Genome-wide association study and Genomic Prediction of spot blotch disease in wheat (Triticum aestivum L.) using genotyping by sequencing

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Abstract

Background Spot blotch caused by Bipolaris sorokiniana is a major constraint in wheat production in tropics and subtropics. There is limited information available on GWAS and study on genomic prediction is completely lacking. To reveal the genetic markers associated with disease resistance, we performed a genome-wide association study (GWAS) for spot blotch disease in 141 spring wheat lines.

Results Based on the testing under natural infection in three years at hot spots location in Pusa, India and Jamalpur, Bangladesh, the genotypes showed significant genetic variation for disease severity. Using Genotyping-by-Sequencing (GBS) based 18637 polymorphic SNP markers and phenotyping from diverse environments, we identified 23 genomic regions across the genome ( p < 0.001) on 14 chromosomes associated with disease scores. Consistent with the previous reports, a most stable genomic region on chromosome 2B, 5B and 7D were detected across the environments. The new genomic region on chromosome 3D was also identified. We performed functional annotation with wheat genome assembly annotation (IWGSC Ref Seq v1.0) and identified NBS-LRR and 35 other plant defense-related protein families across multiple chromosome regions. Using a five-fold cross-validation scheme, we observed moderate prediction accuracy for 3 of 4 environments indicated that our model was able to successfully capture the quantitative variation underlying the SB variation in our population.

Conclusions The GWAS based on the phenotypic data from PUSA India and BARI Bangladesh resulted in a total of 23 genomic regions on 14 chromosomes. The new genomic region appeared on chromosome 3D associated with Zinc finger protein that play important role in plant disease resistance. The genomic prediction model for spot blotch disease resistance in wheat was tested and obtained moderate prediction accuracy.

Background

Wheat (Triticum aestivum L.) is the major staple for more than 35% of the world’s population [1]. The pace of wheat improvement must accelerate to meet the projected global food demand by 2050. Green revolution played a key role in India, Pakistan, Nepal and Bangladesh to ensuring food security in this densely populated region of the world [2]. However, wheat production faces multiple threats
via rapidly evolving pathogen variants, pests and increased climate variability, which considerably jeopardize crop productivity growth [3, 4, 5, 6]. Breeding wheat for climatic resilience and disease resistance combined with good agronomy can potentially improve wheat productivity to meet the future food demands [7].

Spot blotch caused by Bipolaris sorokiniana is a major constraint in wheat production in tropics and subtropics [8, 9]. The pathogen has a worldwide dispersal, but it is predominantly aggressive under conditions of warm, high relative humidity and temperature associated with imbalanced soil fertility. Yield losses are variable but are important in fields with low inputs and under late-sown conditions [2]. Bipolaris sorokiniana act as a causal agent for numerous diseases like head blight, seedling blight, foliar blight/spot blotch, common root rot and black point of wheat, barley, other small cereal grains and grasses [10]. However, spot blotch of wheat is considered as one of the most important diseases caused by this pathogen in the mega environments characterized by high temperature (coolest month greater than 17°C) and high humidity [11].

When desired level of resistance to several diseases is required, it is often difficult to achieve through conventional breeding approaches. Disease resistance can be inherited both qualitatively and quantitatively, as is the case in many wheat diseases [12, 13, 14, 15]. The genetic basis of spot blotch resistance has earlier been recognized to eight major quantitative trait loci (QTLs), namely QSB.bhu-2A, QSB.bhu-2B, QSB.bhu-2D, QSB.bhu-3B, QSB.bhu-5B, QSB.bhu-6D, QSB.bhu-7B and QSB.bhu-7D [12, 13]. Sharma et al (2007b) [16] reported three microsatellite markers (Xgwm67, Xgwm570 and Xgwm469) linked with spot blotch resistance. Lr34 and Lr46, the two broadly used genes conferring leaf rust resistance have also been reported to contribute to spot blotch resistance [17]. Lr34 gene is reported to be the major locus on chromosome 7D and explains up to 55% phenotypic variation for spot blotch disease resistance and this locus was given the gene designation Sb1 [17]. During past few years, several QTLs and genetic markers for spot blotch resistance have been identified in multiple studies in winter Wheat [18, 19, 20, 21].

Due to changes in pathogen populations, resistance genes can lose their effectiveness over time. Given these challenges, identification and mapping of novel resistance genes would aid breeding for
disease resistance in wheat. One approach to identify spot blotch resistance QTLs is through association mapping. This approach leverages historic recombination and generally have better mapping resolution compared to biparental mapping [22]. A promising strategy to identify QTLs for traits of interest is genome wide association study (GWAS) that takes advantage of decreasing sequencing cost and high throughput genotyping assays [23]. A key approach in GWAS is to have enough genome coverage so that functional alleles will be in linkage disequilibrium (LD) with at least one marker [24]. The association studies for disease resistance including spot blotch have been conducted [25, 26, 27, 28, 29, 30].

Limited research was done where same set is exposed to diverse environments in large geographic area for wheat spot blotch. Therefore, the primary objective of this study was to establish marker-trait associations for spot blotch using genotyping by sequencing (GBS) SNP markers in spring wheat in the South Asian wheat growing regions. The significant SNPs can give insights into the biological function and its relationship with resistance more relevant to the South Asian region. This study aims not only to identify novel QTLs but to validate known genomic loci conferring spot blotch resistance. So far, there is no study on genomic predictions for spot blotch disease resistance in wheat, therefore, we made first attempt to test genomic prediction models across environments.

Results
Spot blotch disease was recorded visually on a scale of zero (no symptoms visible) to 100 (completely susceptible) across three years (2016-17, 2017-18 and 2018-19). The populations displayed significant phenotypic variation for spot blotch resistance with nearly continuous distribution of lines in all environments (Fig 1). The mean disease severity ranged from 8.90 to 31.23 in four environments (2016-17 to 2018-19) including Pusa, India and Bangladesh Agricultural Research Institute (BARI), Bangladesh (Table 1). Highest mean disease severity was recorded in BARI 2016-17 (Env4) while lowest was at PUSA 2017-18 (Env2). The analysis of variance revealed highest heritability in Env 4 (0.80) while lowest was in Env 3 (0.50). Based on the combined analysis of all environments, we observed moderate heritability (0.55). There was significant Genotype × Environment interactions (P <0.0001; Table 2).
Genetic linkage mapping

The genetic linkage map was prepared using the most significant SNPs found on 14 hexaploid wheat chromosomes. A total of 70 SNPs were used and clustered in 23 linkage groups (Fig 2). A linkage group was considered to be different if the gap between them is more than 10cM on the same chromosome. We observed a maximum of three linkage groups on chromosome 2B and 5B. Similarly, two linkage groups were found on each of 2A, 5A, 7A, 7B, and 7D. Maximum number of significant SNPs (18 SNP markers) were observed on chromosome 2B (Table 3).

Principal component analysis

Population structure was determined using genotyping information and principal component analysis (PCA) based approach where genotypes clustered in 12 subgroups. The clustering was done using Ward method in JMP v.13 (Fig 3). The Group 1 (G-I) consisted of eight lines including the resistant check HD2733. Maximum number of lines included in G-VII (24 lines) while minimum in G-X and G-XII (6 lines in each). Most of the lines in a group shared alleles descended from common parents. The lines without common parents or less than three sibs per family were classified as ‘others’. The largest group (G-VII) consists lines with mixed pedigrees including SAUAL, WBLL#1, Kachu #1, BAV92//IRENA/KAUZ, FRANCOLIN#1, MUCUY and PBW343. The parental lines with TRCH/SRTU//KACHU cross in their genetic backgrounds dominated G-VIII. The parental line AKURI was a most common parent in G-IX and similar in case of G-XII where sister lines dominated the group. The intra-chromosomal LD was calculated as the pairwise marker correlations ($R^2$) between the GBS markers and plotted against the physical distance for significant marker-trait associations (Fig S1).

Marker -Trait associations

The GWAS of spot blotch resistance was performed based on the data collected at adult plant stages. After false discovery rate, a q-value (corrected p value) <0.05 was used as a threshold to identify significantly associated markers. The GWAS results of spot blotch resistance from the trials conducted at Borlaug Institute for South Asia (BISA) in Pusa, India and BARI in Jamalpur, Bangladesh are given in supplementary tables (Table S1). A total of 70 most significant SNPs markers associated with spot blotch disease resistance belong to 23 linkage groups were detected on chromosomes 1A, 1B, 1D, 2A,
2B, 3A, 3B, 3D, 5A, 5B, 6A, 7A, 7B and 7D (Fig 2). The phenotypic variation explained by the most significant chromosome region ranged from 6.75% (Env2) to 13.65% (Env1) in three years trials. The phenotypic variation explained in individual environments ranged from 7.37-13.65%, 6.75-9.39%, 7.51-12.72% and 7.60-11.68% in Env1, Env2, Env3 and Env4 respectively (Table S1). The largest phenotypic variation explained by the SNP marker located on chromosome 5B in Env1 followed by 2B in Env3 explaining up to 13.65% and 12.72% of phenotypic variation respectively. Most of the SNPs marker regions appeared in more than one environment. We detected seven significant chromosome regions on chromosome 2B while six were on chromosome 5B. A total of 15 SNP markers (S1B_646895451, S2A_31851904, S2B_504717, S2B_525073, S2B_594959, S2B_6253562, S2B_8311062, S2B_90662917, S3B_763230831, S3B_763236179, S3B_763267753, S3B_764192662, S4A_710830493, S6B_719904092 and S7D_181974079) were significant in at least two environments (Table S1).

**Gene functional annotations**

The significant SNPs identified from the GWAS analysis were further studied for the known candidate genes relevant to disease resistance using the recently annotated wheat reference sequence (RefSeq V1.0). We identified NBS-LRR and 35 other plant defense-related protein families across multiple chromosome regions. The significant SNPs S2B_13751999 identified in Env1 on chromosome 2B was located between the GenelDs, TraesCS2B01G030100 and TraesCS2B01G030200. The latter gene play an important role in the resistance of various plant diseases including the downy mildew [31] by producing RPP13 protein while the former gene is involved in synthesis of lectin-receptor kinase which has an important plant immunity function [32]. Similarly, the SNP S2B_699219601 identified in Env4 belongs to the GenID, TraesCS2B01G505200 also involve in downy mildew disease resistance response and other diseases (Table 3). The SNP S2B_8311062 identified in Env1 and Env3 was located close the geneID, TraesCS2B01G018200, which is involved in **NBS-LRR disease resistance protein** synthesis. Similarly, the SNP S2B_28592818 identified in Env4 located close the geneID, TraesCS2B01G058900 also involved in synthesis NBS-LRR disease resistance protein. The annotation of S5B_683352145 revealed that the gene on chromosome 5B identified in Env4 also involved in
synthesis of NBS-LRR disease resistance protein family-1. The SNP S2B_6253562 falls within the GeneID, TraesCS2B01G012400 that encodes Avr9/Cf-9 rapidly elicited protein. The Avr9/Cf-9 protein is involved in early signaling events in the Avr9/Cf-9-dependent plant defense response. The most important SNP S5A_595393566 detected in Env1 on chromosome 5A belongs to the gene TraesCS5A01G402800 which mediates spot blotch (Bipolaris sorokiniana) resistance in wheat (Table 3). The other SNPs S7B_1020705 and S3B_763230831 associated with Leucine-rich repeat receptor-like protein kinase and transmembrane protein contribute to Fusarium resistance in cereals. The SNP (S5B_233586644; geneID TraesCS5B01G128000) explaining highest phenotypic variance (13.65%) was detected in Env1, produces signal recognition particle subunit SRP68 which play a crucial role in targeting secretory proteins to the rough endoplasmic reticulum membrane. The second highest phenotypic variance explained by the SNP (S2B_504717) located on chromosome 2B involved in DON resistance through cytochrome 450. The SNP on 1A, 1B, 1D, 2A, 2B, 3A, 3B, 4B, 5A, 5B, 6A and 7B found to be involved directly in disease resistance mechanism (Table 3).

To further confirm the results, we looked at the common proteins through gene annotation associated with different SNPs, identified independently in different environments. For example, Zinc finger family protein were associated with the SNP on identified on 2B (S2B_15129248) and 3B (S3B_764747435) in Env1, 3D (S3D_610628298) in Env2 and 3A in Env3 (S3A_67348475). Similarly, the kinase protein family were associated with the SNPs on chromosome 2B (S2B_13761590), 2A (S2A_708482943) and 5B (S5B_683514734) in Env4. The results based on the gene annotation and synthesized proteins are presented in supplementary Table S2 and the Vann diagram as Fig 4.

**Evaluation of Prediction Accuracy**

For genomic prediction modeling, the 141 lines were randomly divided into training and testing sets of size 4/5 and 1/5, respectively for each of the four environments. In the initial model training step, both genotypes and the observed phenotypes (i.e., phenotypic BLUPs) were used to estimate the genetic marker effects from the training population. The estimated marker effects were subsequently used to predict the phenotypes of the testing set population. This process was repeated 100 times to sample a random set of training and testing set population during each iteration. The average
correlation between the observed and the predicted phenotypes was calculated. The prediction accuracy distributions, means, and standard errors were reported by each environment. The cross-validation prediction accuracy of the Env4 was 0.33 while for the Env1 and Env2 had a prediction accuracy of 0.29 and 0.24 respectively (Fig 5). In contrast to other environments, the prediction accuracy of the Env3 was negative (-0.16).

**Discussion**

The field trials were conducted at BISA research farm, Pusa, in India for three consecutive years from 2016-17 to 2018-19 and BARI farm, Jamalpur in Bangladesh during 2016-17 crop season. Both the locations fall under the non-traditional, warmer wheat-growing regions belonging to Mega-environment 5 characterized by hot, humid conditions as per CIMMYT’s system for classifying wheat-growing environments in developing countries [11]. The average temperature during the wheat plant reproductive phase at Jamalpur and Pusa is higher than 19°C with a high relative humidity [33] (Table S3).

The spot blotch disease incidence was captured as percentage of infected leaf area at three different growth stages to minimize the chances of disease escape due to environmental factors. However, the scoring date showing highest disease pressure (usually the second one) was used in the analysis. Since the susceptible parent displayed highest disease severity at growth stage 77 (GS77) on Zadoks scale [34], to make better judgment about the level of resistance, disease severity was recorded at this stage (usually second scoring) was used to differentiate each line.

The nearly continuous distribution of lines in all the environments show quantitative nature of resistance. The same has been supported by earlier findings where more than two genes [35, 12, 13, 36, 37] and multiple alleles with minor effect [30, 21] to control spot blotch resistance is reported. It was found that the log transformation improved the data normality which was also reflected by the improved consistency in the GWAS results across locations. We observed significant genetic variation for disease susceptibility in the population. The genetic variances and moderate to high heritabilities for spot blotch were comparable with earlier findings in wheat [37, 38, 36]. Despite significant genotype x environments interactions, we observed moderate to high heritabilities within
environments (Table 2). The environmental interactions might ascribe to difference in the pathogen isolates prevalent in NEPZ of India and Bangladesh in case of locations and weather conditions mainly within location. For example, the maximum mean disease severity of the susceptible lines were up to 43% in Env3 while it was 70% in Env1 (Table 1).

The linkage analysis was based on 18637 filtered SNP markers covering all chromosomes. The redundant SNPs with 0 cM distance and with same gene annotation were removed from the linkage mapping as no additional information is expected. After GWAS analysis, 14 chromosomes harboring significant QTL regions forming 23 linkage groups were used for further analysis and graphical representations. The SNP lies more than 10 cm apart based on linkage mapping, were placed in a separate linkage group. (Fig. 2).

We used genotyping information for the PCA where most of the groups were based on the proportion of genome shared by the parental pool except few exceptions. For example, the subgroup (G-VIII) consisted common parent TRCH/SRTU//KACH while the largest group (G-VII) consists lines with mixed pedigrees dominated by SAUAL, WBLL#1, Kachu #1, BAV92//IRENA/KAUZ, FRANCOLIN#1, MUCUY and PBW343.

Several spot blotch resistance QTLs have been reported on different chromosomes [39, 16, 40, 12, 13, 41, 42, 17, 43, 20, 18, 44, 45]. However, only three major QTLs designated as Sb1 on 7D [17], Sb2 on 5B [41], and Sb3 on 3B [44] are well described. We also observed consistent chromosome regions on 2B and 5B, appeared in more than 2 or all the environments (Table 3). The QTL on 5B, named as Sb2 gene earlier have been studies in detail [44]. The Sb2 gene is known to interact with Tsn1 gene, conferring susceptible reaction to tan spot and Septoria nodorum blotch [46]. The gene ToxA virulent to Tsn1 exists in both Pyrenophora tritici-repentis and Parastagonospora nodorum confer susceptible reaction to tan spot and Septoria nodorum blotch respectively [21]. Friesen et al. 2018 [47] demonstrated major effects of the Tsn1 locus on chromosome 5B. However, the importance of Tsn1 in spot blotch disease resistance under field condition is not known. The QTL on 7D was the first one studied in detail and reported to be associated with Lr46 [17], Lr34 and leaf tip necrosis [36]. Based on the fine mapping studies, it was named as Sb2 gene [41]. It is interesting to note that Ayana et al.
2018 [30] identified six potential QTLs (QSb.sdsu-2D.1, QSb.sdsu-3A.1, QSb.sdsu- 4A.1, QSb.sdsu- 4B.1, QSb.sdsu-5A.1, QSb.sdsu-7B.1) in hard winter wheat using the isolate, SD40 in greenhouse conditions. The chromosome regions on 4A (Env2 and Env4) and 4B (Env4) and 7B (Env1 and Env3) were consistent with the results of [30]. Similarly, four chromosome regions on 1B, 3B, 4B and 5B are validating the finding of [21] which were based on testing in the field condition.

Regardless of % phenotypic variance explained by an allele, almost all wheat chromosomes except 3D and 5D reported to have contributed for spot blotch disease resistance depending on, spot blotch isolate, the breeding material or parents in case of bi-parental population [16, 12, 13, 42, 17, 20, 18, 19, 30, 21]. The minor QTL were reported on, 1BS, 1D, 2D and 3A, 4DS and 6D contributed by ‘CIANO T79’, ‘WUYA’ and ‘BARTAI’ [21]. The broad range of environmental conditions at our field sites allowed us to capture considerable genetic variation underlying spot blotch resistance. We identified 23 QTL regions on 14 chromosomes validating previous results. The new genomic region detected on chromosome 3D associated with the SNP marker S3D_610628298 explained up to 6.94% of the phenotypic variance but detected in Env2 only. Similarly, the SNP (S2B_90662917) on chromosome 2B was most significant, explained only up to 10% of phenotypic variance, while the SNP on 5B explained largest phenotypic variance in Env1 (Table S1). Out of 23, 9 chromosome regions on seven chromosomes (1A, 1B, 2A, 5A, 5B, 7B, 7D) were already mapped in independent studies earlier [16, 13, 42, 17, 20, 18, 19, 30, 21].

So far, based on the consistency in independent QTL mapping studies using different source of resistance, it seems that there is not much genetic variability in spot blotch pathogen across the continent. However, several studies described clear grouping among spot blotch isolates based on the fungal hyphae color, aggressiveness and DNA fingerprinting [48, 49, 50, 51]. Four chromosomal regions on 1B, 2B, 4A and 6B are consistent between Pusa India and Jamalpur Bangladesh. This may be due to prevalence of most aggressive isolate of spot blotch pathogen (isolate No. ICMP 13584, Auckland, New Zealand) common in South Asia [52].

To study the importance of significant SNPs in disease resistance, we annotated all SNPs using wheat genome assembly annotation (IWGSC Ref Seq v1.0) and traced the protein synthesized by the
annotated gene. The literature was mined to look for the putative functions of those proteins. We found that several genes functional annotation strongly associated with disease resistance and observed across the year and environments (Table 3). For example, seven SNPs (S2B_13814702, S2B_533178164, S2B_14809954, S2B_14963432, S2B_15129248, S2B_504717, S2B_78065) on chromosome 2B associated with eight geneIDs, TraesCS2B01G030500, TraesCS2B01G373900, TraesCS2B01G031700, TraesCS2B01G031900, TraesCS2B01G032000, TraesCS2B01G032100, TraesCS2B01G001100 and TraesCS2B01G000400 involved in synthesis of Cytochromosome P450 family protein. The role of Cytochrome P450 family protein in plant defense, secondary metabolite biosynthesis in the classical xenobiotic detoxification pathway is well established by Thapa et al. 2018 [53]. It is involved in resistance to DON which is a trichothecene mycotoxin produced by Fusarium species and increase yield. The Cytochrome P450 family protein may not involve directly in yield increase but to enhanced Fusarium head blight disease resistance.

The SNP (S2B_28592818) detected in Env4 on same chromosome (2B) but at different region synthesizes NBS-LRR disease resistance protein family contribute for disease resistance [54, 55]. Similarly, the SNP S2B_8311062 and S5B_683352145 also associated with the gene synthesize NBS-LRR disease resistance protein family and contribute for fungal disease resistance. The role of NBS-LRR disease resistance protein is disease resistance mechanism is well established [54, 55]. One of the significant SNP located on chromosome 2B, S2B_15129248 is linked to two geneIDs, namely, TraesCS2B01G032100 (synthesize Cytochrome P450 family proteins) and TraesCS2B01G032200 (involved in GRF zinc finger family protein). Both the proteins play an important role in plant disease resistance [53, 56].

It is interesting to note that the most important SNP S5A_595393566 detected in Env1on chromosome 5A belongs to the gene TraesCS5A01G402800 which mediates spot blotch resistance in wheat. This gene is involved in the synthesis of Myb family transcription factor-like protein, found to mediate host resistance to Bipolaris sorokiniana in wheat [57]. The same region has been reported in other independent studies as well [18, 20, 30, 21]. Similarly, the SNP S3A_67065083 associated with geneID TraesCS3A01G103500 involved in synthesis of 1R-MYB Transcription factor which plays an important
role in disease resistance against stripe rust fungus and ear head disease in wheat [45].

The key SNPs on chromosome 1A (MAPK module FgSte50-Ste11-Ste7 in F. graminearum), 1B (stripe rust & powdery mildew), 1D (Serine/threonine-protein kinase), 2B (RPP13, Avr9/Cf-9 rapidly elicited protein, NBS-LRR protein, F-box family protein, pentatricopeptide repeat-containing protein, Peptidylprolyl isomerase, Uroporphyrinogen decarboxylase and resistance to DON), 3A and 3B (1R-MYB TF, wheat NAC protein and interaction with an orphan protein), 4B (Uroporphyrinogen decarboxylase), 5A (Myb family transcription factor-like, Serine/arginine repetitive matrix, NBS-LRR & transmembrane protein), 5B (B3 domain-containing, Mannitol transporter & NBS-LRR family-1 protein) and 7D (implicated in the defense through cell wall modification, degradation, carbohydrate metabolic processes) annotated and found to synthesize different proteins involved in fungal defense mechanism (Table 3) [58, 59, 60, 55, 61, 54, 62, 63, 64, 65, 32, 66, 67, 68, 69, 57, 70, 71]. The consistency in identification of key SNPs involved in resistance mechanism through protein annotation was confirmed where same protein family was identified independently in different environments (Fig. 4).

Maximum number of known proteins involved in fungal defense were based on 14 SNPs on chromosome 2B showing the importance of this chromosome in disease resistance. The earlier independent findings also describe the importance of chromosome 2B in spot blotch disease resistance [12, 19]. Interestingly QTL found in the present study on 1B in Env1, the proteins involved are Pentatricopeptide repeat-containing protein (TraesCS1B01G424000) mRNAs renders more susceptible to pathogenic bacteria and fungi in Arabidopsis thaliana [109] and Homeobox protein (TraesCS1B01G424100) associated with reaction to stripe rust and powdery mildew in common wheat [72]. The SNPs (S1B_646895451 and S1B_647195634) detected in two environments located on chromosome 1B are 18.97 cM apart on genetic linkage map while those belong to the same geneID TraesCS1B01G424000. The gene annotation results indicate role in plant disease resistance [69, 65, 72]. This information obtained from gene annotation could potentially be used in fine mapping and map-based cloning to further characterize the mechanisms of spot blotch disease resistance. The markers with lowest P-values may be converted in to diagnostic markers to validate the SNPs and
used in identification of lines with desired alleles in early generations.

Genomic Prediction
We used the rrBLUP GS model, which includes all marker information to predict a line’s genomic estimated breeding values (GEBV) [73, 74]. rrBLUP requires much lower computational time compared to the other GS models and it is shown to be robust in different GS scenarios [75, 76, 77]. Improving disease resistance in crops via single or a few major genes may be a temporary solution because the effectiveness is limited to only selected races of the pathogen and therefore, have no broad-environment application [78, 79]. The costs connected with the introgression of major genes or QTLs into elite backgrounds is challenging and may unintentionally affect the breeding operations and fast-track the evolution of the pathogen. Here, genomic prediction is well-placed to improve the effectiveness of quantitative disease resistance efforts in wheat breeding [80, 81]. The SB in wheat is shown to be controlled by many small to large effect genes [12, 13, 82, 83]. Therefore, instead of focusing on a few large-effect QTL, the prediction of spot blotch disease infection based on both small- and large-effect QTLs promise a holistic genetic-based approach for broad-spectrum resistance to evade the development and spread of spot blotch contagion.

Our results provide additional evidence in support of the quantitative nature of the disease resistance in wheat. Moderate to high prediction accuracies for 3 of 4 environments indicated that our model was able to successfully capture the quantitative variation underlying the SB variation in our population (Fig. 5). The contrasting prediction accuracies for PUSA19 environments in our study underscore the need for additional research to investigate the stability of genomic predictions across environments [75, 84, 85]. The role of environmental variation in the form of genotype-by-environment and marker-by-environment interactions in genomic selection has been highlighted by others [86, 87]. In view of both the positive and negative findings in this genomic prediction work, this study would provide an important precursor for future wheat breeding research in South Asia which is proven by other researchers also [79, 81].

Conclusions
This study aimed to identify genetic regions underlying spot blotch resistance in the elite spring
wheat genotypes. The variable conditions at four field environments in India and Bangladesh allowed us to capture the considerable phenotypic variation for spot blotch disease in our trials. The GWAS based on the phenotypic data at each site resulted in a total of 23 genomic regions on 14 chromosomes. We were able to validate earlier findings and identified new genomic regions on chromosome 3D contributing up to 6.94% of phenotypic variation. The literature mining of the functional gene annotations identified 36 SNPs encoding single protein or protein family, directly or indirectly involved in disease resistance. The SNPs on chromosome 5A associated with the known gene encodes ‘Myb family transcription factor-like protein’ found to have direct involvement in spot blotch resistance. Using a five-fold cross-validation scheme, we observed moderate prediction accuracies for 3 of 4 environments indicated that our model was able to successfully capture the quantitative variation underlying the SB variation in our population. The results are of importance for the breeders developing spot blotch resistant varieties targeting South Asian region. Given the aggressive pathogen spread and food security concerns, the breeding programs in South Asia could benefit by deploying a genomic selection based breeding scheme for broad-spectrum spot blotch disease resistance in wheat.

Methods

**Plant material and field layout**

The population was a genetically diverse collection comprising 141 advanced breeding lines of spring wheat. It represents 25 years of research at CIMMYT and was carefully assembled to avoid the confounding effects of phenology. The lines were evaluated in two replicates at two field locations: BISA, Pusa, India (25°57'22.8"N, 85°40'13.1"E) in the north India, and BARI, Jamalpur, Bangladesh (24°22'07.7"N, 88°39'42.0"E) during 2016-17, 2017-18 and 2018-19 seasons in a Randomized Block Design. For convenience, the different location-season combinations were termed as Env1 (Pusa 2016-17), Env2 (Pusa 2017-18), Env3 (Pusa 2018-19) and Env4 (BARI 2016-17). Both these locations are known as hot spot for spot blotch disease [3]. The plots of 3.8-meter length were sown in six rows with 0.22 meter spacing at each environment. The trials were timely sown with full irrigation applied through gravity flood-irrigation. The spreader rows of susceptible variety Sonalika were also planted
in alleys for disease build up. In addition, four auxiliary gravity flood-irrigations were also given at regular intervals. All agronomic practices like fertigation and weeding were performed as recommended for each location.

**Screening for spot blotch disease resistance**

The material was evaluated under natural infection conditions in the field. To limit the number of escapes, spot blotch response was evaluated during the mid to advanced-phases of disease development i.e., between heading (GS50 on Zadoks scale) and grain filling stage (GS80) [34]. The disease severity (SEV) were recorded visually on 0 to 100 scale where 0 is complete resistance and 100 is complete susceptibility.

**Genotyping**

Seeds of all lines were obtained from the CIMMYT genetic resources program and genomic DNA was extracted from five bulked leaves using a CTAB procedure [88,] modified as described in Dreisigacker et al. 2013 [89] in CIMMYT, Mexico. The DNA samples were sent to Kansas State University, USA for GBS. The GBS was performed following the protocol of Poland et al 2012 [90]. All lines were sequenced with Illumina Hi Seq2000 or HiSeq2500. GBS-SNP markers were called with TASSEL v5.2 pipeline [91] and aligned to the reference Chinese Spring Wheat Assembly v1.0. The following SNP filtering criteria was applied on raw SNP calls: less than 30% missing markers, minimum 5% minor allele frequency (MAF) and less than 20% heterozygosity. The filtering step yielded 18637 markers and the remaining missing values were imputed using Beagle v4.1 [92].

**Statistical analysis**

The experimental design in each environment was an randomized complete block design with two replications per location. META-R v6.03 developed by CIMMYT [93], Mexico was used to perform multi-environment mixed model analysis. The environments were used as random effects and genotypes as fixed effects. The resulting analysis produced the adjusted trait phenotypic values in the form of best linear unbiased predictions (BLUPs) within and across environments. In addition, the components of phenotypic variance were also extracted to calculate broad-sense line-mean heritability. The genetic linkage map was prepared using IciMapping v.4.0 [94] while principal component and dendograms
were performed using JMP v13 (SAS Inc., Cary, NC, USA). In addition to BLUP, we calculated Log, square root and Arrhenius values from severity percentage using JMP v13. The raw phenotypic distributions of disease scores at each environment were plotted to check normality assumptions. The data was log-transformed for cases where the distributions deviated significantly from normality.

**PCA and linkage disequilibrium**

The PCA was performed using 18637 SNPs and 141 genotypes in Tassel v5.2 [91] for the genetic relationship among genotypes. The first two principal components were drawn to show the clustering among genotypes. The population structure was inferred using the JMP v13. The Kinship was calculated using IBS method in Tassel 5.2. In order to determine the number of SNP marker for GWAS, the LD was estimated in TASSELv5.2 [91] using 18637 markers. The long-distance LD approach was used where critical $r^2$ of 0.2 was drawn in R software v3.5.2. [95] The marker $r^2$ for some chromosomes showed extensive noise, therefore instead of calculating chromosome-wide LD-decay thresholds, we generated LD heat maps of the significant markers for each chromosome separately. The intra-chromosomal LD was calculated as the pairwise marker correlations ($R^2$) between the GBS markers and plotted against the physical distance for significant marker-trait associations.

**Association analysis**

TASSEL v5.2 [91] was used to calculate population Kinship Matrix based on scaled IBS (identity by state) method. Mixed linear model (MLM) was used to test the marker-trait association between the SNP markers and spot blotch disease severity (SEV). MLM has proven useful in controlling for population structure and relatedness within genome wide association studies. Subsequent GWAS and genomic prediction analyses were performed on 16037 SNPs scored on 141 lines from the seasons 2016-17, 2017-18 and 2018-19. Genome-wide significance threshold was established to reduce the false discovery rate due to multiple test comparisons.

**Gene functional annotations**

GWAS results were further analyzed to test if the marker-trait associations fall within the known genic regions and by functional annotation from the reference genome assembly (IWGSC Ref Seq v1.0).
Functional annotation of the genes harboring significant SNPs were retrieved and examined for their association with spot blotch resistance from the genome annotations provided by IWGSC. Subsequently, genes annotated proteins functions were literature mined.

**Genomic Prediction**

The Log transformed BLUPs of spot blotch scores calculated across years were used for genomic prediction modeling. A five-fold cross-validation scheme [96] was implemented. The advanced breeding lines were randomly divided into five subsets (i.e., folds), and four of them were used as the training set. each with approximately the same number of individuals. The random cross-validation runs were repeated for 100 iterations. At each step, the predictive accuracy of the markers was assessed by Pearson’s correlation between the predicted values and the BLUP phenotypes. Overall average of the fifth fold was reported as accuracy of the prediction. All calculations were performed in R software [95] and by using the packages lme4 and rrBLUP [97, 74].

**Declarations**

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**Author contribution**

**VT:** Drafted the manuscript, recorded data in the field and analyzed the data; **RPS:** Provided the breeding material and experiment design; **JP:** Genomic predictions and GBS data of breeding material; **DJ:** Performed genomic predictions; **AKJ:** Spot evaluation and trial management; **PKS:** scientific inputs for manuscript preparation; **PKB:** Experimental design and data recording; **SK:** Recorded data in the field; **MR:** Spot evaluation and trial management at BARI; **GSD:** Linkage mapping analysis; **BST:** revised the manuscript; **UK:** Design the experiment, revised manuscript and overall supervised the research work. All authors read and approved the final manuscript.
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Ethics declarations

Ethics approval and consent to participate
Not applicable

Consent for publication
Not applicable

Competing interests
The authors declared that they had no competing interests

Declaration
The data can be made available on request. It is confirmed that the data may be uploaded into public domain once the manuscript is accepted for publication.

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# Tables

Table 1: Range, mean values, standard deviation and sample variance of 141 advanced lines evaluated for spot blotch disease resistance at growth stage 77 on Zadoks Scale in four environments

| Statistic                  | Env1   | Env2   | Env3   | Env4   |
|----------------------------|--------|--------|--------|--------|
| Mean                       | 21.67  | 8.90   | 15.10  | 31.23  |
| Standard Error             | 0.88   | 0.71   | 0.39   | 0.72   |
| Standard Deviation         | 14.81  | 11.96  | 6.55   | 12.03  |
| Sample Variance            | 219.45 | 143.09 | 42.88  | 144.83 |
| Range                      | 0-70   | 0-50   | 0-43   | 11-66  |
| Confidence Level (95.0%)   | 1.73648152 | 1.402192864 | 0.767555673 | 1.410680491 |

Table 2: Analysis of variance of 141 lines evaluated for spot blotch disease resistance in four environments based on BLUP of disease severity recorded at growth stage 77 on Zadoks scale

| Statistic                  | Env1       | Env2       | Env3       | Env4       | Combined   |
|----------------------------|------------|------------|------------|------------|------------|
| Heritability               | 0.76       | 0.71       | 0.50       | 0.80       | 0.55       |
| LSD                        | 16.49      | 14.16      | 14.41      | 8.74       | 11.04      |
| CV                         | 28.21      | 30.06      | 29.73      | 22.41      | 28.53      |
| Replications               | 2          | 2          | 2          | 2          | 2          |
| Residual variance          | 184.28     | 146.56     | 222.91     | 48.98      | 151.63     |
| Gen variance               | 284.48     | 174.91     | 113.18     | 96.25      | 68.80      |
| Gen significance           | 1.11022E-15| 2.6390E-13 | 4.73255E-05| 0          | 2.82891E-08|
| Gen × Env variance         | -          | -          | -          | -          | 96.00      |
| Gen × Env significance     | -          | -          | -          | -          | 3.69898E-14|
| Env variance               | -          | -          | -          | -          | 104.36     |
| Env significance           | -          | -          | -          | -          | 0.0115412318|

Table 3. List of significant SNPs with the corresponding proteins and possible function elucidated based on the gene annotation using wheat reference sequence (RefSeq V1.0) database

| SNP markers | GeneID | Gene annotation | Possible Function/Description | References |
|-------------|--------|-----------------|------------------------------|------------|
| ENV1 (PUSA17)| | | | |
| S1A_9565 | TraesCS1A01G | Trichome birefringence | Cellulose biosynthesis | [98] |
| 863 | 018700 | | | |
| | TraesCS1A01G | | | |
| | Transmembrane | Regulates fungal development and | | |
| | | | | |
| ID         | Protein Name                             | Description                                                                 | Reference |
|------------|------------------------------------------|-----------------------------------------------------------------------------|-----------|
| S1D_6541  | TraesCS1D01G012500                       | protein C2 calcium/lipid-binding and GRAM domain protein                     | [99]      |
| 259        |                                          | C2 domain protein BAP1 negatively regulates defense responses in *Arabidopsis* |           |
| S1D_6715  | TraesCS1D01G012800                       | Serine/threonine-protein kinase                                              | [66]      |
| 588        |                                          | Confers powdery mildew resistance in wheat                                  |           |
| S2B_1375  | TraesCS2B01G030100                       | Disease resistance protein RPP13, lectin-receptor kinase                     | [31]      |
| 1999       | TraesCS2B01G030200                       | Lectin receptor kinases are involved in plant immunity                      |           |
|            |                                          | Play important roles in the resistance of various plant diseases including the downy mildew |           |
| S2B_1376  | TraesCS2B01G030200                       | Lectin receptor kinase                                                      | [32]      |
| 1590       |                                          | Lectin receptor kinases are involved in plant immunity                      |           |
| S2B_1381  | TraesCS2B01G030500                       | Cytochrome P450 family protein                                              | [100]     |
| 4702       |                                          | Enhances both resistance to deoxynivalenol and grain yield                   |           |
| S2B_1426  | TraesCS2B01G030900                       | Myrcene synthase                                                            | [101]     |
| 1851       |                                          | Interference and disruption of membranes in maize infected by *Fusarium spp* |           |
| S2B_1480  | TraesCS2B01G031700                       | Cytochrome P450                                                             | [100]     |
| 9954       | TraesCS2B01G031800                       | Enhances resistance to deoxynivalenol and increase grain yield              |           |
|            | ATP-dependent Clp protease adapter protein | Essential for early development in *Arabidopsis*                           | [102]     |
| S2B_1496  | TraesCS2B01G031900                       | Cytochrome P450                                                             | [100]     |
| 3432       |                                          | Enhances resistance to deoxynivalenol and increase grain yield              |           |
| S2B_1512  | TraesCS2B01G032100                       | Cytochrome P450                                                             |           |
| 9248       |                                          | Enhances resistance to deoxynivalenol and increase grain yield              |           |
|            | Zinc finger family protein               | Involved in plant disease resistance                                        | [56]      |
| S3B_7632  | TraesCS3B01G519800                       | Transmembrane protein                                                       | [69]      |
| 30831      |                                          | Transmembrane protein FgSho1 regulates fungal development and pathogenicity via MAPK module Ste50-Ste11-Ste7 in *F. graminearum* |           |
|            | TraesCS3B01G519900                       | General regulatory factor 1                                                |           |
| S3B_7632  | TraesCS3B01G520000                       | Serpin family protein                                                       | [103]     |
| 36179      |                                          | Participate in the regulation of complex proteolytic systems                |           |
| S3B_7632  | TraesCS3B01G520100                       | 2-oxoglutarate (2OG) and Fe(II)-dependent oxygenase superfamily protein    | [100]     |
| 67753      |                                          | Enhances resistance to deoxynivalenol and increase grain yield              |           |
|            | TraesCS3B01G520200                       | Serpin family protein                                                       | [103]     |
|            |                                          | Participate in the regulation of complex proteolytic systems                |           |
| S3B_7641  | TraesCS3B01G520800                       | -                                                                          | [58]      |
| 73978      |                                          | Negative regulator of the defense response                                  |           |
| S3B_7647  | TraesCS3B01G521400                       | Glycerol-3-phosphate dehydrogenase                                          | [59]      |
| 47435      |                                          | Contributes to systemic acquired resistance against *P. striiformis* f. sp. *tritici* |           |
| GenBank ID | Description | Function | Reference |
|------------|-------------|----------|-----------|
| S3B_764804887 | TraesCS3B01G 521500 Zinc finger family protein | Plant disease resistance | [56] |
| S3B_764804887 | TraesCS3B01G 521600 Zinc finger family protein | Plant disease resistance | [56] |
| S3B_764804887 | TraesCS3B01G 521700 Tyrosyl-DNA phosphodiesterase | | |
| S5A_593739698 | TraesCS5A01G 400900 Kinase family protein | As initiators of symbiosis or defense | [104] |
| S5A_593739698 | TraesCS5A01G 401000 Keratin, type II cuticular Hb1 | | |
| S5A_595158840 | TraesCS5A01G 402500 Molybdenum cofactor sulfurease | Overexpression of Arabidopsis Molybdenum Cofactor Sulfurase gene confers drought tolerance in maize | [105] |
| S5A_595158840 | TraesCS5A01G 402600 Cytochrome b6-f complex subunit 7 | Oxygenic photosynthesis | [106] |
| S5A_595373332 | TraesCS5A01G 402700 Transmembrane protein | Regulates fungal development and pathogenicity via MAPK module FgSte50-Ste11-Ste7 in F. graminearum | [69] |
| S5A_595393566 | TraesCS5A01G 402800 Myb family transcription factor-like protein | Mediates host resistance to Bipolaris sorokiniana in wheat | [57] |
| S5A_636959783 | TraesCS5A01G 457100 Guanyl-biding family protein | Non-canonical fungal G-protein coupled receptors promote Fusarium head blight on wheat | [107] |
| S5A_636959783 | TraesCS5A01G 457200 Carboxy-terminal peptidase | | |
| S5B_233586644 | TraesCS5B01G 127900 Hippocampus abundant transcript-like protein 1 | | |
| S5B_233586644 | TraesCS5B01G 128000 Signal recognition particle subunit SRP68 | Crucial role in targeting secretory proteins to the rough endoplasmic reticulum membrane | [108] |
| S6A_29496364 | TraesCS6A01G 056000 F-box family protein | Negative regulator of the defense response | [58] |
| S6A_33023854 | TraesCS6A01G 061900 Kinase family protein | As initiators of symbiosis or defense | [104] |
| S6A_33299007 | TraesCS6A01G 062000 F-box family protein | Negative regulator of the defense response | [58] |
| S6A_33299007 | TraesCS6A01G 062300 DNA-directed RNA polymerase II subunit rpb4 | Pol II alone is capable of RNA transcript elongation and of proof reading | |
| S6A_33299007 | TraesCS6A01G 062400 Pigment epithelium-derived factor | | |
| S7B_1020705 | TraesCS7B01G 002400 Leucine-rich repeat receptor-like protein kinase | Pathogen-Responsive Leucine Rich Receptor Like Kinase Contributes to Fusarium Resistance in Cereals and regulates powdery mildew resistance in wheat | [53, 109] |
| S7B_1261577 | TraesCS7B01G 003000 COP1-interacting-like protein | COP1 and HY5 also play essential roles during UV-B signaling | [110, 111] |
| ENV2 (PUSA18) | | | |
| S1B_6471 | TraesCS1B01G Pentatricopeptide | Involved in plant disease resistance | [65] |
| Plate number | Genotype | Accession | Type | Description | Notes |
|--------------|----------|-----------|------|-------------|-------|
| 40           | 95634    | 424000    | repeat-containing protein | Homeobox protein | Associated with Reaction to Stripe Rust and Powdery Mildew [72] |
| S2A_7084     | 82943    | TraesCS2A01G 424100 | Receptor-like kinase 462700 | Glycosyltransferase 462800 | Involves in plant disease resistance [55] |
| S2A_7091     | 73869    | TraesCS2A01G 463300 | Glycosyltransferase 463500 | - | Involves in plant disease resistance [55] |
| S2B_9066     | 2917     | TraesCS2B01G 123200 | Transcription repressor ofp17 | - | Transcriptional Repressor of TaSHY2 and TaIAA7, Enhances Root Length and Biomass in Wheat [112] |
| S3D_610      | 628298   | TraesCS3D01G 537500 | Zinc finger protein (Dof) | - | Zinc Finger Proteins Involved in Plant Disease Resistance [56] |
| S5B_4000     | 97303    | TraesCS5B01G 224500 | Golgi SNAP receptor complex member 1 | - | Involved in plant disease resistance through regulation of Plant Cell Death [113] |
| S5B_6638     | 22910    | TraesCS5B01G 496700 | Chaperone protein dnaJ | - | Leads to thermosensitive gametophytic male sterility in Arabidopsis [114] |
| S7A_7097     | 69485    | TraesCS7A01G 530700 | Pentatricopeptide repeat-containing protein | - | Essential Role in Organelle Biogenesis [115] |
| S7D_1819     | 74079    | TraesCS7D01G 221000 | Hydroxysteroid dehydrogenase | - | Involved in regulating plant growth and development [117] |
| ENV3 (PUSA19) | 7405    | TraesCS7D01G 221100 | Hydroxysteroid dehydrogenase | - | - |
| S2A_3185     | 1904     | TraesCS2A01G 071500 | Flavin-containing monooxygenase | - | Mediate two-step auxin biosynthesis pathway in Arabidopsis [118] |
| S2B_5047     | 17       | TraesCS2B01G 001100 | Cytochrome P450 001120 | Chaperone protein DnaJ | Enhances resistance to deoxynivalenol and increase grain yield [100] |
| S2B_3190     | 90       | TraesCS2B01G 000700 | 3\(^{-}\)N-debenzoyl-2\(^{-}\)deoxytaxol N-benzyoltransferase | - | Leads to thermo-sensitive gametophytic male sterility in Arabidopsis [114] |
| S2B_7806     | 5        | TraesCS2B01G 000400 | Cytochrome P450 000500 | Peptide transporter | Enhances resistance to deoxynivalenol and increase grain yield [100] |
| Gene ID | Description 1 | Description 2 | Reference |
|---------|----------------|---------------|-----------|
| S2B_8311 062 | TraesCS2B01G018100 | Terpene synthase | A potential role of terpenes in drought tolerance [119] |
|         | TraesCS2B01G018200 | NBS-LRR disease resistance protein-like | Disease resistance protein [54, 55] |
| S2B_6253 562 | TraesCS2B01G012400 | Avr9/Cf-9 rapidly elicited protein | Early signaling events in the Avr9/Cf-9-dependent plant defence response [61] |
| S2B_9066 2917 | TraesCS2B01G123200 | Transcription repressor of p17 | Transcriptional repressor of TaSHY2 and TaIAA7, enhances root length and biomass in wheat [112] |
|         | TraesCS2B01G123300 | - | - |
| S2B_8954 0368 | TraesCS2B01G122000 | Mitochondrial transcription termination factor-like | - |
|         | TraesCS2B01G122100 | Expansin protein | - |
| S2B_7485 95375 | TraesCS2B01G552600 | Disease resistance protein RPM1 | Disease resistance protein [120] |
|         | TraesCS2B01G552700 | Signal peptide, CUB and EGF-like domain-containing protein 3 | - |
| S3A_6734 8475 | TraesCS3A01G104000 | helicase with zinc finger protein | - |
|         | TraesCS3A01G104100 | Two-component response regulator ORR24 | Involved in His-to-Asp phosphorelay signal transduction system [121] |
| S3A_6651 3067 | TraesCS3A01G103000 | RING/U-box superfamily protein | Associated with the control of grain size [122] |
|         | TraesCS3A01G103100 | Fasciclin-like arabinogalactan protein | Involved in conidiation and pathogenicity in *Magnaporthe oryzae* [123] |
| S3A_6706 5083 | TraesCS3A01G103400 | Di-glucose binding protein with Kinesin motor domain-containing protein | - |
|         | TraesCS3A01G103500 | Transcription factor-like protein | 1R-MYB Transcription Factor, plays an important role in disease resistance against stripe rust fungus and ear heading in wheat [45] |
| S3A_7207 | TraesCS3A01G | Zinc finger family | Involved in plant disease resistance [56] |
| ID      | Entry      | Description                                           | Function                                                                 | Reference   |
|---------|------------|-------------------------------------------------------|--------------------------------------------------------------------------|-------------|
| 2539    | TraesCS3A01G107400 | Protein NAC domain (GRF)                               | Wheat NAC interacts with an orphan protein and enhances resistance to FHB disease | [124]       |
| S5B_6730 38030 | TraesCS5B01G507700 | Protein plant cadmium resistance                      | -                                                                         | -           |
|         | TraesCS5B01G507800 | Invertase inhibitor                                   | Overexpression of Pectin Methylesterase Inhibitors in Arabidopsis restricts fungal infection by Botrytis cinerea | [125]       |
| S6A_2395 9761 | TraesCS6A01G047300 | Peroxidase                                            | Divergent role in different plant-pathogen systems                       | [126]       |
|         | TraesCS6A01G047400 | -                                                     | -                                                                        | -           |
| S7B_2403 25318 | TraesCS7B01G169400 | Glutathione S-transferase T3                          | Involved in complex stress regulation in Arabidopsis                     | [127]       |
|         | TraesCS7B01G169500 | Cytochrome P450                                        | Enhances resistance to deoxynivalenol and increase grain yield           | [100]       |
| S7D_4964 1314 | TraesCS7D01G082700 | RING/U-box superfamily protein                        | Associated with the control of grain size                                | [122]       |
|         | TraesCS7D01G082800 | Polygalacturonase-1 non-catalytic beta subunit         | Implicated in the defense against fungal pathogens in Plants through cell wall modification, degradation, carbohydrate metabolic processes and as response to stress | [128, 129] |
| S7D_4784 8266 | TraesCS7D01G081100 | Telomerase activating protein                         | -                                                                       | -           |
|         | TraesCS7D01G081200 | Ubiquitin-conjugating enzyme                          | Regulates wheat defence against the phytopathogen Zymoseptoria tritici  | [130]       |
| ENV4 (BARI17) |                     |                                                      |                                                                          |             |
| S1B_6468 95451 | TraesCS1B01G423900 | CXC domain-containing protein                         | Role in reproductive tissues development                                 | [131]       |
|         | TraesCS1B01G424000 | Pentatricopeptide repeat-containing protein           | Involved in plant disease resistance                                    | [65]        |
| S2B_2859 2818 | TraesCS2B01G058800 | Sugar transporter                                     | Role in plant defense responses against fungal pathogens                 | [132]       |
|         | TraesCS2B01G058900 | NBS-LRR disease resistance protein                    | Disease resistance protein                                               | [54, 55]    |
| S2B_5331 78164 | TraesCS2B01G373800 | little nuclei4                                        | -                                                                       | -           |
|         | TraesCS2B01G373900 | Cytochrome P450                                        | Enhances resistance to deoxynivalenol and increase grain yield           | [100]       |
| S2B_6992 19601 | TraesCS2B01G505200 | Disease resistance protein-like                       | Important roles in the resistance of various plant diseases including downy mildew | [31]        |
| S4A_7108 30493 | TraesCS4A01G443100 | Sodium-dependent phosphate transporter                | Importance in the filial tissues during grain filling                    | [133]       |
|         | TraesCS4A01G443200 | ATP-dependent Clp protease ATP-binding subunit        | Essential for early development in Arabidopsis                          | [102]       |
| S4B_3680 23618 | TraesCS4B01G168200 | CBL-interacting Serine/Threonine-                     | Regulates drought stress and ABA responses                               | [134]       |
| Gene ID | Description                                      | Function                                                                                     | Reference |
|---------|--------------------------------------------------|----------------------------------------------------------------------------------------------|-----------|
| TraesCS4B01G 168300 | Uroporphyrinogen decarboxylase                  | Defense responses                                                                         | [135]     |
| SSB_6817 73490 | TraesCS5B01G 518800 | basic helix-loop-helix (bHLH) DNA-binding superfamily protein | Regulate growth and development as well as response to various stresses | [136]     |
| TraesCS5B01G 518900 | Hydroxypyroline-rich glycoprotein family protein | Provides added resistance against pathogen-derived cell wall-degrading enzymes               | [137]     |
| SSB_6833 52145 | TraesCS5B01G 521300 | NBS-LRR disease resistance protein family-1                                                | Disease resistance protein | [54, 55] |
| TraesCS5B01G 521400 | - | -                                                                                           | -         |
| SSB_6835 14734 | TraesCS5B01G 521400 | Kinase family protein                                                                        | As initiators of symbiosis or defense | [104]     |
| TraesCS5B01G 521500 | F-box family protein                             | Negative regulator of the defense response                                                  | -         |
| TraesCS5B01G 522300 | Glycosyltransferase                              | Involved in plant disease resistance                                                       | -         |
| TraesCS5B01G 522400 | ATP-dependent zinc metalloprotease FTSH protein | Overexpression of the ASPG1 gene confers drought avoidance in Arabidopsis                   | [138]     |
| TraesCS5B01G 522500 | ATP-dependent zinc metalloprotease               | Regulates hydrolase, metalloprotease and protease activities -cell division (peptidase/protease) | -         |
| TraesCS5B01G 522600 | Eukaryotic aspartyl protease family protein      | Overexpression of Pectin Methylesterase inhibitors in Arabidopsis restricts fungal infection by Botrytis cinerea | [125]     |
| TraesCS5B01G 522900 | Protein detoxification                           | -                                                                                           | -         |
| TraesCS5B01G 523800 | ATP-dependent zinc metalloprotease               | Associated with the control of grain size                                                   | [122]     |
| TraesCS6B01G 066100 | Invertase/pectin methylesterase inhibitor family protein | Associated with the control of grain size                                                   | [122]     |
| TraesCS6B01G 066200 | PME/invertase inhibitor-like protein             | Overexpression of Pectin Methylesterase inhibitors in Arabidopsis restricts fungal infection by Botrytis cinerea | [125]     |
| TraesCS6B01G 472300 | RING/U-box superfamily protein                   | -                                                                                           | -         |
| TraesCS6B01G 472400 | Integral membrane metal-binding family protein  | -                                                                                           | -         |
| TraesCS7A01G 504700 | Patatin                                          | Pathogen-inducible patatin-like lipid acyl hydrolase facilitates fungal and bacterial host colonization in Arabidopsis | [139]     |
| TraesCS7A01G 504800 | Protein phosphatase 2c                           | Function of the plant PP2Ac genes in plant immune responses                                 | [140]     |
| TraesCS7D01G 067100 | Phosphate-responsive 1 family protein            | Function of the plant PP2Ac genes in plant immune responses                                 | [140]     |

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Figures 1

Distribution of lines for spot blotch resistance and susceptibility evaluated in four environments.
Figure 2

Physical map of QTLs mapped along with already known/mapped loci. The gray and red rectangles represent loci from literature and mapped during this study respectively. The loci marked with (c*) represent clusters of loci mapped at a particular region. Q.S2B.c: Q.S2B.319090, Q.S2B.6253562, Q.S2B.8311062, Q.S2B.13751999, Q.S2B.13761590, Q.S2B.13814702, Q.S2B.14261851, Q.S2B.14809954, Q.S2B.14963432, Q.S2B.15129248, Q.S2B.19097998, Q.S2B.28592818; Q.S3B.c: Q.S3B.763230831, Q.S3B.763236179, Q.S3B.763267753, Q.S3B.764173978, Q.S3B.764747435, Q.S3B.764804887; Q.S5A.c: Q.S5A.593739698, Q.S5A.595158840, Q.S5A.595373332, Q.S5A.595393566, Q.S5A.636959783; and Q.S5B.c: Q.S5B.673038030, Q.S5B.681773490, Q.S5B.683352145, Q.S5B.683514734, Q.S5B.684060726, Q.S5B.684129768, Q.S5B.684230407,
Figure 3
Population structure: PCA and Dendogram

Figure 4
Venn diagram based on the common protein synthesized by the same genes associated with different SNPs over years (The numbers represent the total proteins common between the environments)
Figure 5

Genomic prediction for spot blotch in four different environments using rrBLUP

Supplementary Files
This is a list of supplementary files associated with this preprint. Click to download.

6- Table S2 Common proteins.xlsx
7- Table S3 Weather.docx
5- Table S1 Significant SNPs.xlsx