# GENETIC DIVERSITY OF *PANAX VIETNAMENSIS* VAR. *FUSCIDISCUS* K. KOMATSU, S. ZHU & S.Q. CAI POPULATION IN WESTERN NORTH OF VIETNAM DETECTED BY INTER SIMPLE SEQUENCE REPEAT MARKERS

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# SUMMARY

Panax vietnamensis var. fuscidiscus is a precious medicinal plant which was recently discovered in Sin Ho and Muong Te district, Lai Chau province of Vietnam. Over exploitation of the species in the native habitat poses a serious threat to its existence. Adequate information on the nature and the extent of genetic diversity in this important species is required for developing suitable strategy for its conservation. In this study, inter simple sequence repeat markers were employed to investigate the genetic diversity and variability of 46 individuals belonging to a naturally distributed population of this variety in Vietnam. Genetic diversity at the population level was high ( $H_e = 0.2300$ , I = 0.3665, and PPB = 96.98 %). The group of mature individuals possessed the higher genetic diversity ( $H_{e0} = 0.2291$ ,  $I_0 = 0.3563$ , and  $PPB_{0} = 84.34$  %) as compared to group of young individuals ( $H_{eY} = 0.2086$ ,  $I_v = 0.3291$ , and  $PPB_Y = 81.5$  %). The intergroup gene differentiation was high ( $G_{ST} = 0.0499$ ) with the genetic distance among groups was 0.0298. The similarity coefficient among mature individuals was more moderate (Maximum = 0.873, Minimum = 0.614 and Average = 0.741) than among young individuals (Maximum = 0.916, Minimum = 0.596 and Average = 0.759). Otherwise, the number of discovered individuals was small, distribution area is narrow habitats, and the population showed the reduction in genetic diversity due to the human affects in the habitat and over-exploitation. Results on genetic diversity and variability showed that the investigated population has coped with the risk of decline and needed to be protected.

Keywords: Genetic diversity, inter simple sequence repeat, Panax vietnamensis var. fuscidiscus, Vietnam

# INTRODUCTION

*Panax* L. (ginseng) is a small genus of Araliaceae, which is distinguished from other genera of Araliaceae by slow-growing perennial plants with fleshy roots, stout rootstock. They are the most famous and valuable medicinal plants in the world. In 2003, Zhu *et al.* described a new variety of *Panax vietnamensis* var. *vietnamensis* and named as *Panax vietnamensis* var. *fuscidiscus*. This variety was discovered in Jinping county of Yunnan province, China. In Vietnam, *P. vietnamensis* var. *fuscidiscus* (Lai Chau ginseng) was naturally distributed in Sin Ho district, Lai Chau province, Western North of Vietnam (Phan *et al.*, 2013).

Lai Chau ginseng is found in higher elevations,

between 1600 and 2000 m. The investigated population grows in small groups scattered amongst the herbaceous storey of primary, closed, evergreen, seasonal, tropical, broad-leaved forests on sandy and shale soils (wet and well-drained). This population is very well adapted to the tropical monsoon climate associated with these particular mountainous localities. In the investigated population, mature individuals have been harvested and used by the indigenous people as material for some medical treatments and health enhancement for centuries. And, recently rhizomes of Lai Chau ginseng have been traded as one of P. vietnamensis var. vietnamensis (Ngoc Linh ginseng) adulterants, this increases the harvesting pressure on Lai Chau ginseng. Overexploitation of these medicinal plant's rhizomes for medical uses has led to the loss of genetic biodiversity and as a result the studied species was classified as critically endangered (CR) under the national category with criteria of A2,b,c,d; B2b(ii,iii,v); C2a(i); E (Phan *et al.*, 2013).

Due to the limited distribution, only 80-100 individuals were found in their natural habitat during field investigations in the period 2013 and 2014. Studies on the genetic diversity of this variety have not been conducted in Vietnam. Reduction in genetic diversity is actual risk to P. vietnamensis var. Genetic variation is currently fuscidiscus. understood as a critical variable to the long-term survival of a population or species (Beardmore, 1983; Anatonovis, 1984). Understanding the genetic diversity and variation within and among groups/subpopulations of population of rare and endangered taxa is essential when developing management strategies for both in situ and ex situ conservation activities (Hogbin, Peakall, 1999). Thus. estimating interand intragroups/subpopulations genetic diversity is critical to the protection and long-term availability of P. vietnamensis var. fuscidiscus in both terms of ecological biodiversity and medically-related uses. Current research methods support the use of molecular markers as suitable and accurate tools for population genetic diversity detection. The advantages of inter simple sequence repeat (ISSR) lies within its low-cost use, convenience of use, and high-level of reliability in reproducing results (Zietkiewicz et al., 1994; Nagoaka, Ogihara, 1997; Lu et al., 2009; Roy, Chakraborty, 2009). As such, ISSR methods have established wide spread and accepted use for applications in population genetic studies of both wild and cultivated plants (Roy, Chakraborty, 2009).

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In the current study, the ISSR marker system was employed to induce DNA fingerprints for the estimation of genetic diversity of wild *P*. *vietnamensis* var. *fuscidiscus* population in Lai Chau, Vietnam and investigation of genetic diversity and differentiation in its mature and young groups which distributed in the same natural habitats. The objectives of this study were as follows: (1) to estimate genetic diversity at population and age group levels; (2) to analyze genetic relationships and differentiation among mature and young groups belonging to the population, and (3) to contribute and catalogue the data of this study for the use in the conservation and sustainable utilization of the researched medicinal plants within Vietnam.

# MATERIALS AND METHODS

# **Plant materials**

From March 2013 to May 2014, a total of 46 individuals presenting naturally distributive population of the *P. vietnameensis* var. *fuscidiscus* from Lai Chau Province, Vietnam which corresponded to 2 groups: young and mature, were sampled across their original habitat (Table 1). Twenty four young individuals ( $\leq$ 3-4 years) and 22 mature individuals ( $\geq$  6-10 years) were randomly selected for DNA extraction. Chosen individuals for sampling were separated from each other at least 50 m.

Fresh leaves were collected, kept fresh if DNA extraction within 60 hours or dried in sealed bags with silica gel if DNA extraction executed over 60 hours later and brought to the laboratory where each sample was extracted and preserved at a constant -  $20^{0}$  C for DNA analysis.

**Table 1.** Geographic localities of *P. vietnamensis* var. *fuscidiscus* populations in this study. O: belong to mature group; Y: belong to young group; x: number for sample identification.

Sample quantity and sign	Age group	Geographic localities	Longitude/Latitude range	
22 samples, signed as Ox	> 6 -10 years	Muong Te district and Sin	102 <sup>0</sup> 48'35" to 103 <sup>0</sup> 14'02" E	
24 samples, signed as Yx	≤ 3-4 years	Ho district	22°13'53' to 22°30'51''N	

# DNA extraction purification and quantification

Total genomic DNA was extracted using Cetyltrimethyl Ammonium Bromide (CTAB) protocol I (Weising *et al.*, 2007) with a modification of adding 10 % SDS to the extraction buffer which was then dissolved in water for the subsequent use. The DNA concentration (C) was calculated as follows: C ( $\mu g / \mu L$ ) = OD260 × 50. The OD 260/280 ratio was also calculated to determine DNA purity (Weising *et al.*, 2007).

# **ISSR-PCR** amplification

ISSR primers used in this study were synthesized by Bioneer Corporation (Republic of Korea), according to the primer set published by the University of British Columbia and Zagazig University (Egypt). Sixty ISSR primers were initially screened, and 17 of them, which yielded bright, clear bands and at least possessed one polymorphic band in both populations, were used for the analysis of all 60 samples (Table 2). PCR amplification was repeated for those working primers to check the stability and reproducibility of ISSR DNA fingerprinting. PCRs were performed in 20 µl reactions containing 2 mmol/L MgCl<sub>2</sub>, 0.25 mmol/L each of dNTPs, 1U Tag DNA polymerase (ThermoScientific), 0.2 µmol/L primer and approximately 30 ng DNA templates. The amplifications were performed in a Peqstar 96X Gradient thermocycler (PEQLAB Universal Biotechnologie GmbH, Germany) with the following program: initial denaturation at 94 °C for 5 min; 10 cycles of 94 °C for 45 s, annealing temperature +5  $(T_a+5)$  <sup>0</sup>C (Table 2) for 45 s, decreased 0.5 <sup>0</sup>C/cycle, 72  $^{\circ}$ C for 1 min 30 s; 36 cycles of 94  $^{\circ}$ C for 45 s, annealing temperature for 45 s, 72 °C for 1 min 30 s; Final extension at 72 °C for 15 min; the amplification products were separated in 2 % agarose gel, using TBE buffer at 60 V for 3 hours, stained with ethidium bromide (0.5 µg/ml), and photographed under 254/312 nm wavelength lights using Micro Doc Gel Documentation System (Cleaver Scientific, USA).

# Data analysis

Since ISSR markers were dominantly inherited, each band was assumed to represent the phenotype at a single biallelic locus (Williams *et al.*, 1990). ISSR bands were scored as presence (1) or absence (0) characters, to construct the binary data matrix.

POPGENE software (1.32) was used to calculate genetic diversity parameters: the percentage of polymorphic bands (*PPB*), the average expected heterozygosity (*H*<sub>e</sub>), average effective number of alleles (*ne*), the gene differentiation (*G*<sub>ST</sub>), the genetic distance among investigated sample sets (*D*) and (Yeh *et al.*, 1997). The Nei's genetic distance between pair of sample sets is calculated as:  $D_{XY} =$ ln(I<sub>XY</sub>) is based on the concept of genetic identity (I<sub>XY</sub>): I<sub>XY</sub> = J<sub>yo</sub>/  $\sqrt{(J_X \times I_Y)}$ , where: J<sub>X</sub> = average homozygosity in the first sample set, J<sub>Y</sub>= average calculated by formula J=1- $H_{eT}$ , J<sub>XY</sub>= average intersample set homozygosity = j<sub>XY</sub>/L with j<sub>XY</sub> is homozygosity among two sample sets and L is the number of investigated loci. J<sub>XY</sub> =  $\Sigma_{XYjk}$  (p<sub>XjK</sub> × p<sub>YjK</sub>), where p<sub>XjK</sub> and p<sub>YjK</sub> are frequencies of K<sup>th</sup> allele at locus j<sup>th</sup> in the first and the second sample sets, respectively (Vicente *et al.*, 2003).

Similarity coefficient between pair of samples and UPGMA dendrogram for genetic relationship among all studied samples was calculated and established by using NTSYSpc 2.1 (Numerical Taxonomy and Multivariate Analysis System) software (Rohlf, 2004).

# RESULTS

#### Genetic diversity

The twenty selected primers yielded 166 reproducible bands for total investigated samples. For population level, the number of bands per primer varied between 5 (UBC856C, HB11) and 15 (HB12), with an average of 8.3. For whole population, the *PPB* per primer varied from 83.33 % (ISSR844A, 17899A) to 85.71 % (ISSR 814, 17899B), 87.50 % (UBC807) and 100% (all remaining primers). For young individual group, the *PPB* per primer varied from 42.86 % (17899B) to 100 % (ISSR808, HB8, HB12, HB15, UBC842, UBC856T, UBC873). For mature individual group, the *PPB* per primer varied from 60 % (UBC856T) to 100 % (HB11, UBC842C). There was the reduction of *PPB* from population to age group levels (Table 2).

In the investigated population, the expected genetic heterozygousity was  $H_e = 0.230$ , Shannon information index I = 0.3665, the percentage of polymorphic bands was PPB = 96.98 %, average effective number of alleles Ne = 1.7342. Among the two individual groups separated by age, young group possessed the lower level of genetic diversity ( $H_{eY} = 0.2086$ ,  $I_Y = 0.3291$ ,  $PPB_Y = 81.5$  % and  $Ne_Y = 1.3356$ ), while mature group harbored higher level ( $H_{eO} = 0.2291$ ,  $I_O = 0.3563$ ,  $PPB_O = 84.34$  % and  $Ne_O = 1.8434$ ).

# **Genetic relationship**

In young group, the genetic similarity coefficients among the individuals ranged from 59.96 % ( $Y_{10} - Y_{11}$ ) to 91.60 % ( $Y_{14} - Y_{15}$ ) with a mean of 75.90%, while those of mature group ranged from 61.40 % ( $O_1 - O_7$ ) to 87.30 % ( $O_{13} - O_{14}$ ) with a mean of 74.41 (Table 3 & 4). The gene similarity

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3, 4, 5). The intergroup gene differentiation among

two age groups was  $G_{ST} = 0.0499$  with the genetic distance between them was D = 0.0298.

coefficients among the individuals of population were varied, ranging from 59.00 % ( $Y_8 - O_{20}$ ) to 91.60 % ( $Y_{14} - Y_{15}$ ) with a mean of 74.00 % (Table

 Table 2. ISSR primers used in this study.

	-										
Primer code	Sequence	Ta	Total recorded	PPB (%)							
	(5' <del>→</del> 3')	(°C)	bands for whole	Whole	Young	Mature					
			population	population	group	group					
808	(AG) <sub>8</sub> C	52	9	100.00	100.00	88.89					
814	(CT) <sub>8</sub> TG	51,5	7	85.71	85.71	85.71					
844A	(CT) <sub>8</sub> AC	52	6	83.33	83.33	83.33					
17898A	(CA) <sub>6</sub> AC	54,5	10	100.00	90.00	90.00					
17898B	(CA) <sub>6</sub> GT	54,5	8	100.00	87.50	87.50					
17899A	(CA) <sub>6</sub> AG	54	6	83.33	66.67	66.67					
17899B	(CA) <sub>6</sub> GG	54	7	85.71	42.86	71.43					
HB8	(GA) <sub>6</sub> GG	52	9	100.00	100.00	66.67					
HB9	(GT) <sub>6</sub> GG	52	8	100.00	62.50	87.50					
HB10	(GA) <sub>6</sub> CC	52	13	100.00	61.54	84.62					
HB11	(GT) <sub>6</sub> CC	52	5	100.00	60.00	100.00					
HB12	(CAC) <sub>3</sub> GC	52	15	100.00	100.00	86.67					
HB15	(GTG)₃ GC	52	11	100.00	100.00	90.91					
UBC 807	(AG)8 T	54	8	87.50	50.00	87.50					
UBC 826	(AC) <sub>8</sub> C	54	7	100.00	85.71	85.71					
UBC 842C	(GA) <sub>8</sub> CG	51,5	7	100.00	100.00	100.00					
UBC 842T	(GA) <sub>8</sub> TG	51,5	10	100.00	90.00	90.00					
UBC 856C	(AC) <sub>8</sub> CA	52	7	100.00	85.71	71.43					
UBC 856T	(AC) <sub>8</sub> TA	52	5	100.00	100.00	60.00					
UBC 873	(GACA) <sub>4</sub>	52	8	100.00	100.00	87.50					
Total			166								
Average			8.3	96.98	83.73	84.34					

Table 3. Genetic similarity coefficients among the individuals belong to young group

	Y1	Y2	¥3	Y4	¥5	¥6	¥7	Y8	¥9	Y10	Y11	Y12	Y13	Y14	Y15	Y16	Y17	Y18	Y19	Y20	Y21	Y22	Y23
Y2	0.801																						
Y3	0.783	0.777																					
Y4	0.741	0.663	0.693																				
Y5	0.777	0.687	0.741	0.735																			
Y6	0.801	0.723	0.789	0.783	0.819																		
¥7	0.789	0.735	0.801	0.699	0.807	0.867																	
Y8	0.717	0.747	0.801	0.651	0.675	0.699	0.747																
Y9	0.723	0.693	0.711	0.608	0.777	0.741	0.837	0.681															
Y10	0.699	0.657	0.663	0.596	0.753	0.765	0.801	0.633	0.855														
Y11	0.681	0.614	0.669	0.699	0.651	0.699	0.699	0.614	0.620	0.596													
Y12	0.783	0.741	0.783	0.693	0.789	0.825	0.837	0.693	0.831	0.771	0.705												
Y13	0.783	0.717	0.759	0.657	0.801	0.801	0.873	0.657	0.819	0.795	0.669	0.855											
Y14	0.765	0.711	0.717	0.711	0.843	0.819	0.807	0.639	0.801	0.813	0.651	0.801	0.825										
Y15	0.789	0.723	0.741	0.735	0.855	0.807	0.831	0.663	0.801	0.789	0.711	0.825	0.849	0.916									
Y16	0.801	0.759	0.753	0.723	0.771	0.795	0.795	0.687	0.729	0.741	0.747	0.777	0.813	0.819	0.855								
Y17	0.741	0.747	0.789	0.735	0.759	0.795	0.759	0.723	0.729	0.693	0.663	0.789	0.777	0.759	0.795	0.759							
Y18	0.807	0.753	0.783	0.693	0.729	0.753	0.753	0.729	0.675	0.675	0.681	0.771	0.759	0.705	0.789	0.801	0.789						
Y19	0.765	0.795	0.789	0.675	0.771	0.807	0.819	0.735	0.789	0.777	0.687	0.837	0.813	0.747	0.783	0.819	0.783	0.801					
Y20	0.771	0.837	0.771	0.693	0.693	0.801	0.777	0.765	0.711	0.675	0.633	0.795	0.771	0.741	0.729	0.801	0.789	0.747	0.825				
Y21	0.831	0.753	0.831	0.753	0.789	0.801	0.813	0.765	0.735	0.687	0.693	0.795	0.759	0.717	0.777	0.777	0.789	0.819	0.825	0.759			
Y22	0.783	0.789	0.795	0.657	0.717	0.753	0.789	0.729	0.747	0.723	0.717	0.807	0.783	0.693	0.765	0.825	0.765	0.807	0.825	0.795	0.843		
Y23	0.777	0.795	0.765	0.699	0.711	0.771	0.747	0.723	0.705	0.681	0.747	0.789	0.741	0.699	0.771	0.819	0.759	0.777	0.807	0.813	0.801	0.910	
Y24	0.807	0.729	0.771	0.765	0.837	0.801	0.825	0.681	0.771	0.735	0.693	0.819	0.807	0.825	0.886	0.789	0.813	0.795	0.789	0.759	0.831	0.795	0.789

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	01	02	03	04	05	06	07	08	09	010	011	012	013	014	015	016	017	018	019	020	021
02	0.675																				
03	0.693	0.837																			
04	0.723	0.711	0.813																		
05	0.645	0.729	0.759	0.753																	
06	0.759	0.723	0.741	0.735	0.681																
07	0.614	0.735	0.717	0.675	0.825	0.663															
08	0.669	0.741	0.723	0.729	0.723	0.705	0.693														
09	0.747	0.795	0.813	0.795	0.753	0.747	0.723	0.741													
010	0.717	0.777	0.747	0.729	0.747	0.765	0.729	0.735	0.849												
011	0.735	0.759	0.777	0.819	0.765	0.723	0.699	0.789	0.795	0.753											
012	0.645	0.717	0.735	0.693	0.687	0.681	0.645	0.687	0.753	0.687	0.717										
013	0.693	0.753	0.759	0.777	0.699	0.741	0.705	0.723	0.789	0.747	0.753	0.735									
014	0.735	0.747	0.765	0.771	0.705	0.735	0.711	0.741	0.795	0.717	0.771	0.693	0.873								
015	0.705	0.729	0.783	0.789	0.699	0.765	0.681	0.735	0.765	0.747	0.813	0.699	0.783	0.789							
016	0.747	0.723	0.765	0.807	0.669	0.783	0.663	0.717	0.783	0.777	0.783	0.765	0.789	0.771	0.777						
017	0.657	0.789	0.795	0.777	0.759	0.741	0.753	0.747	0.801	0.771	0.777	0.735	0.807	0.801	0.783	0.753					
018	0.687	0.795	0.741	0.687	0.729	0.723	0.699	0.741	0.783	0.825	0.747	0.729	0.753	0.735	0.741	0.711	0.777				
019	0.717	0.729	0.759	0.789	0.711	0.765	0.717	0.711	0.777	0.783	0.753	0.759	0.783	0.765	0.795	0.801	0.771	0.777			
020	0.633	0.717	0.771	0.741	0.711	0.693	0.693	0.675	0.729	0.687	0.717	0.747	0.771	0.741	0.747	0.705	0.759	0.705	0.783		
021	0.693	0.753	0.759	0.741	0.687	0.741	0.657	0.735	0.777	0.819	0.789	0.687	0.735	0.717	0.735	0.741	0.723	0.765	0.771	0.711	
022	0.699	0.735	0.777	0.771	0.717	0.699	0.651	0.717	0.771	0.693	0.795	0.705	0.789	0.795	0.765	0.735	0.729	0.699	0.693	0.741	0.717

 Table 4. Genetic similarity coefficients among the individuals belong to mature group.

 Table 5. Genetic similarity coefficients between the individuals from young and mature group.

	Y1	Y2	Y3	Y4	Y5	Y6	Y7	Y8	Y9	Y10	Y11	Y12	Y13	Y14	Y15	Y16	Y17	Y18	Y19	Y20	Y21	Y22	Y23	Y24
01	0.687	0.681	0.747	0.705	0.681	0.729	0.705	0.717	0.639	0.639	0.633	0.699	0.663	0.657	0.705	0.705	0.705	0.759	0.729	0.687	0.759	0.711	0.705	0.675
02	0.783	0.729	0.771	0.717	0.801	0.777	0.789	0.693	0.771	0.723	0.729	0.807	0.783	0.765	0.813	0.789	0.765	0.747	0.765	0.747	0.831	0.771	0.753	0.819
03	0.801	0.687	0.741	0.723	0.771	0.759	0.771	0.699	0.705	0.681	0.747	0.741	0.765	0.723	0.771	0.795	0.723	0.765	0.783	0.729	0.837	0.777	0.747	0.801
04	0.783	0.681	0.747	0.741	0.765	0.741	0.729	0.681	0.651	0.602	0.729	0.711	0.723	0.681	0.741	0.741	0.705	0.783	0.717	0.699	0.783	0.747	0.729	0.759
05	0.753	0.699	0.681	0.675	0.831	0.771	0.747	0.590	0.741	0.729	0.651	0.765	0.777	0.855	0.819	0.723	0.711	0.693	0.723	0.681	0.717	0.669	0.663	0.789
06	0.759	0.765	0.771	0.717	0.693	0.777	0.741	0.729	0.687	0.651	0.633	0.735	0.687	0.693	0.681	0.705	0.741	0.747	0.729	0.783	0.795	0.735	0.729	0.735
07	0.711	0.633	0.675	0.669	0.789	0.753	0.741	0.596	0.759	0.795	0.633	0.759	0.771	0.873	0.825	0.729	0.717	0.663	0.705	0.675	0.699	0.651	0.657	0.771
08	0.741	0.687	0.693	0.687	0.735	0.723	0.711	0.590	0.657	0.645	0.663	0.681	0.717	0.735	0.747	0.771	0.687	0.693	0.687	0.705	0.729	0.705	0.699	0.765
09	0.819	0.717	0.759	0.753	0.801	0.837	0.801	0.669	0.687	0.699	0.753	0.771	0.795	0.777	0.837	0.849	0.765	0.831	0.789	0.735	0.831	0.795	0.801	0.819
O10	0.777	0.759	0.765	0.687	0.771	0.831	0.831	0.711	0.765	0.753	0.675	0.813	0.801	0.807	0.819	0.819	0.747	0.789	0.807	0.801	0.789	0.789	0.795	0.825
011	0.819	0.705	0.723	0.753	0.765	0.777	0.741	0.681	0.711	0.687	0.729	0.735	0.747	0.765	0.813	0.789	0.741	0.771	0.741	0.711	0.795	0.735	0.753	0.759
012	0.717	0.627	0.608	0.723	0.699	0.723	0.675	0.602	0.620	0.620	0.687	0.657	0.645	0.687	0.699	0.687	0.687	0.657	0.651	0.681	0.717	0.657	0.675	0.729
013	0.777	0.687	0.741	0.771	0.759	0.771	0.711	0.639	0.633	0.645	0.747	0.717	0.693	0.711	0.747	0.759	0.759	0.753	0.687	0.705	0.753	0.705	0.759	0.813
014	0.759	0.693	0.699	0.801	0.753	0.753	0.705	0.608	0.627	0.651	0.741	0.723	0.711	0.693	0.753	0.765	0.741	0.735	0.693	0.699	0.771	0.735	0.753	0.795
015	0.813	0.687	0.741	0.747	0.699	0.735	0.735	0.675	0.681	0.657	0.723	0.765	0.705	0.699	0.759	0.747	0.699	0.765	0.711	0.693	0.801	0.741	0.735	0.753
O16	0.759	0.693	0.723	0.777	0.705	0.753	0.741	0.753	0.639	0.614	0.705	0.723	0.675	0.681	0.729	0.717	0.729	0.747	0.717	0.747	0.819	0.723	0.765	0.807
017	0.741	0.663	0.681	0.747	0.759	0.807	0.747	0.590	0.705	0.705	0.735	0.741	0.729	0.759	0.783	0.747	0.723	0.729	0.687	0.705	0.777	0.681	0.711	0.789
018	0.759	0.717	0.723	0.693	0.729	0.789	0.789	0.620	0.747	0.747	0.669	0.819	0.795	0.777	0.765	0.753	0.717	0.735	0.753	0.735	0.735	0.723	0.717	0.759
019	0.741	0.711	0.741	0.735	0.711	0.747	0.771	0.687	0.681	0.645	0.735	0.765	0.729	0.723	0.759	0.771	0.735	0.753	0.723	0.729	0.789	0.753	0.783	0.765
O20	0.705	0.639	0.645	0.735	0.675	0.699	0.651	0.590	0.645	0.633	0.759	0.693	0.645	0.687	0.723	0.723	0.699	0.693	0.675	0.645	0.717	0.681	0.723	0.729
021	0.801	0.783	0.777	0.687	0.723	0.819	0.807	0.735	0.717	0.669	0.687	0.789	0.765	0.711	0.747	0.795	0.771	0.765	0.783	0.837	0.801	0.837	0.831	0.765
022	0.747	0.608	0.687	0.789	0.741	0.741	0.717	0.633	0.675	0.627	0.741	0.711	0.699	0.681	0.729	0.729	0.717	0.699	0.669	0.651	0.759	0.711	0.705	0.771

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Figure1. UPGMA dendrogram for genetic relationship of investigated population.

# DISCUSSION

The ISSR markers in this study yielded 139 polymorphic/166 total reproducible bands in 46 individuals which corresponded to two divided by age groups belong to the population of *P. vietnamensis* var. *fuscidiscus* in the Western North of Vietnam. This method provides a highly effective and reliable molecular-level tool for analysis of genetic diversity and genetic relationships within the variety.

This study reports the genetic diversity at population levels and also at individual groups which were classified by age. The extent of genetic variation within two different age groups of each population, the gene differentiation and the genetic distance among them were showed. Studied on the wide ranges of species which possessed the life history traits of dicotyledon, long-lived perennial life form, endemic, outcrossing breeding system and ingested seed dispersal mechanism which were also found in currently investigated variety, Hamrick and Godt (1996) reported that the genetic diversity based on allozyme were PPB = 42 - 46%;  $H_e = 0.10 - 0.14$ ;  $G_{ST} = 0.14 - 0.24$ , and Nymbom (2000) based

on RAPD reported that the genetic diversity were  $H_e = 0.19 - 0.24$ ;  $G_{ST} = 0.17 - 0.23$ . Thus, the results from this study showed that *P. vietnamensis* var. *fuscidiscus* in the Lai Chau province possessed high level of genetic diversity and the gene differentiation between young and mature individual groups was lightly small.

Achieved results showed the higher population genetic diversity related to *PPB* and heterozygousity than that has been reported in previous studies based on RAPD (Artyukova *et al.*, 2004), Allozyme (Jennifer *et al.*, 2004), and AFLP (Zhou *et al.*, 2005; Zhuravlev *et al.*, 2010) in other *Panax* populations. However, the similarity coefficients among the pair of samples in the current study were higher (Bai *et al.*, 1997), which showed the limitations of ISSR markers in individual discrimination.

Using the same technique with current study to induce DNA fingerprinting in *P. ginseng* cultivated in North-East China, Li *et al.* (2011) reported that the genetic diversity was high at the species level ( $H_e$  = 0.2886; *PPB* = 98.96 %) but lower in cultivated types, viz. garden ginseng (h =0.2294, I = 0.3590, *PPB* = 85.42 %), forest ginseng (h = 0.1702, I =

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0.2559, PPB = 57.29 %) and transplanted wild ginseng (h = 0.2021, I = 0.3125, PPB = 76.04 %). These parameters in separated populations of forest ginseng were ranged as h = 0.1065 - 0.1520; I =0.2228 - 0.1854, PPB = 34.38 - 40.62 %). Genetic differentiation  $(G_{ST})$  was also detected among divided by geographic locality subpopulations of forest ginseng 0.2328, garden ginseng 0.3187, and transplanted wild ginseng 0.2540. Comparing to these results, the current study revealed that genetic diversity of the naturally distributed P. vietnamensis var. fuscidiscus population in Western North of Vietnam was as high as compared to P. ginseng garden ginseng population, but it was higher than forest ginseng and transplanted wild ginseng populations in North-East China. The genetic differentiation among divided by age groups of population in this study was significantly lower than that among ginseng populations as previously investigated in the study from Li et al. (2011). The high level of genetic diversity in the study of Li *et al.* (2011) can be attributed to the investigated species' evolutionary development. Due to the long lifespan and overlapping generations of the populations within the prior study, considerable genetic variability has been accumulated and conserved under various selection traits during the evolutionary process (Li et al., 2011) and this may happen to P. vietnamensis var. fuscidiscus population in Western North of Vietnam.

In Western North of Vietnam, near by the habitats of currently investigated population, there is the existence of other Panax taxon, namely P. stipuleanatus with two populations. The genetic diversity of this species was moderate at the species level ( $H_{eT} = 0.254$ ;  $PPB_T = 96.02$  %), and of its populations were lower as ( $H_{eBX} = 0.266$ ;  $PPB_{BX}$ =91.48%) and ( $He_{HT} = 0.235$ ;  $PPB_{HT} = 84.66$ %) for BX and TH populations respectively, the interpopulation gene differentiation was  $G_{ST} = 0.03$ with the genetic distance (D) among them was 0.0103 (Trieu et al., 2016). The intergroup gene differentiation ( $G_{ST} = 0.0499$ ) and the genetic distance between groups (D = 0.0298) were lower than the gene differentiation among subpopulation reported by Li et al. (2011). And it is easy to understand because these groups belong to the same population. However, the values of intergroup gene differentiation and the genetic distance between groups in current study were not small due to without geographic barrier. According to Vicente et al. (2003), the gene differentiation among populations from 0 to 0.05 is considered as small; from 0.05 to 0.15 is considered as moderate.

Because of the low gene differentiation and genetic distance, the two divided by age groups could not separate clearly in the UPGMA dendrogram of whole population (Fig. 1). Instead of this, most individuals belong to the same group had a tendency of grouping to form small clusters, which alternately arranged together. For genetic structure of investigated population, the young group possessed the lower genetic diversity ( $H_{eY} = 0.2086$ ,  $I_Y =$ 0.3291,  $PPB_{\rm Y} = 81.5$  %) than mature group ( $H_{\rm eO} =$  $0.2291, I_0 = 0.3563, PPB_0 = 84.34$  %). And the intergroup gene differentiation among them was high as same as moderate for interpopulation gene differentiation in general as and even higher than P. stipuleanatus species reported in previous study by Trieu et al. (2016).

The similarity coefficient among mature individuals was more moderate (maximum = 0.873, minimum = 0.614 and average = 0.741) than that among young individuals (maximum = 0.916, minimum = 0.596 and average = 0.759). For total population, the maximum genetic similarity occurred in pair of young individuals of  $(Y_{14} - Y_{15})$  and the minimum genetic similarity occurred between a young and a mature individual  $(Y_8 - O_{20})$ . These suggested that even possessed the high genetic diversity, the P. vietnamensis var. fuscidiscus population in Western North of Vietnam has coped with the risk of reduction in genetic diversity through generations. Studied on P. quinquefolius occurred from Georgia to West Virginia, Jennifer and Hamrick (2004) reported that the harvested populations possessed the higher genetic differentiation but lower expected heterozygosity compared to the protected populations due to harvest pressure.

In case of current study, there was a harvest pressure to investigated population as the same situation with *P. quinquefolius* ranged from Georgia to West Virginia. However, the genetic diversity of *P. vietnamensis* var. *fuscidscus* in Lai Chau -Vietnam was still high even as the risk of genetic reduction was evident, this showed that the geographic conditions where the investigated population occurs are suitable for its existence and development. In fact, recently a habitat of studied population has been narrowed by the climate change disaster and human unfavorable activities such as forest clearance, coal making, using the insecticides, and animal hunting. These activities lead to the reduction of pollinators (insects) and dispersive animals (rodents and birds), changes of habitat, especially surface runoff.

The understanding on population genetic variability is essential to effective conservation and sustainable utilization. The relatively high genetic diversity at population and divided by age group levels are the advantages for conservation and development of P. vietnamensis var. fuscidscus in Western North of Vietnam. However, the number of discovered individuals was small, distribution area is narrow habitats, and the population showed the reduction in genetic diversity due to the human affects in the habitat and over-exploitation. Thus, the classification of this variety as critically endangered (CR) is suitable and worthy. Otherwise, because of the heavy harvest pressure, lacking of an actionable conservation strategy may lead to the increased reduction of genetic diversity and reserve of this variety. Thus, it is of critical importance to further investigate and protect this variety for conservation purposes and for sustainable development and harvest these valuable natural resources.

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# REFERENCES

Anatonovis J (1984) Genetic variation within population. In: Perspectives on plant population biology, Dirzor R, Sarukan J (eds) Sinauer, Sunderland pp. 229-241.

Artyukova EV, Kozyrenko MM, Koren OG, Muzarok TI, Reunova GD, Zhuravlev Y N (2004) RAPD and allozyme analysis of genetic diversity in *Panax ginseng* C.A. Meyer and *P. quinquefolius* L. *Russ J Genet* 40(2): 178-185.

Bai D, Brandle J, Reeleder R (1997) Genetic diversity in North American ginseng (*Panax quinquefolius* L.) grown in Ontario detected by RAPD analysis. *Genome 40* 111-115.

Beardmore JA (1983) Extinction, survival and genetic variation. In: Genetics and conservation, Schoenwald-Cox

CM, Chamber SM, Macbryde B, Thomas L (eds) Benjamin-Cummings, Menlo Park pp. 125-151.

Hamrick JL, Godt MJW (1989) Allozyme diversity in plant species. *In:* Plant population genetics, breeding and germplasm resources, Brown AHD, Clegg MT, Kahler AL, Weir BS (eds). Sinauer Associates, Sunderland pp. 43-63.

Hamrick JL, Godt MJW (1996) Effects of life history traits on genetic diversity in plant species. *Philos Trans R Soc Lond Ser B Biol Sci* 351: 1291-1298.

Hogbin PM, Peakall R (1999) Evalution of the contribution of the genetic research to the management of the endangered plant Zieria prostrate. *Conser Biol* 13: 514-522.

Jennifer MC and Hamrick JL (2004) Genetic diversity in harvested and protected populations of wild American ginseng, *Panax quinquefolius* L. (Araliaceae). *American J Bot* 91(4): 540-548.

Li S, Li J, Yang XL, Cheng Z, Zhang WJ (2011) Genetic diversity and differentiation of cultivated ginseng (*Panax ginseng* C. A. Meyer) populations in North-East China revealed by inter-simple sequence repeat (ISSR) markers. *Genet Resour Crop Evol* 58: 815-824.

Lu X, Liu L, Gong Y, Zhao L, Song X, Zhu X (2009) Cultivar identification and genetic diversity analysis of broccoli and its related species with RAPD and ISSR markers. *Sci. Horticult* 122: 645-648.

Nagoaka T, Ogihara Y (1997) Applicability of inter-simple sequence repeat polymorphism in wheat for use as DAN markers in comparison to RFLP and RAPD markers. *Theor Appl Genet* 93: 133-139.

Nei M (1978) Estimation of average heterozygosity and genetic distance from a small number of individuals. *Genetics* 89: 583-590.

Nybom H, Bartish IV (2000) Effects of life history traits and sampling strategies on genetic diversity estimates obtained with RAPD markers in plants. *Perspect Plant Ecol Evol Syst* 3: 93-114.

Rohlf FJ (2004). NTSYSpc - Numerical taxonomy and multivariate analysis system version 2.1 - User guide. Applied Biostatistics Inc.

Roy SC, Chakraborty BN (2009) Genetic diversity and relationships among tea (*Camellia sinensis*) cultivars asrevealed by RAPD and ISSR based fingerprinting. *Indian J Biotechnol.* 8: 370-376.

Schaal BA, Hayworth DA, Olsen KM, Rauscher JT, Smith WA (1998) Phylogeographic studies in plants: problems and prospects. *Mol Ecol* 7: 465-474.

Vicente MCD, Lopez C, Fulton T (2003) Genetic diversity analysis with molecular marker data: Learning module. Journal of Biotechnology 14(4): 619-627, 2016

International Plant Genetic Resources Institute (IPGRI) and Cornell University.

Trieu LN., Mien NT, Tien TV, Ket TV, Duy NV (2016) Genetic diversity of *Panax stipuleanatus* Tsai in North Vietnam detected by inter simple sequence repeat (ISSR) markers. *Biotech & Biotech Equip* 30(3): 506-511.

Weising K, Nybom H, Wolff K, Kahl G (2005) DNA fingerprinting in plants principles, mMethods, and applications (second Edition). Cpc Press Taylor & Fancies Group.

Williams JGK, Kubelik AR., Livak KJ, Rafalski JA, Tingey SV (1990) DNA polymorphisms amplified by arbitrary primers are useful as genetic markers. *Nucl Acids Res* 18: 6531-6535.

Yeh, FC, Boyle TJB (1997) Population genetic analysis of

co-dominant and dominant markers and quantitative traits. *Belgian J Bot* 129-157

Zhou SL, Xiong GM, Li ZY, Wen J (2005) Loss of genetic diversity of domesticated *Panax notoginseng* F.H. Chen as evidenced by ITS sequence and AFLP polymorphism: A comparative study with *P. stipuleanatus* H.T. Tsai et K.M. Feng *Acta Bot Sin* 47(1): 107–115.

Zhuravlev YN, Reunova GD, Kats IL, Muzarok TI, Bondar AA (2010) Research genetic variability and population structure of endangered *Panax ginseng* in the Russian Primorye. *Chinese Medicine* 1-9.

Zietkiewicz E, Rafalski A, Labuda D (1994) Genome fingerprinting by simple sequence repeat (SSR) anchored polymerase chain reaction amplification. *Genomics* 20: 176-183.

# ĐA DẠNG DI TRUYỀN QUÂN THẾ *PANAX VIETNAMENSIS* VAR. *FUSCIDISCUS* K. KOMATSU, S.ZHU & S.Q.CAI Ở TÂY BẮC VIỆT NAM BẰNG CHỈ THỊ PHÂN TỬ INTER SIMPLE SEQUENCE REPEAT

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# TÓM TẮT

*Panax vietnamensis* var. *fuscidiscus* là loài dược liệu quý hiếm được tìm thấy ở huyện Sình Hồ và Mường Tè thuộc tỉnh Lai Châu, Việt Nam. Hiện nay, loài đang trong tình trạng bị đe dọa nghiêm trọng do khai thác quá mức và môi trường sống bị phá hủy. Thông tin đầy đủ về điều kiện sống và đa dạng di truyền làm cơ sở cho việc bảo tồn và phát triển. Trong nghiên cứu này, chỉ thị phân tử inter simple sequence repeat được sử dụng để khảo sát đa dạng di truyền và biến dị của 46 cá thể của quần thể phân bố tự nhiên ở Việt Nam. Đa dạng di truyền ở mức độ quần thể là cao ( $H_e = 0.2300$ , I = 0.3665 và PPB = 96.98 %). Nhóm cá thể trưởng thành có mức độ đa dạng di truyền cao ( $H_{eO} = 0.2291$ ,  $I_O = 0.3563$  và  $PPB_O = 84.34$  %) so với nhóm cá thể có tuổi nhỏ ( $H_{eY} = 0.2086$ ,  $I_y = 0.3291$  và  $PPB_Y = 81.5$  %). Biệt hóa di truyền trong quần thể cao ( $G_{ST} = 0.0499$ ) với khoảng cách di truyền giữa các nhóm tuổi là 0.0298. Hệ số tương đồng giữa các cá thể ở nhóm tuổi nhỏ (Maximum = 0.873, Minimum = 0.614 và Average = 0.741) so với các cá thể ở nhóm tuổi nhỏ (Maximum = 0.916, Minimum = 0.596 và Average = 0.759). Dưới tác động của con người khai thác và phá môi trường sống, số lượng cá thể của loài rất ít, vùng phân bố rất hẹp, khai thác quá mức cũng dẫn đến làm suy giảm da dạng di truyền. Kết quả về đa dạng di truyền và biến dị cho thấy có sự suy giảm mạnh, nên ưu tiên bảo vệ toàn vẹn khu phân bố của quần thể.

Keywords: Đa dạng di truyền, inter simple sequences repeat, Panax vietnamensis var. fuscidiscus, Việt Nam