

Genet Resour Crop Evol (2011) 58:189–207  
DOI 10.1007/s10722-010-9562-8

## RESEARCH ARTICLE

# Detection of genetic integrity of conserved maize (*Zea mays* L.) germplasm in genebanks using SNP markers

Weiwei Wen · Suketoshi Taba · Trushar Shah ·  
Victor H. Chavez Tovar · Jianbing Yan

Received: 10 December 2009 / Accepted: 12 April 2010 / Published online: 7 May 2010  
© Springer Science+Business Media B.V. 2010

**Abstract** Twenty maize landrace accessions regenerated and conserved in five maize genebanks were investigated for genetic integrity using 1,150 Single Nucleotide Polymorphisms (SNPs) and 235 SNP haplotypes. The genetic diversity of three accessions changed significantly in terms of the average number of alleles per locus. Ten out of twenty accessions had significantly different SNP allelic frequencies, either after regeneration or in the same accession held in different genebanks. The proportion of loci with significant changes in SNP allelic frequency was very low (37/1,150). Changes in the major allelic frequency (MAF) for the majority of SNP loci (60.2–75.2%) were less than 0.05. For SNP haplotypes, the genetic diversity of four accessions changed significantly in terms of average number of haplotype alleles and polymorphic information content (PIC) per locus. The proportion of SNP haplotype

alleles lost in the later generations ranged between 0 and 22.6%, and at the same time 0–19.9% of the SNP haplotype alleles appeared in later generations, however, these were absent in the earlier generations. Dynamic changes in genetic integrity, in terms of presence and absence of genes (alleles), by both SNP and SNP haplotype analysis were detected during regeneration. A suboptimum number of ears harvested in one generation can be combined with those from another, repeated regeneration to capture the diversity of the previous generation. Use of molecular markers during regeneration of accessions can help in understanding the extent of genetic integrity of the maize accessions in *ex situ* genebanks and in recommending the best practice for maintaining the original genetic diversity of the genebank accessions.

**Keywords** Conservation · Genebank · Genetic integrity · Maize · Regeneration · SNP · *Zea mays*

**Electronic supplementary material** The online version of this article (doi:[10.1007/s10722-010-9562-8](https://doi.org/10.1007/s10722-010-9562-8)) contains supplementary material, which is available to authorized users.

W. Wen · S. Taba · V. H. Chavez Tovar · J. Yan (✉)  
International Maize and Wheat Improvement Center  
(CIMMYT), Apdo. Postal 6-641, 06600 Mexico,  
DF, Mexico  
e-mail: [j.yan@cgiar.org](mailto:j.yan@cgiar.org)

T. Shah  
International Crops Research Institute for the Semi-Arid  
Tropics (ICRISAT), Patancheru Hyderabad, Andhra  
Pradesh 502324, India

## Introduction

Maize, which originated in southern Mexico, is one of the most important crops in the world. Seed multiplication and regeneration of the original landrace accessions has been performed in *ex situ* genebanks to fulfill seed requests for further characterization and for use in breeding and research. In the course of seed preservation for the active collection, if seed viability

of the accessions drops below 85% germination or the number of seeds falls below 1,500, the accession is regenerated (Taba et al. 2004). The diverse phenotypes and widely differing adaptations of the races of maize and local cultivars are often constraints to regeneration (Taba and Twumasi-Afriyie 2008).

Reducing and managing the loss of genetic integrity of the conserved germplasm during regeneration is an important objective of maize genetic resource conservation programs. The management of seed accessions in different maize genebanks can lead to differential loss of genetic integrity. Identification and rationalization of duplicate accessions in genebanks requires information on the genetic integrity of the accessions. To recommend better practices for maintaining panmictic populations of maize accessions, several studies on genetic integrity during seed multiplication and regeneration using molecular markers from other seed or clonally propagated crops can be useful to compare with those of maize genebank accessions.

The total number of maize and wild species accessions currently held in trust at the International Maize and Wheat Improvement Center (CIMMYT) genebank is 27,441 and this is increased each year with new introductions. Most accessions are maize landraces (24,191); however, breeding lines, gene pools, populations and cultivars, and teosinte are also preserved in this *ex situ* collection. In CIMMYT, collection of germplasm accessions and gene pool enhancement are carried out to promote use of maize genetic resources, towards the ultimate goal of developing improved cultivars. The original maize collections date back to the 1940s, when some 2,000 Mexican accessions were collected and the races of maize in Mexico were studied. Accessions collected from the US National Research Council (NRC) were regenerated several times and the original seed stocks are no longer available. The current seed sources have undergone several generations of seed multiplication. At present, a few seed stocks of only two or three generations are available for comparing genetic integrity. Efforts to regenerate endangered, national seed collections of maize in Latin America started in 1993 and have continued to present (Taba et al. 2005). CIMMYT introduced new accessions and curated some for which seed was in short supply. Seed lots were introduced and regenerated at CIMMYT or at other genebanks. In some instances, the same accession went through several regeneration attempts with

limited success. Different seed sources of the same accessions can be compared if they are genetically different. In this project, the cooperative banks of CIMMYT were the banks of the International Institute of Tropical Agriculture (IITA, Nigeria), the Autonomous National Institute for Agricultural Research (INIAP, Ecuador), the Brazilian Agricultural Research Corporation (EMBRAPA, Brazil), and the National Institute for Investigation in Forestry, Agriculture and Fishery (INIFAP, Mexico).

Previous studies using molecular tools have been performed on the genetic integrity of genebank accessions of some crop species during regeneration. Börner et al. (2000) studied the genetic integrity of eight wheat (*Triticum aestivum* L.) accessions conserved in the Gatersleben genebank and regenerated up to 24 times using 9 wheat microsatellite markers; they concluded that microsatellites can be used as a simple and reliable marker system to verify the integrity and genetic stability of genebank accessions. The genetic integrity of 6 accessions, represented by 14 sub-populations of the open-pollinating species of rye (*Secale cereale* L.), was investigated using 10 rye microsatellites (Chebotar et al. 2003) and found that 4 accessions had significantly different allele frequencies and nearly 50% of alleles identified in the original samples were lost in the regenerated samples. However, some alleles detected in the most recently propagated sub-populations were not observed in the investigated plants of the original seed stocks (Chebotar et al. 2003). Work by van Hintum et al. (2007) used Amplified Fragment Length Polymorphisms (AFLPs) to show that the distribution of genetic diversity in a *Brassica oleracea* collection was related to the effects of regeneration, and argued that accessions with similar levels of differentiation over generations may be combined safely. Also, in a *Brassica oleracea* collection, Soengas et al. (2009) investigated the effect of regeneration on the genetic integrity of 3 accessions based on 25 simple sequence repeats (SSRs) and found that there were significant changes in the population structure and the allelic frequency at individual loci due to the action of genetic drift, directional selection, and possibly assortative mating.

Molecular characterization of germplasm accessions is a useful tool for better management and to study genetic diversity and integrity of conserved germplasm. The new molecular marker system, known as single nucleotide polymorphism (SNP), is

widely used in different crops (Rostoks et al. 2006; Hyten et al. 2008; Muchero et al. 2009), including maize (Buckler et al. 2009; Lu et al. 2009; Yan et al. 2009, 2010). It is highly polymorphic, evenly distributed, co-dominant, accurate, reproducible, high-throughput and cost-effective. SNPs are regarded as the ideal marker system for research which includes genetic diversity analysis, linkage map construction, QTL mapping, and marker-assisted selection (MAS; Yan et al. 2010). In previous studies, a limited number of molecular markers and accessions were employed to analyze the genetic integrity of the crop species mentioned above. Thus, these studies may not reflect the whole picture of genetic changes at a genome-wide level. Few studies of genetic integrity in maize genebank accessions have been reported. In the present study, we genotyped 2–3 comparable generations of twenty maize accessions conserved in five genebanks with a custom GoldenGate assay containing 1,536 SNPs well distributed across the maize genome. The objectives were to detect genetic differences between the comparable generations of the genebank accessions and to investigate the effect of regeneration in different environments and by different genebanks on the genetic integrity of maize accessions.

## Materials and methods

### Plant material and DNA isolation

Seed stocks for the accessions in this study were collected from the CIMMYT maize genebank. The original reference seed sources were the introductions from the cooperative genebanks to the CIMMYT maize genebank. The subsequent generations were the seed accessions regenerated either at CIMMYT or received from the Latin American cooperative genebanks, where regeneration of the same accessions was carried out during the cooperative maize germplasm regeneration project during 1993–2005 (Tabata et al. 2005). The same maize accessions which were regenerated in different times and locations in Latin American maize genebanks were compared for their genetic integrity. The earlier generations (cycles) were recorded as generation 1 and the later generations (cycles) were recorded as generations 2 and 3 (Table 1). Among the accessions included in the

present study, with respect to the genebank accessions (Table 1), were four accessions of the NRC collection from Ecuador in 1954 deposited at the CIMMYT genebank in 1974 after having been regenerated at Carhuas, Peru. They were then sent to the Santa Catalina station of INIAP, Ecuador, for regeneration. CIMMYT received the regenerated seeds sent by the INIAP maize genebank for long term conservation. Both the CIMMYT and EMBRAPA genebanks have preserved the same seed accessions with different seed origins. The EMBRAPA genebank sent them to CIMMYT as the duplicate accessions in 1995. Accessions from the INIFAP genebank were regenerated at Celaya, Mexico, in 2003. The CIMMYT genebank regenerated the same accessions before 2003 from its own seed sources. For the accessions included from the CIMMYT genebank, the seed sources of the earlier generations differ in regeneration sites and years, and the later generations differ in years and different crop seasons at the Tlaltizapán experiment station, in the state of Morelos, Mexico. Regarding materials from the IITA genebank, the CIMMYT genebank received the accessions in 2005 and regenerated them in 2007 and 2009 from the same seed lots, growing them in 8 five-meter rows at Tlaltizapán. Effects on the genetic integrity of the repeated samplings and regenerations were monitored for the seed accessions obtained from different genebanks. Following the recommended procedures for regeneration, 256 plants per plot (60 m<sup>2</sup>) were grown in 16 rows 5 m long, with a spacing of 75 cm between rows. Artificial pollination was carried out by chain crossing, using each plant as a male and female. At least 100 ears were represented in the next generation. The number of seeds of each ear is balanced in making the seed bulk for genebank storage. Seed accessions with records of the number of ears in regeneration were included in this study as much as possible for investigating the effects of sampling size on the genetic integrity of the accessions. The detailed history and passport information of the accessions can be found in Table S1.

In the screenhouse at El Batán, CIMMYT, 200 seeds of each generation were sown during April–May 2009. Depending on seed germination of the accessions, 120–135 seedlings (5–6 leaf stage) per generation were used for DNA extraction. DNA was extracted using a CTAB procedure (CIMMYT Applied Molecular Genetics Laboratory 2003). Each

**Table 1** Accessions and generations of seed origins investigated in this study

Genebank	Accession	Number of regeneration	Generation no.	Generation name	Size of population	Ears harvested	
INIAP	ECUA553	1	1	CA74 -1052-...	135	55	
			2	SC98 -...-44	120	122	
	ECUA937	1	1	CA74 -1174-...	135	88	
			2	SC98 -...-24	120	85	
	ECUA497	1	1	CA74 -1016-...	135	76	
			2	SC97 -...-61	135	86	
	ECUA443	1	1	CA74 -975-...	120	118	
			2	SC97 -...-30	135	136	
	EMBRAPA	BRAZ3881LC		1	TL88A-1901-106	120	67
				2	BRA95 -...-58	135	88
BRAZ3937LC			1	TL88A-1901-139	135	67	
			2	BRA95 -...-60	120	160	
BRAZ3962LC			1	TL88A-1901-153	120	64	
			2	BRA95 -...-61	120	38	
BRAZ4012LC		1	TL88A-1901-185	135	50		
		2	BRA95 -...-62	120	100		
INIFAP	COLI21		1	TL01A-1903-5	135	56	
			2	CE-2003-183	120	70	
	NAYA1		1	TL98B-6903-98	135	46	
			2	CE-2003-189	120	77	
	NAYA 208		1	TL00A-1903-60	135	62	
			2	CE-2003-191	120	66	
	NAYA29		1	TL70B-657-...	135	164	
			2	CE-2003-220	120	28	
CIMMYT	PUEB 42	1	1	BA-69-352	135		
			2	TL94B-6920-93	120	132	
	OAXA265	1	1	TL71B-3421	135		
			2	TL95A-1920-223	120	98	
	CRICA264	2	1	TL73B-4431	135		
			2	AF05A-0903-22	135	108	
			3	TL06A-1903-204	120	30	
GUER 5	2	1	TL-69B-1836	120			
		2	TL94A-1920-214	120	54		
IITA	TANZANIA 87–236	1	1	NIGERIA-05	120		
			2–1	TL07A-1901-11	120	40	
			2–2	TL09A-1903-8	135	24	
	BEN-RB 89–466	1	1	NIGERIA-05	135		
			2–1	TL07A-1901-71	135	30	
			2–2	TL09A-1903-62	135	29	
	BEN-RB 89–84	1	1	NIGERIA-05	120		
			2–1	TL07A-1901-137	120	20	
			2–2	TL09A-1903-123	135	20	

**Table 1** continued

Genebank	Accession	Number of regeneration	Generation no.	Generation name	Size of population	Ears harvested
	BEN-RB 89–428	1	1	NIGERIA-05	120	
			2–1	TL07A-1901-61	135	10
			2–2	TL09A-1903-54	120	59

INIAP = the Autonomous National Institute for Agricultural Research; EMBRAPA = the Brazilian Agricultural Research Corporation; INIFAP = the National Institute for Investigation in Forestry, Agriculture and Fishery; CIMMYT = International Maize and Wheat Improvement Center; IITA = the International Institute of Tropical Agriculture

DNA sample was obtained from the bulked leaves of 15 individuals with equal quantity from each individual. Eight or nine replicated sets of the DNA samples were used for genotyping each generation of accessions. In total, 382 DNA samples were genotyped, including 173 samples from the earlier generations and 209 samples from the comparable generation of later seed origins. One accession from CIMMYT and 4 accessions from IITA had 3 generations of seed origins.

#### SNP genotyping

A custom oligo pool assay (OPA) containing 1,536 SNPs has been developed by Yan et al. (2009) where a total of 943 SNPs with high quality and minor allelic frequency greater than 0.05 were obtained. To develop the OPA used in this study, the 943 high quality SNPs were combined with 593 SNPs selected from the Panzea database ([www.panzea.org](http://www.panzea.org)) on the basis of a designability score higher than 0.5. Designability scores were provided by the Illumina Company, and a score greater than 0.5 indicates that a SNP has a relatively higher probability of success when used in a GoldenGate assay. A total of 382 bulked samples were genotyped using this OPA. SNP genotyping was performed using the Illumina BeadStation 500 G (Illumina, Inc., San Diego, CA, USA) at the Cornell University Life Sciences Core Laboratories Center using the protocol described by Fan et al. (2006). After SNP genotyping, allele calling and further analyses were carried out as described in Yan et al. (2010).

#### Haplotype analysis

The 1,536 SNP reference sequences were used to carry out a BlastN search against the maize accession golden path (AGP) version 1 for B73 (<http://>

[www2.genome.arizona.edu/genomes/maize](http://www2.genome.arizona.edu/genomes/maize); Schnable et al. 2009). The top blast-hit was considered with an  $e$ -value threshold of  $e^{-18}$ . A total of 1,420 unique SNPs out of 1,536 have been mapped *in silico* onto the maize genome, excluding the ones with unknown locations and multiple hits to the genome.

Only 1,150 polymorphic SNPs with good quality and less than 20% missing data were used for final data analysis. A total of 1,116 from 1,150 SNPs were mapped *in silico* onto the maize genome. The relative distance for each SNP was summed and SNPs developed from the same genes were assigned to a uni-locus. A total of 701 uni-loci were identified and 466 SNPs were found unlinked to other SNPs. Remaining SNPs formed the linked groups of two or more SNPs from contiguous DNA sequences, which constituted 235 loci. SNPs from the same locus were grouped into haplotypes that were recorded as alleles; in this way, each locus could have multiple alleles, increasing the information content of the markers. If the genotype of any SNP at a locus was missing in a sample, the locus was regarded as missing an allele in that sample (Yan et al. 2009).

#### Data analysis

Summary statistics for genetic diversity were calculated for each generation of the accessions using the software PowerMarker V3.25 (Liu and Muse 2005). The parameters included average number of alleles per locus, observed heterozygosity, gene diversity (expected heterozygosity), number of heterozygous loci, polymorphic information content (PIC) and pairwise  $F$ -statistics. Gene diversity was calculated at each locus as  $2n(1 - \sum p_u^2)/(2n - 1 - f)$ , where  $n$  is the sample size,  $p_u$  is the frequency of the  $u$ th allele, and  $f$  is the inbreeding coefficient estimated from genotype frequencies (Weir 1996). PIC was used to refer to the

relative value of each marker with respect to the amount of polymorphism exhibited, as described by Botstein et al. (1980). *F*-statistics developed by Wright (1965) are a very useful measure of population subdivision, of which *F*<sub>st</sub> describes the amount of inbreeding-like effects within subpopulations.

Allelic frequencies at each SNP locus were calculated and compared between the comparable generations of the seed origins by year in the same accession. The probability of allelic frequency change of individual loci after regeneration was detected using Fisher's exact test (Mehta et al. 1984). The null hypothesis was rejected at  $P \leq 0.05$ .

Genetic distances, based on allelic frequencies (Nei 1973), were calculated for pair-wise comparisons between generations using PowerMarker V3.25 to detect how far they had diverged from each other. We constructed the dendrogram using the unweighted pair-group method with arithmetic averages (UPGMA) (Sneath and Sokal 1973).

## Results

### Performance and characterization of SNPs

In our OPA, 1,303 SNPs were successfully called in 382 samples with reliable scores and less than 20% missing data, including 153 monomorphic SNPs across all seed accessions. A total of 1,150 polymorphic SNPs were used for further analysis. The number of SNPs per chromosome ranged from 64 (chromosome 9) to 187 (chromosome 1) (Table 2), and 34 were mapped onto contigs of unknown location. Details of all 1,150 polymorphic SNPs are given in Table S2. The table shows 701 loci representing 1,116 SNPs with physical distance information, within which 466 loci contained only one SNP and the remaining 235 loci contained two or more SNPs. Haplotypes were constructed from the loci with more than one SNP (Table 2). In the 382 samples, the 235 SNP haplotypes had a total of 2,116 alleles ranging from 2 to 60 alleles per locus.

### Molecular diversity and genetic distance among generations of each accession

For 20 accessions from 5 different genebanks, the average number of alleles per locus, average observed and expected heterozygosity and the numbers of

**Table 2** Summary of single nucleotide polymorphisms (SNPs) and SNP haplotypes from all chromosomes

Chromosome	SNP number	Uni-locus	
		1 + SNPs <sup>a</sup>	1SNP <sup>b</sup>
1	187	47	61
2	140	27	60
3	128	22	64
4	116	28	44
5	135	30	61
6	67	11	38
7	102	23	36
8	101	21	41
9	64	16	25
10	76	10	36
Unknown	34		
Total	1150	235	466

<sup>a</sup> Represents the uni-locus, contains more than one SNP

<sup>b</sup> Represents the uni-locus, contains only one SNP

heterozygous loci in each generation are shown in Table 3. In general, the average number of alleles per locus, the number of heterozygous loci and the average observed and expected heterozygosity decreased in the later generations of the accessions; however, no significant difference was observed. On the other hand, the average number of alleles per locus between both generations of seed origins was significantly different for the accessions ECUA497, from INIAP, and NAYA1 and NAYA29, from INIFAP.

A dendrogram of genetic distances among the different generations of seed origins was constructed to investigate the divergence (Fig. 1). Those of the same accessions were clustered together in the closest branches. The smallest genetic distances of the same accession were obtained from accessions in INIAP, which were 0.0044, 0.0048, 0.0046 and 0.0056 for ECUA553, ECUA937, ECUA497 and ECUA443, respectively. The greatest genetic distances were obtained in the accessions NAYA29, from INIFAP, GUER 5, from CIMMYT, and BEN-RB 89–428, from IITA, which were 0.0313, 0.0395 and 0.0301, respectively.

Estimation of genetic distance concurred with the pair-wise *F*<sub>st</sub> values (Table 4). The pair-wise *F*<sub>st</sub> between the two generations of ECUA553 from INIAP had the lowest value (0.0083) amongst all pair-wise *F*<sub>st</sub> values. This corresponded with its genetic distance

**Table 3** Genetic diversity of each generation of accessions from five genebanks

Genebank	Accession	Generation no.	Average number of alleles per locus <sup>a</sup>	Average expected heterozygosity	Average observed heterozygosity	Number of heterozygous loci	
INIAP	ECUA553	1	1.33 ± 0.01	0.12 ± 0.01	0.20 ± 0.01	374	
		2	1.32 ± 0.01	0.12 ± 0.01	0.20 ± 0.01	365	
	ECUA937	1	1.32 ± 0.01	0.12 ± 0.01	0.20 ± 0.01	366	
		2	1.31 ± 0.01	0.12 ± 0.01	0.19 ± 0.01	354	
	ECUA497	1	1.34 ± 0.01*	0.12 ± 0.01	0.20 ± 0.01	394	
		2	1.29 ± 0.01*	0.11 ± 0.01	0.19 ± 0.01	340	
	ECUA443	1	1.38 ± 0.01	0.14 ± 0.01	0.23 ± 0.01	434	
		2	1.38 ± 0.01	0.14 ± 0.01	0.22 ± 0.01	436	
	EMBRAPA	BRAZ3881LC	1	1.41 ± 0.01	0.15 ± 0.01	0.25 ± 0.01	470
			2	1.41 ± 0.01	0.15 ± 0.01	0.24 ± 0.01	467
BRAZ3937LC		1	1.34 ± 0.01	0.12 ± 0.01	0.20 ± 0.01	385	
		2	1.31 ± 0.01	0.12 ± 0.01	0.19 ± 0.01	354	
BRAZ3962LC		1	1.36 ± 0.01	0.13 ± 0.01	0.21 ± 0.01	417	
		2	1.39 ± 0.01	0.14 ± 0.01	0.22 ± 0.01	447	
BRAZ4012LC		1	1.41 ± 0.01	0.15 ± 0.01	0.24 ± 0.01	468	
		2	1.37 ± 0.01	0.14 ± 0.01	0.23 ± 0.01	426	
INIFAP	COLI21	1	1.46 ± 0.01	0.16 ± 0.01	0.26 ± 0.01	525	
		2	1.47 ± 0.01	0.17 ± 0.01	0.28 ± 0.01	535	
	NAYA1	1	1.51 ± 0.01*	0.18 ± 0.01	0.29 ± 0.01	581	
		2	1.46 ± 0.01*	0.18 ± 0.01	0.29 ± 0.01	530	
	NAYA 208	1	1.49 ± 0.01	0.19 ± 0.01	0.30 ± 0.01	560	
		2	1.49 ± 0.01	0.18 ± 0.01	0.29 ± 0.01	562	
	NAYA29	1	1.47 ± 0.01*	0.17 ± 0.01	0.26 ± 0.01	535	
		2	1.42 ± 0.01*	0.16 ± 0.01	0.27 ± 0.01	481	
CIMMYT	PUEB 42	1	1.33 ± 0.01	0.12 ± 0.01	0.20 ± 0.01	388	
		2	1.34 ± 0.01	0.12 ± 0.01	0.20 ± 0.01	385	
	OAXA265	1	1.44 ± 0.01	0.16 ± 0.01	0.25 ± 0.01	496	
		2	1.43 ± 0.01	0.16 ± 0.01	0.26 ± 0.01	508	
	CRICA264	1	1.41 ± 0.01	0.15 ± 0.01	0.25 ± 0.01	455	
		2	1.41 ± 0.01	0.15 ± 0.01	0.25 ± 0.01	470	
		3	1.40 ± 0.01	0.15 ± 0.01	0.25 ± 0.01	470	
	GUER 5	1	1.40 ± 0.01	0.16 ± 0.01	0.26 ± 0.01	459	
		2	1.40 ± 0.01	0.15 ± 0.01	0.25 ± 0.01	462	
	IITA	TANZANIA 87–236	1	1.38 ± 0.01	0.15 ± 0.01	0.25 ± 0.01	446
2–1			1.39 ± 0.01	0.14 ± 0.01	0.23 ± 0.01	477	
2–2			1.41 ± 0.01	0.15 ± 0.01	0.25 ± 0.01	441	
BEN-RB 89–466		1	1.37 ± 0.01	0.13 ± 0.01	0.21 ± 0.01	403	
		2–1	1.35 ± 0.01	0.13 ± 0.01	0.21 ± 0.01	382	
		2–2	1.33 ± 0.01	0.12 ± 0.01	0.21 ± 0.01	429	
BEN-RB 89–84		1	1.28 ± 0.01	0.11 ± 0.01	0.18 ± 0.01	366	
		2–1	1.32 ± 0.01	0.11 ± 0.01	0.18 ± 0.01	343	
		2–2	1.30 ± 0.01	0.11 ± 0.01	0.18 ± 0.01	325	

**Table 3** continued

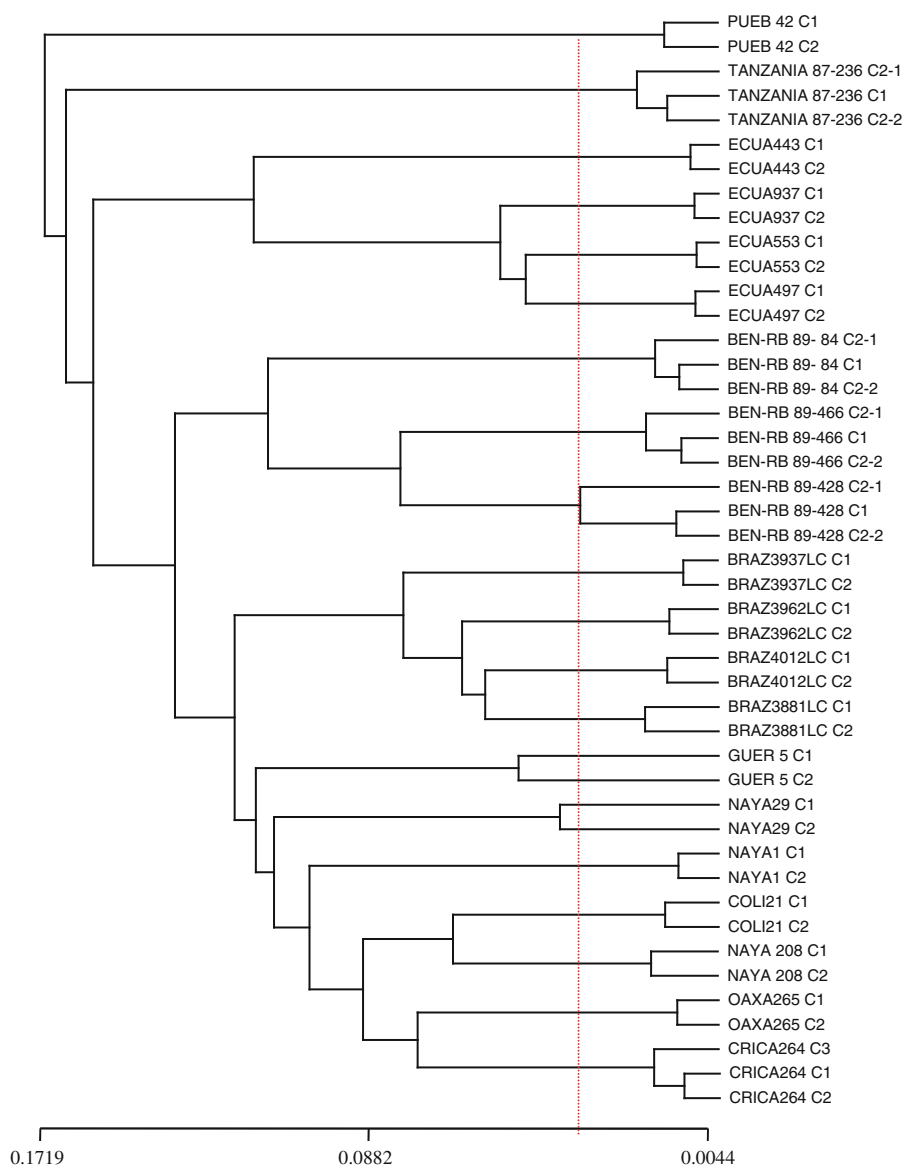
Genebank	Accession	Generation no.	Average number of alleles per locus <sup>a</sup>	Average expected heterozygosity	Average observed heterozygosity	Number of heterozygous loci
	BEN-RB 89–428	1	1.38 ± 0.01	0.14 ± 0.01	0.22 ± 0.01	445
		2–1	1.39 ± 0.01	0.14 ± 0.01	0.21 ± 0.01	4353
		2–2	1.38 ± 0.01	0.14 ± 0.01	0.23 ± 0.01	435

INIAP = the Autonomous National Institute for Agricultural Research; EMBRAPA = the Brazilian Agricultural Research Corporation; INIFAP = the National Institute for Investigation in Forestry, Agriculture and Fishery; CIMMYT = International Maize and Wheat Improvement Center; IITA = the International Institute of Tropical Agriculture

<sup>a</sup> Mean ± Standard Error

\* Significant difference between original and regenerated cycles ( $P = 0.05$ )

**Fig. 1** Unweighted pair group method with arithmetic averages (UPGMA) dendrogram of relationships among generations based on Nei's genetic distance. Generations with genetic distances beyond the red line (genetic distance = 0.0247,  $F_{st} = 0.1221$ ) are significantly different





**Table 4** Genetic distance (top diagonal) and pair-wise  $F_{st}$  (bottom diagonal) between generations of each accession

Genebank	Accession	Generation no.	$F_{st}$ /Genetic distance		
			Generation 1	Generation 2(2–1)	Generation 3(2–2)
INIAP	ECUA553	1	–	0.0044	
		2	0.0083	–	
	ECUA937	1	–	0.0048	
		2	0.0121	–	
	ECUA497	1	–	0.0046	
		2	0.0277	–	
	ECUA443	1	–	0.0056	
		2	0.0094	–	
EMBRAPA	BRAZ3881LC	1	–	0.0145	
		2	0.0574	–	
	BRAZ3937LC	1	–	0.0069	
		2	0.0279	–	
	BRAZ3962LC	1	–	0.0096	
		2	0.0323	–	
	BRAZ4012LC	1	–	0.01	
		2	0.0366	–	
INIFAP	COLI21	1	–	0.0106	
		2	0.0274	–	
	NAYA1	1	–	0.008	
		2	0.0283	–	
	NAYA 208	1	–	0.0133	
		2	0.0367	–	
	NAYA29	1	–	0.0313	
		2	0.1054	–	
CIMMYT	PUEB 42	1	–	0.0106	
		2	0.0092	–	
	OAXA265	1	–	0.0083	
		2	0.019	–	
	CRICA264	1	–	0.0069	0.0115
		2	0.0439	–	0.0143
		3	0.06	0.0168	–
	GUER 5	1	–	0.0395	
2		0.1467	–		
IITA	TANZANIA 87–236	1	–	0.0149	0.0102
		2–1	0.0636	–	0.0175
		2–2	0.0356	0.0767	–
	BEN-RB 89–466	1	–	0.0123	0.0073
		2–1	0.0606	–	0.0164
		2–2	0.0273	0.0916	–
	BEN-RB 89–84	1	–	0.0098	0.0075
		2–1	0.0532	–	0.0149
		2–2	0.0367	0.0915	–
	BEN-RB 89–428	1	–	0.0247	0.0084

**Table 4** continued

Genebank	Accession	Generation no.	<i>Fst</i> /Genetic distance		
			Generation 1	Generation 2(2–1)	Generation 3(2–2)
		2–1	0.1221	–	0.0301
		2–2	0.0206	0.1484	–

INIAP = the Autonomous National Institute for Agricultural Research; EMBRAPA = the Brazilian Agricultural Research Corporation; INIFAP = the National Institute for Investigation in Forestry, Agriculture and Fishery; CIMMYT = International Maize and Wheat Improvement Center; IITA = the International Institute of Tropical Agriculture

between the two generations, which was also the lowest among the genetic distances between the generations in all accessions investigated. The maximum pair-wise *Fst* value (0.1467) was obtained for the accession GUER 5, from CIMMYT, which had the largest genetic distance between two different generations of seed origins.

#### Allelic frequency changes of SNPs among generations

The number of polymorphic loci and the average major allelic frequency (MAF) changes between the two comparable generations of all accessions are shown in Table 5. The smallest MAF changes were obtained for the four accessions from INIAP, all of which were less than 0.04, and the average MAF changes were 0.03 for all accessions in this genebank. The largest MAF changes were detected for GUER 5, from CIMMYT, with an average of 0.093. The INIFAP accessions had the greatest average MAF changes (0.062) amongst the five genebanks. Most of the markers had MAF changes between 0 and 0.05 in all accessions, and the number of markers decreased continuously from the range 0–0.05 to 0.45–0.5 amongst the five genebanks (Fig. 2).

Significant difference ( $P = 0.05$ ) in allelic frequencies of each SNP between the two generations of the accessions were detected and are listed in Table 6. No significant SNP frequency change was detected for nearly half of the accessions studied. They comprised 4 accessions from INIAP, 1 accession from EMBRAPA, 3 accessions from INIFAP, 1 accession from IITA and 1 accession from CIMMYT. A total of 37 SNPs with significant allelic frequency changes were detected for the other 10 accessions. The SNP number within them ranged between 1 and 11 in each accession from four genebanks (Table 6). In 25 out of 37 SNPs, MAF increased to 1 and

heterozygosity decreased at the same time in later generations. The reduction of MAF from 1 to 0.5 was also observed in some cases, which was accompanied by the increase in heterozygosity (Table 6).

#### Allelic frequency changes of SNP haplotypes among generations

SNP haplotypes were also used to characterize the average number of alleles and PIC per locus for each generation (Table 7). In general, no significant difference for the average number of haplotype alleles per locus was identified between the earlier and later generations, except for one accession (ECUA497) from INIAP and two accessions (NAYA1 and NAYA29) from INIFAP. The average number of haplotype alleles per locus ranged between  $1.47 \pm 0.05$  and  $2.03 \pm 0.06$  in the earlier generations of BEN-RB 89–84 and NAYA29, respectively. The average number of haplotype alleles per locus increased in 9 accessions and decreased or remained the same in the other 11 accessions in the later generations, compared to the earlier ones. For two accessions (NAYA1 and NAYA29) from INIFAP, and one accession (PUEB 42) from CIMMYT, the PIC per locus changed significantly between the generations of seed origins. No significant change of PIC was detected in other cases. The increase in PIC value was observed in 8 accessions, whereas for the remaining 12 accessions, it decreased or remained the same in later generations of seed origins. Haplotype allele numbers for the generations ranged from 345 to 478. The earlier generation of the accessions BEN-RB 89–84 and NAYA29 had the minimum and maximum haplotype allele numbers, respectively. The proportion of haplotype alleles lost in later generations ranged from 0 to 22.6%. In addition, 0–19.9% of haplotype alleles was detected in later generations but absent in earlier generations. In 12 cases, the proportion of haplotype allele numbers lost in

**Table 5** Major allelic frequency (MAF) changes between earlier generations and later generations of accessions from five genebanks

Genebank	Accession	Generation	Number of polymorphic loci	Average MAF change <sup>a</sup>	
INIAP	ECUA553	1 vs. 2	415	0.027 ± 0.003	
	ECUA937	1 vs. 2	404	0.028 ± 0.003	
	ECUA497	1 vs. 2	417	0.029 ± 0.003	
	ECUA443	1 vs. 2	494	0.035 ± 0.003	
Average				0.030 ± 0.001	
EMBRAPA	BRAZ3881LC	1 vs. 2	540	0.057 ± 0.004	
	BRAZ3937LC	1 vs. 2	434	0.035 ± 0.003	
	BRAZ3962LC	1 vs. 2	513	0.045 ± 0.004	
	BRAZ4012LC	1 vs. 2	524	0.049 ± 0.004	
Average				0.047 ± 0.002	
INIFAP	COLI21	1 vs. 2	599	0.053 ± 0.003	
	NAYA1	1 vs. 2	621	0.047 ± 0.003	
	NAYA 208	1 vs. 2	634	0.061 ± 0.004	
	NAYA29	1 vs. 2	623	0.087 ± 0.005	
Average				0.062 ± 0.002	
CIMMYT	PUEB 42	1 vs. 2	448	0.043 ± 0.004	
	OAXA265	1 vs. 2	564	0.044 ± 0.003	
	CRICA264	1 vs. 2	528	0.041 ± 0.003	
		1 vs. 3	536	0.050 ± 0.004	
	GUER 5	1 vs. 2	589	0.093 ± 0.006	
Average				0.054 ± 0.002	
IITA	TANZANIA 87–236	1 vs. 2–1	525	0.058 ± 0.004	
		1 vs. 2–2	517	0.046 ± 0.004	
	BEN-RB 89–466	1 vs. 2–1	486	0.050 ± 0.004	
		1 vs. 2–2	468	0.038 ± 0.003	
	BEN-RB 89–84	1 vs. 2–1	413	0.040 ± 0.004	
		1 vs. 2–2	385	0.036 ± 0.004	
	BEN-RB 89–428	1 vs. 2–1	534	0.070 ± 0.005	
		1 vs. 2–2	498	0.041 ± 0.003	
	Average				0.047 ± 0.001

INIAP = the Autonomous National Institute for Agricultural Research; EMBRAPA = the Brazilian Agricultural Research Corporation; INIFAP = the National Institute for Investigation in Forestry, Agriculture and Fishery; CIMMYT = International Maize and Wheat Improvement Center; IITA = the International Institute of Tropical Agriculture

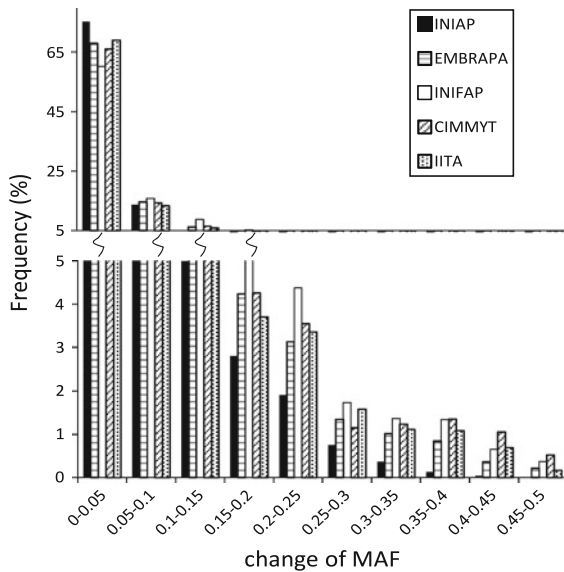
<sup>a</sup> Mean ± Standard Error

later generations was more than those that increased. On the contrary, it was observed that in 11 cases haplotype allele numbers increased in later generations which was more than those that lost haplotype allele numbers. The only accession that did not change its haplotype allele number of each locus was ECUA443 from INIAP (Table 7).

## Discussion

SNP markers can be used to detect the genetic integrity of maize accessions

A large number of SNP markers are available for many plant species for genome-wide fingerprinting.



**Fig. 2** Distribution of major allelic frequency (MAF) changes for single nucleotide polymorphism (SNP) loci detected in accessions from five genebanks. INIAP = the Autonomous National Institute for Agricultural Research; EMBRAPA = the Brazilian Agricultural Research Corporation; INIFAP = the National Institute for Investigation in Forestry, Agriculture and Fishery; CIMMYT = International Maize and Wheat Improvement Center; IITA = the International Institute of Tropical Agriculture

In addition, a wide range of technologies for high-throughput SNP analysis have been developed (as reviewed by Gupta et al. 2008). The GoldenGate assay SNP detection technology of the Illumina Company has been applied widely in genetic analysis, as described in the introduction.

In this study, it has been shown that the 1,536 SNP-OPA was successfully used in genotyping maize landraces. A total of 1,150 polymorphic SNPs of high quality and well-distributed across the genome were observed in 382 samples investigated in the present study. Previous studies had shown SSRs as being highly suitable for studying the genetic diversity of crop germplasm (Liu et al. 2003; Reif et al. 2004), or for verifying identity and genetic integrity of collections in genebanks (Börner et al. 2000; Chebotar et al. 2003; Soengas et al. 2009). Compared with SSRs, SNPs are amenable to very high-throughput genotyping and high-density coverage, and are cost-effective. Furthermore, SNPs are the most abundant type of genetic variation within genomes (Zhu et al. 2003); there is on average one SNP for every 44 bp in maize

(Gore et al. 2009). Hamblin et al. (2007) found that multi-allelic SSR markers are likely to be more informative than bi-allelic SNPs when performing analyses on genetic diversity and relatedness; however, using high-density SNP markers can compensate for this shortcoming (Lu et al. 2009; Yan et al. 2010). Yu et al. (2009) estimated that the power of 1,000 SNPs was similar to that of 100 SSRs for evaluating population structure and relative kinship. On the other hand, haplotypes combining information from several SNPs within the same gene or locus may provide a partial solution to the disadvantage of SNP markers, when used in diversity analyses (Hamblin et al. 2007; Yan et al. 2010). It is suggested that the combined use of both SNP genotypes and haplotypes in genetic diversity analysis can be more powerful than using SNP genotypes alone.

Most recently, Yan et al. (2009) compared the genetic identity of 21 CIMMYT maize lines (CMLs) derived from CIMMYT with the same name but maintained in different labs for more than 20 years using an Illumina GoldenGate assay with 1,536 SNPs. It was found that for some of the “same” lines, the ratio of mismatched SNP markers reached up to 20%. This implies that in maize, an out-crossing crop, it is very difficult to completely maintain the genetic identity and SNPs were a useful tool to detect the identity. In this study, SNP markers and SNP haplotypes were employed successfully to investigate the genetic integrity of maize maintained in different genebanks. The similarities and differences of accessions maintained in different genebanks were estimated and compared, providing some useful information for future maize germplasm conservation and utilization.

Genetic diversity and allelic frequency of individual loci can change significantly after regeneration

In this study, changes in genetic integrity, in terms of average number of alleles per locus, observed heterozygosity, and PIC for almost all accessions, including some significant changes with some accessions in genebanks of INIAP, INIFAP and CIMMYT (Tables 3, 7), were observed during regeneration. In general, regeneration of genebank accessions reduced genetic diversity (Soengas et al. 2009); however, in the present study this did increase in some accessions,

**Table 6** Frequency of major alleles showing significant changes (Fisher's exact test,  $P = 0.05$ ) between generations of each accession

Genebank	Accession	SNP name	Chromosome	Physical position (Mb)	Major allele	Allelic frequency		
						Gen. 1	Gen. 2	Gen. 3
INIAP	ECUA553							
	ECUA937							
	ECUA497							
	ECUA443							
EMBRAPA	BRAZ3881LC	PZA00029.12	2	144.5	A	1	0.5	
		PZA03304.2*	5	4.3	A	0.5	1	
		PZA03711.3	10	121.5	G	0.5	1	
	BRAZ3937LC	PZB00645.1*	6	80.9	G	0.5	1	
	BRAZ3962LC							
	BRAZ4012LC	PZB00228.2*	3	150.8	G	0.5	1	
		PZA03243.3*	1	44.5	A	0.5	1	
INIFAP	COLI21							
	NAYA1							
	NAYA 208							
	NAYA29	PZA03162.1*	7	144.5	A	1	0.5	
		PZA00524.2	5	132.6	A	1	0.5	
		PZB01206.2*	10	99.2	A	1	0.5	
		PZB01389.1	8	134.7	A	0.5	1	
		PZA03710.4	10	121.5	G	0.5	1	
		PZB02227.4	10	123.7	G	0.5	1	
		PZA03255.1*	3	193.6	G	1	0.5	
		PZA03386.1	7	70.4	G	0.5	1	
		PZB01964.5*	3	27.2	A	0.5	1	
		PZA03710.2	10	121.5	A	0.56	1	
	PZA03415.1*	2	21.0	A	1	0.56		
CIMMYT	PUEB 42	PZB01385.3*	8	122.2	C	0.5	1	
		PZA03391.1*	3	219.9	A	0.56	1	
		PZA03612.3*	8	128.6	G	0.93	0.5	
	OAXA265							
	CRICA264	PZA00663.5	4	141.3	A	0.5	0.67	1
	GUER 5	PZA03336.3*	2	11.2	T	0.5	1	
		PZB01083.2	7	4.2	G	0.5	1	
		PZB01358.4	9	106.8	C	0.5	1	
		PZA02869.2*	1	4.4	A	1	0.5	
		PZB00686.2*	5	205.8	C	1	0.5	
PZB00228.6*		3	150.8	G	0.5	1		
PZA03714.2		5	175.5	G	0.5	1		
PZA03625.1*	7	20.2	G	0.5	1			
PZA00933.1	10	71.2	C	0.5	1			
IITA	TANZANIA 87–236	PZB01231.2*	2	39.0	A	1	0.88	
		PZB01569.8*	6	160.7	G	0.5	1	0.83

**Table 6** continued

Genebank	Accession	SNP name	Chromosome	Physical position (Mb)	Major allele	Allelic frequency		
						Gen. 1	Gen. 2	Gen. 3
	BEN-RB 89–466	PZA03503.1	7	0.2	C	0.5	1	
		PZB00093.1	4	122.8	A	0.56	1	0.57
		PHM8549.5	2	NA	T	0.56	1	0.5
	BEN-RB 89–84	PZA02977.4	2	98.7	G	1	0.5	0.88
		PZA00482.10	3	180.3	A	1	0.94	0.94
	BEN-RB 89–428						0.5	

\* Within or close to the QTL regions associated with flowering time; Gen. = generation; SNP = single nucleotide polymorphism; INIAP = the Autonomous National Institute for Agricultural Research; EMBRAPA = the Brazilian Agricultural Research Corporation; INIFAP = the National Institute for Investigation in Forestry, Agriculture and Fishery; CIMMYT = International Maize and Wheat Improvement Center; IITA = the International Institute of Tropical Agriculture

although this increase was not always significant (Tables 3, 7). By using both Nei's genetic distance and pair-wise  $F_{st}$ , a similar pattern of genetic differentiation among generations was detected. Relatively higher genetic distance and  $F_{st}$  values among the different seed generations of NAYA29, GUER 5 and BEN-RB 89–428 may indicate significant loss of genetic integrity.

In the present study, changes detected in genetic integrity, in terms of presence and absence of genes (alleles) by both SNP and SNP haplotype analysis, were dynamic during regeneration. The largest change of average MAF was more than twice that of the smallest change. The majority of loci had MAF changes less than 0.05 between the two generations of all accessions from the five genebanks. Ten of the twenty accessions investigated showed significantly different allelic frequencies between the two generations. Compared to studies in other species, loci with significant changes were very few in the present study. Previous studies found that changes in allelic frequencies after regeneration were substantial in barley (Parzies et al. 2000) and rye (Chebotar et al. 2003). In *B. oleracea*, the proportion of alleles in some regenerated accessions showed significant differences, up to 34%, compared to the original ones (Soengas et al. 2009).

The different heterozygosity for some loci and number of heterozygous loci between the earlier and later generations in this study may be due to inbreeding or genetic drift. The effect of inbreeding can be reduced by maintaining an equal or larger effective population size during regeneration. The initial level of heterozygosity of maize landrace accessions would influence the success in maintaining their genetic integrity during regeneration. Genetically uniform accessions may be impacted less by a small effective population size. On the other hand, highly-diverse germplasm accessions would need a large effective population size: a large number of plants (more than 100) and paired crosses, using one plant as male or female to increase effective population size (Crossa 1989; Crossa et al. 1994).

The observed differences of allelic frequencies in this study may be due to several factors. Firstly, a rare allele can be lost in either the sampling or propagating stages of the regeneration cycle. The small size of the regenerated population can result in bottlenecks, coupled with a loss of alleles. Secondly, allele detection, especially for rare alleles, largely depends on the sample size analyzed. In the present study, we assayed only 8 or 9 bulked samples of 120 or 135 plants. This may mean that rare alleles are possibly not detected. Furthermore, even though the bulking of

**Table 7** Genetic diversity and allele number changes of the single nucleotide polymorphism (SNP) haplotypes between generations of each accession

Genebank	Accession	Gen. no.	Average number of haplotype alleles per locus <sup>a</sup>	Average PIC per locus <sup>a</sup>	Haplotype allele numbers	Number and percentage of haplotype alleles		
						Lost <sup>b</sup>	Increased <sup>b</sup>	
INIAP	ECUA553	1	1.66 ± 0.05	0.17 ± 0.01	389	46	34	
		2	1.60 ± 0.05	0.16 ± 0.01	377	(11.8)	(8.7)	
	ECUA937	1	1.55 ± 0.05	0.15 ± 0.01	365	40	35	
		2	1.53 ± 0.05	0.15 ± 0.01	360	(11.0)	(9.6)	
	ECUA497	1	1.68 ± 0.05**	0.17 ± 0.01	395	62	18	
		2	1.49 ± 0.04**	0.14 ± 0.01	351	(15.7)	(4.6)	
	ECUA443	1	1.71 ± 0.05	0.20 ± 0.01	403			
		2	1.71 ± 0.05	0.20 ± 0.01	403	0	0	
	EMBRAPA	BRAZ3881LC	1	1.81 ± 0.05	0.22 ± 0.01	425	59	70
			2	1.86 ± 0.06	0.22 ± 0.01	436	(13.9)	(16.5)
BRAZ3937LC		1	1.69 ± 0.06	0.17 ± 0.01	396	66	33	
		2	1.54 ± 0.05	0.15 ± 0.01	363	(16.7)	(8.3)	
BRAZ3962LC		1	1.71 ± 0.05	0.19 ± 0.01	401	46	58	
		2	1.76 ± 0.05	0.21 ± 0.01	413	(11.1)	(14.0)	
BRAZ4012LC		1	1.79 ± 0.05	0.22 ± 0.01	421	65	38	
		2	1.68 ± 0.05	0.18 ± 0.01	394	(15.4)	(9.0)	
INIFAP		COLI21	1	1.95 ± 0.06	0.25 ± 0.01	458	56	67
			2	2.00 ± 0.06	0.26 ± 0.01	469	(12.2)	(14.6)
	NAYA1	1	2.01 ± 0.06**	0.26 ± 0.01*	473	86	36	
		2	1.80 ± 0.05**	0.21 ± 0.01*	423	(18.2)	(7.6)	
	NAYA 208	1	1.94 ± 0.05	0.24 ± 0.01	455	57	67	
		2	1.98 ± 0.06	0.26 ± 0.01	465	(12.5)	(14.7)	
	NAYA29	1	2.03 ± 0.06**	0.27 ± 0.02**	478	108	49	
		2	1.78 ± 0.05**	0.21 ± 0.01**	419	(22.6)	(10.3)	
	CIMMYT	PUEB 42	1	1.61 ± 0.05	0.15 ± 0.01*	378	36	64
			2	1.73 ± 0.05	0.19 ± 0.01*	406	(9.5)	(16.9)
OAXA265		1	1.87 ± 0.05	0.22 ± 0.01	440	69	48	
		2	1.78 ± 0.05	0.21 ± 0.01	419	(15.7)	(10.9)	
CRICA264		1	1.79 ± 0.05	0.20 ± 0.01	421			
		2	1.85 ± 0.06	0.21 ± 0.01	435	51(12.1)	65(15.4)	
		3	1.70 ± 0.05	0.19 ± 0.01	400	72(17.1)	51(12.1)	
GUER 5		1	1.83 ± 0.06	0.22 ± 0.01	430	80	74	
		2	1.80 ± 0.05	0.22 ± 0.01	424	(18.6)	(17.2)	
IITA		TANZANIA 87–236	1	1.67 ± 0.05	0.18 ± 0.01	393		
	2–1		1.72 ± 0.05	0.19 ± 0.01	405	46(11.7)	58(14.8)	
	2–2		1.84 ± 0.06	0.22 ± 0.01	432	39(9.9)	78(19.9)	
	BEN-RB 89–466	1	1.79 ± 0.05	0.19 ± 0.01	420			
		2–1	1.68 ± 0.05	0.17 ± 0.01	395	69(16.4)	44(10.5)	
		2–2	1.63 ± 0.05	0.16 ± 0.01	384	70(16.7)	34(8.1)	
	BEN-RB 89–84	1	1.47 ± 0.05	0.12 ± 0.01	345			
		2–1	1.57 ± 0.05	0.15 ± 0.01	370	39(11.3)	64(18.6)	
		2–2	1.55 ± 0.05	0.15 ± 0.01	364	34(9.9)	53(15.4)	

**Table 7** continued

Genebank	Accession	Gen. no.	Average number of haplotype alleles per locus <sup>a</sup>	Average PIC per locus <sup>a</sup>	Haplotype allele numbers	Number and percentage of haplotype alleles	
						Lost <sup>b</sup>	Increased <sup>b</sup>
	BEN-RB 89-428	1	1.80 ± 0.06	0.21 ± 0.01	423		
		2-1	1.86 ± 0.06	0.22 ± 0.01	437	64(15.1)	78(18.4)
		2-2	1.72 ± 0.06	0.20 ± 0.01	405	68(16.1)	50(11.8)

Gen. = generation; PIC = polymorphic information content; INIAP = the Autonomous National Institute for Agricultural Research; EMBRAPA = the Brazilian Agricultural Research Corporation; INIFAP = the National Institute for Investigation in Forestry, Agriculture and Fishery; CIMMYT = International Maize and Wheat Improvement Center; IITA = the International Institute of Tropical Agriculture

<sup>a</sup> Mean ± Standard Error

<sup>b</sup> Numbers in parentheses are the percentage of lost and increased alleles in the original cycles compared to the regenerated cycles

\* Significant difference between original and regenerated cycles ( $P = 0.05$ )

\*\* Significant difference between original and regenerated cycles ( $P = 0.01$ )

DNA from 15 plants to make up one SNP determination is labor- and cost-effective, it requires only 1 of the 15 plants to be heterozygous for the result to indicate that all 15 plants are heterozygous. If all 15 plants have two types of SNP alleles, the result will be the same as one heterozygous plant in the bulked sample, thereby inducing deviations in the detection of allele frequency in the population under study.

In addition, significant changes in allelic frequency of individual loci can be caused by genetic drift, as explained by van Hintum et al. (2007) and Soengas et al. (2009). We found cases where frequencies of major alleles increased to 1 and lost heterozygosity at the same time. This indicated possible genetic drift or directional selection on these loci at pollination that could have been performed among the plants flowering at the same time. To investigate the effect of pollination during regeneration, we checked the loci for significant allelic frequency changes to confirm their relationship with functional genes or QTLs associated with flowering time. Buckler et al. (2009) used joint linkage analysis of a maize NAM (Nested Association Mapping) population, which is based on about 5,000 recombinant inbred lines (RILs) from 25 diverse inbred lines crossed with B73. Thirty-six and 39 QTLs (occupying 47 genome-wide QTL regions using a window of 10 Mb) were identified that explained 89% of the total variation for male flowering and female flowering, respectively. A total of 19 SNP loci (occupying 16 QTL regions) with significant allelic frequency changes in our study were found in or close to (<10 Mb, ~10 cM) the 47

QTL regions associated with flowering time reported in Buckler et al. (2009) (Table 6). More than 57% (16/28) of the SNP loci with significant allelic frequency changes detected in the present study were within the flowering time QTL regions, which is significantly greater than the ratio of random QTL distribution in chromosomes (20%; 47/230). Further studies are needed to identify the potential genes causing the changes; however, present results provide some evidence that the mating system in regeneration may affect the maintenance of genetic integrity. The reduction of MAF from 1 to 0.5 was also observed in our results. This may be due to contamination with foreign pollen and/or seed or handling errors.

#### Strategies for genetic resources conservation and regeneration

The frequency of regeneration should be minimized, as it can cause the loss of genetic integrity of an accession. Several theoretical studies have been carried out to provide simple, efficient sampling schemes and optimal sample size for regeneration and maintenance of crop species based on the statistical model and population genetics theory (Crossa 1989; Crossa et al. 1994; Wang et al. 2004). Regeneration proceeded well at CIMMYT. The accessions which were poorly adapted to the experimental facilities will require appropriate regeneration sites at cooperative genebanks. Artificial pollination is the most common method for accession regeneration and multiplication in maize genebanks.



Chain crosses use a plant as female and male while paired crosses use a plant only once as female or male. When a large number of accessions are regenerated, chain crosses are usually used as they are cost effective. However, it should be noted that the effective population size is near half that of a paired cross (Crossa et al. 1994). To avoid genetic drift and inbreeding, an equal and large effective population size (at least 100 ears at harvest) throughout generations is recommended. In recent years, all landrace regeneration has been conducted at CIMMYT with a goal of 100 ears by chain crosses (half sibs). If a generation of the accession represents fewer than 100 ears at harvest, a repeated regeneration is attempted for the same accession in the following season to reach the required number. IITA accessions were regenerated at CIMMYT repeatedly from the same seed origins (Table 1). A suboptimum number of ears harvested in one generation can be combined with another repeated generation to capture the diversity of the previous generation.

We observed significant losses of genetic diversity in some accessions (i.e. ECUA 497, NAYA 1 and NAYA 29) after regeneration. For these accessions, the number of ears harvested in some generations was less than 100, sometimes even less than 50. This may be one of the reasons for significant changes in genetic diversity. The population genetics theory of sampling a practical number of ears at regeneration (Crossa 1989; Crossa et al. 1994) ensures that the genetic integrity of the accessions in genebanks will be maintained. The SNP loci, with significant changes between generations in relation to flowering time, could indicate that there may have been assortative mating (e.g. early flowering plants mate with early flowering plants; late flowering plants mate with late flowering plants) which can cause a loss of genetic integrity. Further molecular studies are suggested to inspect the on-going practice of regeneration of maize genebank accessions. Regeneration of a landrace accession or other heterogeneous populations in different maize genebanks over many generations can lead to genetically different accessions of the same name due to the change in genetic integrity in time and location. It is recommended to keep a record of duplicate accessions among the genebanks and it is suggested that perhaps the genetic diversity of the original collection cannot be maintained by only one genebank, but rather should be

conserved by collaborative genebanks. During the *ex situ* maintenance of an open pollinating species like maize, the division of the genebank collection into base and active storage can reduce the risk of genetic changes. Core subsets are formed to maximize the efficiency of germplasm evaluation as well as genetic diversity in a collection. A target of good regeneration practice should be to not lose useful alleles during regeneration. When considering the size of the maize genome, the number of loci investigated in the present study is quite limited. With next-generation DNA sequencing technology (Shendure and Ji 2008), it will be feasible to sequence any maize genotype of interest and the sequence of the whole genebank collection may be accessible. Enhanced sequencing throughput available in the near future may uncover the huge diversity present in the genebank accessions. At that time, the real “gene” or allele bank will be constructed. Fingerprinting of genebank accessions can help manage genetic integrity of the germplasm accessions as well as the molecular diversity of maize cultivars.

**Acknowledgments** The authors would like to thank Ramu Punna, contract editor Debra Eaton Mullan, and CIMMYT editor Mike Listman for their help in improving the manuscript. We acknowledge the full financial support received from The International Rice Research Institute, Philippines (IRRI) in conjunction with The Consultative Group on International Agricultural Research (CGIAR) genebank rehabilitation projects of The World Bank, coordinated by System-wide Genetic Resources Programme (SGRP). We also would like to express our sincere thanks to the cooperators of Latin American maize regeneration projects and International Institute of Tropical Agriculture, Nigeria (IITA) for germplasm introductions at CIMMYT maize genebank.

## References

- Börner A, Chebotar S, Korzun V (2000) Molecular characterization of the genetic integrity of wheat (*Triticum aestivum* L.) germplasm after long-term maintenance. *Theor Appl Genet* 100:494–497
- Botstein DR, White RL, Skolnick M, Davis RW (1980) Construction of a genetic linkage map in man using restriction fragment length polymorphisms. *Am J Hum Genet* 32: 314–331
- Buckler ES, Holland JB, Bradbury PJ, Acharya CB, Brown PJ, Browne C, Ersoz E, Flint-Garcia S, Garcia A, Glaubitz JC, Goodman MM, Harjes C, Guill K, Kroon DE, Larsson S, Lepak NK, Li H, Mitchell SE, Pressoir G, Peiffer JA, Rosas MO, Rocheford TR, Cinta Romay M, Romero S, Salvo S, Villeda HS, da Silva HS, Sun Q, Tian F,

- Upadaya N, Ware N, Yates H, Yu J, Zhang Z, Kresovich S, McMullen MD (2009) The genetic architecture of maize flowering time. *Science* 325:714–718
- Chebotar S, Röder MS, Korzun V, Saal B, Weber WE, Börner A (2003) Molecular studies on genetic integrity of open-pollinating species rye (*Secale cereale* L.) after long-term genebank maintenance. *Theor Appl Genet* 107:1469–1476
- CIMMYT Applied Molecular Genetics Laboratory (2003) Laboratory protocols. 3rd edn. Mexico, pp 7–11
- Crossa J (1989) Methodologies for estimating the sample size required for genetic conservation of outbreeding crops. *Theor Appl Genet* 77:153–161
- Crossa J, Taba S, Eberhart SA, Bretting P, Vencovsky R (1994) Practical considerations for maintaining germplasm in maize. *Theor Appl Genet* 89:89–95
- Fan JB, Gunderson KL, Bibikova M, Yeakley JM, Chen J, Wickham Garcia E, Lebruska LL, Laurent M, Shen R, Barker D (2006) Illumina universal bead arrays. *Methods Enzymol* 410:57–73
- Gore MA, Chia JM, Elshire RJ, Sun Q, Ersoz ES, Hurwitz BL, Peiffer JA, McMullen MD, Grills GS, Ross-Ibarra J, Ware DH, Buckler ES (2009) A first-generation haplotype map of maize. *Science* 326:1115–1117
- Gupta PK, Rustgi S, Mir RR (2008) Array-based high-throughput DNA markers for crop improvement. *Heredity* 101:5–18
- Hamblin MT, Warburton ML, Buckler ES (2007) Empirical comparison of simple sequence repeats and single nucleotide polymorphisms in assessment of maize diversity and relatedness. *PLoS ONE* 12:e1367
- Hyten DL, Song Q, Choi IY, Yoon MS, Specht JE, Matukumalli LK, Nelson RL, Shoemaker RC, Young ND, Cregan PB (2008) High-throughput genotyping with the GoldenGate assay in the complex genome of soybean. *Theor Appl Genet* 116:945–952
- Liu KJ, Muse SV (2005) PowerMarker: an integrated analysis environment for genetic marker analysis. *Bioinformatics* 21:2128–2129
- Liu K, Goodman M, Muse S, Smith JS, Buckler E, Doebley J (2003) Genetic structure and diversity among maize inbred lines as inferred from DNA microsatellites. *Genetics* 165:2117–2128
- Lu Y, Yan J, Guimara CT, Taba S, Hao Z, Gao S, Chen S, Li J, Zhang S, Vivek BS, Magorokosho C, Mugo S, Makumbi D, Parentoni SN, Shah T, Rong T, Crouch JH, Xu Y (2009) Molecular characterization of global maize breeding germplasm based on genome-wide single nucleotide polymorphisms. *Theor Appl Genet*. doi: [10.1007/s00122-009-1162-7](https://doi.org/10.1007/s00122-009-1162-7)
- Mehta CR, Patel NR, Tsiatis AA (1984) Exact significance testing to establish treatment equivalence with ordered categorical data. *Biometrics* 40:819–825
- Muchero W, Diop NN, Bhat PR, Fenton RD, Wanamaker S, Pottorff M, Hearne M, Cisse N, Fatokun C, Ehlers JD, Roberts PA, Close TJ (2009) A consensus genetic map of cowpea [*Vigna unguiculata* (L.) Walp.] and synteny based on EST-derived SNPs. *Proc Natl Acad Sci USA*. doi: [10.1073/pnas.0905886106](https://doi.org/10.1073/pnas.0905886106)
- Nei M (1973) Analysis of gene diversity in subdivided populations. *Proc Natl Acad Sci USA* 70:3321–3323
- Parzies HK, Spoor W, Ennos RA (2000) Genetic diversity of barley landrace accessions (*Hordeum vulgare* ssp. *vulgare*) conserved for different lengths of time in *ex situ* gene banks. *Heredity* 84:476–486
- Reif JC, Xia XC, Melchinger AE, Warburton ML, Hoisington DA, Beck D, Bohn M, Frisch M (2004) Genetic diversity determined within and among CIMMYT maize populations of tropical, subtropical, and temperate germplasm by SSR markers. *Crop Sci* 44:326–334
- Rostoks N, Ramsay L, MacKenzie K, Cardle L, Bhat PR, Roose ML, Svensson JT, Stein N, Varshney PK, Marshall DF, Graner A, Close TJ, Waugh R (2006) Recent history of artificial outcrossing facilitates whole-genome association mapping in elite inbred crop varieties. *Proc Natl Acad Sci USA* 103:18656–18661
- Schnable PS, Ware D, Fulton RS, Stein JC, Wei F, Pasternak S, Liang C, Zhang J, Fulton L, Graves TA et al (2009) The B73 maize genome: complexity, diversity, and dynamics. *Science* 326:1112–1115
- Shendure J, Ji H (2008) Next-generation DNA sequencing. *Nat Biotechnol* 26:1135–1145
- Sneath PHA, Sokal RR (1973) Numerical taxonomy. Freeman, San Francisco
- Soengas P, Carrea E, Lema M, Velasco P (2009) Effect of regeneration procedures on the genetic integrity of *Brassica oleracea* accessions. *Mol Breeding* 23:389–395
- Taba S, Twumasi-Afriyie S (2008) Regeneration guidelines: maize. In: Dullo ME, Thormann I, Jorge MA, Hanson J (eds) Crop specific regeneration guidelines [CD-ROM]. CGIAR System-wide Genetic Resource Programme, Rome, p 10
- Taba S, van Ginkel M, Hoisington D, Poland D (2004) Wellhausen-Anderson plant genetic resources center: operations manual, 2004. CIMMYT, Mexico
- Taba S, Shands HL, Eberhart SA (2005) The growth of CIMMYT's maize collection with the introduction of Latin American maize landrace accessions through the cooperative regeneration project, p 1–8. In: S. Taba (ed) Latin American maize germplasm conservation: regeneration, *in situ* conservation, core subsets and prebreeding. Proceedings of a workshop held at CIMMYT 7–10 April 2003. CIMMYT, Mexico
- van Hintum TJJ, van Wiel CCM, Visser DS, van Treuren R, Vosman B (2007) The distribution of genetic diversity in a *Brassica oleracea* gene bank collection related to the effects on diversity of regeneration as measured with AFLPs. *Theor Appl Genet* 114:777–786
- Wang J, Crossa J, Ginkel M, Taba S (2004) Statistical genetics and simulation models in genetic resource conservation and regeneration. *Crop Sci* 44:2246–2253
- Weir BS (1996) Genetic data analysis II. Sinauer Associates, Sunderland
- Wright S (1965) The interpretation of population structure by *F*-statistics with special regard to systems of mating. *Evolution* 19:395–420
- Yan J, Shah T, Warburton M, Buckler ES, McMullen MD, Crouch JH (2009) Genetic characterization and linkage disequilibrium estimation of a global maize collection using SNP markers. *PLoS ONE* 4(12):e8451. doi: [10.1371/journal.pone.0008451](https://doi.org/10.1371/journal.pone.0008451)

- Yan J, Yang X, Shah T, Héctor Sánchez H, Li J, Warburton M, Zhou Y, Crouch JH, Xu Y (2010) High-throughput SNP genotyping with the GoldenGate assay in maize. *Mol Breeding* 25:441–451
- Yu JM, Zhang ZW, Zhu CS, Tabanao DA, Pressoir G, Tuinstra MR, Kresovich S, Todhunter RJ, Buckler ES (2009) Simulation appraisal of the adequacy of number of background markers for relationship estimation in association mapping. *The Plant Genome* 2:63–77
- Zhu YL, Song QJ, Hyten DL, van Tassell CP, Matukumalli LK, Grimm DR, Hyatt SM, Fickus EW, Young ND, Cregan PB (2003) Single-nucleotide polymorphisms in soybean. *Genetics* 163:1123–1134