



Full Length Article

Population Structure, Linkage Disequilibrium and Association Mapping Study through SSR Markers in Sunflower (*Helianthus annuus*)

Danish Ibrar^{1*}, Rafiq Ahmad², Muhammad Ayub Khan¹, Asif Javaid¹, Zubair Ahmed¹, Munir Ahmad² and Talat Mahmood²

¹Pakistan Agricultural Research Council, Islamabad, Pakistan

²Pir Mehr Ali Shah Arid Agriculture University, Rawalpindi, Pakistan

*For correspondence: danish.uar@gmail.com

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Abstract

Sunflower is one of the most important oilseed crops of world. Exploitation of heterosis has been very successful in cross pollinated crops and sunflower being cross pollinated is ideal for hybrid breeding. Exploitation of hybrid vigor with precision saves time and resources, and this can be achieved by using new genetic based crop improvement techniques like association mapping. Association mapping has proved very successful in designing molecular marker based speedy crop improvement. Agro-morphological traits are important for sunflower breeders in selection of genotypes with high performance and other yield contributing traits. Ten agro-morphological characters *i.e.*, days to flower initiation and completion, plant height, stem curvature, head diameter, number of leaves per plant, leaf area, hundred seed weight, seed yield per plant and oil contents were used to study the genetic diversity in 109 sunflower lines at morphological level. A high genetic variability was observed among sunflower lines in field conditions. Population structure and linkage disequilibrium was estimated after genotyping through 40 SSR (simple sequence repeats) markers, that amplified a total of 65 DNA bands to locate the QTLs (quantitative trait loci) controlling the important yield contributing agro-morphological traits. Based on population structure analysis, sunflower genotypes showed two sub-populations. For identification of important QTL's controlling the morphological traits, genotypic and phenotypic data combination showed eleven SSR bands had a significant association ($P \leq 0.05$) with 6 out of 10 tested agro-morphological characters. These identified marker trait associations are expected to be helpful for sunflower breeders in designing a successful breeding strategy. © 2020 Friends Science Publishers

Keywords: Sunflower breeding; Microsatellites; Mixed linear model; Genetic diversity; Association mapping; Linkage disequilibrium

Introduction

Marker assisted selection in plant breeding has been found to be very effective in devising more focused and robust crop improvement programs. As this technique provided more precision and accuracy in selection thus saves time, resources and efforts required for the development of new variety/hybrid (Filippi *et al.* 2015). An important part of this methodology is to locate the markers in the genome that are tightly linked to the quantitative trait loci (QTLs) controlling phenotype of the plant (Darvishzadeh 2016). QTL mapping in crop plants is normally achieved by two frequently utilized techniques *i.e.*, QTL mapping and association mapping (Ilyas *et al.* 2018). In conventional QTL mapping, two parents are crossed in an organized way and the association between phenotypic traits and mapped marker loci allows the identification of QTLs. Because of few recombination events, the QTL of interests may not be tightly associated with the marker identified

(Myles *et al.* 2009). Whereas, association mapping which is a relatively new approach of mapping QTLs, identifies relationship between gene polymorphism and phenotypic variation in existing germplasm collections without the development of mapping populations (Fusari *et al.* 2012). It proves to be efficient in detecting the markers that are tightly linked to a specific QTL (Abdurakhmonov and Abdurakarimov 2008). Association mapping utilizes the population structure and linkage disequilibrium (LD) information, hence, also known as LD mapping (Thornsberry *et al.* 2001). Association mapping provides high resolution mapping by arresting all the meiosis occurring in the breeding history of the crop. Moreover, it is cost effective and time saving technique as compared to QTL mapping or linkage analysis (Ambreen *et al.* 2018).

Microsatellites/SSR markers are frequently being employed for population structure analysis in association mapping studies because of their proven ability for generation of more information content as compared to

biallelic markers (Fusari et al. 2012; Filippi et al. 2015). In Pakistan, sunflower (*Helianthus annuus* L.) was introduced during 1960's along with safflower (*Carthamus tinctorius*) and soybean (*Glycine max*) to increase the local edible oil production. During 2015–16 sunflower was cultivated on an area of 866003 hectares in Pakistan and 35000 tons of vegetable oil was extracted (Ibrar et al. 2018). Worldwide, sunflower is the fourth biggest vegetable oil producing crop after palm oil (*Elaeis guineensis*), soybean and canola (*Brassica napus*) (Rauf et al. 2017). From hybrid breeding perspective, it is considered as second most important crop after maize (*Zea mays*) (Seiler et al. 2017). Domestication of sunflower was started in Pre-Colombian times, but the breeding efforts for oil types were started in 18th century (Mandel et al. 2011). Heterosis on commercial scale was exploited in sunflower after the incorporation of CMS (cytoplasmic male sterility) genes by Leclercq (1969) and discovery of male fertility restoration genes by Kinman (1970) and Lochner (2011). Assessment of genetic diversity for various agro-morphological traits is a pre-requisite for manipulating and introgression precisely for achieving the crop improvement objectives under optimum and less than optimum conditions (Hussain et al. 2018; Noble et al. 2018)

In this study 109 sunflower genotypes panel were evaluated for genetic diversity, linkage disequilibrium and population structure so that to detect the SSR loci associated with nine important agro-morphological traits through association mapping.

Despite of considerable progress being made in plant breeding by conventional approaches; the need to save time and resources, increase accuracy and pace of improvement had urged plant breeders to use new and improved breeding strategies by combining new advancement made in the field of genetics and phenomics. Association mapping has been proved an effective approach being utilized globally for marker assisted breeding program. Sunflower is among the most important oilseed crops but only few association mapping studies has been conducted on this crop so far (Filippi et al. 2015). Therefore, it is needed to characterize sunflower genotypes to upsurge the level of understanding regarding sunflower worldwide genetic map. In this study 109 sunflower genotypes were used to document the level of their genetic diversity, linkage disequilibrium and population structure through microsatellite markers. The SSR loci located in proximity of the genes controlling morphological traits studied could be highlighted at chromosomal level through association mapping analysis. The information gathered in this study will be helpful for plant breeders working on the improvement of morphological traits through directed and precise breeding approaches.

Materials and Methods

Plant material and phenotyping

Present study was performed on 109 sunflower lines

(Table 1) maintained by Oilseeds Research Program (ORP) of National Agricultural Research Center (NARC), Islamabad. For phenotypic evaluation, these sunflower lines were planted in open field conditions at NARC, Islamabad during spring 2016 following augmented block design. Each sunflower line was planted in a 5 m row with row to row distance of 75 cm and plant to plant distance of 25 cm. NPK fertilizers was applied @ 150:60:60 kg/ha. Complete dose of phosphorus and potassium was applied as basal along with half of the nitrogen and remaining half of nitrogen was applied at first irrigation. Thinning was done after 12–15 days to ensure proper plant population. Weeding was done manually twice to keep crop weed free during the growth period. Ten morphological parameters viz. days to flower initiation (days taken from date of sowing till 5% of the plant of an entry starts flowering), days to flower completion (days taken from sowing till 95–100% plants of an entry initiate flowering), days to maturity (days taken from sowing till 95% of the plants turn their brackets brown), plant height (height of the plant from soil surface to base of head at maturity), stem curvature (plant height subtracted from height of the head from the ground surface), number of leaves per plant, leaf area, hundred seed weight, seed yield and oil content were recorded for the phenotyping of sunflower material.

DNA extraction and genotyping with SSR markers

Total genomic DNA was extracted from 10–12 old sunflower seedlings following CTAB (Cetyl trimethylammonium bromide) DNA extraction protocol (Murray and Thompson 1980). DNA extracted was diluted in 50 μ L of TE buffer for working solution and stored at -4°C. Purity and concentration of genomic DNA was checked by running it on 1% agarose gel.

Overall, 40 SSR markers were employed for genotyping (Table 3). These microsatellites were selected from the 17 linkage groups in sunflower identified by Yu et al. (2002) so that a uniform representation of sunflower genome could be ensured. For PCR analysis 20 μ L of reaction mixture was prepared from 1–1.2 μ L DNA solution, 2 μ L Taq Buffer, 2.5 μ L MgCl₂, 2 μ L dNTP's mixture, 0.2 μ L Taq Polymerase enzyme, 10.8 μ L dd.H₂O and 0.8 μ L each of forward and reverse primers (Primers were first diluted with dd.H₂O for making their working solution). A touchdown cycling program was employed to reduce the spontaneous amplification of the PCR product. Cycling protocol comprises of initial denaturation at 94°C for 5 min, followed by 30 cycles of 94°C for 30 s, annealing temperature for 30 sec (it ranges from 55°C to 62°C for different primers), extension temperature of 72°C for 40 sec, with final extension at 72°C for 5 min. The PCR products then obtained were run on a 2% agarose gel for visualization of the amplified segments.

Linkage disequilibrium

Pairwise LD among the SSRs was calculated using the TASSEL program (v.3.0.174) based on the allele frequency correlation (r^2) and LD was drawn to represent the pairwise LD measurements graphically.

Population structure

For population structure analysis Bayesian clustering approach was followed for the SSR genotyping data in the Structure program (v.2.3.4) (Falush *et al.* 2003). For calculation of ancestry fractions of each cluster an admixture model and correlated allele frequencies were applied to each accession, 10 independent runs for each K-value (1–10) were completed with burn-in period of 10,000 followed by 10,000 Markov Chain Monte Carlo repetitions. The delta K method was implemented in Structure Harvester Program (Earl 2012) to determine the most suitable K-value. An unrooted neighbor joining phylogenetic tree was drawn using TASSEL program (v.3.0.174).

Association mapping

Association mapping among the phenotypic data of nine morphological traits and the genotypic data of 40 SSR markers was performed using the program TASSEL (v 3.0.174). SSR markers with known linkage groups and their corresponding positions were used. MLM (mixed linear model) of association mapping that uses both population structure and kinship matrix was employed so that to minimize the probability of false association that may arise in GLM (general linear model) based method. The association between marker and trait was considered significant at $P \leq 0.05$ (Ambreen *et al.* 2018).

Results

Phenotypic variability and population structure analysis

High level of phenotypic variability was observed in the field condition among the sunflower genotypes for all the studied traits (Table 2). Phenotypic data collected exhibited a broad variation among the sunflower studied panel making it an ideal population for documenting the genotypic variability. The data of ten morphological traits was then later combined with the genotypic data revealed by the SSR markers genotyping to highlight the underlying genes controlling these traits. Forty SSR markers were used for the population structure analysis that amplified a total of 65 bands. The admixture model-based analysis provided information about the optimal number of sub-populations. As the value of K increased from 1 to 10 (Fig. 1) LnP(D) also increases continuously and maximum inflection was noticed as the value of K changed from 1 to 2 (Fig. 2). This

optimum number of k was further validated by ΔK , a second order statistics. ΔK value also showed a peak at $K=2$ (Fig. 3). This shows that there were two sub-populations in our samples, based on SSR genotyping data. Further analysis of these 2 sub-populations revealed that these two populations contained maintainer and CMS lines separately.

Neighbor-joining tree

An unrooted neighbor joining phylogenetic tree diagram was generated in TASSEL (v.3.0.174) to compute the level of relatedness among the sunflower accessions. Sunflower accessions study panel was divided into three clusters namely CI, CII and CIII (Fig. 4). CI contains A and B lines mainly with some mixture of open-pollinated and few R-lines as well. Likewise, CIII mainly consists of R-lines, whereas, C-II contains few R-lines along-with some open-pollinated lines. This tree diagram validated that female lines are quite distinct from male (R) lines. However, diversity within the clusters was limited.

Linkage disequilibrium analysis

Linkage disequilibrium was assessed among the entire forty SSR markers over 109 sunflower accessions. A linkage disequilibrium graph was generated on the basis of squared correlation of allele frequencies. The distribution assembly of squared correlation of allele frequencies (r^2) is shown in Fig. 5. Loci in red, green and blue dots exhibited high level of LD having their p-value less than 0.0001, 0.001 and 0.01 respectively.

Association mapping analysis

Linked markers along-with their P -values are shown in Table 4. In this study 11 markers showed a significant association with the underlying QTLs controlling six studied traits i.e., head diameter, leaf area, seed yield, days to flower initiation, days to flower completion and hundred seed weight while no significant marker trait association was detected for plant height, stem curvature, oil content and number of leaves per plant. More than one marker was found to have strong correlation with head diameter and days to flower initiation. A scatter plot diagram was drawn to illustrate the markers expressing strong association with the traits studied with threshold value at $P \leq 0.05$ (Fig. 6).

Discussion

Rapid domestication and urge for more productive cultivars housed with superior quality attributes had resulted in more yields but at the cost of loss in the genetic diversity. To address the threats faced by the crops with narrow genetic base like sunflower, scientists are establishing and maintaining large and genetically divergent germplasm material. In this present research, 109 diverse sunflower

Table 1: List of Sunflower Accessions used in present study

S. No.	Accession No.	Source	S. No.	Accession No.	Source
1	CMS-HAP-12	NARC, Islamabad	56	RHP-38	NARC, Islamabad
2	CMS-HAP-56	NARC, Islamabad	57	RHP-77	NARC, Islamabad
3	CMS-HAP-101	NARC, Islamabad	58	RHP-82	NARC, Islamabad
4	CMS-HAP-54	NARC, Islamabad	59	RHP-42	NARC, Islamabad
5	CMS-HAP-103	NARC, Islamabad	60	RHP-73	NARC, Islamabad
6	CMS-HAP-24	NARC, Islamabad	61	RHP-74DN	NARC, Islamabad
7	CMS-HAP-110	NARC, Islamabad	62	RHP-7485	NARC, Islamabad
8	CMS-HAP-112	NARC, Islamabad	63	RHP-7490	NARC, Islamabad
9	CMS-HAP-25	NARC, Islamabad	64	RHP-7495	NARC, Islamabad
10	CMS-HAP-111	NARC, Islamabad	65	RHP-7498	NARC, Islamabad
11	CMS-HAP-10	NARC, Islamabad	66	RHP-74100	NARC, Islamabad
12	CMS-HAP-114	NARC, Islamabad	67	RHP-74105	NARC, Islamabad
13	CMS-HAP-115	NARC, Islamabad	68	RHP-74107	NARC, Islamabad
14	CMS-HAP-03	NARC, Islamabad	69	RHP-74108	NARC, Islamabad
15	CMS-HAP-99	NARC, Islamabad	70	RHP-74110	NARC, Islamabad
16	CMS-HAP-125	NARC, Islamabad	71	RHP-74112	NARC, Islamabad
17	CMS-HAP-118	NARC, Islamabad	72	RHP-74115	NARC, Islamabad
18	CMS-HAP-116	NARC, Islamabad	73	RHP-74120	NARC, Islamabad
19	CMS-HAP-121	NARC, Islamabad	74	RHP-74125	NARC, Islamabad
20	CMS-HAP-117	NARC, Islamabad	75	RHP-74128	NARC, Islamabad
21	CMS-HAP-122	NARC, Islamabad	76	RHP-74130	NARC, Islamabad
22	CMS-HAP-120	NARC, Islamabad	77	RHP-71	NARC, Islamabad
23	CMS-HAP-123	NARC, Islamabad	78	SFP-14	NARC, Islamabad
24	CMS-HAP-102	NARC, Islamabad	79	SFP-12	NARC, Islamabad
25	CMS-HAP-08	NARC, Islamabad	80	SFP-10	NARC, Islamabad
26	CMS-HAP-119	NARC, Islamabad	81	SFP-40	NARC, Islamabad
27	HAP-12	NARC, Islamabad	82	SFP-42	NARC, Islamabad
28	HAP-56	NARC, Islamabad	83	SFP-38	NARC, Islamabad
29	HAP-101	NARC, Islamabad	84	SFP-18	NARC, Islamabad
30	HAP-54	NARC, Islamabad	85	SFP-36	NARC, Islamabad
31	HAP-103	NARC, Islamabad	86	SFP-31	NARC, Islamabad
32	HAP-24	NARC, Islamabad	87	SFP-37	NARC, Islamabad
33	HAP-110	NARC, Islamabad	88	SFP-24	NARC, Islamabad
34	HAP-112	NARC, Islamabad	89	SFP-09	NARC, Islamabad
35	HAP-25	NARC, Islamabad	90	SFP-41	NARC, Islamabad
36	HAP-102	NARC, Islamabad	91	SFP-19	NARC, Islamabad
37	HAP-10	NARC, Islamabad	92	SFP-22	NARC, Islamabad
38	HAP-114	NARC, Islamabad	93	SFP-25	NARC, Islamabad
39	HAP-116	NARC, Islamabad	94	SFP-43	NARC, Islamabad
40	HAP-123	NARC, Islamabad	95	SFP-33	NARC, Islamabad
41	HAP-111	NARC, Islamabad	96	SFP-46	NARC, Islamabad
42	HAP-99	NARC, Islamabad	97	SFP-08	NARC, Islamabad
43	HAP-122	NARC, Islamabad	98	SFP-07	NARC, Islamabad
44	HAP-120	NARC, Islamabad	99	SFP-16	NARC, Islamabad
45	HAP-03	NARC, Islamabad	100	SFP-26	NARC, Islamabad
46	HAP-08	NARC, Islamabad	101	SFP-13	NARC, Islamabad
47	RHP-68	NARC, Islamabad	102	SFP-35	NARC, Islamabad
48	RHP-72	NARC, Islamabad	103	SFP-20	NARC, Islamabad
49	RHP-53	NARC, Islamabad	104	SFP-32	NARC, Islamabad
50	RHP-73-1	NARC, Islamabad	105	RHP-83	NARC, Islamabad
51	RHP-46	NARC, Islamabad	106	RHP-84	NARC, Islamabad
52	RHP-76	NARC, Islamabad	107	RHP-88	NARC, Islamabad
53	RHP-41	NARC, Islamabad	108	RHP-86	NARC, Islamabad
54	RHP-81	NARC, Islamabad	109	RHP-89	NARC, Islamabad
55	RHP-69	NARC, Islamabad			

lines that included A, B R and OPV's were studied. Ten morphological traits studied showed a high level of genetic variability in field conditions (Table 2) making this panel of sunflower genotypes an ideal fit for determining the marker-trait association by combining the phenotypic and genotypic data. For genotyping 40 SSR markers that yielded 65 scorable DNA bands. SSR markers have been found to be very effective as they showed more resolution power than SNPs (Emanuelli *et al.* 2013).

SSR based genomic data is the most common method of studying the genetic diversity and population structure analysis. Genomic SSR's are useful marker types because of their abundance in genome, higher polymorphic content and reproducibility (Filippi *et al.* 2015). It has also been reported in recent studies that SSR markers produce same results as obtained through SNPs from GBS (Souza *et al.* 2018).

In depth knowledge of population structure is important to avoid any spurious or false associations

Table 2: Mean, range and standard deviation studied traits among sunflower genotypes

Crop traits	Range	Mean	Standard deviation	Genotypes with highest value	Genotype with lowest value
DFI	69 –94	79.86	5.85	RHP-7485	CMS-HAP-102
DFC	72 – 102	91.45	8.12	RHP-7495	CMS-HAP-102
PH	121.8 – 246.25	185.46	27.13	SFP-42	CMS-HAP-111
SC	15.5 – 36.7	27.68	5.45	SFP-12	SFP-31
HD	9.25 – 20.9	14.50	2.93	HAP-10	RHP-69
L/P	22 – 4189	30.84	3.99	SFP-33	HAP-111
L/A	139.08 – 276.48	202.31	39.13	SFP-37	CMS-HAP-12
HSW	2.76 – 6.94	4.92	0.98	CMS-HAP-111	RHP-42
SY/P	23.9 – 62.7	37.14	7.31	CMS-HAP-54	RHP-41
OC	35.6 – 59.93	44.18	4.99	CMS-HAP-12	SFP-33

DFI: Days to flower initiation; DFC: Days to flower completion; PH: Plant height; SC: Stem curvature; L/P: Leaves per plant; L/A: Average leaf area; HSW: 100-seed weight, SY/P: Seed yield per plant; SY/ha: Seed yield per hectare OC: Oil contents

Table 3: List of SSR markers used to study the association mapping and population structure in sunflower

Primer Name	Linkage Group	Forward Sequence	Reverse Sequence
ORS-605	1	CGCGTGATGTGACGATTATT	ACGGAGCAAAGTTTCGAGGT
ORS-543	1	CCAAGTTTCAGTTACAATCCATGA	GGTCATTAGGAGTTTGGGATCA
ORS-371	1	CACACCACCAAACATCAACC	GGTGCCTTCTTCCCTTGTG
ORS-453	2	CCTGTGAGCTACAATACTCCACA	GATTCTGATTAGGCGGTGGT
ORS-1053	2	TTTCATCACATTAGACCATAGACCA	GGCTTTCCTTCGTGGTTTGTAT
ORS-752	3	CACTGATGAACAAGTGGGAGA	ATGATTCCTATACCCACCAA
ORS-924	3	TAAATCGCCATAACCACTCCATC	TATCAGCAGGAAGAACGCCTAAT
ORS-366	4	AACCAACTGAGCATTCTGTGA	GCGCTAGGTTAAGAGGACAAA
ORS-1068	4	AATTTGTCGACGGTGACGATAG	TTTTGTCAATTCATTACCCAAGG
ORS-337	4	TGGTTCATTATCCTTGGTC	GGTGTGGTGGTTAATTCGTC
ORS-1024	5	GGGAAGTGGGCTTGTCTATGTAT	AACACACCGAAATCACCTATGAA
ORS-533	5	TGGTGGAGTCACTATTGGA	AGGAAAGAAGGAAGCCGAGA
ORS-608	6	CATGGAAAGCCAGTCTCT	CGTGCGTGATTAACATACCC
ORS-1256	6	GATGTTGATGTTGGTGAAGTTGC	CTCCGTCACCTTAAGCACTTGTA
ORS-400	7	CGAACCCTGTGTACCGTTT	ACTTCGTTACAAGGCACAA
ORS-700	7	GTACCCACCACGCTTAACCA	AGTCTCCACAGCAACGTCA
ORS-830	8	CAAGTGCAATTAGGTGTTCTAACA	GCCCTCTGACTGTTGTATGACTG
ORS-599	8	TTCCCTATCACACGCCTCTC	GAAAGGAAGTAGCGGTGGTG
ORS-882	9	AAACCCGCATGTAAGATATTCCG	ATCGGGAGCAGAAGAAGAGTATG
ORS-617	9	GGTACTTGGTATTCATGGGTCAT	GACACCGCAACTTAACACTT
ORS-795	9	CGTAGTTACACCCAGATCG	TGTCCACAGGTTGAAGATCG
ORS-613	10	GTA AACCTTAGGTCAATTTGCAG	ATCTCCGAAAACATTCTCG
ORS-1088	10	ACTATCGAACCTCCTCCAAAC	GGATTCTTTCATCTTTGTGGTG
ORS-433	10	CCGAGGTTTGATCGCTATTT	AGCGTTTGTGATTTGATTACGA
ORS-769	11	GTTTATTTATGTAGAAATGTTCTGGAA	ATGTGGTGGTAAGGTTTGGT
ORS-697	11	TTGGGCTGTGGTTCTTAAAC	AAGAGATGGGAGTGTGTATGC
ORS-1085	12	GACCTCAAGGCATGCTAACACTC	ACTAAGTGTGTGGACGGGGAAA
ORS-1040	12	CTGCTGATCGTTTCTTGGATAGA	TGTAATCCTTCTAATCAACTCCAC
ORS-879	13	GAACCTCCCTTTGTCTGCATATC	CTCCGTTGCTGTTGATGTCT
ORS-781	13	GTCAACCCATGACCCAAACC	GATGTGGAGGAGAGGGGTGT
ORS-511	13	TGGCTCAGATTAAGTTCACACAG	CGGTTGCGAGTAACAGGTA
ORS-307	14	CAGTTCCTGAAACCAATTCA	GCAGTAGAAGATGACGGGATG
ORS-1086	14	TTGTTTGTGCGCACTCAAGATT	ATTATCGGCACATCTTTGGATTT
ORS-857	15	ACATCCGAACGAAGGACAATC	CAAGAAAGTATGTCACCCAATAGCA
ORS-562	15	CACACACAAAACCCTAGCTCT	CAATCATATCGAGCACACATCA
ORS-768	16	CCACTCATCATCAAGCCTAACA	AGTGGTGCTGTTGTAGGT
ORS-1064	16	TGAATGATCTATGAGTGGTGTAGG	ACTCGAGTGGTAAGTCTGTTAGG
ORS-495	16	CCAGGATTAGGTAGCTTAGTTCG	GCGATCTGAGGTTGACTCGT
ORS-811	17	CCTTCTCCTCAATCTTTGGCTA	AGGAATGAAATGGGTGTGTGT
ORS-845	17	GGTGCCCTATCTTCATCTCTG	CTAAAGGGTATCACACATTTGACATT

(Flint-Garcia *et al.* 2005). Based on the diagrams of ΔK and $\ln P(D)$, the sunflower accessions were divided into two sub-populations as highest peak was observed at $K = 2$. Mandel *et al.* (2013) studied 433 sunflower lines and found two sub-populations based on the optimum K value. In sunflower population studies CMS and R lines tend to group separately from each other and this trend has been

observed previously by (Lochner 2011; Ibrar *et al.* 2018)

Linkage disequilibrium is an important analysis while performing marker assisted selection and association mapping analysis. It is considered as a non-random association of alleles at different loci present on the same chromosome. It assumes the co-segregation of a specific trait and DNA marker and by using this information locates

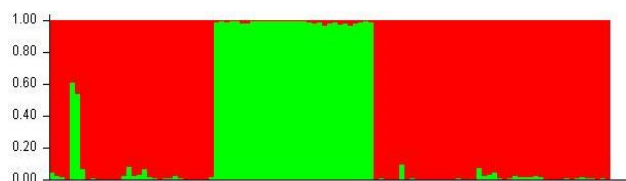


Fig. 1: Population structure of 109 sunflower accessions based on SSR genotyping data at K=2 (Structure 2.3.4)

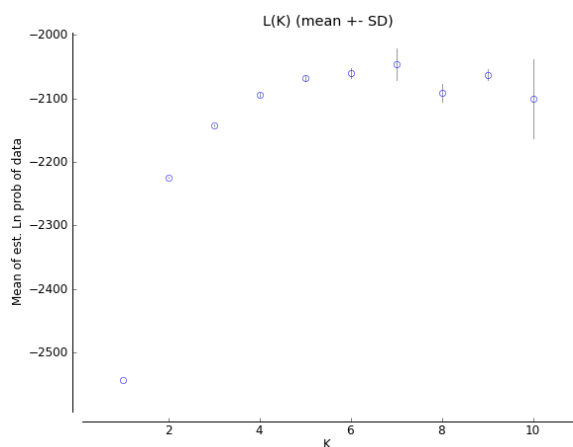


Fig. 2: Diagram of LnP(D) of possible clusters (K) from 1 to 10

the QTLs or major genes (Darvishzadeh 2016). Resolution of association mapping depends on the structure of LD across the genome. LD is dependent on various factors like outcrossing, inbreeding, population size, physical separation and recombination frequency between loci, mutation rate, selection, historical sub-division and admixture of populations and genomic rearrangements (Ilyas *et al.* 2018). Presence of linkage disequilibrium is a pre-requisite for any association mapping studies as LD determines the significance of association among QTL's and the phenotype (Maulana *et al.* 2018). Mandel *et al.* (2011) also reported two distinct populations of sunflower one primarily composed of R lines and the other of B or female lines. This distinction among R and B lines is may be due to their separate breeding history and origin (Lochner 2011). The rare mixing of B lines in R line group may arise because of continuous breeding efforts as sunflower breeders had introgressed superior traits from one genotype to the other for the development of superior inbred lines for hybrid development (Fick and Miller 1997).

This study provided us to detect QTLs controlling some important morphological traits like head diameter, days to flower initiation, leaf area, 100 seed weight, days to flower completion and seed yield per plant. These traits expressed a strong correlation with seed yield in sunflower (Arshad *et al.* 2010; Jalil *et al.* 2014); thus these markers can be used for designing sunflower breeding programs for increased seed yield alongwith short duration. However for determination of marker-trait association with other

Table 4: List of traits along with significantly associated markers ($P \leq 0.05$)

Sr. No.	Trait	Marker	P- value
1	Head diameter	ORS-1088	0.0054
2	Head diameter	ORS-371	0.0054
3	Leaf area	ORS-1085	0.0079
4	Seed yield per plant	ORS-769	0.019
5	Head diameter	ORS-608	0.025
6	Head diameter	ORS-608	0.025
7	Head diameter	ORS-608	0.025
8	Days to flower initiation	ORS-433	0.027
9	Days to flower completion	ORS-433	0.027
10	Hundred seed weight	ORS-811	0.028
11	Days to flower initiation	ORS-605	0.033

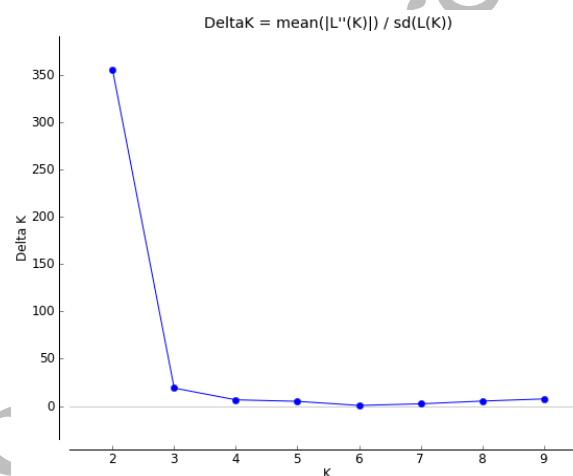


Fig. 3: ΔK based on the rate of change of LnP(D) between successive K values

important characters like oil content and plant height further evaluation should be carried out. Neighbor joining tree shows two sub-populations in the sunflower studied panel along-with a minor cluster containing OPVs. However, out of two major clusters one contains mostly A and B lines while other predominantly contains R lines. Similar clustering pattern has previously been reported by Lochner (2011) and Ibrar *et al.* (2018).

Conclusion

As very few association mapping studies have been conducted on this crop; therefore, in this effort forty SSR markers was utilized to identify marker-trait association. MLM based approach of association mapping coupled with Q+K model identified eleven SSR markers that have been found to be in proximity ($p \leq 0.05$) of the genes controlling six phenotypic traits i.e., head diameter, days to flower initiation and completion, leaf area, seed yield and 100-seed weight. The marker-trait association identified could be used in designing sunflower improvement/breeding programs with more precision and efficacy thus saving the time and resources needed for new cultivar/hybrid development.

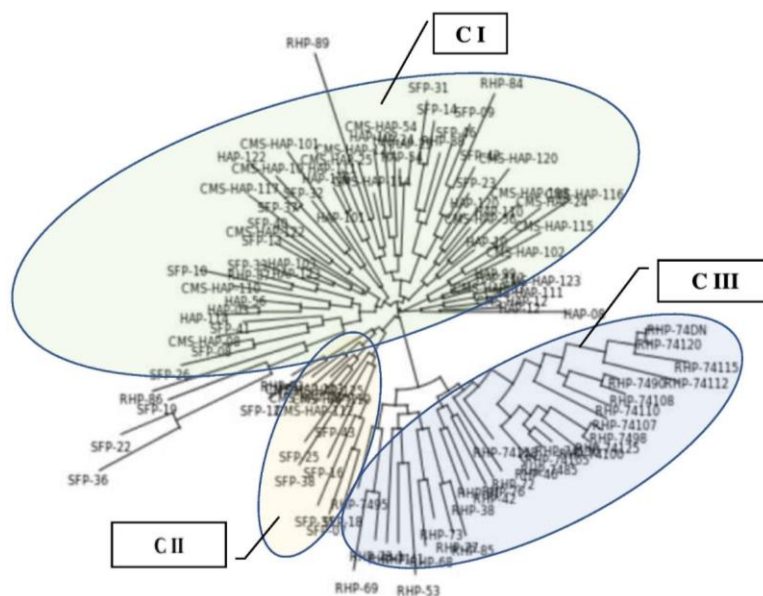


Fig. 4: An unrooted neighbor joining phylogenetic tree of 109 sunflower accessions based on SSR data. C I (Cluster I) consists of mainly CMS and maintainer lines, C II (Cluster II) consists of OPV's and C III (Cluster III) contains restorer lines

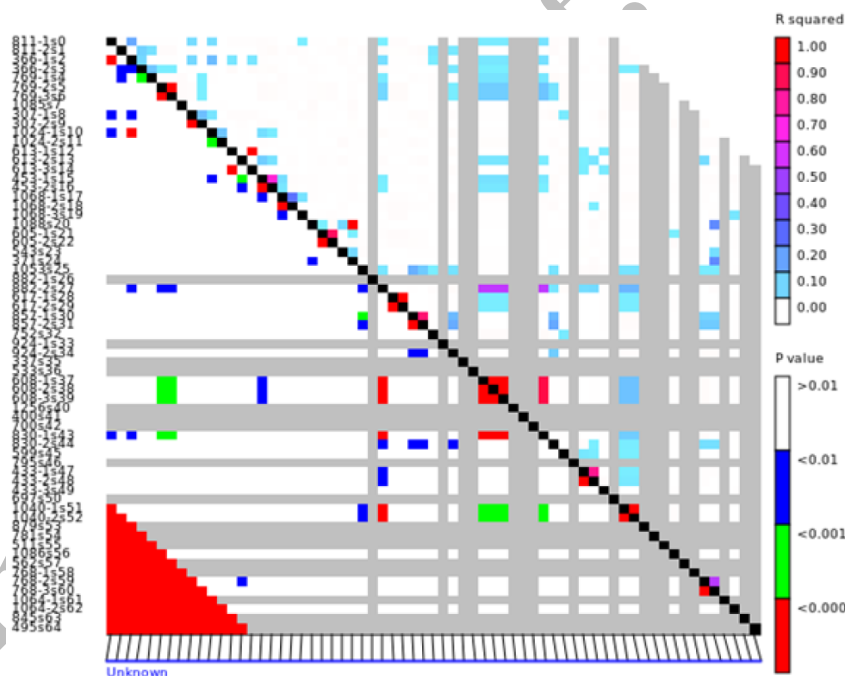


Fig. 5: LD distribution pattern based on squared correlation of allele frequencies (r^2). Each cell represents the comparison of two pairs of marker sites with color codes for the presence of significant linkage disequilibrium

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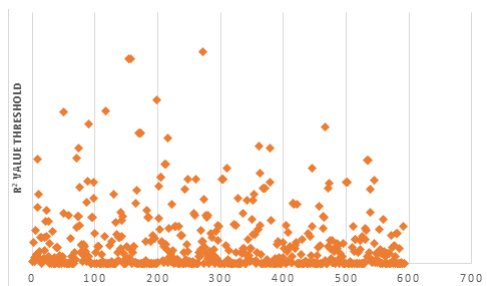


Fig. 6: A scatter plot depicting the marker-traits association

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