

## Genetic Diversity and Geographic Differentiation in *Tacca chantrieri* (Taccaceae): an Autonomous Selfing Plant with Showy Floral Display

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Received: 25 December 2005 Returned for revision: 27 March 2006 Accepted: 12 April 2006 Published electronically: 21 June 2006

• **Background and Aims** Despite considerable investment in elaborate floral displays, *Tacca chantrieri* populations are predominantly selfing. It is hypothesized that this species might possess considerable spatial or temporal variation in outcrossing rates among populations. To test this hypothesis, genetic variability and genetic differentiation within and among *T. chantrieri* populations were investigated to find out if they are in agreement with expectations based on a predominantly inbred mating system.

• **Methods** Genetic diversity was quantified using inter-simple sequence repeats (ISSR) in 303 individuals from 13 populations taken from known locations of *T. chantrieri* in China, and from one population in Thailand.

• **Key Results** Of the 113 primers screened, 24 produced highly reproducible ISSR bands. Using these primers, 160 discernible DNA fragments were generated, of which 145 (90.62%) were polymorphic. This indicated considerable genetic variation at the species level. However, there were relatively low levels of polymorphism at population levels, with percentages of polymorphic bands (PPB) ranging from 8.75% to 55%. A high level of genetic differentiation among populations was detected based on different measures (Nei's genetic diversity analysis:  $G_{ST} = 0.5835$ ; AMOVA analysis:  $F_{ST} = 0.6989$ ). Furthermore, based on levels of genetic differentiation, the 14 populations clustered into two distinct groups separated by the Tanaka Line.

• **Conclusions** High levels of differentiation among populations and low levels of diversity within populations at large spatial scales are consistent with earlier small-scale studies of mating patterns detected by allozymes which showed that *T. chantrieri* populations are predominantly selfing. However, it appears that *T. chantrieri* has a mixed-mating system in which self-fertilization predominates, but there is occasional outcrossing. Significant genetic differences between the two distinct regions might be attributed to vicariance along the Tanaka Line. Finally, possible mechanisms of geographic patterns based on genetic differentiation of *T. chantrieri* are discussed.

**Key words:** *Tacca chantrieri*, floral display, population genetic structure, gene flow, ISSR markers, mating system, geographic differentiation, Tanaka Line.

### INTRODUCTION

The spatial distribution of genetic diversity in plant populations, which is characterized by the genetic variability and genetic differentiation within and among populations, is primarily determined by the life history, including its reproductive traits, of the plant (Schoen, 1982a, b; Schoen and Clegg, 1985; Hamrick and Godt, 1996), but population history is also a determinant factor of the genetic variation within species (Schaal *et al.*, 1998).

Population history, represented as fluctuations in the number and size of populations, and the evolutionary and biogeographic histories of species, may have played critical roles in determining its current genetic composition (Schaal *et al.*, 1998). Contemporary biogeographic patterns of genetic variation are determined by historical patterns of gene flow and vicariance among populations (Hewitt, 1996; Soltis *et al.*, 1997; Avise, 2000). This history should be reflected in the genetic structure and phylogeography of extant populations, which should provide information enabling evolutionary processes to

be inferred and for biogeographical scenarios that underlie patterns of genetic differentiation to be tested. As such, if the genetic structure of a population of a species is understood, its evolutionary history can be elucidated (Bauert *et al.*, 1998).

Plant reproductive traits also determine the population genetic structure via the influence of the plant's mating system (Hamrick and Godt, 1990; Schoen *et al.*, 1996). The close relationship between the mating system and the level of genetic variation and genetic structure has been documented in many studies using different methods (Brown *et al.*, 1989; Hamrick and Godt, 1990). Inbreeding species are expected to have less genetic diversity and heterozygosity within populations, as well as more genetic differentiation among populations, than outcrossing species (Charlesworth and Charlesworth, 1995; Hamrick and Godt, 1996). Therefore, genetic diversity within and among populations can reflect, in a certain extent, the relative rates of inbreeding versus outcrossing in a species.

As an important reproductive trait, floral design and display affect plant mating systems by attracting animal pollinators and thereby promoting pollen dispersal and cross-pollination (Harder and Barrett, 1996; Emms *et al.*,

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1997). It is believed that plants with greater investment in floral structures attractive to pollinators will benefit from increased fitness via cross-pollination. In this case, species with a high investment in extravagant floral displays are expected to be largely outcrossing (Charlesworth and Charlesworth, 1987).

*Tacca chantrieri* is a widespread species that occurs in humid tropical regions of south-east Asia. Its distribution has contracted due to overexploitation, habitat destruction, and forest fragmentation. *Tacca chantrieri* has extravagant floral displays and a high investment in reproductive structures that lead to the expectation that it is largely outcrossing (Drenth, 1972; Saw, 1993). Furthermore, dark floral colours, the presence of long filiform appendages or bracts, and the absence of nectar are commonly associated with fly pollination by deceit (sapromyophilous pollination syndrome) (Proctor *et al.*, 1996). Surprisingly, in field pollination experiments and a mating system study, it was found that, despite considerable investment in extravagant floral display, most seeds produced by plants in four populations in south-west China resulted from autonomous self-pollination (Zhang *et al.*, 2005). This mismatch between floral morphology and mating system might cause *T. chantrieri* to exhibit spatial and temporal variation in population genetic structure. To test this hypothesis, larger surveys of its population genetic structure are required.

Here, inter-simple sequence repeat (ISSR) markers are employed to determine the levels and distribution of genetic differentiation within and among populations of *T. chantrieri*. These markers have recently become widely used in population genetic studies because they require less investment in time, money and labour than other methods, are highly variable (Wolfe and Liston, 1998; Harris, 1999) and exhibit Mendelian inheritance (Gupta *et al.*, 1994; Tsumura *et al.*, 1996). Inter-simple sequence repeats also generate higher percentages of polymorphic loci than other methods (Esselman *et al.*, 1999). The most recent review that evaluated among- and within-population diversity in wild angiosperms and gymnosperms using different nuclear DNA markers concluded that estimates of genetic variation derived by the dominantly inherited markers RAPD, AFLP and ISSR are very similar and may be directly comparable to codominant marker types; however, ISSRs tend to produce somewhat higher estimates of within-population variation (Nybohm, 2004).

The main objective of this study was to quantify genetic variation in *T. chantrieri* from a range of populations and to compare it with expectations for selfing species. The following specific questions are addressed: (1) What are the levels of genetic variation within and among populations of *T. chantrieri*? (2) Are patterns of genetic diversity consistent with expectations based upon the highly selfing mating system estimates from previous work? (3) Are there any geographical patterns of genetic differentiation in this species? Following the presentation of the results, the implications of observed genetic patterns on the possible areas of origin and diversification of *T. chantrieri* are discussed.

## MATERIALS AND METHODS

### *Plant species*

*Tacca chantrieri* André (also called the ‘bat flower’ or ‘bat plant’) is one of the most widespread species in the genus *Tacca*, which is distributed on the Thailand–Indo-Chinese Peninsula and on the Malay Peninsula to southern China. Its distribution in China extends from southern Yunnan to Guangxi and Hainan, and it inhabits moist and shady understorey habitats in tropical forests (Drenth, 1972; Wu *et al.*, 2003) (Fig. 1). Plants are 50–100 cm tall, with tubers or creeping rhizomes and alternate elliptic entire leaves. It has a curious inflorescence which is bat-like, both in shape and colour, with wide-spreading wing-like bracts of rich maroon-black accompanied by long trailing filaments or ‘whiskers’. If pollinated, the small black flowers develop into large black berries. In the Yunnan Province, south-west China, where the present studies were conducted, the species flowers from April to July, and by September–October the berries are ripe.

### *Study sites and sampling*

From 2003 to 2004, leaves of *T. chantrieri* were collected from 14 populations in south China and Thailand. The populations were chosen because they represented the overall distribution of the species in China (Fig. 1), as well as one population from Thailand. The fourteen populations are geographically separated with each other by at least 60 km, and the size of the populations ranged from 600 to 3500 m<sup>2</sup>, with the exception of the CZ population which is much smaller (about 600 m<sup>2</sup> with some 50 plants) (Table 1). In each of the 14 populations, a random sample of 15–35 plants was taken. Considering the sampling strategy, a Pearson’s correlation test was used to analyse the effect of sample size on genetic diversity estimates. The total sample included 303 individuals. Distances between adjacent samples were at least 10 m to increase the possibility of detecting genetic variation within each population. Leaves were collected in the field and dried directly with silica gel.

### *DNA extraction and ISSR–polymerase chain reaction (PCR) amplification*

DNA was extracted using a modified CTAB method (Doyle, 1991). Total DNA was dissolved in 1× TE for subsequent use. ISSR–PCR amplifications were performed in a GeneAmp PCR System 9700 DNA Thermal Cycler (PerkinElmer, USA) with two cycling profiles: (1) 5 min at 94 °C, followed by 35 cycles of 30 s at 94 °C, 45 s at 48 °C, 1.5 min at 72 °C, and ending with 7 min at 72 °C; and (2) 5 min at 95 °C, followed by 35 cycles of 45 s at 94 °C, 30 s at 50 °C, 1.5 min at 72 °C, and ending with 7 min at 72 °C. One hundred and thirteen primers (UBC primer set no. 9, Biotechnology Laboratory, University of British Columbia) were screened initially to identify well-amplified polymorphic bands among the populations. Of the 113 primers tested, 24 produced strong, clear and reproducible bands. These were selected for further study of the 303 *T. chantrieri* individuals. PCR was carried out

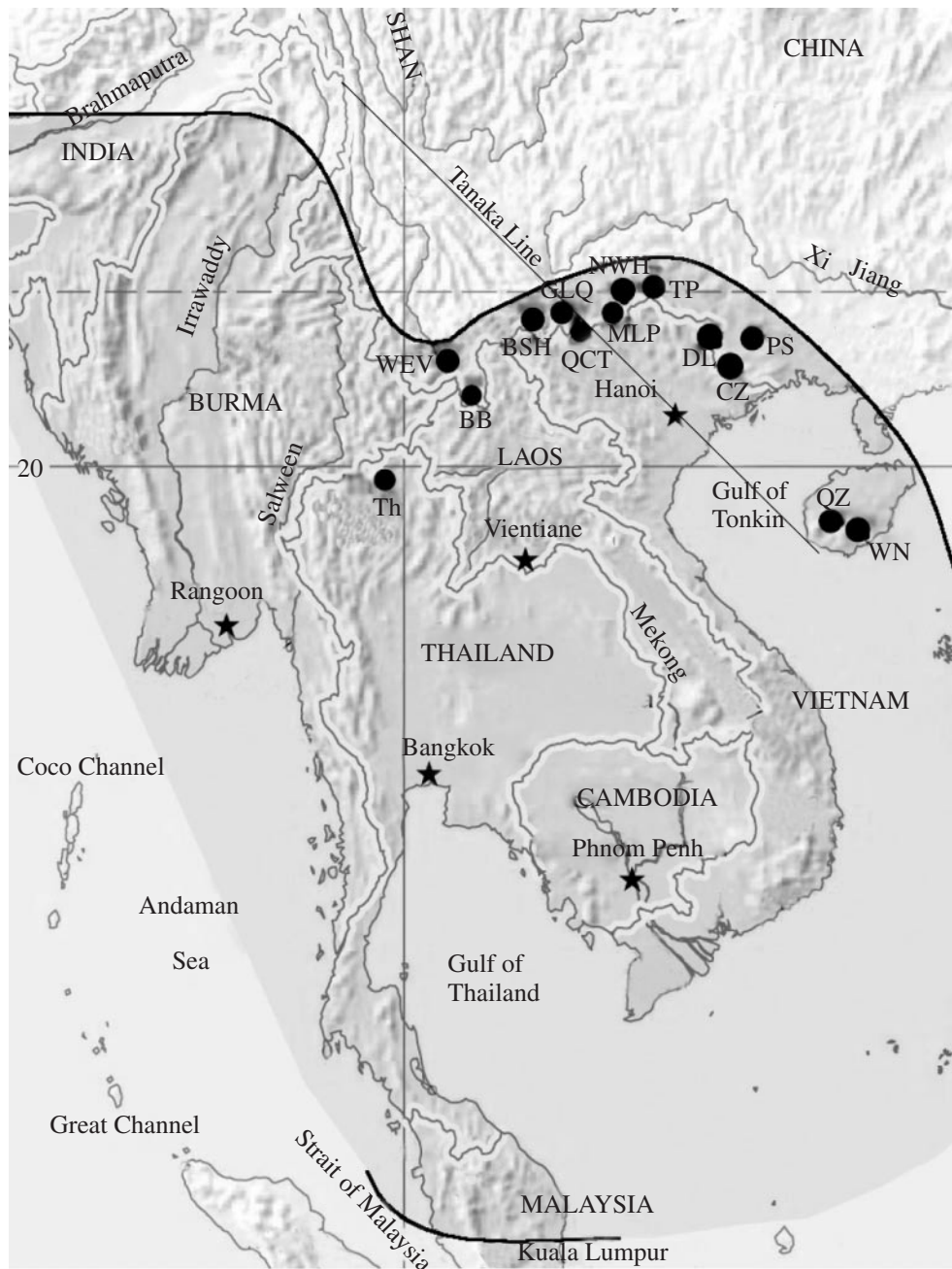


FIG. 1. A map of the known distribution of *Tacca chantrieri* (shaded area between two thick black lines) and locations of sampled populations with the abbreviations used in this study (see Table 1 for explanations of population abbreviations). This map was based on data from Drenth (1972).

in a total volume of 20  $\mu$ L, which included 100 ng of template DNA, 2  $\mu$ L 10 $\times$  PCR buffer ( $Mg^{2+}$  Plus), 0–1  $\mu$ L  $MgCl_2$  (25 mM), 0.8  $\mu$ L dNTPs Mixture (2.5 mM), 1.5 U of Taq polymerase (TaKaRa Biotechnology Dalian Co., Ltd, Dalian, China), 0.75 mM primers (Shanghai Sangon Biological Engineering Technology and Service Co. Ltd, Shanghai, China) and double-distilled water. Amplification products were separated via electrophoresis on 1.5 % (w/v) agarose gels with 0.5 $\times$  TBE buffer at 120 V for 3–4 h along with a GeneRuler100 ladder (Fermentas UAB, Inc.), stained with ethidium bromide (0.1 mg  $\mu$ L<sup>-1</sup>). They

were then photographed with an Epson digital still Af camera. Negative controls, which lacked template DNA, were included in each PCR set to test for possible contamination.

#### Genetic diversity analysis

Because ISSR markers are dominant, it was assumed that each band represented the phenotype at a single biallelic locus (Williams *et al.*, 1990). Amplified fragments were scored for the presence (1) or absence (0) of

TABLE 1. Details of all the populations of *Tacca chantrieri* sampled

Abbreviation	Sample size	Locality	Position	Altitude (m)	Size (individuals)
QCT	22	Qincaitang, Hekou County, Yunnan Province	22°40'N, 104°01'E	680–780	~120
GLQ	21	Gulingqing, Maguan County, Yunnan Province	22°45'N, 103°58'E	700–1200	~200
BSH	20	Bashahe, Lvchun County, Yunnan Province	22°53'N, 101°56'E	680–960	~600
BB	20	Wangtianshu, Mengla County, Yunnan Province	21°37'N, 101°35'E	680	~180
WEV	20	Yexianggu, Jinghong County, Yunnan Province	22°10'N, 100°51'E	760	~150
MLP	21	Balihe, Malipo County, Yunnan Province	22°58'N, 104°51'E	800–1000	~100
NWH	15	Nanwenhe, Malipo County, Yunnan Province	22°59'N, 104°44'E	720–920	~50
TP	20	Tianpeng, Funing County, Yunnan Province	23°12'N, 105°32'E	900–1000	~100
DL	35	Delong, Napo County, Guangxi Province	23°17'N, 105°50'E	520–680	~180
CZ	16	Ecological park, Chongzuo County, Guangxi Province	22°24'N, 107°30'E	100	~50
PS	25	Pingshan, Longan County, Guangxi Province	22°53'N, 107°39'E	200	~300
WN	23	Xinlong tropical flower garden, Wanning County, Hainan Province	18°41'N, 110°12'E	80	~100
QZ	25	Baihualing, Qiongzong County, Hainan Province	19°00'N, 109°49'E	350	~1000
Th	20	Chiangmai, Thailand	19°15'N, 98°55'E	500	~100

homologous bands. The resulting presence/absence data matrix of the ISSR phenotypes was analysed using POPGENE version 1.31 (Yeh *et al.*, 1997) to estimate the following genetic diversity parameters at the species level: the percentage of polymorphic bands (PPB), expected heterozygosity ( $H_E$ ) and genetic diversity measures ( $H_T$ , total population gene diversity;  $G_{ST}$ , coefficient of gene differentiation). An analysis of molecular variance (AMOVA) was used with ARLEQUIN version 2000 (Schneider *et al.*, 2000) to estimate the partitioning of ISSR phenotypic variation within populations, among populations and between two regions (south Yunnan–Thailand and south-east Yunnan–Guangxi–Hainan). The AMOVA variance components were used as estimates of genetic diversity within and between populations. The significance of this  $F$ -statistic analogue was tested with 1000 random permutations. A dendrogram was generated from pairwise Nei's genetic distances among the populations with the neighbour-joining algorithm using MEGA v. 2.1 (Kumar *et al.*, 2001).

Gene flow was estimated indirectly using the formula:  $Nm = 0.5(1 - G_{ST})/G_{ST}$  (McDermott and McDonald, 1993). Test for a correlation between Nei's genetic distance and geographical distances (in kilometres) between populations, a Mantel test was performed using tools for population genetic analysis (Miller, 1997) (computing 999 permutations). To detect the geographical pattern of genetic differentiation, significance correlation between the genetic divergence of populations with geographic distance was also tested.

## RESULTS

### Genetic diversity released by ISSR markers

The 24 selected primers generated 160 bands that ranged in size from 2 kb to 320 bp in *T. chantrieri*, which corresponded to an average of 6.67 bands per primer. Of these bands, 90.62% (145 in total) were polymorphic among the 303 individuals. The PPB for a single population ranged from 8.75% (Thailand) to 55% (TP), with

an average of  $32.7 \pm 12.8\%$ , and the average effective number of alleles per locus was 1.19. Assuming Hardy–Weinberg equilibrium, the average gene diversity was estimated as 0.165 within populations ( $H_E$ ), and 0.264 at the species level ( $H_T$ ) (Table 2). Although the sample size varied among populations (15–35 individuals per population), there was no relationship between the sample size and genetic diversity (Pearson's correlation test;  $n = 14$ ,  $r = 0.455$ ,  $P = 0.102$ ). Among the 14 populations, populations TP and DL exhibited the greatest level of variability (PPB, 55% and 50%;  $H_E$ , 0.181 and 0.165, respectively). Populations Th and WEV exhibited the lowest level of variability (PPB, 8.75% and 15%;  $H_E$ , 0.034 and 0.041, respectively).

### Population genetic structure

Most  $H_T$  in *T. chantrieri* was distributed among populations. The mean  $G_{ST}$  for all populations was estimated as 0.5835, which indicated that 58.35% of the genetic variability was distributed among populations. The AMOVA analysis is consistent with the results of Nei's genetic structure in that there is a high degree of population differentiation. Populations of *T. chantrieri* were grouped into two geographic regions: the south-east Yunnan–Guangxi–Hainan region (NWH, PS, CZ, DL, TP, MLP, WN and QZ); and the south Yunnan–Thailand region (BSH, BB, WEV, Th, QCT and GLQ). Highly significant ( $P < 0.001$ ) genetic differences were detected between regions, among populations (within regions) and among individuals (within both populations and regions) (Table 3). Of the total molecular variance, 33.44% was attributable to regional divergence, 36.46% to population differences within regions and 30.10% to individual differences within populations. When the total variance was partitioned without considering the regional distribution of populations, 69.89% was attributable to populations ( $F_{ST}$ ) and 30.1% to individual differences within populations. The number of migrants ( $Nm$ ) was estimated as 0.3568 individuals per generation between populations, and the strong genetic differentiation in

TABLE 2. Pooled values and mean genetic variabilities within populations of *T. chantrieri* detected by ISSR analysis

Population		<i>n</i>	<i>H<sub>E</sub></i>	PPB (%)	<i>G<sub>ST</sub></i>	<i>N<sub>m</sub></i>	<i>F<sub>ST</sub></i>
West region	BSH	20	0.098 (0.164)	27.5			
	QCT	22	0.111 (0.174)	35			
	BB	20	0.079 (0.159)	25			
	Th	20	0.034 (0.116)	8.75			
	WEV	20	0.041 (0.117)	15			
	GLQ	21	0.085 (0.164)	27.5			
	Mean		0.0747 (0.031)	23.125			0.6678
	Pooled	123	0.168	71.25	0.563	0.388	
East region	MLP	21	0.156 (0.204)	41.25			
	NWH	15	0.09 (0.164)	27.5			
	TP	20	0.181 (0.20)	55			
	DL	35	0.165 (0.191)	50			
	CZ	16	0.104 (0.181)	28.12			
	PS	25	0.151 (0.201)	40.62			
	WN	23	0.16 (0.201)	45			
	QZ	25	0.095 (0.159)	31.88			
	Mean		0.1378 (0.036)	39.92			0.5601
	Pooled	180	0.234	77.5	0.4168	0.6997	
	Average species level	21.64	0.165 (0.068) 0.264	32.72 (12.82) 90.62	0.5835	0.3568	0.6829

*n*, Sample size; *H<sub>E</sub>*, expected heterozygosity; PPB, percentage of polymorphic loci; *G<sub>ST</sub>*, genetic differentiation between populations estimated by using POPGENE 1.31; *N<sub>m</sub>*, estimated gene flow; *F<sub>ST</sub>*, genetic differentiation between populations estimated by using Arlequin. Standard deviations are shown in parentheses.

TABLE 3. Analysis of molecular variance (AMOVA) within/among populations and between geographic regions in *Tacca chantrieri*

Source of variation	d.f.	SSD	Variance component	Variation (%)	<i>F</i> statistics	<i>P</i> *
Between regions <sup>†</sup>	1	1289.84	7.56	33.44	<i>F<sub>CT</sub></i> = 0.3344	<0.001
Among populations within regions	12	2204.49	8.24	36.46	<i>F<sub>SC</sub></i> = 0.5477	<0.001
Within populations	288	1960.54	6.81	30.1	<i>F<sub>ST</sub></i> = 0.6989	=0.00196
Total	301	5454.86	22.61			
Analysis of the populations within west region						
Among populations	5	854.06	9.052	66.78	<i>F<sub>SC</sub></i> = 0.6082	<0.001
Within populations	116	526.93	4.504	33.22	<i>F<sub>ST</sub></i> = 0.6678	<0.001
Analysis of the populations within east region						
Among populations	7	1359.41	10.673	56.01	<i>F<sub>SC</sub></i> = 0.4337	<0.001
Within populations	172	1441.9	8.383	43.99	<i>F<sub>ST</sub></i> = 0.5601	<0.001

d.f., Degrees of freedom; SSD, sums of squares; *F<sub>CT</sub>*, total deviation from Hardy–Weinberg expectations; *F<sub>SC</sub>*, among-population deviations from Hardy–Weinberg expectations; *F<sub>ST</sub>*, deviation from Hardy–Weinberg expectations due to population subdivision.

\**P* values are the probabilities of having a more extreme variance component than the observed values by chance alone. Probabilities calculated by 1000 random permutations of individuals across populations.

<sup>†</sup>Geographic regions for *T. chantrieri* are those in the west (BSH, BB, WEV, QCT, GLQ, Th) and those in east (MLP, NWH, TP, DL, PS, CZ, WN, QZ).

*T. chantrieri* suggests that the two regions examined are isolated and that gene flow between the two regions is limited.

The neighbour-joining dendrogram based on the genetic distance between populations revealed a similar pattern: the genetic distances among the populations showed a spatial pattern that corresponded to their geographic locations (Fig. 2). Moreover, all 14 populations were clustered into two geographical groups: a clear geographical pattern of genetic diversity was identified along the Tanaka Line (Fig. 1), and a phytogeographical Line was identified between Sino-Japanese and Sino-Himalayan

genera of East Asian flora (Tanaka, 1954; Li and Li, 1997). On the west side of the Line (south Yunnan–Thailand region), 66.78 % of the variance was attributable to populations (*F<sub>ST</sub>*), and 33.22 % to individual differences within populations. On the east side (south-east Yunnan–Guangxi–Hainan region), 56.01 % of the variance was attributable to populations (*F<sub>ST</sub>*) (Table 3).

The result of a Mantel test with 1000 permutations revealed that the genetic divergence of populations (Nei's genetic distance) was significantly correlated with geographic distance in *T. chantrieri* ( $r = 0.5862$ ,  $P = 0.001$ ) (Fig. 3).

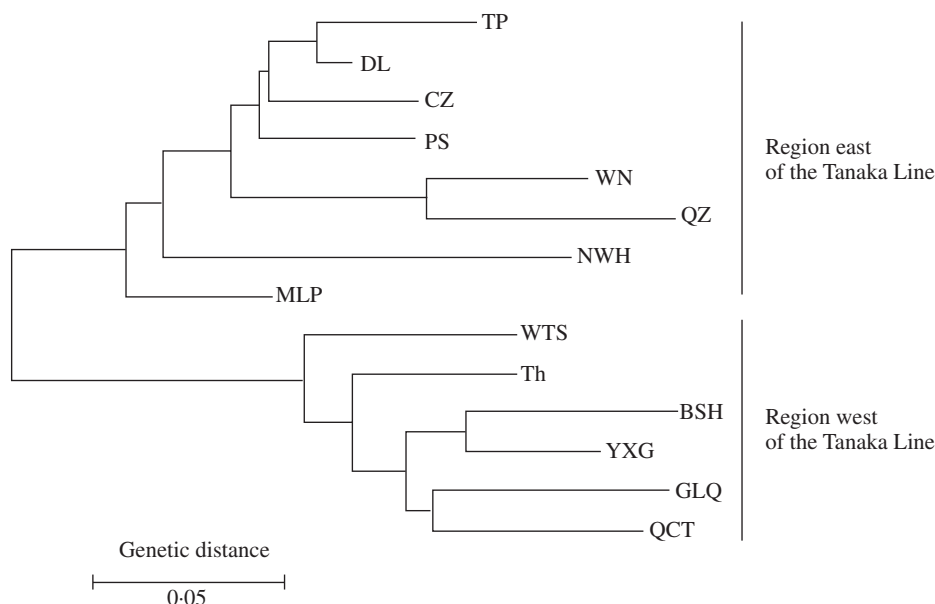


FIG. 2 Neighbour-joining tree of *T. chantrieri* based on pairwise Nei's genetic distance between populations (for explanation of codes see Table 1).

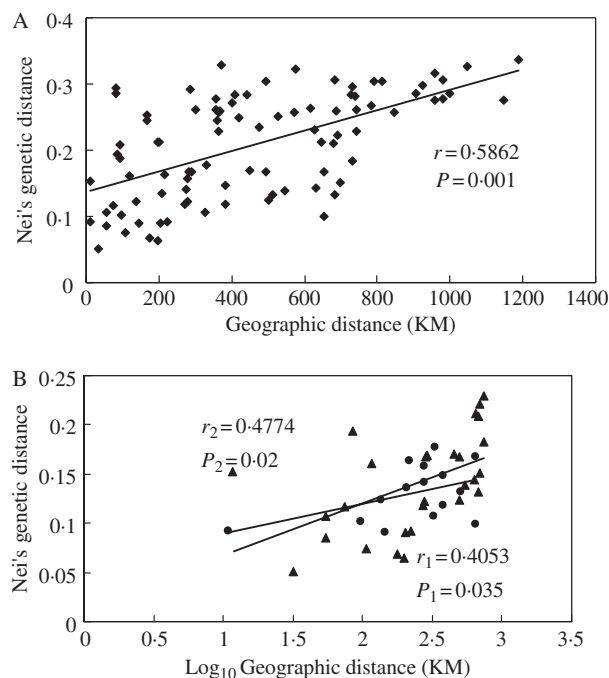


FIG. 3 Correlation between geographical distance and Nei's genetic distance revealed by the Mantel test (TFPGA; Miller, 1997): (A) all populations; (B) considering separation by the Tanaka Line (circles, populations located on the west side of the Tanaka Line; triangles, populations located on the east side of the Tanaka Line).

## DISCUSSION

### *Genetic variation in T. chantrieri and its correlation with the mating system*

The ISSR survey of 14 populations of *T. chantrieri* revealed a high level of genetic variation at the species

level, with 90.62 % of bands displaying polymorphism. However, there was considerable variation in PPB, with values ranging from 8.75 % to 55 %, and an average of  $32.72 \pm 12.82$  %. This implied that a large proportion of genetic variation was partitioned among populations. In general, selfing species usually possess lower genetic diversity within populations and higher genetic differentiation among populations relative to outcrossing species (Hamrick and Godt, 1996). Therefore, the present data on the population genetic structure in *T. chantrieri* at large spatial scales are consistent with the highly selfing mating system documented at smaller spatial scales (Zhang *et al.*, 2005). Among these 14 populations, the mating systems of WEV and BB have been quantified previously by allozyme markers. The WEV population had the highest selfing rate ( $S_m = 0.941$ ), and exhibited the lowest genetic diversity (PPB = 15 %,  $H_E = 0.041$ ). The BB population also had a quite high selfing rate ( $S_m = 0.859$ ) and contained very low genetic variation (PPB = 25 %,  $H_E = 0.079$ ) (Zhang *et al.*, 2005). Among the four populations of *T. chantrieri* examined previously, estimates of the population level maternal selfing rate ( $S_m$ ) averaged 0.86 (range 0.76–0.94). Such a figure is high, and is similar to that of other obligately selfing species. Consistent with this, the average genetic diversity in the south Yunnan–Thailand region was very low (PPB = 23.125 %,  $H_E = 0.075$ ). Similarities between the conclusions based on ISSR markers in this investigation and previous studies of reproductive biology and the mating system of *T. chantrieri* illustrate the potential of ISSR markers for population genetic studies.

Another feature of the population genetic variation of *T. chantrieri* is that populations maintain quite a large amount of ISSR variation, but it is not correlated with population size. The highest genetic variation was in population TP (PPB = 55 %,  $H_E = 0.181$ ), which had quite

a small population size (only about 100 reproductive plants). Population QZ possessed the greatest size (>1000 reproductive individuals), but maintained only a medium level of genetic variation (PPB = 31.88 %,  $H_E$  = 0.095). This result indicated that current population size cannot be a criterion for population genetic variation in this species.

*Genetic structure patterns among populations of T. chantrieri and their possible causes*

The present analyses of the data obtained from ISSR markers using different approaches (Nei's genetic diversity analysis and AMOVA) demonstrated similar patterns of genetic structure for populations of *T. chantrieri*. The AMOVA indicated that 69.89 % of the total genetic variation was partitioned among populations. In comparison with genetic variation and structure based on RAPD analyses of other wild plant populations (Nybom and Bartish, 2000; Nybom, 2004), the amount and pattern of genetic variation in *T. chantrieri* is more comparable to selfing or mixed mating taxa than to outcrossing species. The  $G_{ST}$  among populations was 0.5835, which was similar to the average for selfing plant species (0.51) in the analysis by Hamrick and Godt (1990).

A high level of population differentiation may be explained by several factors, such as the species breeding system, genetic drift, demographic fluctuations, or the genetic isolation of populations (Hogbin and Peakall, 1999). When populations are small and geographically and genetically isolated from one another, genetic drift influences the genetic structure and increases differentiation among populations (Barrett and Kohn, 1991; Ellstrand and Elam, 1993). Estimates of the effective gene flow per generation ( $Nm$ ) of *T. chantrieri* were lower (0.3568) than one successful migrant per generation. This indicated limited gene flow among populations, which may be insufficient to counteract the effect of genetic drift. While inferences of the migration rate from estimates of  $Nm$  are not definitive for populations that do not exhibit metapopulation dynamics or large demographic shifts (Whitlock and McCauley, 1999), the method is still a reasonable guide to levels of gene flow among populations. The low estimates of migration among *T. chantrieri* populations correspond well with the geographic isolation of the populations, in which genetic differentiation among populations appears to be correlated with geographic distance between populations. For example, TP and DL are geographically close, and the genetic distance between them is also relatively small.

*Geographical patterns of genetic differentiation of T. chantrieri and their implications*

The neighbour-joining dendrogram based on the genetic distance between populations revealed a similar pattern to that of the genetic distances among populations: both showed a spatial pattern that corresponded to their geographic location (Fig. 2). From the neighbour-joining tree, significant genetic differences were found between the south Yunnan–Thailand and south-east Yunnan–Guangxi–Hainan regions, and coincidentally separated by a presumed

biogeographic boundary, The Tanaka Line (Tanaka, 1954; Li and Li, 1997). It is suggested that current genetic diversity distribution pattern of *T. chantrieri* populations might be due to a possible evolutionary event under vicariance from a single common ancestor through fragmentation of its original geographic range, and this vicariance could be explained by the different history of the geological structure on each side of the Tanaka Line.

The Tanaka Line is considered to be a boundary between the Sino-Japanese plate/biogeographic region in the east and the Sino-Himalayan plate/biogeographic region in the west. The approximate position of the Tanaka Line can be shown as a straight Line starting at the intersection of 28°N, 98°E southward to approximately 18°45' or 19°N, 108°E (Fig. 1). In general, the floral components of the Sino-Japanese region are relictual, and the elements of the Sino-Himalayan are evolved. The south-east Yunnan–Guangxi–Hainan area is part of an important floristic region in China called the Dian–Qian–Gui biogeographic region, and is located on the east side of the Tanaka Line. This region is noted for species abundance, endemism and historically high rates of speciation. The results of the present study indicated that expected heterozygosity of *T. chantrieri* was higher to the east ( $H_E$  = 0.234) than to the west ( $H_E$  = 0.168) of the Tanaka Line (Table 2). However, genetic differentiation among populations was greater to the west ( $G_{ST}$  = 0.563,  $F_{ST}$  = 0.6678) than to the east of the Line ( $G_{ST}$  = 0.417,  $F_{ST}$  = 0.5601) (Table 2). All these observations are consistent with an evolutionary origin for *T. chantrieri* in the Dian–Qian–Gui region, with a relatively recent range expansion to the west, resulting in reduced diversity and a higher population differentiation in the western region.

The genetic structure of plant populations is also influenced by the long-term evolutionary and ecological history of the species, which would include shifts in distribution, habitat fragmentation and population isolation (Schaal *et al.*, 1998). Wu *et al.* (2003) proposed that *Tacca* originated from the southern marginal area of the Palaearctic continent when Pangaea expanded to the Pacific Ocean for the first time. Later, this genus became differentiated in a succession of nearby environments. Moreover, they also hypothesized that the northern part of the Indo-Chinese peninsular, stretching from Yunnan to Tibet, might be the site of ancient differentiation of this genus. The pattern of genetic structure of *T. chantrieri* is consistent with this: the genetic diversity was quite high south-east of Yunnan (TP and MLP populations), Guangxi (DL and PS populations) and Hainan (WN population) areas. This geographic area exactly corresponds to one of the centres of endemism in China (south-east of the Yunnan, Guizhou and Guangxi regions) (Li, 1996) and Hainan, which became separated from the south of China five million years ago (Zhu and Roos, 2004). The present results might support Wu's hypothesis to a certain extent: the original geographic region of this species lies from Vietnam to the northern edge of the subtropics in south-east Yunnan, Guizhou, Guangxi and Hainan (the Dian–Qian–Gui region) (Wu *et al.*, 2003). At the very least, the

populations on the east side of the Tanaka Line originated earlier than those on the west side.

## CONCLUSIONS

For ISSR markers, *T. chantrieri* exhibits low levels of diversity within populations, and significant genetic variation among populations. This genetic structure is unexpected for a species with an extravagant floral display, but corresponds with the mating system of this species as previously quantified. Geographical patterns of genetic differentiation of *T. chantrieri* provide strong evidence for both its evolutionary and ecological history, and its vicariance among populations. Additional studies measuring genetic variation in other populations in southern distribution areas of this species, and in other species sympatric to it, would be especially helpful in determining if the population genetic structure detected in *T. chantrieri* is unique among other *Tacca* species.

## ACKNOWLEDGEMENTS

We thank Dr Zhi-Yong Zhang, Mr Jie Cai for assistance in some laboratory work, Dr Wan-Jin Liao and Dr Xue-Jun Ge for assistance in ISSR analysis, as well as Mr Jiang-Yun Gao, Mr Zhi-Lin Bai and Mr Fan Chen for their help during fieldwork and collection of samples. Special thanks to Dr Spencer Barrett and Dr David Erickson for their comments and suggestions to help improve early drafts of the manuscript, and Dr David Westcott for improving the English. The research was supported by a grant from the Ministry of Science and Technology (2004 DKA30430) to D.-Z. Li, the Yunnan Provincial Natural Science Foundation grant 2003C0024Q to L. Zhang, and the National Natural Science Foundation of China (30225007) to Q.-J. Li.

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