

RESEARCH ARTICLE

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Genetic characterization and genome-wide association mapping for stem rust resistance in spring bread wheat

Elias Shewabbez^{1*} , Endashaw Bekele¹, Admas Alemu², Laura Mugnai³ and Wuletaw Tadesse⁴

Abstract

Background: Emerging wheat stem rust races have become a major threat to global wheat production. Finding additional loci responsible for resistance to these races and incorporating them into currently cultivated varieties is the most economic and environmentally sound strategy to combat this problem. Thus, this study was aimed at characterizing the genetic diversity and identifying the genetic loci conferring resistance to the stem rust of wheat. To accomplish this, 245 elite lines introduced from the International Center for Agricultural Research in the Dry Areas (ICARDA) were evaluated under natural stem rust pressure in the field at the Debre Zeit Agricultural Research Center, Ethiopia. The single nucleotide polymorphisms (SNP) marker data was retrieved from a 15 K SNP wheat array. A mixed linear model was used to investigate the association between SNP markers and the best linear unbiased prediction (BLUP) values of the stem rust coefficient of infection (CI).

Results: Phenotypic analysis revealed that 46% of the lines had a coefficient of infection (CI) in a range of 0 to 19. Genome-wide average values of 0.38, 0.20, and 0.71 were recorded for Nei's gene diversity, polymorphism information content, and major allele frequency, respectively. A total of 46 marker-trait associations (MTAs) encompassed within eleven quantitative trait loci (QTL) were detected on chromosomes 1B, 3A, 3B, 4A, 4B, and 5A for CI. Two major QTLs with $-\log_{10}(p) \geq 4$ (EWYP1B.1 and EWYP1B.2) were discovered on chromosome 1B.

Conclusions: This study identified several novel markers associated with stem rust resistance in wheat with the potential to facilitate durable rust resistance development through marker-assisted selection. It is recommended that the resistant wheat genotypes identified in this study be used in the national wheat breeding programs to improve stem rust resistance.

Keywords: Markers, *Puccinia graminis* f. sp. *tritici*, QTL, GWAS, SNP

Background

Wheat (*Triticum aestivum* L.) is a leading crop, both in terms of economic value and area of production worldwide [1, 2]. Developing countries account for nearly 77% of total global wheat imports [3]. Wheat provides nearly 20% of daily world human caloric requirements [4] and

demand is expected to increase to 60% by 2050 [5]. However, various challenges have hindered meeting this demand, with recurrent emerging fungal pathogens proving to be one of the leading problems worldwide [6].

Wheat stem (black) rust, caused by *Puccinia graminis* Pers. f. sp. *tritici*, Eriks. & E. Henn (*Pgt*), has been recognized as a major threat to global food security [7, 8]. Concerns regarding this disease have increased significantly, especially following the 1998 outbreak of the novel virulent race Ug99 which originated in Uganda. Since then, this race has produced 13 different variants throughout

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East Africa [9, 10]. The race can infect 90% of the wheat varieties grown worldwide [11] and yield losses can reach up to 100% in susceptible cultivars under conducive environmental conditions [12]. Races other than Ug99 were also reported in different parts of Western Europe. In 2013, a stem rust epidemic arose in Germany and spread to Denmark, Sweden, and the UK [13]. In 2016/2017, Italy chronicled two epidemics of wheat stem rust caused by race TTRTF, which destroyed tens of thousands of hectares of cultivated wheat [14]. All these reports indicate that the disease is re-emerging as a threat to wheat production globally.

Ethiopia is considered to be a hotspot for the development and evolution of new *Pgt* races [15]. Many new variants of *Pgt*, which were first identified in this country, have spread to different parts of the world. TTKSK, TKTTF, TRTTF, JRCQC, and TTTTF are the current major wheat stem races that are threatening wheat productivity in Ethiopia [16]. In 2013/2014, severe stem rust epidemics were caused by *Pgt* race TKTTF (not a member of Ug99 lineage), resulting in almost total yield loss on widely grown wheat cultivars. Since then, this race has spread widely and has been found in 10 different countries, including Western Europe [17].

To overcome this problem, host plant resistance developed through molecular marker technology is the most sustainable, cost-effective, and environmentally friendly approach for controlling rust diseases [7, 18]. Accordingly, many molecular markers linked with *Pgt* resistance were discovered throughout the wheat genome during the past couple of decades using genome-wide association mapping (GWAS). GWAS has been the most effective tool to detect several quantitative trait loci (QTLs), with moderate to minor effects against *Pgt* disease [19]. However, factors such as population structure and kinship similarity should be controlled properly to avoid false-positive QTLs. To overcome this, several models, including the mixed linear model (MLM) have been implemented. Since the first report in 2007 [20], various GWAS studies were carried out successfully and high numbers of QTLs have shown *Pgt* resistance in wheat [21–24]. So far, more than 80 genes conferring resistances to *Pgt* have been cataloged in common wheat and wheat relatives [24]. However, only a few genes are effective against all pathogen strains. Of these, *Sr2*, *Sr13*, *Sr22*, *Sr25*, *Sr26*, *Sr35*, *Sr39*, and *Sr40* were reported to be the most effective against Ug99 [18].

The frequent co-evolution of host and pathogen remains a big challenge in the durability of the released resistant cultivars [25]. The narrow genetic diversity of cultivated wheat cultivars [22, 26] and the impact of climate change [12] are the major cause of this problem. Thus, additional sources of resistant QTLs, followed by

marker-assisted gene pyramiding, are required to produce durable resistant varieties. Therefore, this study aimed to characterize the genetic diversity and to identify novel QTLs associated with resistance to stem rust of wheat through GWAS.

Results

Phenotypic variation and heritability

The performance of genotypes towards stem rust resistance varied greatly. For instance, the disease severity score was ranged between 10 and 80%. The majority (46.7%) had a disease severity (DS) score of 15–30% whilst 8.5% had a DS score of 0% (Fig. 1, Additional file 1). The best linear unbiased estimates (BLUP) values of DS and coefficient of infection (CI) were calculated from adjusted means of each accession across two years, and are summarized in Fig. 1.

The data of disease severity (DS) and infection response (IR) were combined to define the disease response as the coefficient of infection (CI) and 71% of lines had less than 30 (Fig. 1C). Of these, the top twenty resistance lines (presented in Table 1) ranged with the average CI values of 4.5 for pedigree SERI.1B//KAUZ/HEVO/3/AMAD/4/CHAM-6/FLORKWA-2 to 12 for pedigree SERI.1B//KAUZ/HEVO/3/AMAD/4/WEAVER/JACANA. Additional genotypes scored between 6 to 80 of CI and are presented in Additional file 1. On the other hand, all local controls (i.e. Digelu, Kubssa, Hidasse, Honqolo, and Ogolcho) were susceptible, with average CI ranging from 60 for HIDASSE to 80 for OGOLCHO and HONQOLO.

The ANOVA analysis revealed highly significant variation among genotypes ($P < 0.001$) and genotype \times year interactions ($P < 0.001$) for all parameters. Heritability (H) values for DS were 79% and IR 72%, suggesting that all parameters had a strong genetic basis. In addition, the disease distribution of the breeding lines was high between seasons, with average correlations of 0.76, 0.85, and 0.78 for DS, IT, and CI, respectively (Table 2).

Population structure and genetic diversity analysis

The population structure of the panel was inferred through the Bayesian clustering model, principal component analysis (PCA), and neighbor-joining (NJ) tree. The Bayesian clustering model applied on STRUCTURE software and subsequent application of STRUCTURE HARVESTER showed a delta K peak value of two (Fig. 2A). As a result, accessions were classified into two sub-populations composed of 106 and 139 lines in sub-populations 1 and 2, respectively (Fig. 2C). The scree plot of PCA showed that weak kinship existed among the lines. For the first 10 principal components (PCs), variances of SNP markers were changed from 7.5% (PC1) to 2% (PC10) and between 0 and 2% after PC10 (Fig. 2B).

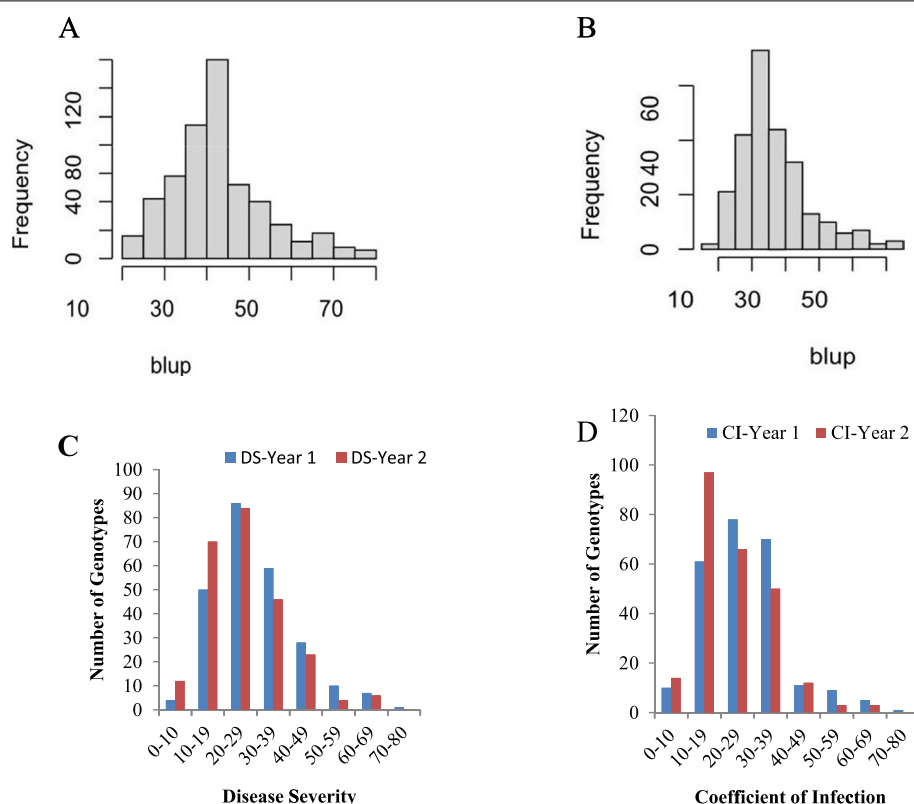


Fig. 1 Distribution of adult plant stage disease resistance (APR) response and best linear unbiased estimates (BLUPs) as disease severity (DS), and coefficient of infection (CI). (A) BLUPs of DS; (B) BLUPs of CI; (C) Frequency of genotypes for disease severity (DS); (D) frequency of genotypes for the coefficient of infection

Phylogenetic tree analysis of the genetic relationship between the populations was carried out based on the distance-based neighbor-joining tree on TASSEL software v5.2.35 followed by web-based visualization software iTOL. The resulting dendrogram shows three phylogenetic groups color-coded with a STRUCTURE probability distribution. This is not consistent with the STRUCTURE result (which was two groups). Since the genotypes are elite lines, passed by complex breeding history, such inconsistency is expected. However, the majority of lines were still grouped in the same group as the STRUCTURE result and some lines were grouped in the mixed group. For instance, 78 (56%) of the lines in the first group were composed of a sub-population 1, whereas 61 lines (44%) were categorized in subpopulation 2. The second group was composed mainly of subpopulation 2, which consists of 49 (70%) lines; whereas 21 (30%) lines were classified in subpopulation 1. The third group was composed of 58 (76%) lines from subpopulation 2, and 21 (30%) lines from subpopulation 1 (Fig. 3).

Genetic data and linkage disequilibrium

Once sub-optimal quality markers had been filtered out, 9523 SNP markers were retained from 245 lines. The distribution of SNPs across the A, B, and D sub-genomes was 50, 39, and 11%, respectively. The maximum number of SNP markers was recorded on chromosome 2B (930) and the minimum number was on chromosome 4D (48) (Fig. 4). The mean genome-wide heterozygosity, genome-wide polymorphic information content (PIC), and gene diversity were 0.006, 0.2, and 0.38, respectively. The PIC scores of SNPs varied, with only 1% being highly informative (>0.5), while 75 and 24% of markers had moderate (0.25–0.5) and least (<0.2) PIC scores, respectively.

Linkage disequilibrium decay based on SNP markers of each chromosome was calculated as the Pearson correlation coefficient (r^2) between marker pairs as a function of genetic distance (cM). The LOESS curve intercepted the line of critical value at 6 cM in A genome, 8 cM in B genome, and 5 cM in D genome, indicating that all markers within these ranges were considered as a single locus (Fig. 5).

Table 1 Lists of top resistant lines and their pedigree during 2017/2018 main season at Debre Zeit Agricultural Research Center, Ethiopia

No	Pedigree	Disease severity and response to Sr	
		2018	2019
1	SERI.1B//KAUZ/HEVO/3/AMAD/4/CHAM-6/FLORKWA-2	10MR	15MR
2	SERI.1B//KAUZ/HEVO/3/AMAD/4/MO88/MILAN	15MR	10MR
3	SERI.1B//KAUZ/HEVO/3/AMAD/4/TNMU/MILAN/5/WATAN-12	15MR	10MR
4	PBW343*2/KUKUN//22SAWSN – 97	10MRMS	15MR
5	SERI.1B//KAUZ/HEVO/3/AMAD/4/ESDA/SHWA//BCN	10MR	10MR
6	SERI.1B*2/3/KAUZ*2/BOW//KAUZ/4/SHIHAB-7	10MR	10MR
7	CROC-1/AE.SQUARROSA (224)//OPATA/3/FLAG-7	10MR	10MR
8	TRACHA-2/SHUHA-3/3/SHUHA-7//SERI 82/SHUHA'S'	15MRMS	10MRMS
9	SERI.1B//KAUZ/HEVO/3/AMAD/4/PFAU/MILAN	15MR	10MR
10	WATAN-7/SEKHRAH-2	10MR	15MRMS
11	WEAVER/WL 3928//SW 89.3064/3/SOMAMA-3	15MRMS	10MR
12	SERI.1B//KAUZ/HEVO/3/AMAD/4/SHUHA-7//SERI 82/SHUHA'S'	15MR	15MS
13	KAUZ'S'/SERI/3/TEVEE'S'//CROW/VEE'S'	15MRMS	15MR
14	ATTILA*2/CROW/3/VEE#5/SARA//DUCULA	15MR	15MR
15	TILILA/MUBASHIIR-1	15MR	15MR
16	QAFZAH-27/SEKSAKA-6	15MR	15MR
17	SERI.1B*2/3/KAUZ*2/BOW//KAUZ/4/SHIHAB-7	15MR	15MR
18	STAR*3/LOTUS-5/3/CHUM//7*BCN/4/FLAG-2	15MR	10MR
19	HADIAH-14/ANGI-2	10MRMS	15MRMS
20	SERI.1B//KAUZ/HEVO/3/AMAD/4/WEAVER/JACANA	15MR	15MR

Table 2 Mean response, variance component estimates and heritability for IR, DS, and CI variables

	DS (%)	IR(0–1)	CI
Range	10–80	0.3–1	3–80
Grand mean	33.90	0.8	28.27
BLUEs	32.60	0.8	26.902
σ^2_G	385.3***	0.013***	374.5***
σ^2_E	21.05*	0.000 ^{ns}	15.7 ^{ns}
$\sigma^2_{G \times E}$	20.70**	0.005 ^{ns}	24.1**
σ^2_{error}	166.10	0.004	200.3
H	78.95	72.2	75.7
CV	37.93 0.76	8.52 0.85	50.06 0.78
r			

Disease severity (DS); infection response (IR); coefficient of infection (CI); BLUEs, best linear unbiased estimate; σ^2_G estimate of genotypic variance; σ^2_E estimate of environmental variance; $\sigma^2_{G \times E}$ is the genotype by environment interaction variance; σ^2_{error} is the residual error variance; heritability (H); r Pearson's correlation coefficients among stem rust DS, IT, and CI between two seasons. *, **, *** and ns represents significance at $P < 0.05$, $P < 0.01$, $P < 0.001$, and not significant, respectively

Marker-trait associations

A mixed linear model (MLM) was implemented for MTA, including the population structure and kinship similarity matrix (Q + K) and the BLUPs estimated values

of CI of genotypes and quality checked SNP markers. The model appropriately discovered valuable MTAs with neither inflation (false-positive/type I error) nor overcorrection (false-negative/type II error) problems as depicted from the Q-Q plot (Fig. 8).

A total of 46 MTAs included in 11 QTLs were discovered for CI with the considered exploratory significant threshold ($-\log_{10}(p) \geq 3$). The highest number of MTAs (44) was detected on the B sub-genome, of which 36, 4, 3, and 1 MTAs were located on chromosomes 1B, 3B, 4B, and 5B, respectively (Table 3, Fig. 7). The remaining 2 QTLs were identified from chromosomes 3A and 4A. The explained phenotypic variance of QTLs was taken from the most significant SNP marker (QTL tag-SNP). The three QTLs with high explained phenotypic variance (PV) of tag SNPs were EWYP1B.4 (8.8%), EWYP1B.5 (8.37%), and EWYP1B.3 (8.16%). The other PV of QTLs tag-SNPs ranged from 7.48 to 4.76% (Table 3, Fig. 7). Details of markers distribution in each accession are presented in Additional file 1.

Discussion

Stem rust has been increasing in severity and incidence and now poses a serious threat to global wheat production [8]. To overcome this threat, efforts are ongoing

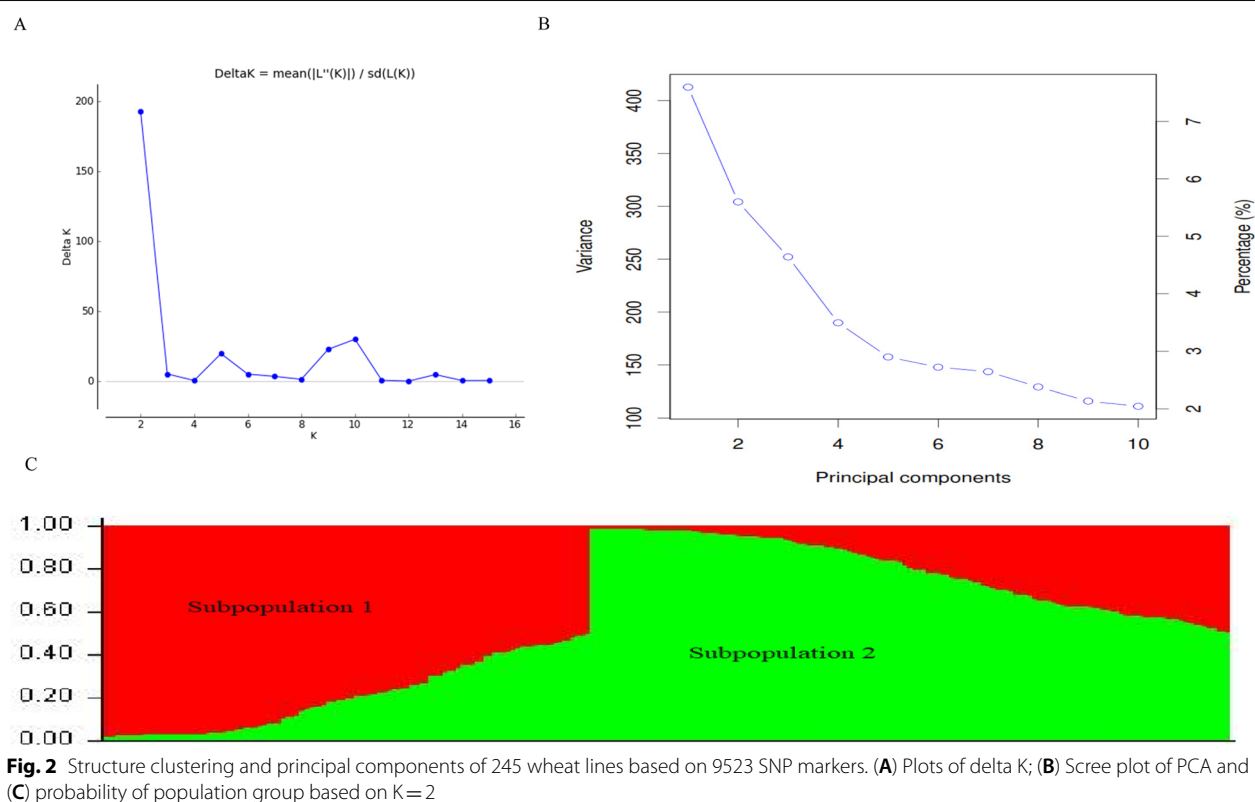


Fig. 2 Structure clustering and principal components of 245 wheat lines based on 9523 SNP markers. **(A)** Plots of delta K; **(B)** Scree plot of PCA and **(C)** probability of population group based on K=2

worldwide to monitor rust diseases, identify rust pathotypes, and evaluate wheat germplasm for rust resistance [36]. As part of the global effort, this study was designed to quantify the existing allelic variation of breeding lines and to search for sources of resistant QTL for *Pgt* resistance. Consequently, 245 elite bread wheat lines were evaluated in the field condition to identify QTLs for adult plant resistance to wheat stem rust. A significant variation was observed between breeding lines for adult plants' resistance to stem rust. This study detected several MTAs included in 11 different QTLs with different effects that could potentially play an important role in future marker-assisted pyramiding against the disease.

Field evaluation of wheat germplasm for resistance to stem rust

Disease response characterization under high disease pressure in field conditions remains the best stem rust management strategy in breeding for developing stem rust-resistant cultivars [37]. Ethiopia is considered to be a hotspot for the development of *Pgt* race diversity and frequent disease epidemics. Studies carried out in Ethiopia showed that most previously identified races such as TTKSK, TKTFE, TTTTE, TRTTE, RRTTE, and others were virulent on most varieties grown in the country

[38]. Accordingly, many field evaluation studies for *Pgt* response have been carried out in different wheat-growing regions of the country [22, 39, 40]. Most elite breeding lines skewed towards moderate resistance, although some differences were observed between individual genotypes. The two parameters (i.e. DS and IR) showed moderate to high heritability with significant variation among lines and genotype x year interactions, indicating that most of the existing variation was due to genetic bases. CI has been used as the most efficient trait to discover QTLs of stem rust resistance in wheat via GWAS analysis [23, 41].

Population structure and genetic diversity

Systematic characterization of population structure and genetic diversity provides a foundation for efficient exploitation of genetic resources and can enhance breeding for durable stem rust resistance in wheat. For the population structure study, three different approaches were applied. Although there is considerable overlap between the three techniques for population analysis, the overall conclusion suggests that there is no clear and substantial separation between individual genotypes. This could be owing to the panel's complicated evolutionary and breeding history. It is suggested that more researches

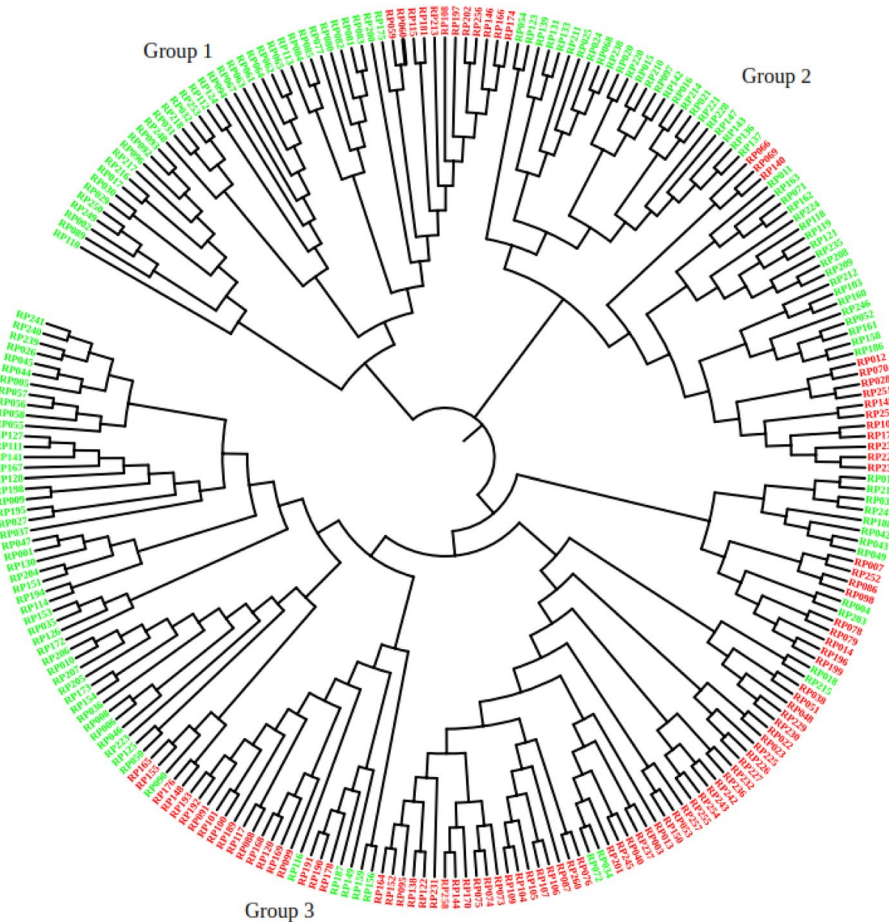


Fig. 3 A dendrogram illustrating the clusters of wheat lines based on Nei's genetic distance. The lines were color-coded with STRUCTURE probability distribution. Clusters with similar pedigrees and genetic backgrounds were named by their common parent

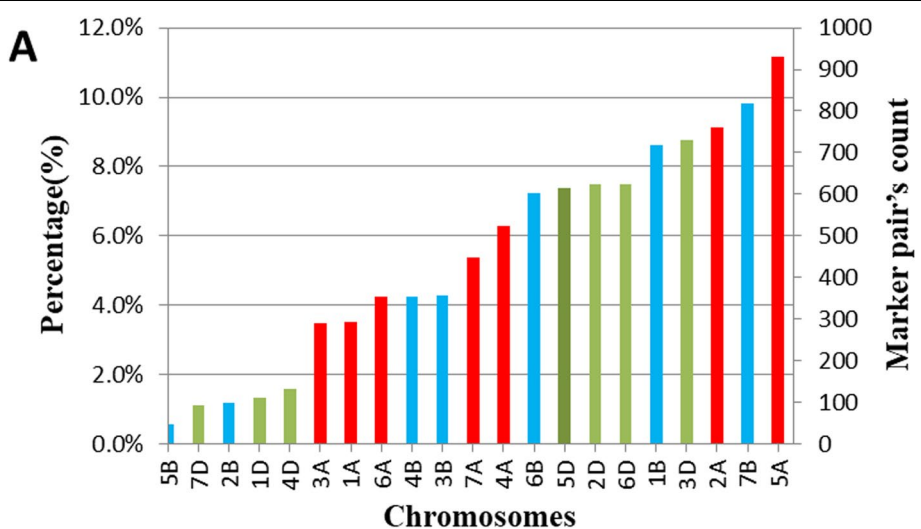


Fig. 4 Genome-wide distributions of single nucleotide polymorphisms (SNPs) based on 15K genotyping results

need to be done to better understand the relationships between genotypes from different groups.

The mean PIC and gene diversity of the genome was 0.25 and 0.3, respectively. Several studies have previously reported various rates of Nei's gene diversity and PIC in different wheat populations [22, 34, 42–44].

Linkage disequilibrium and MTAs

The LD of the genome and sub-genomes of the current panel was estimated using SNP markers. The fastest LD decay was observed on the D sub-genome, which agreed with the previous report [45]. The LOESS curve intercepted the line of critical value at 6 cM in A genome, at 8 cM in B genome, and 5 cM in the D genome, indicating that all markers within these ranges are considered as a single locus (Fig. 5). Since many significant SNP markers (36 MTAs) were identified in the present study, the LD pattern in chromosome 1B was analyzed independently and detected five LD blocks (Fig. 6). Similar large LD blocks in 1B chromosomes have been reported previously [46].

The current study unveiled 46 SNP significant markers encompassed within 11 QTLs. Of these, only two QTLs (EWYP1B.1, EWYP1B.2) were identified as major QTLs ($-\log_{10}(p) \geq 4$). EWYP1B.1, EWYP3A, EWYP3B.1, and EWYP5B, respectively, were found near genomic areas of *Sr31*, *Sr27*, *Sr2*, and *Sr56* [33]. The remaining seven QTLs (EWYP1B.2, EWYP1B.3, EWYP1B.4, EWYP1B.5, EWYP3B.2, EWYP4A, and EWYP4B) were newly discovered in the current study. These new QTLs could play paramount importance in enhancing *Pgt* resistance through marker-assisted selection or introgression.

We found five QTLs on chromosome 1B (EWYP1B.1, EWYP1B.2, EWYP1B.3, EWYP1B.4, and EWYP1B.5) that encompassed 36 MTAs ranging in size from 30.34 cM (*wsnp_Ku_c13229_21142792*) to 114 cM (*Tdurum_contig10036_977*). On this chromosome, three resistance genes (*Sr14*, *Sr31*, and *Sr58*) were cataloged previously [32]. Of these, only *Sr31* has been reported in association with wheat stem rust disease at locus EWYP1B. 1 [27, 28]. The remaining four QTLs (EWYP1B.2, EWYP1B.3, EWYP1B.4, and EWYP1B.5) were likely novel resistance loci identified in the current study. Four of the 36 MTAs found on this chromosome have previously been associated to other wheat diseases: *snp_BE442716B-Ta_2_1* and *wsnp_Ex_rep_c69266_68192766* with stripe rust [47], *wsnp_Ex_c38116_45719983* with Fusarium head

blight [48], and *BS00070139_51* with crown rot resistance [49].

On chromosomes 3A, 3B, 4A, 4B, and 5B, the additional six QTLs containing 10 MTAs were discovered. The marker *Tdurum_contig777_260* (IWB73429) designated as EWYP3A QTL was adjacent to the all-stage resistance gene *Sr27* which is transferred from *Secale cereale* and *Sr35* gene which is transferred from *Triticum monococcum* [32]. Because EWYP3A is so close to the *Sr27* gene area, *Sr27* is most likely the underlying gene for this region. On the short arm of chromosome 3B, *Sr2* came from *Triticum dicoccum* and *Sr12* originated from *Triticum turgidum* ssp. were cataloged previously [32]. On this chromosomal, we discovered the EWYP3B.1 QTL, which consists of three markers (*Tdurum_contig12899_342*, *Excalibur_c20277_483*, and *Tdurum_contig12008_803*). The nearest *Sr* gene to these markers was the *Sr2* gene [30, 31]. This *Sr2* gene has been extensively used in breeding as a source of durable and broad-spectrum adult plant resistance. Individual genotypes carrying the favorable allele of these SNP markers have shown an apparent difference in the CI score (Additional file 1). On chromosome 5B, the SNP marker *RAC875_c30011_426* (IWB56412) explained 5.7% of the total phenotypic variation. This marker is found near the chromosome region previously discovered the *Sr56* gene that confers the APR to wheat stem rust [32, 33]. To the best of our knowledge, the other three QTLs, EWYP3B.2, EWYP4A, and EWYP4B, which were found on chromosomes 3B, 4A, and 4B, respectively, have never been reported and could potentially be novel QTL sources for stem rust resistance breeding programs.

Conclusions

This study characterized the genetic diversity of elite ICARDA breeding lines and performed GWAS based on the evaluation of field stem rust. As a result, substantial genetic variability and field disease response to *Pgt* was observed among the lines. The study detected several potentially novel loci associated with *Pgt* resistance. These markers could provide useful genetic information to unlock the genetic basis of resistance to *Pgt* in wheat. Furthermore, the result will accelerate the introgression of identified resistance QTLs in the wheat breeding program through marker-assisted introgression. The identified resistant lines could also be used as crossing parents in stem rust-resistant breeding programs.

(See figure on next page.)

Fig. 5 Scatterplots showing genome-wide linkage disequilibrium (LD) decays based on 15 K genotyping results in 245 wheat breeding lines. R^2 as a function of genetic distance (cM) between pairs of SNP markers estimated for A, B, and D sub-genomes. (A) LD for A sub-genome; (B) LD of B sub-genome; (C) LD of D sub-genomes. The LOESS representing the decay of R^2 along genetic distance is illustrated for each genome. LD critical threshold estimated from LD distribution of pairs of unlinked SNP markers is indicated by the dashed horizontal red line

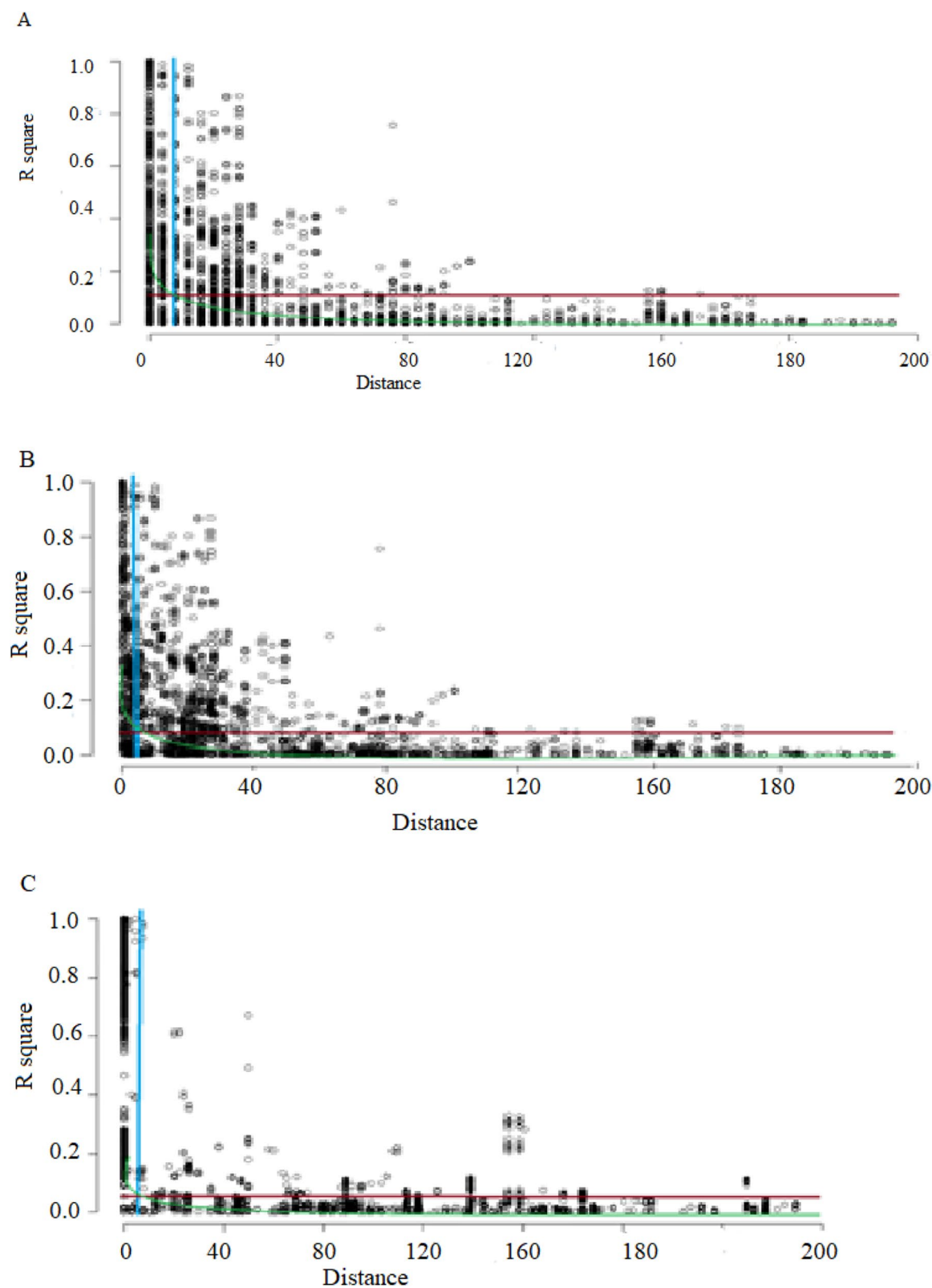


Fig. 5 (See legend on previous page.)

Table 3 Lists of QTLs identified for adult plant resistance (APR) to wheat stem rust

QTLs	SNPs	Markers Name	Alleles	Chr ^a	Pos ^b	Pos ^c	MAF	−log ₁₀ P	R ²	FDR ^d	Sr genes	References
EWP1B.1	IWA6489.1	wspn_Ku_c13229_21142792	T/C	1B	30.34	unknown	0.213439	4.398146	7.14	0.0192	Sr31	Mettin et al., 1973 [27]; Zeller, 1973 [28]
	IWB29508	Excalibur_c95327_51	A/G	1B	43.86	9.37986	0.418972	3.000182	5.91	0.0265		
EWP1B.2	IWB29475	Excalibur_c94756_540	T/C	1B	57.6	unknown	0.41502	3.398788	6.63	0.0163		new
	IWB1569	BobWhite_c22266_315	C/T	1B	60.62	329.48968	0.213439	4.17831	8.16	0.0050		
EWP1B.3	IWB43258	Kukri_c26168_423	C/A	1B	60.62	1.417758	0.462451	3.07204	6.06	0.0171		new
	IWB23446	Excalibur_c20228_135	A/G	1B	64.1	305.2701	0.094862	3.717446	7.27	0.0130		
IWB54643	RAC875_c18282_1390		T/C	1B	64.1	unknown	0.090909	3.676459	7.15	0.0171		
	wspn_Ex_rep_c69266_68192766		T/C	1B	64.1	299.97785	0.094862	3.668026	7.13	0.0130		
IWB6405	BS00011450_51		T/C	1B	64.1	318.23318	0.13834	3.473183	6.74	0.0080		
IWA131	wspn_BE443531B_Ta_1_1		C/T	1B	64.32	156.68751	0.114625	3.490435	6.77	0.0231		
IWB48689	Kukri_rep_c101799_95		C/A	1B	64.46	unknown	0.114625	3.490435	6.77	0.0231		
IWA3631	wspn_Ex_c38116_45719983		C/T	1B	64.89	340.37296	0.079051	3.957779	7.72	0.0775		
IWB51549	Ra_c23839_884		T/C	1B	65.42	367.44011	0.146245	3.61409	7.02	0.0080		
IWB31732	GENE-0193_197		A/G	1B	65.42	367.44095	0.146245	3.471958	6.74	0.0080		
IWA6890	wspn_Ku_c30982_40765254		T/G	1B	66.07	336.20988	0.075099	4.081555	7.97	0.1850		
IWB58051	RAC875_c44575_561		C/T	1B	66.07	325.79901	0.075099	3.973875	8.40	0.0266		
IWB461	BobWhite_c1318_691		T/C	1B	66.73	387.64464	0.150198	3.618488	7.03	0.0080		
IWB37720	JD_c64600_281		G/A	1B	67.14	409.62805	0.893281	3.718149	7.48	0.0208		
IWA106	wspn_BE442716B_Ta_2_1		T/G	1B	67.14	403.156221	0.102767	3.67975	7.15	0.0208		
IWB38394	Ku_c13515_171		C/T	1B	67.14	unknown	0.102767	3.67975	7.15	0.0208		
IWB35871	IACX2701		C/T	1B	67.38	413.08886	0.098814	3.795799	7.43	0.0208		
IWB10444	BS00070139_51		A/C	1B	68.04	418.16246	0.150198	3.618055	7.03	0.0080		
IWB27852	Excalibur_c59016_839		A/G	1B	68.04	426.07288	0.150198	3.618055	7.03	0.0080		
IWB56778	RAC875_c32894_1038		C/T	1B	68.04	unknown	0.150198	3.354313	8.83	0.0080		
IWB59327	RAC875_c5796_424		A/G	1B	68.04	IWB59327	0.150198	3.618055	7.03	0.0080		
IWB8148	BS00038929_51		T/C	1B	68.04	417.85674	0.150198	3.618055	7.03	0.0080		
EWP1B.4	IWB74145	tbl0023b14_704	G:T	1B	70.08	438.2896	0.339921	3.35714	6.59	0.0245		new
	IWB47566	Kukri_c73734_175	C/T	1B	76.89	532.565453	0.213439	4.268153	8.37	0.0282		
IWB60433	RAC875_c7674_634		G/A	1B	76.89	532.56545	0.27668	3.545613	7.40	0.0282		
IWB6504	BS00011973_51		T/G	1B	76.89	531.855670	0.280632	3.485333	6.78	0.0282		
IWA775	wspn_CAP11_c543_375403		A/G	1B	76.89	542.43163	0.217391	3.044683	5.89	0.0282		
IWA5228	wspn_Ex_rep_c66389_64588992		A/G	1B	79.77	561.70458	0.110672	3.731492	7.28	0.0080		
IWB74900	tbl0048b10_1365		A/G	1B	79.77	552.5332	0.110672	3.731492	7.28	0.0080		

Table 3 (continued)

QTLs	SNPs	Markers Name	Alleles	Chr ^a	Pos ^b	Pos ^c	MAF	−log ₁₀ P	R ²	FDR ^d	Sr genes	References
EWYP1B.5	IWB10621	BS00072791_51	A/G	1B	105.83	629.26299	0.27668	3.778586	7.61	0.0080		new
	IWB70380	Tdurum_contig32775_78	A/G	1B	112.07	unknown	0.335968	3.732007	7.26	0.0080		
	IWB66198	Tdurum_contig10036_977	C/A	1B	114.13	634.6536	0.094862	3.49178	6.81	0.0080		
EWYP3A	IWB73429	Tdurum_contig777_260	G/A	3A	20.74	102.20749	0.339921	3.609065	7.12	0.0080	Sr27	McIntosh et al, 1995 [29]
EWYP3B.1	IWB67769	Tdurum_contig12899_342	T/C	3B	9.7	5.585837	0.379447	3.484232	6.90	0.0210	Sr2	Ausemus et al, 1946 [30]; Knott, 1968 [31]
	IWB23457	Excalibur_c20277_483	A/G	3B	9.7	5.58572	0.371542	3.042077	5.88	0.0210		
	IWB67389	Tdurum_contig12008_803	T/C	3B	9.7	5.584656	0.395257	3.016518	5.83	0.0210		
EWYP3B.2	IWB75222	tplb0059m03_622	C/T	3B	20.14	7.373615	0.434783	3.022107	5.95	0.0208		new
EWYP4A	IWB72664	Tdurum_contig59603_74	G/A	4A	26.5	9.92939	0.150198	3.240453	6.40	0.260		new
EWYP4B	IWB24798	Excalibur_c29127_552	G/A	4B	64.58	232.88033	0.094862	3.73037	7.26	0.0208		new
	IWB48189	Kukri_c8973_1986	T/C	4B	62.22	Unknown	0.098814	3.126412	4.76	0.073		
	IWB53758	RAC875_c13639_2159	T/C	4B	62.92	272.17459	0.098814	3.126412	4.76	0.0190		
EWYP5B	IWB56412	RAC875_c30011_426	C/T	5B	104.55	571.47521	7.08548	2.9914	5.77	0.0210	Sr56	Park 2016 [32]; Yu et al, 2014 b[33]

QTLs Quantitative trait loci. SNPs Single nucleotide polymorphism. Chr^a Chromosome position, Pos^b Marker's genetic position mapped in the wheat 90KSNP consensus map [34] in centimorgans (cM); Pos^c, marker's physical position produced by the International Wheat Genome Sequencing Consortium (IWGSC RefSeq v1.0) [35] in megabase pairs (Mbp); FDR^d, The false-discovery rate adjusted P-values; (MAF), minor allele frequency; R², phenotypic variance explained by the markers



Broad-sense heritability (H^2) was calculated using the following formula:

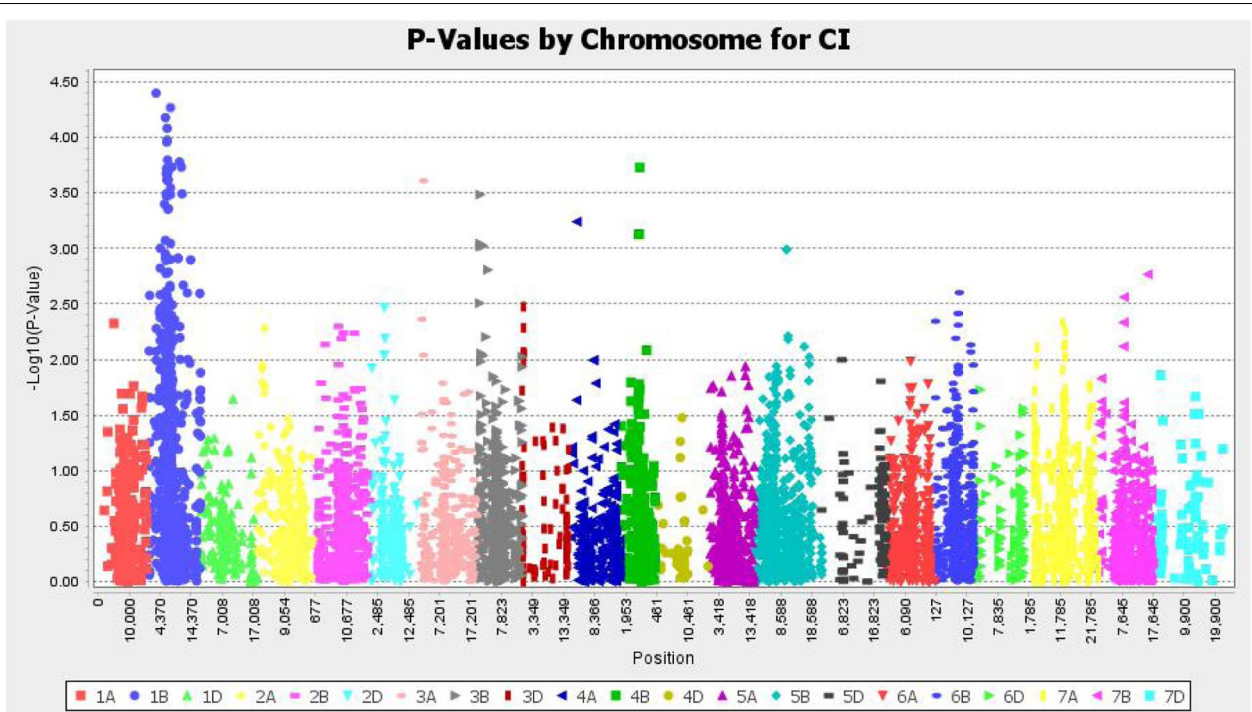


Fig. 7 GWAS results of the Manhattan plot along with the 21 chromosomes showing significantly associated markers with adult plant stem rust resistance. The position of each marker was based on the wheat consensus SNP map [34]

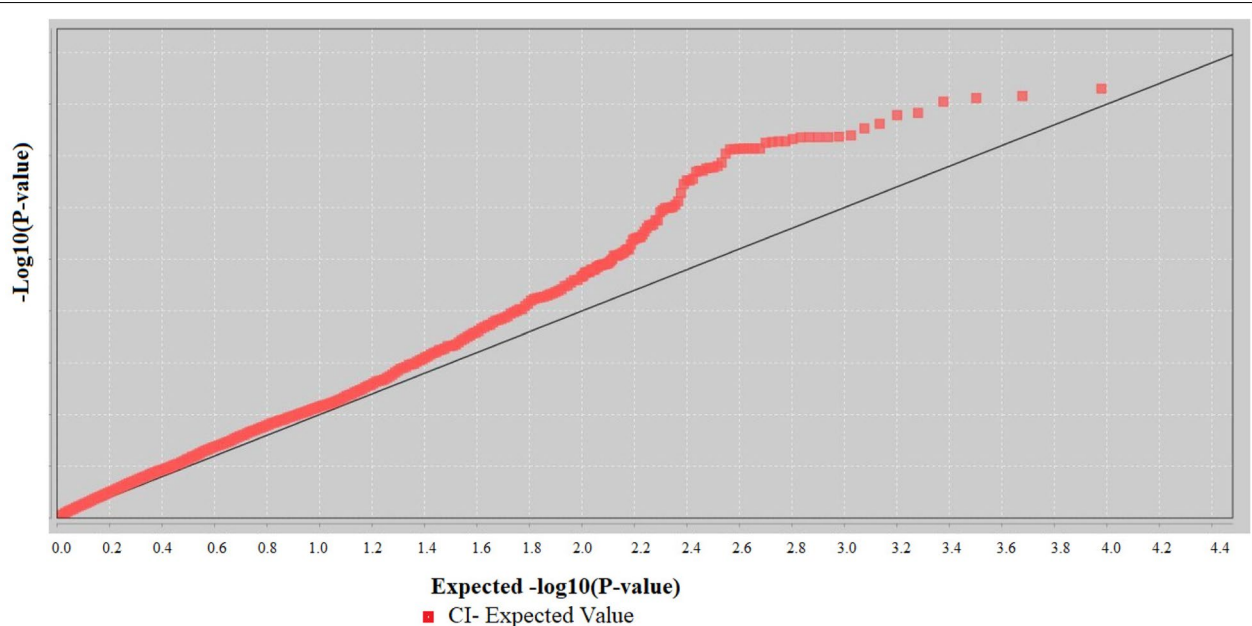


Fig. 8 Q-Q plot for stem rust resistance in a panel of 245 wheat breeding lines using the MLM model. The plots show the observed p -values (p) for the association between CI and each tested marker expressed as $-\log_{10}(P\text{-value})$ of p (y -axis) plotted against $-\log_{10} P$ of the expected p -values (x -axis) under the null hypothesis of no association for the analyses

$$H^2 = \frac{\sigma^2_G}{\sigma^2_G + (\sigma^2_{G \times E})/n + (\sigma^2_{error})/n}$$

Where σ^2_G is the genotypic variance, σ^2_E is the environmental variance, $\sigma^2_{G \times E}$ is the genotype by environment interaction variance, σ^2_{error} is the residual error variance and n is the number of years.

To reduce false-positive associations, best linear unbiased predictors (BLUPs) for CI were calculated using a mixed model in *lme4* package implemented in R environment [51] according to the following model where y is the response variable:

$$y = \text{lmer}(\text{Trait} \sim (1|\text{Genotype}) + (1|\text{Year}))$$

Population structure and genetic diversity

The optimal sub-populations of the panel were estimated based on three different approaches. The Bayesian model-based population structure was estimated from 100 unlinked SNP markers located at least 10 cM apart across the genome using STRUCTURE 2.3.1 software [52, 53]. To execute this, three independent runs were performed for each hypothetical K value run from 2 to 15 with the length of the burn-in period of 10,000 steps followed by 100,000 Monte Carlo Markov Chain (MCMC). The results obtained from this procedure were used in a web-based informatics tool namely, "Structure Harvester" [54] to define the optimal K value, based on ΔK method Evanno, 2005 [55]. Each genotype was assigned to one subpopulation based on its membership probability. The second approach used to determine the optimal subpopulation was based on a marker-based kinship matrix (K matrix) on a scaled identity-by-state method using the whole set of SNP markers from TASSEL 5 software [56]. Finally, the principal components analysis (PCA) of genetic relatedness was performed with the same software and added to the regression model as a covariant.

Genetic diversity was estimated based on polymorphic information content (PIC), heterozygosity, and Nei's gene diversity using the whole set of SNP markers from PowerMarker 3.25 software [57]. Phylogenetic analysis based on distance-based neighbor-joining method was calculated with TASSEL 5 software and visualized through web-based program iTOL (v 4.3.2) [58].

Genotyping, linkage disequilibrium, and genome-wide association analysis

DNA extraction of lines was carried out on one-week-old seedlings following the protocol described by Allen et al. (2006) [59] using Cetyltrimethylammonium bromide

(CTAB). Genotyping was performed by Illumina iSelect 15K single nucleotide polymorphism (SNP) wheat array and called by GenomeStudio V2011.1 software. The resulting 13,006 SNPs were further screened using those only minor allelic frequency (MAF) > 5%, and missing data percentage of < 10%. Five lines were excluded as a result of this screening. Finally, 9523 quality SNP markers were generated from 245 lines that were used for further analysis.

The resulting SNP data were subjected to linkage disequilibrium (LD) analysis as squared allelic frequency correlations (R^2) between each pair implemented in TASSEL v5.2 and GAPIT (Genomic Association and Prediction Integrated Tool) R package [60]. The critical R^2 value (where the LD is due to the physical linkage) was determined by taking the 95% of R^2 data of unlinked markers as the threshold, according to Brescaghello and Sorrells (2006) [61].

Marker-trait association analysis (MTAs) between the BLUP value of CI and SNPs markers were analyzed using a mixed linear model (MLM) in TASSEL 5.2 software. Using the formula: $y = X\alpha + Q\delta + K\mu + e$; where y = phenotypic values, X is SNP marker genotypes, α is a vector containing fixed effects as a result of the genotype, Q is population structure as PCA, δ is a vector containing fixed effects resulting from population structure, K is the relative kinship matrix, μ is a vector of random additive genetic effects and e is a vector of residuals. Marker trait associations were declared significant at a threshold value of $-\log_{10}(p) \geq 3$ (corresponding p value ≤ 0.001) [62].

Abbreviations

ANOVA: Analysis of variance; BLUP: Best linear unbiased prediction; CI: Coefficient of infection; DARC: Debre Zeit Agricultural Research Center; GWAS: Genome-wide association study; DS: Disease severity; ICARDA: International Center for Agricultural Research in the Dry Areas; IR: Infection response; LD: Linkage disequilibrium; MAF: Minor allele frequency; MAS: Marker-assisted selection; MCMC: Markov chain Monte Carlo; MLM: Mixed Linear Model; MTA: Marker-trait association; PC: Principal components; PCA: Principal component analysis; *Pgt*: Wheat stem rust fungus *Puccinia graminis*; PIC: Polymorphism information content; PV: Phenotypic variance; QQ: Quantile-quantile; QTL: Quantitative trait locus; SNP: Single nucleotide polymorphism.

Supplementary Information

The online version contains supplementary material available at <https://doi.org/10.1186/s12863-022-01030-4>.

Additional file 1: List of pedigree stem rust response and associated SNPs for wheat stem rust.

Acknowledgments

This project is a part of ICARDA's shuttle breeding program which has been designed for performing stem rust resistance screening. Authors acknowledge Debre Zeit Agricultural Research Center for facilitating field experiments.

Authors' contributions

ES, WT, and EB conceived and designed the study. WT designed the study and provided the germplasm and the genotypic data. EB guided the project development. ES wrote this manuscript. ES and AA performed data analysis. LM edited the manuscript. All authors read and approved the final manuscript.

Funding

This research did not receive any specific grant from funding agencies in the public, commercial, or not-for-profit sector.

Availability of data and materials

The datasets generated and/or analyzed during the current study are available in the figshare data repository, <https://doi.org/10.6084/m9.figshare.17711150.v3>.

Declarations

Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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Received: 22 December 2020 Accepted: 31 January 2022

Published online: 14 February 2022

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