

## Genetic diversity among tropical maize inbred lines as revealed by SSR markers

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### Abstract

Genetic diversity and phenotypic superiority are important attributes of parental inbred lines for use in hybrid breeding programs. In this study, genetic diversity among 30 maize (*Zea mays* L.) inbred lines comprising of 28 introductions from the International Maize and Wheat Improvement Center (CIMMYT), one from Indonesia and a locally developed, were evaluated using 100 simple sequence repeat (SSR) markers, as early screening for potential parents of hybrid varieties. All markers were polymorphic, with a total of 550 unique alleles detected on the 100 loci from the 30 inbred lines. Allelic richness ranged from 2 to 13 per locus, with an average of 5.50 alleles ( $n_a$ ). Number of effective alleles ( $n_e$ ) was 3.75 per locus, indicating their high effectiveness in revealing diversity among inbred lines. Average polymorphic information content (PIC) was 0.624, with values ranging from 0.178 to 0.874, indicating high informativeness of the markers. High gene diversity was observed on Chromosomes 8 and 4, with high number of effective alleles, indicating their potential usefulness for QTL analysis. The UPGMA dendrogram constructed identified four heterotic groups within a similarity index of 0.350, indicating that these markers were able to group the inbred lines. The three-dimensional PCoA plot also supports the dendrogram grouping, indicating that these two methods complement each other. Inbred lines in different heterotic groups have originated from different backgrounds and population sources. Information on genetic diversity among the maize inbred lines are useful in developing strategies exploiting heterosis in breeding programs.

**Keywords:** genetic distance; heterotic group; molecular estimation; polymorphism; SSR; *Zea mays* L.

**Abbreviations:** CIMMYT\_International Maize and Wheat Improvement Center; CML\_CIMMYT Maize Lines; CTAB\_cetyl trimethylammonium bromide; PCoA\_Principal coordinate analysis; PCR\_Polymerase chain reaction; PIC\_Polymorphic information content; QTL\_Quantitative trait loci; SSR\_Simple sequence repeats; TE\_Tris- ethylenediaminetetraacetic acid.

### Introduction

In maize (*Zea mays* L.) hybrid breeding, success in the exploitation of heterosis depends on genetic diversity of the parental lines (Falconer and Mackay, 1996), as means to select lines that exhibit maximum heterosis without making all possible crosses among them (Makumbi et al., 2011). Lack of genetic diversity may lead to limitation in breeding progress and genetic advance from selection (Cornelius and Sneller, 2002; Onaga et al., 2012).

Various approaches have been employed to examine genetic diversity and inter-relationship among crop genotypes including morpho-agronomic, pedigree, biochemical and DNA-based molecular means (Mohammadi and Prasanna, 2003; Govindaraj et al., 2014). A range of molecular tools has been successfully utilized for assessment of genetic diversity with good reliability. The most common being Restricted Fragment Length Polymorphisms (RFLPs), Amplified Fragment Length Polymorphisms (AFLPs), Random Amplified Polymorphic DNA (RAPD), Inter-Simple Sequence Repeats (ISSR) and Simple Sequence Repeats (SSRs). However, in the last decade, SSRs has been preferred because of the advantages offered.

SSRs also known as microsatellites, are short run of tandemly repeated sequences, which are evenly distributed in the

eukaryotic genome (Taramino and Tingey, 1996). Being co-dominant, SSRs markers are mainly employed in plant analysis due to their hyper variable nature, wide genome coverage, high reproducibility, reliability and discriminating ability, and ability to give high allelic variations (Ko et al., 2016), besides highly specific and polymorphic (Jones et al., 1997).

Numerous studies have utilized SSR markers to assess genetic diversity and relationships among maize populations (Cömertpay et al., 2012; Aci et al., 2013; Nyaligwa et al., 2015; Shayanowako et al., 2018; Adeyemo and Omidiji, 2019). Li et al. (2006) studied 22 CIMMYT subtropical QPM and 24 Chinese inbred lines using 64 SSR markers and obtained two main clusters. Makumbi et al. (2011) studied diversity, heterosis and combining ability among 15 tropical maize lines using 32 SSR markers and found that most lines were grouped together among four clusters according to their pedigrees. Kanagarasu et al. (2013) utilized 10 SSR markers on 27 exotic and indigenous maize inbred lines and obtained five major clusters representing different heterotic groups. Singh et al. (2018) were able to differentiate 18 maize inbred lines into four heterotic groups using SSR markers, while Wende et al. (2013) found that separation of lines into distinct clusters was a result

of prior selection of polymorphic SSR markers, showing high genetic distances.

The objectives of the present study were to determine genetic diversity among 28 introduced tropical maize inbred lines obtained from the International Maize and Wheat Improvement Center (CIMMYT). One inbred line originated from Indonesia and one locally developed. We used SSR markers to assess genetic diversity among them and subsequently assign them into appropriate heterotic groups for potential use as parents in future maize hybrid breeding programs. These CIMMYT inbred lines have previously been identified phenotypically as potential parents for forage use. They have not been subjected to molecular diversity assessment in the past.

## Results

### *Allelic variability and polymorphism*

All the 100 SSR markers utilized were found to be polymorphic, comprising of dinucleotide repeat motifs (63%), trinucleotide repeat motifs (18%), tetranucleotide repeat motifs (7%), and pentanucleotide repeat motifs (7%), while 5% were unidentified. Among the 2866 bands amplified, 550 were unique. The highest percentage of polymorphic bands (PPB) was obtained from SSR marker bnlgl1444 (PPB = 43.33%) (Supplementary Table 1), which also revealed the highest number of alleles (13 alleles), with the dinucleotide type of repeat located in Bin 4.08, followed by SSR marker bnlgl1917 (PPB = 40.00%) located in Bin 4.10, bnlgl1152 (PPB = 40.00%) located in Bin 8.06 and umc1029 (PPB = 36.67%) located in Bin 7.04 (all were dinucleotide type of repeats). In contrast, the lowest PPB was obtained from nine SSR markers, namely umc1467 (a trinucleotide repeat motif in Bin 1.02), umc1604 (a trinucleotide repeat motif in Bin 2.08), umc1639 (a pentanucleotide repeat motif in Bin 3.10), umc1963 (an unspecific number repeat motif in Bin 4.04), umc2406 (a trinucleotide repeat motif in Bin 5.04), umc1348 (a dinucleotide repeat motif in Bin 5.05), umc1143 (a pentanucleotide repeat motif in Bin 6.00), umc1695 (a dinucleotide repeat motif in Bin 7.00) and umc1279 (a trinucleotide repeat motif in Bin 9.00), all having the same PPB value of 6.67. Results of the evaluation revealed high genetic variations, indicating that the inbred lines possessed high genetic variability. Allelic richness ranged from two alleles per locus (for SSR marker umc1467, umc1604, umc1639, umc1693, umc2406, umc1348, umc1143, umc1695 and umc1279) to 13 alleles per locus (for SSR marker bnlgl1444), with a mean of 5.50 alleles per locus, while number of effective alleles per locus ranged from 1.25 to 8.67 alleles, with a mean of 3.75 alleles per locus. The highest number of effective bands per locus was obtained from SSR marker bnlgl1444, with 8.67 alleles, followed by umc1278, with 8.53 alleles per locus and umc1031, with 8.45 effective alleles per locus. Average PIC value among the markers was 0.624, with values ranging from 0.178 (umc1639) to 0.874 (bnlgl1444), where 77% of those values were higher than 0.500. This describes the usefulness of these markers in measuring genetic diversity among the inbred lines.

Average observed homozygosity and heterozygosity among the inbred lines were 0.983 and 0.017, respectively. High homozygosity and low heterozygosity values are indications of the presence of Hardy-Weinberg disequilibrium caused by selfing and selection imposed on the inbred lines over generations, reducing heterozygosity among them. The SSR

marker bnlgl1444 was found to reveal the highest Shannon's information index ( $I = 2.339$ ) and highest Nei's expected heterozygosity (Nei's = 0.885) (Supplementary Table 1). In contrast, SSR marker umc1639 revealed the lowest Shannon's Information Index ( $I = 0.349$ ) and lowest Nei's expected heterozygosity (Nei's = 0.198). The average genetic diversity for the two SSR markers based on Shannon's information index ( $I$ ) and Nei's gene diversity estimated were 1.354 and 0.671, respectively. The information on gene diversity from Shannon's index and Nei's gene diversity indicates high level of within-line genetic diversity. This information could be utilized to investigate diversity revealed by the 30 maize inbred lines studied. The high genetic differentiation ( $F_{ST} = 0.989$ ) and low level of gene flow ( $Nm_f$ ) (0.003) revealed manifestations of high efficiency of the SSR markers.

Nine SSR markers were obtained from Chromosomes 2 and 10, 10 from Chromosomes 1, 4, 5, 6, 7, 8 and 9, and 12 from Chromosome 3. Information from each chromosome was calculated based on the alleles detected by the SSR markers. Results have shown that, on the average, Chromosome 8 had the highest allele number per locus ( $n_a$ ) (7.10) and number of effective alleles ( $n_e$ ) (4.55), indicating that this chromosome had the richest gene diversity (Table 2). This was also reflected by the highest Nei's expected heterozygosity (0.744) revealed by the chromosome, inferring its high genetic diversity. This is an evidence that this chromosome possessed the most diverse genes on the loci involved. Besides Chromosome 8, Chromosome 4 also showed high gene diversity, with observed number of alleles of 6.80 per locus and number of effective alleles of 4.38. In addition, Chromosomes 8 and 4 also had high Shannon's information indices (1.601 and 1.460, respectively), and high Nei's heterozygosity coefficients (0.744 and 0.673, respectively), further emphasizing their gene richness and informativeness. Genetic information of each of the inbred lines is presented in Table 3, which was subsequently used in the analysis on genetic similarities.

Based on results of genetic identities of the 30 inbred lines studied, inbred line SM7-11, IPB 8 and CML498 were found to be the most heterogeneous, revealing high Nei's heterozygosity estimates of 0.0803, 0.0306 and 0.0275, respectively. They also portrayed the highest number of alleles, number of effective alleles and Shannon's information index (Table 3). In contrast, the least heterogeneous were inbred lines CML147, CML311, CML312, CML322, CML331, CML478 and CML495, revealing very low values for these genetic parameters.

### *Genetic similarities among inbred lines for SSR loci*

A dendrogram established from similarity coefficients among the inbred lines using the UPGMA method is presented in Fig 1. Result of Mantel test giving cophenetic value of  $r = 0.818$  is the evidence for the good agreement between result of the genetic similarity matrix and what is portrayed by the dendrogram. The cluster analysis has clearly separated the lines into different groups. Genetic similarities ranged from 0.114 (between CML 383 and CML 479) to 0.813 (between CML 383 and CML 384). At a vertical cut of 0.352 of Dice's coefficient on the UPGMA dendrogram (Fig 1), four diverse groups were identified as follows: Group I represented by nine CIMMYT inbred lines, CML269, CML311, CML312, CML322, CML331, CML376, CML373, CML383 and CML384, with an average similarity coefficient of 0.369; Group II represented by six other CIMMYT inbred lines CML71, CML78, CML146, CML147, CML152 and CML160, with coefficient of 0.404;

Group III consisted of five CIMMYT inbred lines, CML491, CML494, CML495, CML498 and CML503, the inbred line from Indonesia, IPB8, and the line from Malaysia, SM7-11, with coefficient of 0.386. Group IV comprised of eight CIMMYT inbred lines, CML428, CML448, CML447, CML449, CML451, CML476, CML478 and CML479), with an average similarity coefficient of 0.389. The most genetically similar inbred lines were CML383 (standard pedigree: P502-C1-771-2-2-1-1-B) and CML 384 (standard pedigree: P502-C1-771-2-2-1-3-B) placed in Group I, which were both originated from the same source population (P502), and with a related pedigree (CIMMYT Global Maize Program, 2015). The second highest similarity was shown between CML311 (standard pedigree: S89500-F2-2-2-2-B\*5) and CML312 (standard pedigree: S89500-F2-2-2-1-1-B\*5), which were also placed in Group I. As seen from the pedigrees, they were both originated from the same source population and shared 56.5% similarity based on the SSR markers. Although some inbred lines showed different pedigrees but were genetically related and grouped into the same cluster. Inbred line CML71 and CML78 were both developed from two different source populations (Antigua and G32, respectively) but were genetically related (0.455 of similarity coefficient), as they are placed in the same cluster. Based on the dendrogram established, Group IV consists of several lowland inbred lines developed from seven different population sources (P145, P43, P21, P32, REC, G24 and P24).

#### **Genetic distance by PCoA**

A three-dimensional principal coordinate analysis (PCoA) plot was constructed (Fig 2) to depict the distance among the maize inbred lines. Their distribution based on information from the PCoA were derived from the genetic distances from the three main component factors of the eigenvalues (data are not shown). Interestingly, it is clearly seen that the three-dimensional projection supports the dendrogram constructed by UPGMA grouping method (Fig 1), indicating that these two methods complement each other towards the group separation. The only obvious exception is that inbred line SM7-11 seems to fit in the cluster corresponding to Group I of the plot (Fig 2), not Group III as previously depicted in the dendrogram (Fig 1).

#### **Discussion**

A total of 2866 bands amplified by the 100 SSR markers distributed on 10 chromosomes. They were used as the basis of the analysis of genetic diversity among the 30 maize inbred lines. Among the polymorphic bands amplified, 550 alleles were considered adequate for the assessment.

The extent of diversity was assessed using PIC, gene diversity and genetic distances. The presence of high polymorphism and high number of rare alleles were observed among these inbred lines, which is an explanation to the presence of high intra-group diversity, as described by Glaszmann et al. (2003). The magnitude of mean PIC value (0.624) was higher than that previously reported by Smith et al. (1997), which involved 58 maize inbred lines and four maize hybrids (PIC = 0.62). Sserumaga et al. (2014) reported (PIC = 0.61) on 42 tropical maize inbred lines; Senior et al. (1998) on 94 elite maize inbred lines (PIC = 0.59), Belalia et al. (2018) on 56 maize landraces (PIC = 0.57) and Nyaligwa et al. (2015) on 79 elite maize inbred lines (PIC = 0.51). However, lower values were reported by Li et al. (2006) from a study on 22 CIMMYT QPM maize inbred lines and 24 Chinese maize inbred lines (PIC =

0.66), Shayanowako et al. (2018) reported (PIC = 0.58) on 37 diverse maize genotypes and Adeyemo and Omidiji (2019) on 19 diverse maize varieties (PIC = 0.75). The high PIC values obtained is an indication of the high informativeness of the markers, as reflected by the scale proposed by Botstein et al. (1980), where PIC value of  $> 0.5$  accounts for a highly informative marker,  $0.5 > \text{PIC} > 0.25$  for informative marker, and  $\text{PIC} \leq 0.25$  for a slightly informative marker. High estimates of PIC obtained were probably because primers with low discriminatory ability have been eliminated during the initial screening. The distribution of loci involved all the 10 chromosomes, representing the actual diversity of the inbred lines. The high level of polymorphism indicates that variations among the inbreds were considerably high, where exploitation of heterosis in hybrid crosses could be effective. Shiri et al. (2014) observed a high Nei's expected heterozygosity (0.52) as well as a high Shannon's information index (0.94) in their study on maize, supporting our findings. However, these values were lower than those reported by Xiang et al. (2010) and Belalia et al. (2018).

Average number of alleles detected was 5.5 per locus, slightly lower than that reported from a previous study involving 86 white and 69 yellow lowland tropical CIMMYT maize inbred lines (7.4 alleles per SSR locus) (Xia et al., 2004), and that reported on 73 subtropical, 11 tropical highland and 10 tropical lowland CIMMYT maize inbred lines (7.2 alleles per SSR locus) (Xia et al., 2005). A lower level of allelic richness was found in the studied maize inbred lines, indicating that the use of a small number of inbred lines has contributed to the narrow gene diversity within the population. Average observed homozygosity and observed heterozygosity found on the inbred lines were slightly higher than that reported by Kashiani et al. (2012b) ( $\text{Hom}_o = 0.9396$  and  $\text{H}_o = 0.0604$ ). High gene variation in the chromosomes of the inbred lines studied showed that they possess the information required in genetic diversity. Similarly, result on genetic differentiation (F) indicates that there was a high genetic variability among the studied inbred lines. High F values revealed that a large portion of total genetic diversity could be attributed to the high genetic variation among the inbred lines. Previous workers have reported different values of observed F (Aci et al., 2013; Ferreira et al., 2018), depending on genetic materials assessed.

Based on results of the individual genetic identities of the inbred lines, the high variation observed among the inbred lines was due to different pedigrees of these lines that originated from different breeding programs utilizing different source populations, giving a reflection that these inbred lines belong to different genetic groups. This diversity could be utilized to develop superior hybrid varieties with high level of heterosis by involving parental lines from different groups. It is interesting to note that the local inbred line SM7-11, and the one introduced from Indonesia IPB8, revealed high diversity among loci, probably due to the diverse sources involved in their development, making them more heterogeneous.

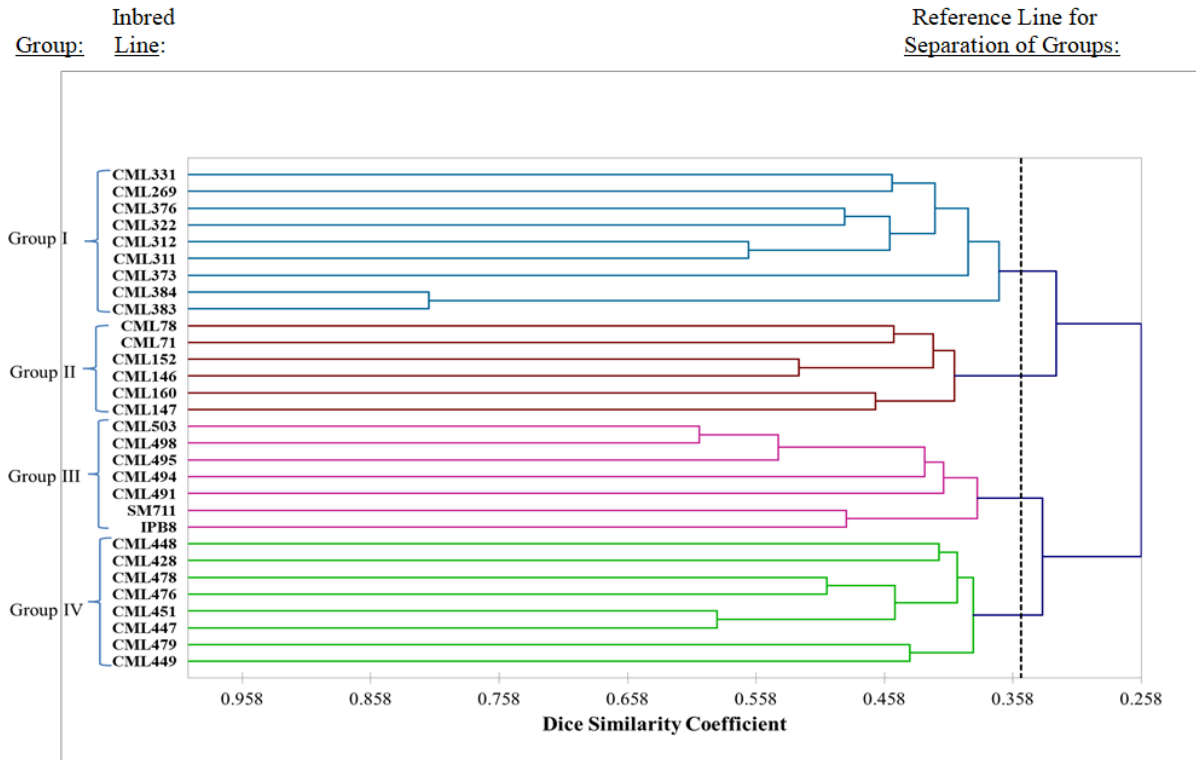
From the dendrogram constructed based on SAHN clustering and from the three-dimensional PCoA projection, four heterotic groups were formed separating the inbred lines.

Eight subtropical lines were clustered together in one group. Two of them (CML383 and CML384) were also found to be clustered in the same group in another study reported by Xia et al. (2005). These two highly related inbred lines had highest genetic similarity coefficient between them, because they have similar pedigrees, since they originated from the

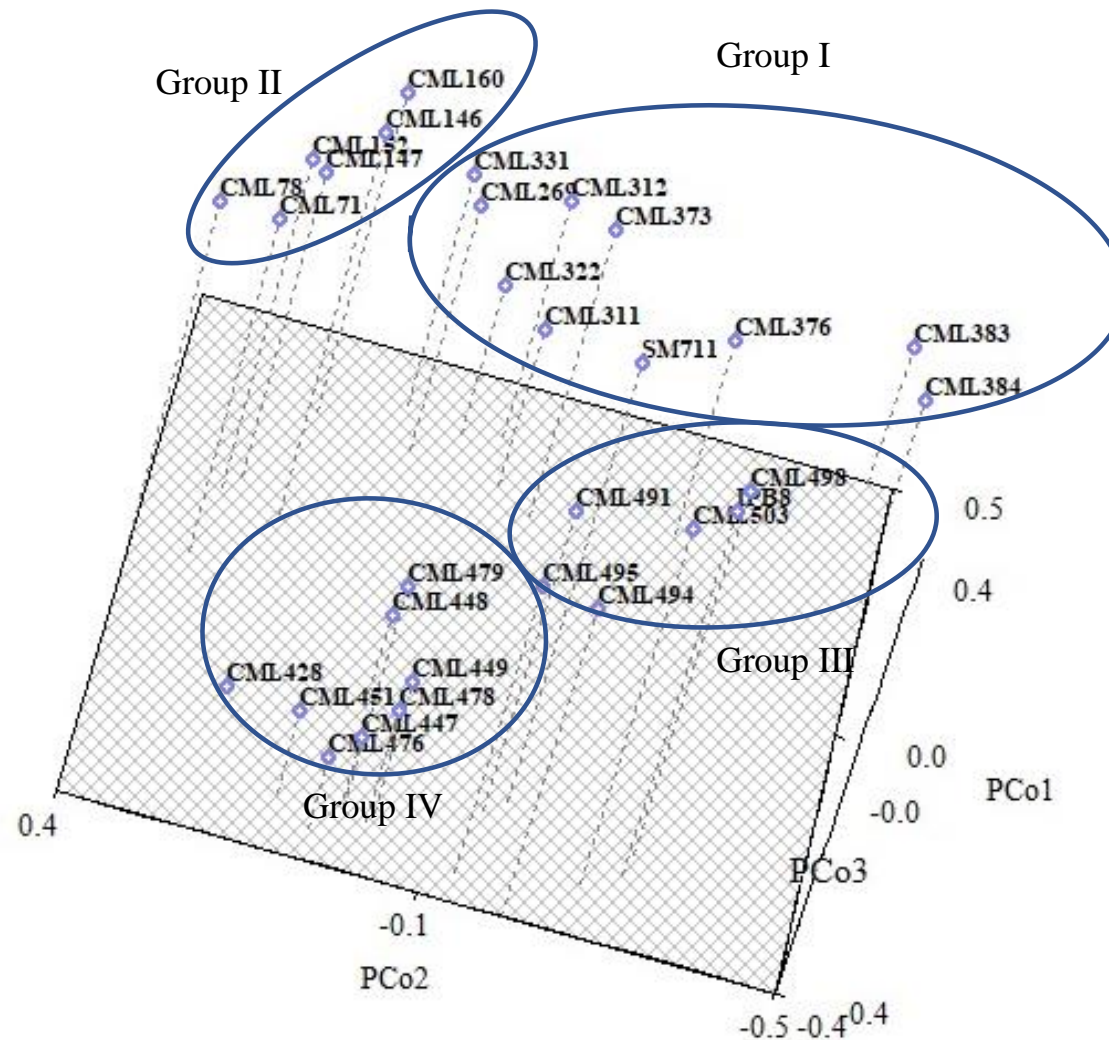
**Table 1.** List of maize inbred lines evaluated, and their standard pedigrees, origins, adaptations and source populations

Inbred line	Standard pedigree	Origin	Adaptation	Source population
CML 71	ANTGP2-5-#-1-2-1-1-5-5-7-B	CIMMYT	Lowland	Antigua
CML 78	G32-C19-HS32-1-#-2-B-#*3-3-B	CIMMYT	Subtropical	G32
CML 146	AC8563-HS35-3-1-B-2-1-B-B-1-B	CIMMYT	Lowland	P63
CML 147	P63-C2-FS53-1-1-B*3-9-B	CIMMYT	Lowland	P63
CML 152	S8662Q-1-4-4-5-B	CIMMYT	Lowland	P62
CML 160	P63-C2-FS6-2-1-1-B-2-1-B	CIMMYT	Lowland	P63
CML 269	P25STE-C1-FS13-6-1-1-B	CIMMYT	Lowland	P25
CML 311	S89500-F2-2-2-2-B	CIMMYT	Subtropical	P500
CML 312	S89500-F2-2-2-1-1-B	CIMMYT	Subtropical	P500
CML 322	LLMBR-17-B-5-3-1-4-B	CIMMYT	Subtropical	REC
CML 331	((SUWAN8422)/(P47/MP78-518)-#-183-1-2-1-2-2-B	CIMMYT	Subtropical	REC
CML 373	P43SR-4-1-1-2-1-B-8-1-B	CIMMYT	Subtropical	P43SR
CML 376	SLWHGA-F118-2-1-1-B-1-B-1-B	CIMMYT	Subtropical	SLWHG
CML 383	P502-C1-771-2-2-1-1-B	CIMMYT	Subtropical	P502
CML 384	P502-C1-771-2-2-1-3-B	CIMMYT	Subtropical	P502
CML 428	SW91145-2-3-B-B-#-#-3-B	CIMMYT	Asia Lowland	P145
CML 447	(P43-F95/P21-F219)-1-B*3-1-B	CIMMYT	Lowland	P43
CML 448	P21MRRS-C1-430-1-B	CIMMYT	Lowland	P21
CML 449	P32MRRS-F2-C2-23-2-B	CIMMYT	Lowland	P32
CML 451	((NPH-28-1/G25)/NPH-28-1)-1-2-1-1-3-1-B	CIMMYT	Lowland	REC
CML 476	P21MRRS-C1-525-1-B	CIMMYT	Lowland	P21
CML 478	(G24-F119/G24-F54)-6-3-2-1-B*3-1-B	CIMMYT	Lowland	G24
CML 479	(P24STE-5/P24STE-17)-B*4-#*3-B-3-B-1-B	CIMMYT	Lowland	P24
CML 491	(6207QB/6207QA)-1-4-#-2-2-B	CIMMYT	Lowland	REC
CML 494	LAPOSTASEQ-C4-F7-1-2-2-2-2-B	CIMMYT	Lowland	P43
CML 495	(PNVABCOSD/NPH-28-1)-F32-B-1-B-1-2-B	CIMMYT	Lowland	RCW
CML 498	((CML265/CL00303)/CML264)-B-8-1-3-B	CIMMYT	Lowland	RCW
CML 503	((((CML176/CML264)-B)/CML264)-7-1/CML264)/CML264)-F2-65-B	CIMMYT	Lowland	POP21
IPB8	N.A	Indonesi	Tropical	N.A.
SM7-11	UPM Selection	Malaysi	Tropical	SMC317

Abbreviations used in pedigree: # = sibbing; B = Sefled and Bulked; -1, -2, -3 = ear-to-row; HS = Half sibs; FS = Full sibs; \* = backcross of a specific parent; C = Cycle; F = filial; ( ) = denotes pedigrees of one parent in a cross; / = cross between parents; P = Pool. N.A. = information not available, introduced from Institut Pertanian Bogor (IPB), Indonesia.



**Fig 1.** Dendrogram constructed using Unweighted Pair Group Method using Arithmetic Averages (UPGMA), separating 30 tropical maize inbred lines into four diverse heterotic groups. The four groups are separated at the Dice's similarity coefficient of 0.352 (Dice, 1945). The CML lines were introduced from CIMMYT, while SM7-11 and IPB 8 are from Indonesia and Malaysia, respectively.



**Fig 2.** Plot from three-dimensional principal coordinate analysis (PCoA) projected using the first three principal coordinates obtained from amplification of 100 SSR markers, measured using Dice similarity indices (Dice, 1945), on 30 maize inbred lines. PCo1, PCo2 and PCo3 are projected based on the first three principal coordinates obtained from molecular data of the 30 maize inbred lines. CML lines were introduced from CIMMYT, while SM7-11 and IPB 8 are from Indonesia and Malaysia, respectively. Four heterotic groups are indicated by the circles containing the inbred lines within the group.

**Table 2.** Chromosomal genetic variations revealed by 100 SSR markers amplified on 30 maize inbred lines.

Chromosome	$n_a$	$n_e$	$Hom_o$	$H_o$	$I$	Nei's	$\hat{H}$
1	5.10	3.52	0.997	0.340	1.268	0.647	0.0017
2	4.56	3.41	0.993	0.326	1.251	0.663	0.0037
3	5.50	3.42	0.940	0.364	1.289	0.625	0.0264
4	6.80	4.38	0.976	0.315	1.460	0.673	0.0117
5	3.90	2.78	1.000	0.372	1.104	0.617	0.0000
6	5.50	3.75	1.000	0.303	1.380	0.683	0.0000
7	5.30	3.64	0.993	0.308	1.343	0.680	0.0033
8	7.10	4.55	0.958	0.242	1.601	0.744	0.0200
9	5.70	4.00	0.990	0.306	1.418	0.681	0.0050
10	5.44	4.09	0.992	0.275	1.438	0.712	0.0037
Mean	5.50	3.75	0.983	0.316	1.354	0.671	0.0333
SD	2.60	1.75	0.055	0.156	0.472	0.152	0.0080

$n_a$  = observed number of alleles,  $n_e$  = observed number of effective alleles,  $Hom_o$  = observed homozygosity,  $H_o$  = observed heterozygosity,  $I$  = Shannon's information index, Nei's = Nei's expected heterozygosity  $\hat{H}$  = average heterozygosity, SD = standard deviation.

**Table 3.** Molecular genetic identities of 30 maize inbred lines amplified using 100 microsatellite DNA markers.

Inbred line	$n_a$	$n_e$	$Hom_o$	$H_o$	$I$	Nei's	$\hat{H}$
CML 71	1.0112	1.0112	0.9888	0.0112	0.0078	0.0056	0.0090
CML 78	1.0220	1.0220	0.9780	0.0220	0.0152	0.0110	0.0075
CML 146	1.0213	1.0213	0.9787	0.0213	0.0147	0.0106	0.0085
CML 147	1.0000	1.0000	1.0000	0.0000	0.0000	0.0000	0.0063
CML 152	1.0104	1.0104	0.9896	0.0104	0.0072	0.0052	0.0083
CML 160	1.0104	1.0104	0.9896	0.0104	0.0072	0.0052	0.0083
CML 269	1.0110	1.0110	0.9890	0.0110	0.0076	0.0055	0.0071
CML 311	1.0000	1.0000	1.0000	0.0000	0.0000	0.0000	0.0081
CML 312	1.0000	1.0000	1.0000	0.0000	0.0000	0.0000	0.0082
CML 322	1.0000	1.0000	1.0000	0.0000	0.0000	0.0000	0.0080
CML 331	1.0000	1.0000	1.0000	0.0000	0.0000	0.0000	0.0084
CML 373	1.0106	1.0106	0.9894	0.0106	0.0074	0.0053	0.0082
CML 376	1.0106	1.0106	0.9894	0.0106	0.0074	0.0053	0.0085
CML 383	1.0102	1.0102	0.9898	0.0102	0.0071	0.0051	0.0082
CML 384	1.0104	1.0104	0.9896	0.0104	0.0072	0.0052	0.0083
CML 428	1.0215	1.0215	0.9785	0.0215	0.0149	0.0108	0.0070
CML 447	1.0211	1.0211	0.9789	0.0211	0.0146	0.0105	0.0082
CML 448	1.0111	1.0111	0.9889	0.0111	0.0077	0.0056	0.0070
CML 449	1.0104	1.0104	0.9896	0.0104	0.0072	0.0052	0.0083
CML 451	1.0105	1.0105	0.9895	0.0105	0.0073	0.0053	0.0075
CML 476	1.0109	1.0109	0.9891	0.0109	0.0075	0.0054	0.0074
CML 478	1.0000	1.0000	1.0000	0.0000	0.0000	0.0000	0.0079
CML 479	1.0222	1.0222	0.9778	0.0222	0.0154	0.0111	0.0089
CML 491	1.0104	1.0104	0.9896	0.0104	0.0072	0.0052	0.0068
CML 494	1.0105	1.0105	0.9895	0.0105	0.0073	0.0053	0.0084
CML 495	1.0000	1.0000	1.0000	0.0000	0.0000	0.0000	0.0065
CML 498	1.0549	1.0549	0.9451	0.0549	0.0381	0.0275	0.0088
CML 503	1.0206	1.0206	0.9794	0.0206	0.0143	0.0103	0.0082
IPB 8	1.0612	1.0612	0.9388	0.0612	0.0424	0.0306	0.0082
SM7-11	1.1158	1.1158	0.8842	0.1158	0.0579	0.0803	0.0084
Mean	1.0170	1.0170	0.9830	0.0170	0.0110	0.0092	0.0079
SD	0.0234	0.0234	0.0234	0.0234	0.0132	0.0152	0.0007

$n_a$  = observed number of alleles,  $n_e$  = effective number of alleles,  $Hom_o$  = observed homozygosity,  $H_o$  = observed heterozygosity,  $I$  = Shannon's information index, Nei's = expected heterozygosity,  $\hat{H}$  = heterozygosity, SD = standard deviation.

same germplasm source (Table 1). We also found that, although CML71 and CML78 were genetically related (with 45.5% genetic similarity), they were derived from two different source populations (Table 1). As reported by Xia et al. (2005), small divergence among the inbred lines was probably due to previous intermingling among the source populations in

CIMMYT. Reid et al. (2011) emphasised that inbred lines should not be grouped together based on similarity of germplasm sources, but rather on similarity of their genetic background. Past workers have also verified that separation by heterotic groups represents the pedigrees of the lines (Barata and Carena, 2006; Reid et al., 2011).

Assessment of genetic relationship among genetic materials is an important step in a crop breeding program as it would determine genetic diversity of the breeding populations or genotypes. Accurate assessment of genetic diversity in breeding programs would genetically identify the most diverse parental combinations in crosses to form hybrids or among segregating progenies for further utilization in breeding programs. Molecular markers are important and complementary to evaluation of phenotypic traits in assessing genetic diversity and relationships among corn genotypes for efficient selection of parents in breeding programs (Nyaligwa et al., 2015). From the cluster analysis on the inbred lines, the subtropical and lowland lines were clearly grouped together according to their related pedigrees. With a very slight exception of line SM7-11, results of the PCoA and the cluster analysis were in good agreement with each other, and both were used in complement for interpreting the relationships among the inbred lines. The distribution of the inbred lines based on three-dimensional PCoA was derived from genetic distances among the three main component factors of the eigenvalues (26.56% of total genetic variance). As described by Messmer et al. (1992), the PCoA graph constructed from the first three eigenvectors, which accounted for more than 25% of the original variation in the populations. A high level of heterogeneity was reported among the studied inbred lines. Formation of four heterotic groups among the inbred lines explains their diverse genetic backgrounds at the DNA level, indicating their high potential for use in exploitation of heterosis. Cross combinations among inbred lines with diverse genetic backgrounds could produce high heterosis, resulting from favorable dominant gene effects and interactions in the progenies. Srdic et al. (2008) found that the magnitude of genetic distances revealed by SSR markers was in good agreement with the extent of specific combining ability in a series of sweet corn inbred lines.

## Materials and Methods

### Plant materials

A total of 30 maize inbred lines adapted to the lowland subtropical and tropical environments were utilized in this study. Twenty-eight inbred lines were introduced from the International Maize and Wheat Improvement Center (CIMMYT), Mexico, one was introduced from Indonesia and the other was locally developed at Universiti Putra Malaysia (UPM) (Table 1).

### DNA extraction and quantification

Young leaves from seedlings of one to two weeks old of each inbred line were collected separately for DNA extraction following the standard modified CTAB method described by Hoisington et al. (2005). DNA concentration was then measured using Thermo scientific NanoDrop 2000c spectrophotometer (Thermo scientific™, USA), following the standard protocol as outlined by Joseph (2010). The DNA extracted from each inbred line was diluted and suspended in TE (Tris-EDTA) buffer at the concentration of 40-60 ng/μl. The quality of DNA extracted was determined by running it on 0.5% agarose gel.

### PCR and gel electrophoresis

Amplification of DNA was carried out in a 15 μL reaction mix using T100 Thermal Cycler (Bio-Rad Laboratories, Inc., CA, USA), with reaction mixture containing 1X Biorad MyTaq™ Red Mix PCR buffer (KCl, (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>) (contains dNTPs, MgCl<sub>2</sub> and Taq polymerase), 0.5 μM of primers (for each forward and reverse), and 30 ng of genomic DNA. The touchdown PCR program steps included an initial denaturing step of 3 min at 95°C, and 20 cycles, each with 1 min DNA denaturation at 94°C, 1 min annealing at 68°C~58°C (the annealing temperature was decreased by 0.5°C in each cycle to 58°C), 1 min extension at 72°C, and 10 cycles of 1 min at 94°C, 1 min at 58°C, 1 min at 72°C, and a final extension of 10 min at 72°C. Products of PCR reaction were separated using 4% Lonza MetaPhor® agarose gel that was stained using GelRed DNA stain (Biotium, Inc., CA, USA), and run using horizontal gel electrophoresis (Bio-Rad Laboratories, Inc., CA, USA) at 90 volts for 90 min.

### Data analyses

One hundred microsatellite regions evenly distributed throughout the maize genome were selected from the database MaizeGDB (Andorf et al., 2010), based on their PIC, as well as from information on QTLs reported in previous investigations (Xie et al., 2009; Li et al., 2010; Kashiani et al., 2012a). Gel images were captured using the Molecular Imager Gel DocTM XR+ (Bio-Rad Laboratories, Inc., CA, USA) and processed with Image LabTM Software, and the amplified bands were scored manually using molecular ladders (Qiagen, Netherlands) of 25 bp and 50 bp as reference for data scoring. A binary data was created by denoting presence as "1" and absence as "0" for all fragment band sizes. Bands that were too faint and thus unable to be scored were considered as missing value by denoting as "9". The PIC was calculated for each SSR locus and each inbred line using the genetic analysis software PowerMarker version 3.25 (Liu and Muse, 2005) following the formula by Anderson et al. (1993).  $PIC = 1 - (\sum P_{ij}^2)$ , where  $P_i$  is the frequency of  $j^{th}$  alleles for the  $i^{th}$  locus. Total number of alleles ( $n_a$ ), number of effective alleles ( $n_e$ ) (Anderson et al., 1993), observed homozygosity ( $Hom_o$ ), observed heterozygosity ( $H_o$ ) (Levene, 1949), Shannon's Information index (I), average heterozygosity ( $\bar{H}$ ) (Lewontin, 1972), Nei's expected heterozygosity (Nei's) (Nei, 1973) and coefficient of inbreeding (F) (Lukas and Donald, 2002), were calculated using the software POPGENE version 1.31 (Yeh et al., 1999). Similarly, in the same way the above molecular genetic differentiations were also determined for each inbred line.

Genetic similarities (GS) among the inbred lines were subsequently estimated based on Dice coefficients (Dice, 1945), with the formula  $Dc = 2n_{xy} / (n_x + n_y)$ , where  $n_x$  and  $n_y$  represent the number of putative SSR alleles for X and Y, respectively, and  $n_{xy}$  represents the number of putative SSR alleles shared between X and Y. A dendrogram tree was constructed by applying the Unweighted Pair Group Method using Arithmetic Averages (UPGMA) following the Sequential Agglomerative Hierarchical and Nested (SAHN) method, utilizing the software NTSYS-pc version 2.1 (Rohlf, 2000). The same data matrix of genetic distance was utilized to construct the Principal Coordinate Analysis (PCoA) plots using the same software. Two-way Mantel test (Mantel, 1967) was performed

to construct the matrix of correlation for comparison between genetic distance and genetic similarity among the genotypes for 1000 permutations.

## Conclusion

The SSR markers utilized in this study were able to reveal high variations among the maize inbred lines. Assignment of inbred lines into four heterotic groups was in close agreement with the genetic background of their source populations. Most SSR markers that showed high polymorphism were dinucleotide SSR markers. They revealed high percentage of polymorphic bands. The markers used were also identified as having high polymorphic information content and high gene diversity. Chromosome 8 revealed high gene diversity with a high number of effective alleles, indicating the presence of diverse genes on its loci in the inbred lines. Therefore, they can be used to identify QTLs for useful traits including yield. Since the CIMMYT inbred lines had previously been identified phenotypically as good potential parents for forage utilization, the high genetic diversity revealed by the SSR markers could be utilized for development of superior forage hybrid varieties in the future work of exploiting heterosis.

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