

**MORPHOLOGICAL AND GENETIC ANALYSIS OF *Triplaris guayaquilensis* Wedd  
(POLYGONACEAE): ONE NATIVE TREE OF ECUADOR**

ANÁLISIS MORFOLÓGICO Y GENÉTICO DE *Triplaris guayaquilensis* Wedd (POLYGONACEAE):  
UN ÁRBOL NATIVO DE ECUADOR

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**ABSTRACT**

In this paper, we assessed six native populations (55 trees) of *Triplaris guayaquilensis* Wedd (Fernan Sanchez), one of the major forest species from Ecuador, using morphological and AFLP (Amplified Fragment Length Polymorphisms) data. The populations were collected through two macro-sites (Central coastals: Quevedo, Ventanas, la Guayas; Andean surroundings: la Maná, Patricia Pilar, Pichincha). The populations showed the following traits: straight shaft (66 %); round, irregular top shape (50 %); and branch insertion angle 0° - 30° (86 %). Four qualitative (straight shape, type of leaf edge, leaf width and leaf pubescence) and four quantitative (commercial tree height, basal area, commercial volume and total volume) traits were the most explicative traits present after Principal Component Analysis (PCA). PCA separated populations into two groups: one group included populations from Central Coastals which showed morphological traits highly and positively correlated with wood production, and the other group included populations with lower tree growth from the Andean surroundings. Populations from Central Coastals showed the highest values of genetic diversity indexes, AFLP markers separated populations based on the macro site of origin. For  $K = 2$  Bayesian analysis separated FS populations into two groups; two populations from Central Coastals region and the other four the Andean surroundings region (3) and 1 from Central Coastals (La Guayas). For greater  $K$  values, the genetic fragmentation of populations by origins was evident since for  $K = 5$  four groups were performed: one including populations from Quevedo and Ventanas and other from La Guayas (Coastals), the third group included trees from La Mana and Pichincha and the fourth, from Patricia Pilar (Andean surroundings). Results suggested the constant and effective genetic recombination or the genetic flow among and within Fernan Sanchez populations with a clear tendency towards genetic differentiation.

**Keywords:** Fernan Sanchez; forest genetic resources; genetic variability; molecular markers.

**RESUMEN**

En éste trabajo, seis poblaciones nativas (55 árboles) de *Triplaris guayaquilensis* Wedd (Fernán Sánchez), una de las principales especies forestales de Ecuador, se sometieron al análisis morfológico y genético con AFLPs (polimorfismos en la longitud de los fragmentos amplificados). Las poblaciones se colectaron a través de dos macro-sitios (Litoral Central: Quevedo, Ventanas, la Guayas; Estribaciones de los Andes: la Maná, Patricia Pilar, Pichincha). Las poblaciones exhibieron las siguientes características: fuste recto (66 %); forma de la copa irregular y redonda (50 %); ángulo de inserción de las ramas de 0° a 30° (86 %). Cuatro características cualitativas (forma del fuste, tipo de terminación de hojas, ancho de hojas y

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pubescencia de las hojas) y cuatro cuantitativas (altura comercial del árbol, área basal, volumen comercial y volumen total) fueron las más explicativas de acuerdo con el análisis de componentes principales (ACP). El ACP separó las poblaciones en dos grupos: uno incluyó poblaciones del Litoral Central, con características morfológicas alta y positivamente correlacionadas con la producción de madera y, el otro, con poblaciones con crecimiento de árboles menor provenientes de las estribaciones de los Andes. Las poblaciones del Litoral Central mostraron los mayores valores de diversidad genética y los marcadores AFLP separaron dichas poblaciones con base en el macrositio de origen. Para un valor  $K = 2$  el análisis Bayesiano separó las poblaciones de FS en dos grupos: dos poblaciones de la región Litoral Central y las otras cuatro de la región de las estribaciones de los Andes (3) y una del Litoral Central (La Guayas). Para valores  $K$  mayores fue evidente la fragmentación genética de las poblaciones de acuerdo con el origen pues para  $K = 5$  se formaron cuatro grupos: uno incluyó poblaciones de Quevedo y Ventanas y otro de La Guayas (Litoral), un tercer grupo incluyó árboles de La Mana y Pichincha y un cuarto grupo, de Patricia Pilar (estribaciones de los Andes). Los resultados sugieren la constante y efectiva recombinación genética o flujo genético entre y dentro de poblaciones de Fernán Sánchez con una tendencia clara hacia la diferenciación genética.

**Palabras clave:** Fernán Sánchez; recursos genéticos forestales; variabilidad genética; marcadores moleculares.

## INTRODUCTION

Ecuador has native resources with high productive potential for several purposes. Among the forest genetic resources outstands *Triplaris guayaquilensis* Wedd commonly known as Fernán Sánchez (FS), is exceptional (LITTLE and DIXON, 1969). FS is a forest species that belongs to Polygonaceae, and it is native of Ecuador where it is well distributed throughout coastal regions and near the Andes. FS shows fast vegetative growth, and it is highly demanded for the construction, agroforestry and furniture industries.

The coastal region of Ecuador is one of the more endangered regions in terms of biological diversity due deforestation and advancement of agriculture despite conservation strategies, such as the establishment of biological reservations and protected areas. Highly valuable woody species, such as *Clorophora tinctoria*, *Cordia machranta*, *Tectona grandis* and *Triplaris guayaquilensis*, have been indiscriminately used. Currently, original forestry populations are poor and have a risk of genetic erosion. Throughout the coastal region, forests and agriculture farms coexist (VALVERDE, 1991). Little knowledge has been gathered about *Triplaris* with the exception of botanical and biological studies (JORGESEN and LEON, 1999). Several studies have promoted a slow advancement of botanical, genetic and molecular analyses of native populations in Ecuador (AGUIRRE and LARS, 2005).

Some advances have been obtained in the genetic analyses of major forest resources of Ecuador, such as *Schizolobium parahybum* (CANCHIGNIA-MARTINEZ et al., 2007) and *Tectona grandis*

(NIETO-RODRIGUEZ, 2010), where molecular marker strategies have been successfully used to analyze genetic relationships among and within native populations. In addition to the improved knowledge of genetic diversity patterns and population genetics, the identification of promising trees should increase the potential of breeding and seed production of major forest resources (NAMKOOG and KOSHY, 2001) and optimize conservation. In this study, we analyzed *Triplaris guayaquilensis* native populations from two macro sites of Ecuador to determine the genetic relationships among and within populations based on morphological and genetic markers.

## MATERIALS AND METHODS

### Germplasm

Young leaves of each-one of 55 trees from six Fernan Sanchez (FS) populations were collected through two macro-sites of Ecuador (Table 1). One macro-site (Central Coastal) includes populations from Quevedo, Ventanas and Guayas with tropical, humid forest conditions (bh-T). The other macro-site (Andean surroundings) includes plants from La Mana, Pichincha and Patricia Pilar, and these regions have tropical, humid and warm forest conditions (bh-PMtc) (HOLDRIGE, 1977; CAÑADAS et al., 1986, VALLEJO and MALDONADO, 1987). In each site, geographical (latitude, longitude and altitude) and weather data were recorded. Seed collection, manipulation, and storage were conducted as previously described by ORDOÑEZ et al. (2005) for conservation in one local germplasm bank of the 'Universidad Técnica Estatal de Quevedo',

Quevedo, in Ecuador. In each site of collection trees is naturally growing and they were randomly chosen taking care that all trees were representative of each population and similar height and age; the minimum distance between trees was 8 m and maximum of 50 m. The size of surface in each forest was variable (1 -3 ha).

### Morphological analysis

FS plants were morphologically analyzed based on descriptors for tropical woody species as previously described by LEOPOLD et al. (2001) and modified by ORDÓÑEZ et al. (2005) (Table 2). For this study, 16 qualitative traits and 6 quantitative traits were registered as follows: 11 traits corresponded to tree timber, 2 traits to top tree, and 9 to leaves.

### Genetic analysis

Genomic DNA was isolated and purified from 100 mg of fresh young leaves from each plant/tree using a DNeasy® Plant Mini kit (Qiagen®). Samples were stored at 4 °C. The AFLP procedure was performed as previously described by VOS et al. (1995). Approximately 200 ng of genomic DNA was digested with 50 U of EcoRI and 15 U of *Tru91* endonucleases at 37 °C for 210 min followed by incubation at 70 °C for 15 min in a thermocycler (Gene Amp 9700, Applied Biosystems). The DNA fragments were linked to EcoRI and MseI adapters (EcoRI adapter, 5'-CTCGTAGACTGCGTACC-3'/3'CTGACGCATGGTTAA-5'; and MseI adapter, 5'-GACGATGAGTCCTGAG-3'/3'-TACTCAGGACTCAT-5') at 20 °C for 2 h. After preselective amplification by PCR using nucleo-

TABLE 1: Geographical origin of *Triplaris guayaquilensis* populations from Ecuador.

TABLA 1: Origen geográfico de poblaciones de *Triplaris guayaquilensis* de Ecuador.

Location	n.	Geographical origin			Climatic conditions
		South latitude	West longitude	Altitude (masl)	
Macro-site: Central Coastals					
La Guayas	10	01° 27'	79° 08'	54	Dryland-tropical forest with 16-32 °C of mean temperature and > 1000 mm of annual precipitation; clayish-loam soils with neutral pH and sedimentary origin.
Quevedo	9	01° 05'	79° 04'	74	Sub-humid tropical forest with 23 – 32 °C of mean temperature and > 1500 mm of annual precipitation; silt-loam or sandy-loam soils with slightly acid-neutral pH and volcanic origin.
Ventanas	8	01° 25'	79° 27'	60	Sub-humid tropical forest with 23 – 32 °C of mean temperature and > 1500 mm of annual precipitation; sandy-clay soils with neutral-slightly alkaline pH and sedimentary origin.
Macro-site: Andean surroundings					
Patricia Pilar	8	0° 02'	79° 02'	90	Sub-humid tropical forest with 23 – 28 °C of mean temperature and > 1500 mm of annual precipitation; loam or silt-loam soils with acid pH and volcanic origin.
La Mana	10	0° 56'	79° 13'	315	Highly-humid tropical forest with 18 – 30 °C of mean temperature and > 2000 mm of annual precipitation; loam or silt-loam soils with acid pH and volcanic origin.
Pichincha	10	1° 04'	79° 08'	57	Humid tropical forest with 12 – 26 °C of mean temperature and > 2000 mm of annual precipitation; loam or silt-loam soils with acid pH and volcanic origin.

TABLE 2: Morphologic traits measured in *Triplaris guayaquilensis* from Ecuador.TABLA 2: Características morfológicas medidas en *Triplaris guayaquilensis* de Ecuador.

Tree Organ	Traits	Classes/Scales	
Shaft	Shaft shape	Straight (1), curved (2), very curved (3)	
	Height of bifurcation	1/3 upper (1), 1/3 middle (2), 1/3 bottom (3)	
	Principal axis dominance	Total (1), partial (2), null (3)	
	Branch insertion angle	0 to 30° (1), 30 to 60° (2), 60 to 90° (3)	
	Un-bark degree	High (1), low (2)	
	Shaft diameter to the breast height	cm	
	Total height	m	
	Commercial height	m	
	Trunk base area	m <sup>2</sup>	
	Shaft total volume	m <sup>3</sup>	
	Shaft commercial volume	m <sup>3</sup>	
	Crown	Crown diameter	> 10 m (1), 10 to 5 m (2), < 5 m (3)
		Crown shape	Round (1), round-irregular (2), ovoid (3), enlarged (4)
Leaf	Edge shape	Serrated (1), undulating (2), smooth (3)	
	Adaxial color	Light green (1), Dark green (2)	
	Abaxial color	Light green (1), Dark green (2)	
	Pubescence	Present (1), absent (2)	
	Leaf length	Large (1), medium (2), small (3)	
	Leaf shape	Round (1), enlarged (2), ovoid (3)	
	Leaf width	Wide (1), narrow (2)	
	Leaf texture	Smooth (1), rough (2)	
	Venation pattern	Close to the edge (1), dispersed (2)	

<sup>a</sup> Classes were described on Table 2. Numbers in brackets indicate the number of accessions per class.

tide A, a selective amplification by PCR was performed with four combinations of EcoRI + 3 and MseI + 3 primers with the following PCR specifications: 94 °C for 30 s; 20 cycles of 56 °C for 60 s and 72 °C for 60 s (pre-amplification); 94 °C for 30 s; 6 cycles of 56 °C for 30 s and 72 °C for 60 s; 94 °C for 30 s; and 23 cycles of 56 °C for 30 s and 72 °C for 60 s (selective amplifications). Amplifications used the EcoRI + 3 oligonucleotide (5'-GACTGCGTACCAATTC/NNN-3'), which was marked with IRDye<sup>TM</sup>-800, and the MseI + 3 oligonucleotide (5'-GATGAGTCCTGAGTAA/NNN-3'). Amplified products were stored at -20 °C until use. Amplified fragments were electrophoresed in an automate sequencer IR<sup>2</sup> (model 4200-02G, LICOR, Lincoln, NE). Acrylamide gels were prepared according to the AFLP LI-COR Bioscience's manual. AFLP data were collected in real-time from

the sequencer.

### Data analysis

The mean, variance, rank, standard deviation and coefficient of variation of each morphological trait were calculated using 'Statistica' version 5 for Windows (STATSOFT, 2004). The frequencies of predetermined classes of qualitative data were also calculated. The data were subjected to principal component analysis (PCA). We then used the most explicative variables to perform a cluster analysis by estimating Nei distances and a dendrogram was constructed based on UPGMA (HAIR et al., 1992) using 'Statistica'. AFLP bands were visually numerated according their migration on a gel and we assigned the number one to the band with the highest molecular weight and so on to the band with the lowest molecular weight. We assumed that

two bands with the same molecular weight in different individuals or plants as identical. We denoted the presence of one band with the number one and we denoted the absence of a band with zero. Binary matrices of ones and zeros were used to estimate similarity coefficients (NEI and LI, 1979), and the genetic distances among plants and populations were then calculated. Genetic distance matrices were used to construct one dendrogram based on the UPGMA algorithm (NEI and KUMAR, 2000) and the FreeTree software (HAMPL et al., 2001). Cluster analysis was corroborated by bootstrap analysis using 1,000 permutations, and the consensus dendrogram was obtained with TreeView version 1.6.6 (PAGE, 2000). In addition, we estimated the diversity indexes based on the formula previously described by POWELL et al. (1996), and we also estimated the percentage of polymorphic loci (IP)  $\pm$  standard errors per population taking into account the four AFLP primer combinations using Excel 2000. We used the IP instead of the Nei index (NEI, 1972), which has been previously reported for cedar *C. odorata* (GILLES et al., 1997; CAVERS et al., 2003; DE LA TORRE et al., 2008), pine *Pinus* sp. (THOMAS et al., 1999; DÍAZ et al., 2001) and mahogany *S. macrophylla* (LOWE et al. 2003), because IP measures the degree of polymorphism in the population whereas the Nei index measures the heterozygosity of individuals. It is important to note that one dominant marker system, such as AFLP, is not capable to directly measure heterozygosity. Similarity matrices were used to calculate the hierarchical analysis of molecular variance (AMOVA) (EXCOFFIER et al., 2005) as previously described by HUFF et al. (1993) using Arlequin 2.0 (SCHNEIDER et al., 1997). The hierarchies were macro-sites, populations, and accessions. The population genetic structure was inferred by Bayesian clustering model implemented by STRUCTURE version 2.3.1 (PRITCHARD et al., 2000) using AFLP data. To run the program, a number of genetic clusters ( $K$ ) characterized by the matrices of allele frequencies was first assumed. For each individual, the proportion of its genome derived from each genetic cluster (proportion of ancestry) was then estimated. Seven independent runs for each value of  $K$  ranging from 1 to 6 were performed using 30,000 Markov Chain Monte Carlo repetitions and 300,000 burn-in periods. To calculate a global rate of assignment, individuals were arbitrarily deemed as assigned to a single genetic cluster when the proportion of ancestry in that cluster was greater

than 0.8. The ideal  $K$  value was the one with the highest Ln P(D) values as described by Evanno et al. (2005).

## RESULTS

### Morphological variability

From the 22 morphological traits measured in the FS plants, we classified 16 as qualitative and six as quantitative. Five qualitative traits showed three or more classes (Table 3), and all quantitative traits exhibited high variation coefficients (i.e. shaft diameter to the breast height and commercial height with values up 50 %). The germplasm had the following traits: straight shaft (66 %); round, irregular top shape (50 %); and branch insertion angle from 0° to 30° (86 %). However, quantitative descriptors depend on location of each tree within forest areas, and they can be significantly influenced by environmental conditions. PCA explained 86.7 % of total variability with the former three principal components (PC).

TABLE 3: Summary of qualitative traits of *Triplaris guayaquilensis* populations from Ecuador.

TABLA 3: Resumen de las características cualitativas de poblaciones de *Triplaris guayaquilensis* de Ecuador.

Trait	Classes (frequencies) <sup>a</sup>	
	Shaft	
Shaft shape	1 (34), 2 (19), 3 (2)	
Height of bifurcation	1 (32), 2 (18), 3 (5)	
Principal axis dominance	1 (34), 2 (20), 3 (1)	
Branch insertion angle	1 (45), 2 (10), 3 (0)	
Un-bark degree	1, (28), 2 (27)	
Crown		
Crown diameter	1 (29), 2 (25), 3 (1)	
Crown shape	1 (4), 2 (27), 3 (2), 4 (22)	
Leaf		
Edge shape	1 (0), 2 (0), 3 (55)	
Adaxial color	1 (55), 2 (0)	
Abaxial color	1 (0), 2 (55)	
Pubescence	1 (0), 2 (55)	
Leaf length	1 (53), 2 (2), 3 (0)	
Leaf shape	1 (0), 2 (55), 3 (0)	
Leaf width	1 (0), 2 (55)	
Leaf texture	1 (55), 2 (0)	
Venation pattern	1 (55), 2 (0)	

According to the PCA, four qualitative (straight shape, type of leaf edge, leaf width and leaf pubescence) and four quantitative (commercial tree height, basal area, commercial volume and total volume) traits were the most explicative traits in FS. The most explicative traits corresponded to straight and trunk descriptors and all of the traits were positively associated with morphological variability in FS germplasm (Table 4). Using the data from the two major PCs (Figure 1), the FS populations were dispersed into four quadrants where quadrant I included individuals from Ventanas, and it had high values of commercial height, basal area, commercial volume and total volume but low values of straight shape, bifurcation height, principal axis dominance and total height. Quadrant II included plants from La Guayas and Quevedo, and it had high values of straight shape, bifurcation height, principal axis dominance, total height, commercial height, basal area, commercial and total volume, which are all desirable traits for commercial exploitation of FS. Quadrant III included trees from Patricia Pilar with high values of straight shape, bifurcation height, principal axis dominance, type of leaf edge, leaf width and total height, but had low values of commercial height, basal area, commercial volume and total volume. Finally, quadrant IV included plants from La Maná and Pichincha with low values for all previously described traits. Cluster analysis separated populations into two groups based on their geographical origin. Group I included populations from Quevedo, Ventanas and La Guayas (Central Coastals) with traits highly and positively correlated with wood production as described by PCA. Group II had populations with lower tree growth, which belonged to the Andean surrounding regions (Figure 2).

**AFLP analysis**

The four AFLP primer combinations produced 348 bands, and 276 of the bands were polymorphic (79 %). The EcoRI+ACG/MseI+AAC combination showed the highest informativeness according to the diversity indexes calculated (data not shown). Populations from Central Coastals had the highest values of genetic diversity index (Quevedo = 0.78±0.13; Ventanas = 0.77±0.13; La Guayas = 0.69±0.14), and populations with the lowest genetic diversity were from Andean surroundings (Pichincha = 0.58±0.06; Patricia Pilar = 0.55±0.09; la Mana = 0.52±0.08). As the morphological analysis, AFLP analysis separated FS populations

based on geographical origin with high levels of robustness. One group included populations from the Andean surroundings with low genetic diversity (DI = 0.55±0.03). The other three populations (Quevedo, La Guayas and Ventanas) belonging to the Central Coastal region had high genetic diversity (DI = 0.74±0.05) (Figure 3). AMOVA indicated significant differences among macro-sites and populations as well as within populations. The highest proportion of molecular variance (46 %) was found

TABELA 4: Eigenvectors of most descriptive traits of *Triplaris guayaquilensis* from Ecuador as indicated the Principal Component Analysis of morphological data.

TABLE 4: Vectores característicos de las características más descriptivas de *Triplaris guayaquilensis* de Ecuador de acuerdo con el Análisis de Componentes Principales de datos morfológicos.

Traits	Principal Component		
	1 <sup>a</sup>	2	3
Shaft shape	0.90*	-0.20	0.30
Height of bifurcation	0.84*	-0.02	0.21
Principal axis dominance	0.87*	-0.27	0.22
Branch insertion angle	0.51	-0.11	0.20
Un-bark degree	-0.44	0.29	0.05
Crown diameter	0.47	-0.25	-0.28
Crown shape	-0.58	0.51	0.48
Edge shape	0.88*	0.16	-0.19
Adaxial color	0.13	0.13	0.20
Abaxial color	0.18	0.44	-0.28
Pubescence	0.87*	-0.41	0.05
Leaf length	0.70*	-0.14	0.01
Leaf shape	0.10	-0.07	0.95*
Leaf width	0.90*	0.16	-0.19
Leaf texture	0.65	-0.50	0.16
Venation pattern	0.34	-0.32	0.16
Shaft diameter to the breast height	-0.13	0.92	-0.02
Total height	0.85*	0.36	-0.22
Commercial height	-0.34	0.87*	-0.17
Trunk base area	0.01	0.98*	0.01
Shaft total volume	0.02	0.98*	0.01
Shaft commercial volume	0.08	0.98*	0.00
Total explained variance (%)	52.70	24.49	9.52
Accumulated variance (%)	52.70	77.19	86.71

<sup>a</sup> Asterisks (\*) indicate the most descriptive traits (p<0.01).

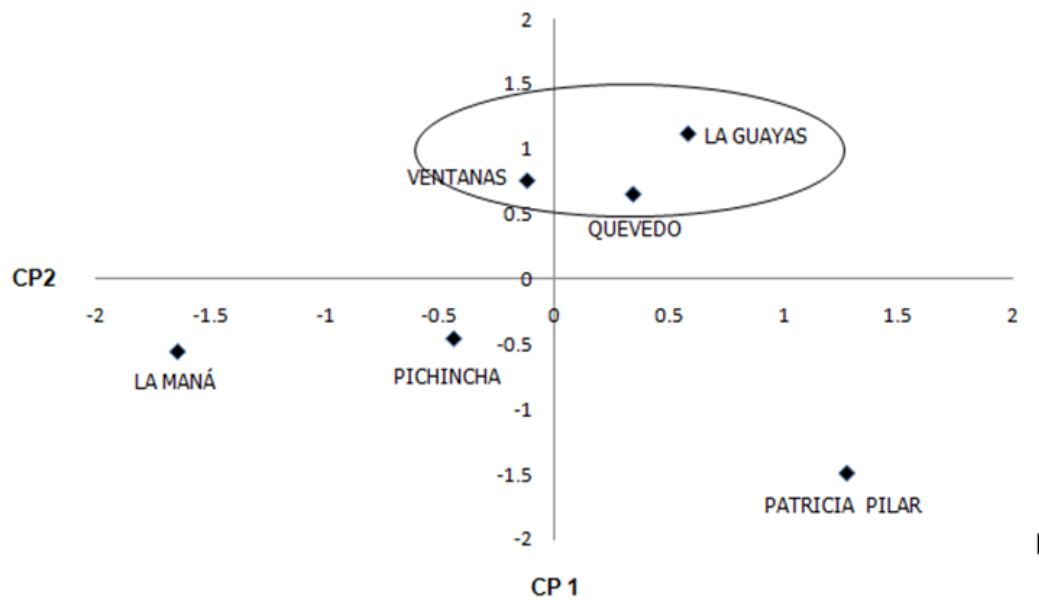


FIGURE 1: Dispersion of *Triplaris guayaquilensis* populations from Ecuador based on the data of two major principal components of PCA of morphological data.

FIGURA 1: Dispersión de las poblaciones de *Triplaris guayaquilensis* de Ecuador con base en datos de los dos principales componentes del ACP de datos morfológicos.

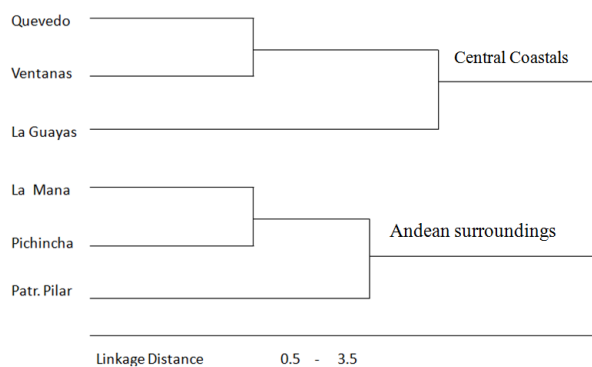


FIGURE 2: Dendrogram of *Triplaris guayaquilensis* populations from Ecuador by using morphological data.

FIGURA 2: Dendrograma de poblaciones de *Triplaris guayaquilensis* de Ecuador con base en datos morfológicos.

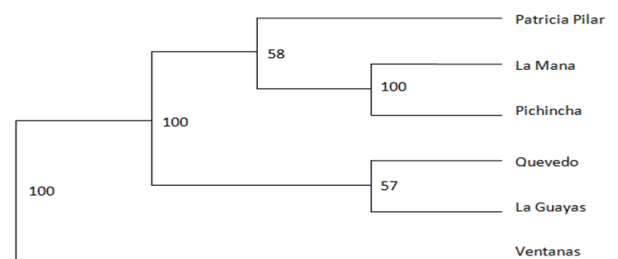


FIGURE 3: Dendrogram of *Triplaris guayaquilensis* populations from Ecuador based on AFLP data. Numbers of each node indicate repeatability values (%) from bootstrap analysis.

FIGURA 3: Dendrograma de poblaciones de *Triplaris guayaquilensis* de Ecuador con base en datos AFLP. Los números en cada nodo indican los valores de repetibilidad (%) del análisis de robustez.

within populations, and the lowest proportion of molecular variance (25 %) was found among populations (Table 5). The genetic differentiation index ( $F_{ST}$ ) was 0.74 indicating high differentiation among all hierarchies analyzed. As mentioned above in Materials and Methods, the population structure was determined with the software STRUCTURE v. 2.3.1 (Pritchard et al., 2000) without pre-identification of individuals within any particular group.

Therefore, the program itself assigned genotypes to populations. Furthermore, the optimum number of populations was decided on with the  $\Delta K$ -based test of Evano et al. (2005). The  $\Delta K$  value constantly decreased from 2351 ( $K = 2$ ) to 24 ( $K = 6$ ), the highest peaks were found for  $K = 2$  (2351),  $K = 3$  (623) (data not shown) and then both were used in this

TABLE 5: AMOVA of six *Triplaris guayaquilensis* populations from Ecuador by using AFLP data.

TABLA 5: AMOVA de seis poblaciones de *Triplaris guayaquilensis* de Ecuador con base en datos AFLP.

Source of variation	d.f.	Squared means	Explained variance (%)	P
Among macro sites	1	938.96	27.65	0.0133
Among populations	4	216.62	25.52	< 0.0001
Within populations	46	34.6	46.8	< 0.0001
Total	54	62.87	100	

study because of our interest in subdividing the full population into groups. For  $K = 2$  the FS populations were divided into two groups; two populations from Central Coastals region and the other four the Andean region (three) and one from Central Coastals (La Guayas). For greater  $K$  values, genetic fragmentation of populations by origins was evident since for  $K = 5$  four groups were performed: one including the populations from Quevedo and Ventanas and other from La Guayas (Coastals) as well as one third group from La Mana and Pichincha and other from Patricia Pilar (Andean surroundings) (Figure 4).

## DISCUSSION

The Fernan Sanchez (*Triplaris guayaquilensis*) is one of the native forest species of Ecuador, and it is currently endangered due to several factors including indiscriminate fell, selective exploitation, no management plans and no conservation of genetic resources. Despite the negative factors, our results emphasize the great genetic variability of FS throughout Ecuador. PCA was highly explicative of morphological variability of *Triplaris guayaquilensis* (> 85 % within the three former PCs), which has previously been described by LI et al. (2008) for *Paramichelia baillonii* and by MIRANDA et al. (2000) for *P. halepensis*. The following traits were the most explicative traits: six traits from the tree shaft, two traits from the tree crown and four traits from the leaves. All of these traits positively associated with genetic variability of FS. Further studies in FS should take into account the most explicative traits as indicated below.

Based on the values for the two major PCs, *Triplaris guayaquilensis* populations were

grouped according to common morphological traits. Populations from Ventanas, La Guayas and Quevedo were in quadrants I and II. All of them had high tree growth and wood production due to their location in geographical regions with climatic conditions for tree growth. The other FS populations had low values for tree growth and wood production due to their location near the Andes where weather conditions are not favorable for reproduction and tree growth. LI et al. (2008) used 45 descriptors to analyze morphological variations in *P. baillonii* and they reported that traits, such as length of wood fibers or wood relative density, clearly grouped similar populations. Our morphological descriptors were powerful and reliable enough to group and to discriminate *T. guayaquilensis* based on previous geographical origin data. In addition, the descriptors were advantageous because of the easy registering method.

The polymorphism registered for *Triplaris guayaquilensis* (79.5 %) was similar to the polymorphism previously reported by CAVERS et al. (2003) in *C. odorata* (84 %) using AFLPs but was lower than the values reported by GILLIES et al. (1997) using RAPDs and DE LA TORRE et al. (2008) using AFLPs (93.8 % and 98.8 %, respectively) in *C. odorata*. Moreover, the polymorphism registered for *T. guayaquilensis* was higher than the polymorphism reported by DÍAZ et al. (2001) (43.8 %) in *P. oocarpa*. Our data demonstrated the high efficiency of AFLPs to detect polymorphisms, to produce high numbers of amplified products per reaction and their efficiency of AFLPs to detect genetic variations in forest species, such as FS. The genetic diversity index values of FS were lower than the values reported for *S. parahybum* in Ecuador (CANCHIGNIA-MARTÍNEZ et al., 2007) but similar to values found for *Tectona grandis* (teak) (NIETO-RODRÍGUEZ, 2010).

While the AFLP marker system was only capable of detecting and clearly differentiating native Ecuadorian *S. parahybum* ecotypes from commercial or foreign populations (CANCHIGNIA-MARTÍNEZ et al., 2007), molecular markers in this study were able to propose a probable site of dispersion of *Triplaris guayaquilensis* based on the criteria previously described by RIVERA-OCASIO et al. (2002). Coastal region (Quevedo and Ventanas) and FS populations were dispersed and then colonized throughout Ecuador's territory. RIVERA-OCASIO et al. (2002) analyzed *Pterocarpus officinalis* populations from South America, Central America and



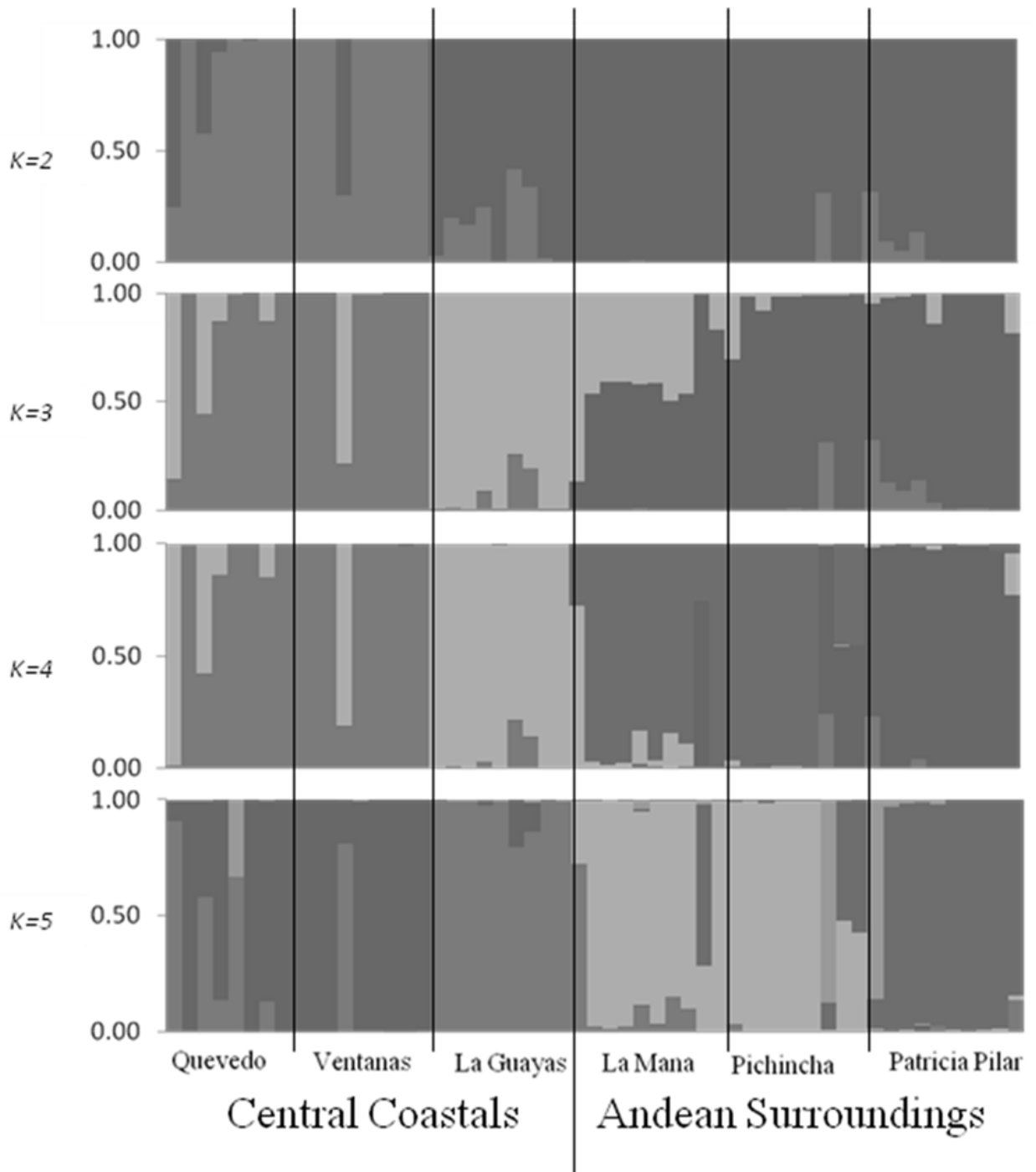


FIGURE 4: Estimated *Triplaris guayaquilensis* genetic structure. Each tree is represented by a thin vertical line which is partitioned into  $K$  segments representing the individual estimated membership fractions in  $K$  clusters. Black lines separated individuals from different populations labeled by population and origin.

FIGURA 4: Estructura genética estimada en *Triplaris guayaquilensis*. Cada árbol se representa por una línea vertical que se particiona en  $K$  segmentos que representan las fracciones de membresía estimada de cada individuo en los  $K$  conglomerados. Las líneas negras separan los individuos de las diferentes poblaciones etiquetadas por población y origen.

the Caribbean. The highest genetic diversity values were found in continental populations (Venezuela) and decreased in the Caribbean islands. The variation in genetic diversity values suggested an introduction site of any exotic species to a new environment and the most probable dissemination pattern. Thus, genetic variability found in FS from Ecuador reflects the degree of perturbation and fragmentation due to anthropic activity, which is greater near urban regions and may cause genetic erosion of the species (DE LA TORRE et al., 2008).

The germplasm of *Triplaris guayaquilensis* was divided into two groups for  $K = 2$ , one group included two populations from Central Coastals region and the other four ones from the Andean region (three) and Central Coastals (La Guayas). For greater  $K$  values genetic fragmentation of populations by origins, it was evident since for  $K = 5$  four groups were performed: one including the populations from Quevedo and Ventanas and other from La Guayas (Coastals) as well as one third group from La Mana and Pichincha and another one from Patricia Pilar (Andean surroundings). Data suggested a genetic mixture among populations and probable fragmentation for larger  $K$  values. Bayesian cluster analysis is based on the statistical models that use allele frequencies of each locus (PRITCHARD et al., 2000; BONIN et al., 2007), and the statistical methods that use genetic distances are based on the presence or absence of each locus (similarity or dissimilarity among individuals). Our data suggest a constant and effective genetic recombination or genetic flow among and within FS populations with a clear tendency towards genetic differentiation.

## CONCLUSIONS

The morphologic analysis of *Triplaris guayaquilensis* indicated that populations can be clearly separated by using both qualitative (straight shape, type of leaf edge, leaf width and leaf pubescence) and quantitative (commercial tree height, basal area, commercial volume and total volume) traits. Populations from Central Coastals showed morphologic traits highly and positively correlated with wood production and they can be used for Fernan Sanchez genetic improvement or direct propagation and further re-planting of eroded or degraded areas.

Genetic analysis indicated the genetic fragmentation of populations based on each origin suggesting the constant and effective genetic recom-

bination or genetic flow among and within Fernan Sanchez populations with a clear tendency towards genetic differentiation.

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