGTCCAAACCGGATA)/Em1(GACTGCGTACGAATTAAT)	12	12	100.0	0.314	3.768	5.32	0.316	0.486
Me1(TGAGTCCAAACCGGATA)/Em19(GACTGCGTACGAATTACG)	16	15	93.8	0.286	4.292	7.56	0.303	0.458
Me1(TGAGTCCAAACCGGATA)/Em20(GACTGCGTACGAATTTAG)	9	9	100.0	0.284	2.556	3.00	0.248	0.395
Me2(TGAGTCCAAACCGGAGC)/Em26(GACTGCGTACGAATTCGG)	14	13	92.9	0.257	3.343	5.48	0.254	0.391
Me3(TGAGTCCAAACCGGAAT)/Em17(GACTGCGTACGAATTATG)	13	11	84.6	0.289	3.178	5.72	0.294	0.442
Me4(TGAGTCCAAACCGGACC)/Em5(GACTGCGTACGAATTAAC)	15	15	100.0	0.291	4.365	6.92	0.305	0.464
Me4(TGAGTCCAAACCGGACC)/Em21(GACTGCGTACGAATTTCCG)	17	16	94.1	0.276	4.415	7.28	0.291	0.444
Me9(TGAGTCCAAACCGGACA)/Em14(GACTGCGTACGAATTCTC)	15	13	86.7	0.270	3.511	5.72	0.273	0.419
Me10(TGAGTCCAAACCGGACG)/Em20(GACTGCGTACGAATTTAG)	13	12	92.3	0.301	3.612	5.48	0.305	0.465
Me10(TGAGTCCAAACCGGACG)/Em21(GACTGCGTACGAATTTCCG)	13	12	92.3	0.271	3.252	5.00	0.266	0.410
Me10(TGAGTCCAAACCGGACG)/Em26(GACTGCGTACGAATTCGG)	16	15	93.8	0.304	4.562	7.32	0.324	0.489
Me11(TGAGTCCAAACCGGACT)/Em5(GACTGCGTACGAATTAAC)	18	16	88.9	0.271	4.337	8.24	0.293	0.436
Me11(TGAGTCCAAACCGGACT)/Em13(GACTGCGTACGAATTCTA)	15	14	93.3	0.300	4.199	7.32	0.319	0.477
Me11(TGAGTCCAAACCGGACT)/Em20(GACTGCGTACGAATTTAG)	16	14	87.5	0.253	3.542	5.76	0.255	0.391
Me11(TGAGTCCAAACCGGACT)/Em21(GACTGCGTACGAATTTCCG)	10	8	80.0	0.278	2.224	3.36	0.251	0.390
Me12(TGAGTCCAAACCGGAGG)/Em1(GACTGCGTACGAATTAAT)	16	15	93.8	0.269	4.037	6.40	0.276	0.425
Me12(TGAGTCCAAACCGGAGG)/Em19(GACTGCGTACGAATTACG)	15	15	100.0	0.341	5.115	8.20	0.373	0.555
Me13(TGAGTCCAAACCGGAAA)/Em5(GACTGCGTACGAATTAAC)	12	10	83.3	0.261	2.609	4.00	0.243	0.379
Me13(TGAGTCCAAACCGGAAA)/Em18(GACTGCGTACGAATTAGC)	16	13	81.3	0.217	2.823	4.48	0.204	0.324
Me15(TGAGTCCAAACCGGAGA)/Em1(GACTGCGTACGAATTAAT)	13	10	76.9	0.282	2.819	5.32	0.284	0.424
Me15(TGAGTCCAAACCGGAGA)/Em26(GACTGCGTACGAATTCGG)	15	10	93.3	0.273	3.821	6.48	0.284	0.426
Me16(TGAGTCCAAACCGGATA)/Em1(GACTGCGTACGAATTAAT)	15	14	93.3	0.271	3.793	5.80	0.272	0.424
Me16(TGAGTCCAAACCGGATA)/Em19(GACTGCGTACGAATTACG)	18	16	88.9	0.235	3.760	6.04	0.236	0.369
Me17(TGAGTCCAAACCGGTAG)/Em19(GACTGCGTACGAATTACG)	12	11	91.7	0.291	3.202	5.00	0.289	0.440
Me17(TGAGTCCAAACCGGTAG)/Em21(GACTGCGTACGAATTTCCG)	17	17	100.0	0.291	4.947	7.64	0.310	0.472
Me18(TGAGTCCAAACCGGCAT)/Em19(GACTGCGTACGAATTACG)	16	13	81.3	0.239	3.109	5.12	0.234	0.364
Me19(TGAGTCCAAACCGGTTG)/Em1(GACTGCGTACGAATTAAT)	13	9	69.2	0.236	2.123	4.04	0.220	0.337
Me19(TGAGTCCAAACCGGTTG)/Em18(GACTGCGTACGAATTAGC)	16	14	87.5	0.301	4.214	7.60	0.325	0.480
Me19(TGAGTCCAAACCGGTTG)/Em19(GACTGCGTACGAATTACG)	14	13	92.9	0.303	3.941	6.92	0.319	0.479
Me19(TGAGTCCAAACCGGTTG)/Em22(GACTGCGTACGAATTGTC)	9	7	77.8	0.310	2.171	4.28	0.299	0.439
Me20(TGAGTCCAAACCGGTGT)/Em1(GACTGCGTACGAATTAAT)	14	13	92.9	0.310	4.032	6.76	0.330	0.487
Me20(TGAGTCCAAACCGGTGT)/Em17(GACTGCGTACGAATTATG)	13	11	84.6	0.254	2.793	4.52	0.243	0.376
Me20(TGAGTCCAAACCGGTGT)/Em19(GACTGCGTACGAATTACG)	18	17	94.4	0.254	4.316	6.64	0.261	0.406
Me20(TGAGTCCAAACCGGTGT)/Em26(GACTGCGTACGAATTCGG)	13	12	92.3	0.278	3.336	5.00	0.274	0.423
Me21(TGAGTCCAAACCGGTCA)/Em19(GACTGCGTACGAATTACG)	11	10	90.9	0.247	2.470	2.92	0.209	0.349
Total	498	445	89.4	-	-	-	-	-
Mean	14.2	12.7	89.9	0.277	3.560	5.79	0.279	0.427
SD	2.4	2.6	7.3	0.026	0.792	1.44	0.038	0.050

Me15, Me16, Me17, Me18, Me19, Me20 and Me21) and 11 reverse primers (Em1, Em5, Em13, Em14, Em17, Em18, Em19, Em20, Em21, Em22 and Em26) (Table 3). Each 25 μ l PCR reaction mixture consisted of 0.5 μ l of genomic DNA (about 50 ng), 0.5 μ l forward primer (10 μ mol/l), 0.5 μ l reverse primer (10 μ mol/l), 12.5 μ l 2 \times Taq Plus PCR MasterMix (TIANGEN, Beijing, China) and 11 μ l double distilled water. The thermal cycling condition was the same as that of Li and Quiros (2001). The PCR products were separated by electrophoresis through 2% agarose gels with DL2000 DNA markers (TIANGEN, Beijing, China) as references. The gels were then stained with ethidium bromide and visualized under UV light using a Universal Hood II imaging system (Bio-Rad, Hercules, CA, USA).

Statistical analysis

Phenotypic variability was displayed by the parameter of F and CV (coefficient of variation) as assessed by one-way analysis of variance (ANOVA) process in Statistical Analysis System (SAS V 8.1) (SAS Institute, Cary, North Carolina). Repeatability (R) for each trait was evaluated with following formula: $R = 1 - 1/F$. The phenotypic correlation coefficient for each pair of traits was calculated by SAS PROC CORR program.

The SRAP bands were scored manually and recorded as a binary matrix, with codon of 1 and 0 representing presence and absence of a band at a particular location in each lane respectively. The polymorphism information content (PIC) for each primer set was calculated with software PowerMarker V 3.25 (Liu and Muse 2005). The marker index (MI) could be estimated with the formula: $MI = PIC \times NPB$, where NPB is the number of polymorphic bands per primer set (Powell et al. 1996). The primer resolving power (RP) was evaluated using the formula: $RP = \sum I_i$, where $I_i = 1 - (2 \times |0.50 - p|)$, and p is the proportion of the 50 genotypes containing the I band (Prevost and Wilkinson 1999). The POPGENE V 1.31 was employed to estimate the parameter of h (Nei's gene diversity) and I (Shannon's Information Index) (Yeh et al. 1999). The program GenAlEx V 6.5 incorporated with Microsoft Excel 2010 software was implemented to conduct the AMOVA (analysis of molecular variance) procedure (Peakall and Smouse 2012).

For cluster analysis, the Numerical Taxonomy Multivariate Analysis System (NTSYSpc V 2.10e) (Rohlf 2000) was applied to estimate the genetic similarities of the genotypes with the simple matching coefficient (SM) and Dice coefficient for phenotype and SRAP data respectively, and then separately generate a dendrogram using the unweighted pair

group method with arithmetic average (UPGMA) with the SAHN module. The goodness-of-fit of the dendrogram to the original genetic similarity matrix was validated by measuring the Cophenetic Value (COPH) and Matrix Comparison Plot (MXCOMP) using NTSYSpc packages.

Marker-trait association assessment was performed by software TASSEL V 5.0 with general linear model (GLM) and mixed linear model (MLM) (Bradbury et al. 2007). The identified markers displaying a statistic value of $p < 0.05$ in both models were regarded as the significant markers associated with traits.

Results

Genotypic variation in growth and wood property traits

All the growth and wood property traits varied significantly among genotypes ($n = 50$), as evidenced by ANOVA results (F value; Table 1). It was also observed that the coefficient of variation (CV) spanned from 6.8 (tracheid diameter) to 31.3% (stem volume) across traits. While the repeatability for the traits of height (H), DBH (diameter at breast height), stem volume (V), and wood basic density (WBD), hygroscopicity (Hy), and tracheid diameter (TD) seemed to be relatively high (≤ 0.6).

Further analysis revealed that the tree height, DBH, V and HR (heart-wood ratio) were positively correlated to each other at a significant level ($p < 0.01$), as shown in Table 2. Positive correlations were also found between Hy and DBH and V , and for HR with TL (tracheid length) and TD with TLDR (tracheid length-diameter ratio) ($p < 0.05$ or 0.01). Typically, negative correlations were consistently observed between WBD and growth and Hy ($p < 0.05$ or 0.01).

Genotypic variation in SRAPs

Thirty-five SRAP primer combinations consistently resulted in a detectable PCR profile with clear, stable and rich polymorphism bands as represented by Fig. 1. These primer sets produced 498 scorable bands including 445 polymorphic loci with a PPB (percentage of polymorphic bands) of 89.4% across the present Chinese fir genotypes (Table 3). The number of bands for each assay ranged from 9 (Me1/Em20 and Me19/Em22) to 18 (Me11/Em5, Me16/Em19 and Me20/Em19) with an average of 14.2 bands and 12.7 polymorphic loci per primer combination. The PPB varied from 69.2 (Me19/Em1) to 100.0% (Me1/Em1, Me1/Em20, Me4/Em5, Me12/Em19 and Me17/Em21) with an average of

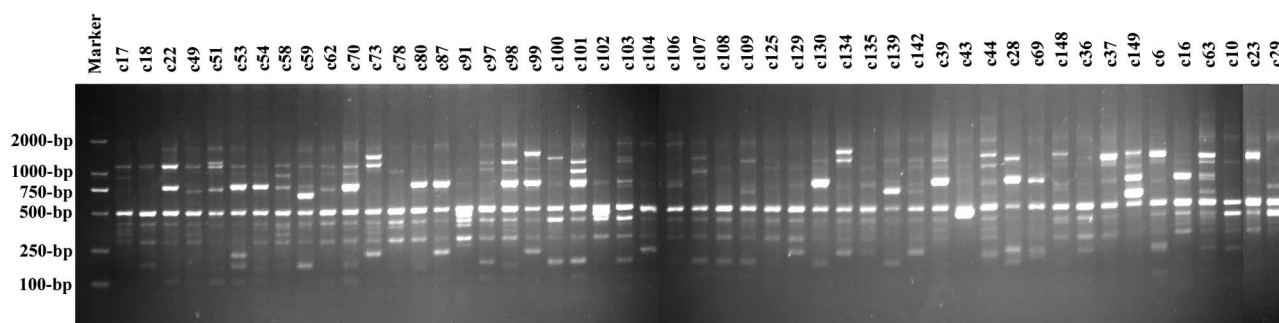


Fig. 1. The represented SRAP profile with primer combination Me20/Em19 for the present 50 Chinese fir genotypes

89.9% per primer set. Each primer combination was found to express a reasonably informative PIC (polymorphic information content), MI (marker index) and RP (resolving power) for the genotypes with a stretching of 0.217 – 0.314 (mean = 0.277), 2.123 – 5.115 (mean = 3.560) and 2.92 – 8.24 (mean = 5.79) respectively. While the Nei's gene diversity (h) ranged from 0.204 to 0.373 (mean = 0.279), and the Shannon's Information Index (I) presented an average value of 0.427 (0.324 – 0.555).

Significance ($p < 0.01$) of the variability of SRAP polymorphism among genotypes was further demonstrated by AMOVA (analysis of molecular variance) (Table 4). When considering the provenance aspect, it was also found that a total of 93.0% variation resided within provenances (Guangdong, Guangxi, Guizhou and Hunan, Jiangxi and Fujian), while there had low (7.0%) but significant genetic variation among provenances. Typically, significant variation of the genotypes within Lechang provenance was observed at SRAPs.

Clustering of genotypes

For genotype clustering, we preliminarily constructed a dendrogram based on the growth and wood property data (matrix correlation $r = 0.98$) (Fig. 2 A). It was found that the present 50 genotypes could be classified into 23 groups with a simple matching coefficient (SM) lower than 0.56. All the Groups seemed to harbor a limited number of members as less than five.

To further our understanding of the relationship of the genotypes at DNA level, UPGMA dendrogram was reconstructed with SRAP markers using DICE's coefficients (matrix correlation $r = 0.73$) (Fig. 2 B). In general, the 50 Chinese fir genotypes

could be grouped into 7 clusters (I to VII). Cluster I, II (except c28 from Guizhou), III, V and VI were composed by the genotypes derived from Lechang (Lechang provenance, Guangdong), while cluster IV harbor a complex origin (provenance) including those from Lechang (Guangdong, e.g. c125, c129, c130, c135 and c142), Guangxi (c39, c43 and c44), Guizhou (c69 and c148), Hunan (c36, c37 and c149), Jiangxi (c6, c16 and c63) and Fujian (c10). The genotypes of c23 and c29 were grouped into a separate cluster (VII) reflecting their closed relationship in provenance (Fujian).

SRAP markers associated with growth and wood property traits

In the next step, we aimed to determine the association of the SRAP markers with growth and wood property traits in Chinese fir. A total of 99 statistically significant ($p < 0.05$) marker-trait associations were detected using TASSEL as confirmed by both GLM and MLM (Table 5). These marker-trait associations corresponded to 77 different markers with R^2 (percentage of the phenotypic variation explained by marker) ranging from 8.3 to 26.4%; 59 of these were found to be associated with only one trait, while the other 18 SRAP markers were linked to more than one traits (e.g. Me16/Em1₁₀₀₀ significantly associated with DBH, V, WBD and Hy).

Discussion

In this study, we evidenced that the present 50 Chinese fir genotypes varied significantly in growth and wood property traits and SRAP polymorphisms suggesting a comprehensive diversity in phenotype

Table 4. Analysis of molecular variance (AMOVA)

Source	df	SS	MS	Est. Var.	%	Stat	Value	P value
Within genotypes	49	6981.760	142.485	71.242	100.0%	PhiPT	1.000	0.001
Among provenances	5	462.156	92.431	4.796	7.0%	PhiPT	0.065	0.001
Within provenances	44	3028.724	68.835	68.835	93.0%	PhiPT	0.065	0.001
Within Lechang provenance	34	4732.114	139.180	69.590	100.0%	PhiPT	1.000	0.001

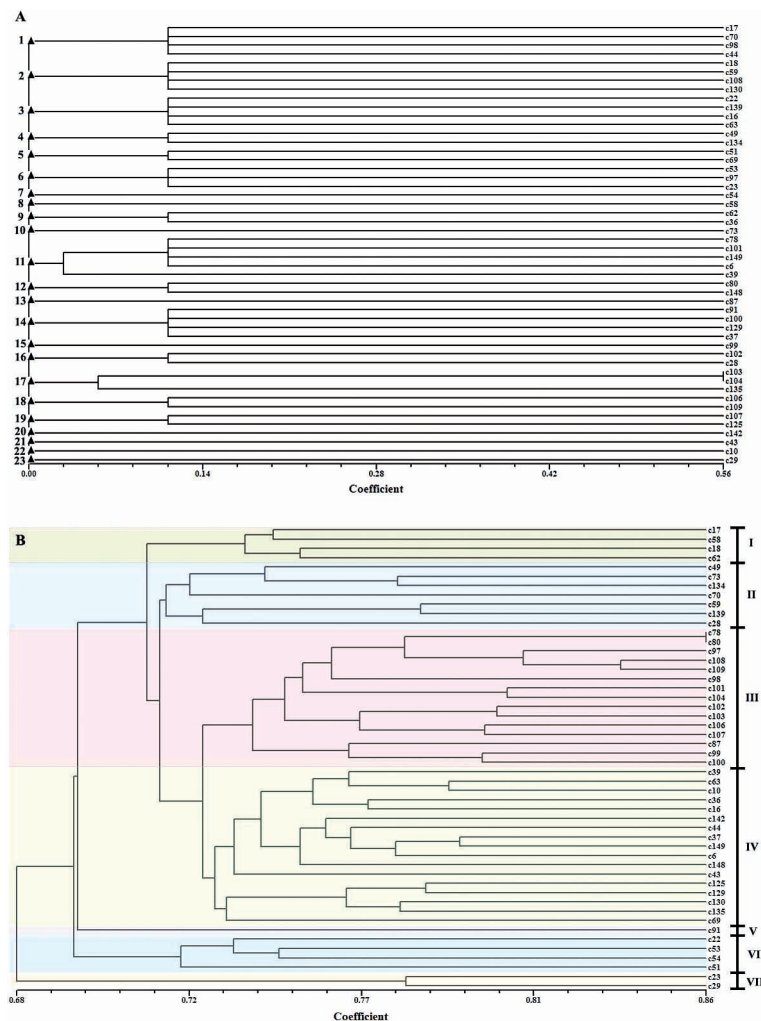


Fig. 2. UPGMA dendrograms of the Chinese fir genotypes based on growth and wood property data (A), and SRAP markers (B) respectively. A: Preliminary clustering is indicated with Arabic numerals followed by a black triangle; the coefficient was obtained by the simple matching method. B: Clusters are shown in different background, and indicated with scale and upper roman numerals; the coefficient was obtained by Dice method

and genotype. SRAP' clustering analysis further revealed that these genotypes could be split into 7 groups that help us to select the breeding parents with more distant genetic relationships. Herein, we also identified a set (77) of growth and wood property associated SRAP markers. These markers may be useful for the next marker-assisted selection (MAS) programs in Chinese fir.

To rapidly obtain the Chinese fir genetic information in genotypes, PCR-based SRAP marker technique was herein employed. In fact, several other PCR-based DNA marker systems have also been used for the germplasm evaluation of Chinese fir. Li et al. (2007) applied the RAPD markers to decipher the genetic diversity and relationships of 182 second-generation elite genotypes of Chinese fir based on 29 primers that generated a total of 311 fragments with a PPB of 78.8% and an average of 8.4 polymorphic bands per primer. Yang et al. (2009) reported a molecular polymorphic study of different geographic

provenance germplasm ($n = 24$) of Chinese fir by ISSR markers. In their study, a total of 173 polymorphic bands were produced by 22 primers with an average of 7.9 polymorphic bands per primer. Recently, Wen et al. (2013) examined 27,666,670 Chinese fir transcriptome and finally identified 28 polymorphic useful EST-SSR markers. While Ouyang et al. (2014) developed 52 SSR markers from Chinese fir, and then successfully detected 254 polymorphic loci from a germplasm collection (1st breeding population) with an average of 4.9 alleles per loci. In this report, we observed that the SRAP system enabled to produce 498 bands with a PPB of 89.4% and an average of 12.7 polymorphic bands per primer combination in the tested genotypes. Furthermore, the marker attributes including PIC, MI and RP appeared to be rather informative for each primer set (Table 3). These performances suggested that the present SRAP system was also efficient for the detection of DNA variability of Chinese fir. Application of SRAP

Table 5. SRAP markers associated with growth and wood property traits in Chinese fir. H, height; DBH, diameter at breast height; V, stem volume; WBD, wood basic density; Hy, hygrosopicity; HR, heart-wood ratio; TL, tracheid length; TD, tracheid diameter; TLDR, tracheid length-diameter ratio. Only the markers with significant marker-trait associations ($p < 0.05$) in both GLM (general linear model) and MLM (mixed linear model) are mentioned. The maker is named with its original primer set followed by size (bp). R^2 indicated the percentage of the phenotypic variation explained by marker. The value of R^2 was calculated by MLM

Traits	The significantly associated SRAP markers ($p < 0.05$)	Range of R^2 (%)
H	Me10/Em26 ₁₆₀ , Me11/Em5 ₃₈₀ , Me11/Em5 ₅₅₀ , Me11/Em21 ₇₅₀ , Me12/Em19 ₉₀₀ , Me13/Em18 ₁₅₀ , Me15/Em26 ₂₅₀ , Me19/Em19 ₇₅₀ , Me21/Em19 ₆₀₀	9.1–12.8
DBH	Me10/Em26 ₁₆₀ , Me11/Em5 ₅₅₀ , Me11/Em13 ₁₁₀₀ , Me16/Em1 ₁₀₀₀ , Me18/Em19 ₇₀₀ , Me19/Em19 ₇₅₀ , Me20/Em19 ₄₀₀ , Me20/Em19 ₈₅₀	8.8–12.8
V	Me1/Em20 ₂₅₀ , Me10/Em26 ₁₆₀ , Me13/Em18 ₁₅₀ , Me16/Em1 ₁₀₀₀ , Me17/Em21 ₂₀₀₀ , Me20/Em19 ₄₀₀	8.4–16.1
WBD	Me2/Em26 ₄₇₀ , Me3/Em17 ₂₀₀₀ , Me10/Em20 ₂₅₀ , Me11/Em20 ₂₀₀ , Me12/Em19 ₄₅₀ , Me16/Em1 ₁₀₀₀ , Me19/Em22 ₁₅₀ , Me20/Em1 ₇₅₀	9.3–26.4
HR	Me1/Em1 ₉₀₀ , Me1/Em19 ₉₀₀ , Me1/Em19 ₁₀₀₀ , Me10/Em21 ₉₃₀ , Me10/Em26 ₁₆₀ , Me10/Em26 ₅₀₀ , Me11/Em13 ₄₅₀ , Me11/Em20 ₁₂₀₀ , Me13/Em5 ₅₅₀ , Me18/Em19 ₇₀₀ , Me20/Em19 ₂₇₀ , Me20/Em19 ₈₅₀ , Me20/Em26 ₃₀₀ , Me21/Em19 ₃₅₀	8.3–17.5
Hy	Me2/Em26 ₄₇₀ , Me3/Em17 ₂₀₀₀ , Me10/Em20 ₂₅₀ , Me11/Em20 ₂₀₀ , Me12/Em19 ₄₅₀ , Me16/Em1 ₁₀₀₀ , Me19/Em22 ₁₅₀ , Me20/Em1 ₇₅₀	9.0–24.6
TL	Me1/Em20 ₃₆₀ , Me4/Em5 ₃₂₀ , Me11/Em5 ₁₇₀₀ , Me11/Em20 ₃₃₀ , Me11/Em20 ₁₂₀₀ , Me12/Em19 ₁₁₀ , Me12/Em19 ₂₀₀ , Me13/Em18 ₈₈₀ , Me17/Em19 ₂₀₀ , Me17/Em19 ₁₀₀₀ , Me19/Em19 ₃₀₀ , Me20/Em19 ₇₅₀ , Me20/Em19 ₁₄₀₀ , Me20/Em26 ₄₅₀ , Me21/Em19 ₁₀₀₀	8.3–17.5
TD	Me10/Em20 ₁₆₅₀ , Me10/Em21 ₅₀₀ , Me11/Em20 ₁₅₀₀ , Me13/Em18 ₃₅₀ , Me16/Em19 ₈₀₀ , Me17/Em19 ₂₀₀₀ , Me17/Em21 ₂₀₀ , Me17/Em21 ₁₇₀₀ , Me19/Em19 ₁₈₀₀ , Me19/Em22 ₆₀₀	8.3–14.8
TLDR	Me1/Em1 ₂₅₀ , Me3/Em17 ₅₀₀ , Me11/Em5 ₂₅₀ , Me11/Em5 ₁₇₀₀ , Me11/Em13 ₉₅₀ , Me11/Em21 ₃₀₀ , Me11/Em21 ₁₈₀₀ , Me12/Em19 ₅₀₀ , Me12/Em19 ₁₀₀₀ , Me13/Em5 ₂₀₀ , Me13/Em5 ₇₀₀ , Me13/Em18 ₁₀₀ , Me13/Em18 ₆₀₀ , Me13/Em18 ₇₀₀ , Me13/Em18 ₇₅₀ , Me13/Em18 ₈₀₀ , Me15/Em1 ₄₉₀ , Me15/Em1 ₅₀₀ , Me15/Em26 ₁₄₀₀ , Me17/Em21 ₇₀₀ , Me21/Em19 ₇₅₀	8.9–15.7

technique in conifer was additionally reported by Feng et al. (2009). In their study, nine SRAP primer pairs yielded 249 bands with a PPB of 55.42% from a total of 480 *Pinus koraiensis* samples that belonged to 24 difference provenances. Albeit so, there have few studies on the application of the SRAP technique in conifer.

Genetic diversity refers to the level of genetic differentiation within a species or population, which also reflects the ability of the germplasm to adapt to the changeable environments and its genetic potential for breeding (Cravero et al. 2007; Yu et al. 2014). In terms of the measurements of PPB (89.4%), h (0.204–0.373, mean = 0.279) and I (0.324–0.555, mean = 0.427), we found that the genetic diversity of the present Chinese fir genotypes was relatively high. This diversity furthered our understanding of the phenotypic variation in growth and wood property traits, and strongly suggested that it was possible to select the divergent germplasms (genotypes) for the breeding programs.

With respect to the genetic relationship, it is not always the best way to define the genetic similarity using the phenotypic traits because of the degree of divergence between genotypes at the phenotypic level is not necessarily correlated with a similar degree of genetic difference (Hamrick and Loveless 1989; Mwase et al. 2010). As shown in Fig. 2, the phenotypic dendrogram had low correspondence to the DNA tree. For example, the genotypes of c17, c70, c98 and c44 were considered to be classified into a

group (group 1) in the phenotypic dendrogram, but according to the molecular (SRAP) distance matrix, they were distributed into 4 different clusters (I – IV). Albeit the clustering of c18, c59, c108 and c130 in the phenotypic tree and their consistent Lechang origin, the SRAP dendrogram revealed that these genotypes belonged to different clusters at DNA level. This divergence can be largely explained by: (1) genetic interactions, i.e., two or more different combinations of genes may lead to same phenotype (Dillman et al. 1997; Oliveira et al. 2013), (2) limited phenotypic/morphological characters tested (Oliveira et al. 2013), i.e. only 9 quantitative traits (growth and wood property traits) were involved in this study, while higher amount of SRAP markers (TNB = 498, NPB = 445) was used, and (3) volatility of the phenotype influenced by the environmental factors. Indeed, when comparing the morphological and molecular clustering patterns it was commonly reported that these two methods were inconsistent and highly variable (Rana et al. 2005; Mamunur Rahman et al. 2011). Discrepancy of the genetic distances between morphological traits and molecular markers (AFLP) could also be found in the tree species *Uapaca kirkiana* müell. Årg (Mwase et al. 2010). In our study, the SRAP dendrogram definitely had higher discriminating capacity over that of phenotype, and it could be employed as an authentic reference for the assessment of the genetic relationship when these germplasms (genotypes) were considered to be used.

Success of MAS depends on the availability of marker-trait associations. In this report, a total of 99 significant marker-trait associations corresponding to 77 different SRAP markers were detected (Table 5). These markers could explain 8.3 to 26.4% variation (R^2) of the growth and wood property traits at a significant level ($p < 0.05$). Strikingly, 18 of which were found to be associated with more than one phenotypic traits. For example, Me19/Em19₇₅₀ was linked to H and DBH, while Me16/Em1₁₀₀₀ was accounted for both growth (DBH and V) and wood property (WBD and Hy) traits. Such associations may be caused by pleiotropism or QTL interactions (Wang et al. 2013), and also reflected a correlation between traits, in accordance with the results of Table 2. Collard et al. (2005) proposed that a QTL explaining more than 10.0% of total phenotypic variation (R^2) could be considered as major QTL. Referring to this, 41 SRAP markers ($R^2 > 10.0\%$) corresponding to 53 marker-trait associations could be regarded as pivotal genetic loci for growth and wood property traits. One such marker, Me2/Em26₄₇₀, even explained more than 20.0% of the phenotypic variation for wood property traits (26.4% and 24.6% for WBD and Hy respectively). Since the SRAP technique aimed to detect the functional genomic regions (Li and Quiros 2001), the present marker-trait associations seemed to be particularly useful for Chinese fir MAS programs.

Acknowledgments

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