Genetic Evaluation and Screening of Diverse Wheat Genotypes for Spot Blotch Resistance

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Abstract
Production of wheat (Triticum aestivum L.) the main food source of South Asian countries including India faces several constraints including spot blotch caused by Bipolaris sorokiniana resulting in yield loss of 25–43 % depending upon the stage of infection. Fifty genotypes were evaluated for nine quantitative characters and area under disease progress curve (AUDPC) to identify superior genotype with spot blotch resistance. High heritability coupled with moderate to high genetic advance as percent of mean was registered for grains per spike, tillers per square meter, days to 50% heading and days to 50% flowering indicating the characters to be governed by additive genes. Correlation and path coefficient analysis favored days to 50% heading, days to 50% flowering and grains per spike since they had significant positive correlation with yield and simultaneous negative correlation with AUDPC and also conferring highest positive direct effect towards yield. Multiple linear regression (MLR) analysis indicated days to 50% heading to be most sensitive with negative influence on AUDPC. D² analysis grouped the 50 genotypes into 10 clusters suggesting presence of diversity among the genotypes. Frequency distribution of AUDPC among the genotypes showed more or less normal distribution of the character. Low AUDPC score with acceptable level of yield performance were recorded for the genotypes 29882, 29610, 29473, 29940, 29477, 29748 and 30081. Identification of high yielding and less susceptible genotypes for spot blotch disease in the present investigation offered an opportunity for wheat improvement through selective breeding.

Introduction
Bread wheat (Triticum aestivum L.) one of the oldest cereal crop is regarded as the ‘King of Cereals’ since it shares a large area under production, high productivity and holds a prominent position in the international food grain trade (Hazra et al. 2019a). It is the main food source of South Asian countries (Singh et al. 2016) and in India it is the principal cereal crop next to rice (Kumari et al. 2020). The overall production of wheat in India has gone up tremendously from 12.26 million tonnes in 1964-65 to 103.6 million tonnes in 2018-19. Recently, an estimate of United States Department of Agriculture (USDA) states that India is expected to touch the new level of wheat production about 107.0 million tonnes in 2020-21 (USDA 2021).

Production of bread wheat in South Asia during countries including India still faces various constrains like raising temperature, unexpected hailstorms, erratic and unusual precipitation during February-March (Duveiller 2004), exposing the crop to several diseases and pests including spot blotch or foliar blight of wheat. West Bengal is categorized as a hotspot for the disease because of its mild and short winter, humid climate and late sowing due to delay in harvesting of kharif rice and sometimes excessive soil moisture after rice harvest. Warm and humid climate of this region aggravates the disease which seriously hampers the production of intensive cropping system (Singh et al. 2016). The yield loss may range from 25–43% depending upon the stage of infection and the national yield loss is recorded to be around 18–22% (Acharya et al. 2011).

Spot blotch is caused by a hemi biotrophic fungal pathogen Bipolaris sorokiniana (Sacc.) Shoem syn. Drechslera sorokiniana (Sacc.) Subrm and Jain (syn. Helminthosporium sativum) and its teleomorph is Cochliobolus sativus (Singh et al. 2016). Typical symptoms of the spot blotch disease appear on the leaves, sheath, nodes and glumes with brown lesions of oval to oblong or elliptical in shape measuring 5 to 10 mm long and 3 to 5 mm wide (Gupta et al. 2018). Other symptoms include darkening of the sub crown region, dark brown lesions on culm, coleoptile, crowns and roots. This pathogen first attacks the older leaves at the base of the plant and then progresses upward (Joshi et al. 2002). Lesions on leaves may start from few mm and later it turns into dark brown spots and can extend up to 1–2 cm (Chand et al. 2002). As the disease progresses, the lesions get scattered throughout the leaves and subsequently their size increase to coalesce with each other to form large necrotic spots which result in loss chlorophyll that causes reduction in photosynthetic area of the leaf (Gupta et al. 2018). Sometimes yellowing can be seen due to toxin production from the lesion (Chowdhury et al. 2013). In its severe form the fungal pathogen attacks the spikes forming dark brown to black discoulouration around the gerninating point of the seed known as “Black Point” (Gupta et al. 2018).

The genetic base of cultivated wheat genotypes has become narrow due to continuous inbreeding (Rehman et al. 2018) and the present agricultural scenario has led to rapid decline in both inter and intra varietal variability as a result of continuous breeding of the elite genotypes. Complete resistance in bread wheat against spot blotch or foliar blight is still lacking, although low to high levels of resistance have been reported (Rosyara et al. 2007; Singh et al. 2020). In high yielding varieties of wheat resistance to spot blotch is poor and requires rigorous investigation to improve the resistance along with good yield (Joshi et al. 2007; Meena et al. 2014). The existing genetic variability can be exploited by intercrossing among diverse genotypes to isolate superior transgressive segregants (Baranwal et al. 2012; Shah et al. 2020). Identification of superior diverse genotypes with desirable traits and their consequent use in breeding program and establishment of successful selection criteria can be helpful for successful varietal improvement (Hazra et al. 2019b).

The present investigation has been aimed to assess the interrelation between the disease score and other important agro-morphological traits of fifty diverse wheat genotypes grown naturally under a hotspot region for the disease in India for formulating an effective selection criterion for spot blotch resistance wheat breeding.

Materials And Methods
Experimental material
Fifty bread wheat genotypes (Table 1) collected from International Center for Agricultural Research in Dry Areas (ICARDA), Aleppo, Syria and AICRP-Indian Institute of Wheat and Barley Research (ICAR-IIWBR), Karnal, India was used in the present study to screen for the resistance to spot blotch disease and other agronomic traits. Out of the fifty genotypes, HD2967, HD3086 and DBW107 were used as yield check as suggested by Gupta et al. (2017) while Sonalika, vulnerable to spot blotch, was used as a susceptible check variety for spot blotch disease screening according to Turan et al. (2017).
Table 1  
Fifty wheat genotypes used in the present investigation

| Sl. No. | Code no. of genotypes | Parentage | Source |
|--------|-----------------------|-----------|--------|
| 1      | TERBOL                | TERBOL    | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 2      | ATLAS                 | ATLAS     | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 3      | TESFA                 | TESFA     | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 4      | 30140                 | ABU-REYAA-1/LEITH-1 | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 5      | 30081                 | ABUZIG-10/2*PFAU/MILAN | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 6      | 29882                 | ALMAZ-19/ETBW 4919/3/NING MAI 9558//CHIL/CHUM18 | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 7      | 30053                 | ANBER-6//WORRAKATTA/PASTOR | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 8      | 29889                 | ASEEL-1//MILAN/PASTOR/3/SHAMISS-3 | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 9      | 29760                 | ATILIA*2/AMAD//ENKOY/3/PFAU/MILAN | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 10     | Sonalika (C)          | SONALIKA (C) | AICRP-Indian Institute of Wheat and Barley Research (ICAR-IIWBR) |
| 11     | 29761                 | ATILIA*2/AMAD//ENKOY/3/PFAU/MILAN | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 12     | 29872                 | ATILIA-1/NS732/HER//PARUS/PASTOR/3/TEMPORALERAM 87*2/KONK | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 13     | 29610                 | BACANORA T 88/RUTH-2 | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 14     | 29821                 | BACANORA T 88/RUTH-2//PFAU/MILAN | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 15     | 29690                 | BAOBAB-1//MILAN/PASTOR | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 16     | 29672                 | CHAM-6//SHUHA-14/5/KAUZ/3/MYNA/VUL//BUC/FLK/4/MILAN | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 17     | 29748                 | CHAM-8/FLAG-3//MILAN/PASTOR | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 18     | 29612                 | CHAM-8/RUTH-3 | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 19     | 29824                 | CHAMRAN/4/OPATA/BOW//BAU/3/OPATA/BOW/5/SAMIRA-9 | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 20     | 29506                 | CROC-1/AE.SQUARROSA (205)//MILAN/KAUZ/3/MILAN/PASTOR | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 21     | 29509                 | CROC-1/AE.SQUARROSA (205)//MILAN/KAUZ/3/MILAN/PASTOR | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 22     | HD 2967 (C)           | HD2967 (C) | AICRP-Indian Institute of Wheat and Barley Research (ICAR-IIWBR) |
| 23     | 29865                 | DAJAJ-5/4/CHEN/AEGILOPSSQUARROSA (TAUS)//BCN/3/KAUZ/5/WBLL1*2/KIRITATI | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 24     | 29522                 | DURRA-2/TAZA-2 | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 25     | 30001                 | FAYEQ-2/3/NESMA*2/14-2//2*SAFI-3 | International Center for Agricultural Research in Dry Areas (ICARDA) |

The genotypes designated as (C) has been used as check.
| Sl. No. | Code no. of genotypes | Parentage | Source |
|--------|-----------------------|-----------|--------|
| 26     | HD3086 (C)            | HD3086 (C) | AICRP-Indian Institute of Wheat and Barley Research (ICAR-IIWBR) |
| 27     | 29784                 | GEMMEIZA-10/SHAMISS-3 | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 28     | 29782                 | HUBARA-1/ACHTAR/INRA1764/7/CHAM-8/6/SAKER’S/5/RBS/ANZA/3/KVZ/HYS//YM/H/TOB/4/BOW’S’ | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 29     | 29641                 | HUBARA-1/5/KAUZ/3/MYNA/VUL//BUC/FLK/4/MILAN | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 30     | 29502                 | INQALAB91*2/TUKURU//MILAN/PASTOR | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 31     | 29703                 | JAWAHIR-6/ETBW 4921 | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 32     | 29490                 | KATILA-13/PFAU/MILAN | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 33     | 29473                 | PBW343*2/KUKUN//ANBER-9 | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 34     | 29769                 | PBW343/ETBW 4921//QAMAR-6 | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 35     | 29752                 | PFAU/MILAN//ABIER-2/3/SHUHA-3//TURACO/CHIL | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 36     | 29858                 | PFAU/MILAN/FLAG-3/3/NEJMAH-9 | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 37     | 29992                 | QADANFER-4//ACHTAR/INRA 1764/3/SHAMISS-3 | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 38     | 30098                 | RABHI-10/ETBW 4922//KAUZ’S/FLORKWA-1 | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 39     | 29526                 | RABHI-3/5/KAUZ/3/MYNA/VUL//BUC/FLK/4/MILAN | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 40     | 29945                 | TEMPORALERA M 87*2/KONK//FAYEQ-1 | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 41     | 29988                 | TEVEE’S/SHUHA’S//ACHTAR/INRA 1764/3/CHIL-1/SHUHA-1 | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 42     | 29903                 | TRAP#1/BOW//PFAU/3/MILAN/4/ETBW 4922/5/PFAU/MILAN | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 43     | 29812                 | WATAN-6/ETBW 4919//ZAKIA-14 | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 44     | 29798                 | WBLL1*2/BRAMBLING/3/OPATA/RAYON//KAUZ | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 45     | 29940                 | WEAVER/TSC//WEAVER/3/WEAVER/4/WAXWING/5/DURRA-8 | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 46     | 29477                 | WHEATEAR//ACHTAR/INRA 1764 | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 47     | 29476                 | WHEATEAR//ACHTAR/INRA 1764 | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 48     | 29671                 | ZAIN-4/QADANFER-11 | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 49     | 29913                 | ZERBA-6/FLAG-6/3/TAM200/PASTOR//TOBA97 | International Center for Agricultural Research in Dry Areas (ICARDA) |
| 50     | DBW107 (C)            | DBW107 (C) | AICRP-Indian Institute of Wheat and Barley Research (ICAR-IIWBR) |

The genotypes designated as (C) has been used as check.
The present investigation was conducted at ‘AB’ block farm BCKV (Bidhan Chandra Krishi Viswavidyalaya), Kalyani (22°59’ N, 88°48’ E, and 9.75 m above mean sea level) under the new alluvial zone of West Bengal, India during the Rabi season of two consecutive years 2018–2019 and 2019–2020. This area is primarily considered as hot spot area of spot blotch disease due to prevalent sub-tropical humid climatic condition with annual mean temperature range of 12.5° C – 36.3° C and rainfall of 1120–1500 mm with relative humidity of 50–80%, which is ideal for the development of disease. Fifty test genotypes were sown following randomized block design with three replications. In all the replications each genotype was planted in three rows of 3 m length keeping 18 cm distance between rows. The susceptible check Sonalika was included after every 20 test entries and along the borders to provide the chance of creating equal disease pressure to all the test genotypes. The sowing was done in the first week of December so that the post anthesis stage is exposed to warm and humid environment which is conducive for disease development (Chaurasia et al. 2000). During both the years, the crop was raised with recommended package of practices. A fertilizer dose of 120-60-40 kg/ha N-P₂O₅-K₂O was applied to the experimental plot. Half nitrogen and full amount of phosphorous and potassium was provided as basal during field preparation, a quarter of nitrogen was top dressed at 21 days after sowing (DAS) and another quarter at 40 DAS. Five irrigations were applied as recommended in the critical growth stages of the crop (crown root initiation stage, tillering stage, late jointing stage, flowering stage and dough stage) although the soil contained sufficient organic matter content (0.78%) to retain moisture.

**Assessment of agro-morphological traits**

The agro-morphological traits assessed in this experiment were plant height (cm), days to 50% heading, days to 50% flowering, tillers per square meter, days to maturity, spike length (cm), grains per spike, test weight (g), yield/plant (g). Plant height was measured from the base at ground level to the tip of spike of main tiller excluding awns at maturity. Days to 50% heading and days to 50% flowering were counted as the number of days from sowing until 50% of the ear emerges fully from the boot of flag leaf and anthesis occurred in 50% of the ear in each plot respectively. Days to maturity were counted as the number of days from sowing till the grains became hard enough and contained moisture levels near 12%. Test weight and yield/plant was recorded for each test genotype separately by taking weight in electric balance. Grains per spike were counted manually and panicle length was measured by a 30 cm long scale bar with 0.1 cm interval markings after harvest.

**Artificial inoculation of pathogen**

To impose an optimum disease pressure for thorough screening of test genotypes, artificial epiphytotic condition was created beside natural disease occurrence. A pure culture of aggressive *Bipolaris sorokiniana* isolate was obtained from Department of Plant Pathology, Bidhan Chandra Krishi Viswavidyalaya and maintained on potato dextrose agar (PDA) medium at 24 ± 1°C. For bulk propagation of spores, culture was multiplied on autoclaved sorghum (*Sorghum bicolor* L. Moench) grains for 20 days at 24 ± 1°C under 12 hours alternate dark and light cycle (Duveiller and Altamirano 2000). The spore concentration in the suspension was kept roughly 10^4 spores/ml. The suspension was sprayed uniformly as a mist in the field using hand atomizer at three different growth stages (GS) of the crop on Zadoks scale (Zadoks et al. 1974), namely- tillering (GS20), flag leaf emergence (GS37) and anthesis (GS65) during evening hours. The field was irrigated immediately after inoculation to maintain a high level of moisture for successful disease development.

**Assessment of spot blotch disease reaction**

The spot blotch disease score was recorded using double digit scale (Singh and Kumar 2005) at three growth stages viz, GS73 (early milking), GS77 (late milking) and GS83 (soft dough) of Zadoks scale. The first digit and second digit of double-digit score represent the percent leaf area covered by spot blotch infection on the flag leaf and the penultimate leaf respectively. The disease severity percent of each genotype was estimated by following formula (Duveiller et al. 2005):

\[ \text{Disease severity} \% = \frac{(D1/9) \times (D2/9)}{100} \]

Where,

- D1- first digit, refers to the vertical disease progress in accordance with the plant height
- D2- second digit, refers to severity measured as the extent of diseased leaf area

To have an idea about the progress of disease with time, area under disease progress curve (AUDPC) based on disease severity score was calculated using the following expression (Das et al. 1992):

\[ \text{AUDPC} = \sum_{i=1}^{n-1} \left[ \frac{1}{2} \left( Y_i + Y_{i+1} \right) \times \left( t_{i+1} - t_i \right) \right] \]

Yi is the disease severity measured on the ith date and \((t_i, t_{i+1})\) is the number of days in between two consecutive dates of disease scoring and n is the number of dates on which spot blotch was recorded.

**Statistical analysis**
The data obtained from the quantitative parameters of the 50 genotypes grown for two years were subjected to pooled analysis to consider the consistency of response of the test genotypes over the years. Analysis of variance (ANOVA) was performed for the characters under study. Estimates of genetic parameters like genotypic coefficient of variance (GCV) and phenotypic coefficient of variance (PCV) as per Burton (1952) and Burton and De Vane (1953), broad sense heritability ($h^2$) as per Hanson et al. (1956) and genetic advance as a percent of mean as per Johnson et al. (1955) were evaluated using the R-packages, version 3.6.1. Character association expressed in terms of correlation coefficient as per Al-Jiboari et al. (1958) and path coefficient analysis as suggested by Wright (1921) and discussed by Dewey and Lu (1959) were determined by Statistical Package for Agricultural Research (SPAR-I). Character with maximum influence on AUDPC was determined through multiple linear regression (MLR) using SPSS version 23.0. Genetic divergence among the genotypes was determined by the Mahalanobis’ generalized distance (Mahalanobis 1936) as per Rao (1952) using Genres software version 7.01. Skewedness for AUDPC among the genotypes was calculated using SPSS version 23.0.

### Results

Analysis of variance (Table 2) revealed significant differences among the fifty genotypes for all the ten quantitative characters. Heat map analysis (Fig. 1) arranged the genotypes and the characters into hierarchical clustering simultaneously based on similarity and distances between them and the pattern of colour mosaic indicated the association between the genotypes and the characters. High values (> 20%) of genotypic coefficient of variation (GCV) and phenotypic coefficient of variation (PCV) could not be recorded for any characters. However, among the characters the GCV and PCV values (Fig. 2) were highest for yield per plant followed by tillers per meter square and grains per spike. High heritability in broad sense (> 60%) was observed for all the characters excepting test weight, which was moderate (51.50%). The characters like plant height and days to maturity showed high heritability but their genetic advance (Fig. 2) as percentage of mean was low (< 10%). The characters, tillers per square meter, grains per spike and yield per plant showed high heritability and high genetic advance as percent of mean. Very high heritability and moderate genetic advance was observed for days to 50% heading (94.50 % and 14.43 % respectively) and days to 50% flowering (91.60 % and 12.11 % respectively) Moderate heritability (51.50 %) and low genetic advance as percent of mean (9.42 %) was recorded for test weight.

| Sources of variation | Degree of freedom | Plant height (cm) | Days to 50% heading | Days to 50% flowering | Tillers/sq.m | Days To maturity | Spike length (cm) | Grains per spike | Test weight (g) | Yield/plant (g) | AUDPC Score |
|----------------------|-------------------|------------------|---------------------|-----------------------|-------------|-----------------|------------------|-----------------|----------------|----------------|-------------|
| Genotype             | 49                | 43.86**          | 71.11**             | 64.24**               | 210.32**    | 39.29**         | 1.53**           | 80.90**         | 20.19**        | 1.86**         | 290442.4**  |

** significant at $P \leq 0.01$.

For character association study using correlation analysis (Fig. 3a and 3b), area under disease progress curve (AUDPC) was taken into consideration to understand its interrelation with yield per plant and yield components. Yield per plant showed significantly negative correlation (0.323* and 0.280*) with AUDPC at both genotypic and phenotypic levels. Significant positive correlation with yield per plant at both genotypic and phenotypic levels were observed for days to 50 % heading (0.394** and 0.331*), days to 50 % flowering (0.383** and 0.319*), tillers per square meter (0.530** and 0.343*), spike length (0.432** and 0.449**) and grains per spike (0.884** and 0.834**). Days to maturity revealed significant positive correlation with yield per plant at only genotypic level although the correlation was positive it was not significant. Among the characters registering significant positive correlation with yield per plant, only days to 50 % heading (-0.493** and - 0.464**), days to 50 % flowering (-0.455** and - 0.409**), tillers per square meter (-0.323* and 0.281*) and grains per spike revealed significant negative correlation with AUDPC at both genotypic and phenotypic level. Path coefficient analysis at both genotypic and phenotypic level (Fig. 4a and 4b) revealed that grains per spike had maximum direct positive effect on yield per plant (0.794 and 0.758 respectively) followed by days to 50 % heading (0.080 and 0.041 respectively) and days to 50 % flowering (0.079 and 0.040 respectively). The other characters having significant positive correlation with yield per plant had negative direct effect. The residual effect was low for both genotypic (0.02) and phenotypic (0.05) level. Multiple linear regression (Table 3) revealed days to 50 % heading to be most sensitive towards AUDPC. The predictability (in terms of adjusted $R^2$) of the MLR was highly significant being 0.661.

| AUDPC Score = $2883.350 - 30.867 \text{Days to 50\% heading}$ | $R^2$ | Adj $R^2$ | SE(est) |
|---------------------------------------------------------------|-------|-----------|---------|
| 0.698                                                         | 0.661 | 275.27    |         |

The genotypes could be grouped into 10 clusters (Table 4) based on $D^2$ square analysis. Cluster VI accommodated maximum genotypes (9), followed by cluster I accommodating 7 genotypes. Cluster II and cluster IX comprised of 6 genotypes each, while cluster X accommodated 5 genotypes and cluster VIII consisted of 3 genotypes while two genotypes grouped in cluster III and cluster IV, each. Maximum intra cluster distance was observed for cluster IV (9.23), followed by cluster VI (9.22), cluster X (8.93) and cluster I (8.26). The intra cluster distance of cluster II and cluster VI were also on a higher side being 7.29 and 7.91 respectively, while minimum intra cluster distance was recorded for cluster III (2.61) followed by...
cluster IV (2.73). Inter cluster distance was recorded to be maximum between cluster VII and cluster VIII (13.26), followed by cluster III and cluster V (12.03), cluster V and cluster VII (11.98) and cluster VIII and cluster X (11.68). Minimum inter cluster distance was registered between cluster II and cluster IV (6.53) followed by cluster II and cluster VII (7.86). Out of the 10 quantitative characters studied (Fig. 6), yield per plant contributed maximum (26.20%) towards total divergence, followed by days to 50% heading (23.92%), grains per spike (15.18%) and days to 50% flowering (14.94%). The contribution of plant height, tillers per square meter and spike length were not evident while the contribution of days to maturity and spike length were also less. The cluster mean analysis (Table 5) revealed the maximum cluster mean for days to 50% heading (73.00) and days to 50% flowering in cluster VIII while cluster IV exhibited highest mean for grains per spike (50.17) and yield per plant (6.18g). Minimum cluster mean for days to 50% heading (61.56) and days to 50% flowering (69.50) were registered in cluster VII and minimum cluster mean for grains per spike (41.78) and yield per plant were recorded in cluster VIII and cluster V respectively.

| CLUSTER | GENOTYPES |
|---------|-----------|
| I       | TERBOL    | ATLAS   | TESFA   | 30140   | 30081   | 29782   | 29641   |
| II      | 29882     | 30053   | 29889   | 29760   | 29761   | 29784   |
| III     | HD 2967 (C) | 29473   |         |         |         |         |
| IV      | 29824     | 29858   |         |         |         |         |
| V       | Sonalika (C) | 29872   | 29612   | 29703   |         |         |
| VI      | 29610     | 29821   | 29690   | 29672   | 29748   | 29506   | 29509   | 29522   | 29798   |
| VII     | 29865     | 30001   | HD3086 (C) | 29502   | 29752   | 30098   |
| VIII    | 29490     | 29477   | 29476   |         |         |         |
| IX      | 29769     | 29992   | 29526   | 29945   | 29988   | 29913   |
| X       | 29903     | 29812   | 29940   | 29671   | DBW107 (C) |         |

The genotypes designated as (C) has been used as check.

Table 4. Clustering of the genotypes based on D² statistics (pooled data of two years)

| CLUSTER | Plant height (cm) | Days to 50% heading | Days to 50% flowering | Tillers/sq.m | Days To maturity | Spike length (cm) | Grains per spike | Test weight (g) | Yield/plant (g) |
|---------|-------------------|---------------------|-----------------------|--------------|-----------------|-------------------|-----------------|----------------|----------------|
| I       | 90.78             | 66.05               | 73.10                 | 64.76        | 96.91           | 9.25              | 45.29           | 35.07          | 5.29           |
| II      | 94.66             | 69.56               | 77.44                 | 75.06        | 97.39           | 9.14              | 47.67           | 36.72          | 5.82           |
| III     | 86.90             | 70.67               | 78.50                 | 84.50        | 101.83          | 9.80              | 49.00           | 35.03          | 5.70           |
| IV      | 97.75             | 68.00               | 74.83                 | 81.83        | 98.00           | 9.08              | 50.17           | 37.07          | 6.18           |
| V       | 95.94             | 63.00               | 70.67                 | 68.58        | 95.75           | 8.22              | 40.83           | 34.51          | 4.67           |
| VI      | 95.05             | 68.59               | 75.96                 | 71.59        | 98.59           | 9.02              | 42.70           | 35.16          | 5.00           |
| VII     | 92.48             | 61.56               | 69.50                 | 70.78        | 96.00           | 9.38              | 44.28           | 35.37          | 5.17           |
| VIII    | 96.58             | 73.00               | 78.78                 | 77.44        | 105.67          | 8.59              | 41.78           | 36.56          | 5.05           |
| IX      | 94.95             | 67.44               | 74.11                 | 78.67        | 101.33          | 9.57              | 48.33           | 34.68          | 5.56           |
| X       | 89.89             | 65.60               | 73.20                 | 73.00        | 98.73           | 9.97              | 45.07           | 36.62          | 5.53           |

Table 5. Cluster mean analysis (pooled data of two years)

Analysis of variance (Table 1) of the set of test genotypes not only depicted the variation for agro-morphological traits but also for their disease severity. Frequency distribution of AUDPC among the genotypes showed slight positive skewness (0.052) for the variable (Fig. 7) which was not statistically significant suggesting more or less normal distribution of the character. No genotype showed complete resistance (disease score 00 according to double digit score classification by Kumar et al. 2016) against Bipolaris sorokiniana. Lowest AUDPC value was recorded in genotype 29882 (245) followed by 29610 (250), 29473 (265), 29940 (410). The AUDPC value of the susceptible check variety Sonalika was very high (961.67). Two genotypes 29872 (1378.33) and 30001 (1375) showed higher AUDPC value than that of Sonalika. Low AUDPC score with acceptable level of...
yield performance (Fig. 8) with respect to the yield checks HD 2967, HD3086 and DBW 107 were recorded for the genotypes 29882, 29610, 29473, 29940, 29477, 29748 and 30081.

Discussion

Significant variation among the genotypes for all the quantitative characters was the testimony of varied parentage of the genotypes taken under study as suggested earlier by Arya et al. (2017a) and different agro-climatic conditions from where the genotypes were obtained (Yadav et al. 2014). Significant variation was further reflected by heat mapping, which not only hierarchically clustered the genotypes according to their distance and similarity but also simultaneously clustered the characters under study and revealed the interaction between the genotypes and characters. Hierarchical clustering is a powerful tool to agglomerate a group of individuals into a cluster based on their similarity and distance. Heat map consists of rectangular tiling, with each tile shaded on a colour scale to represent the value of the corresponding element of the data matrix and high and low values are depicted with complementary colour code (Wilkinson and Friendly 2009). Rectangular matrix of similar colour along the row helps to identify the traits that appear to be characteristic for corresponding genotype cluster. Phenotypic coefficient of variation (PCV) was greater than genotypic coefficient of variation (GCV) for all the characters suggesting environmental influence on the genotypes for the expression of the characters (Kumar et al. 2013; Hazra et al. 2019b). Low to moderate values of GCV and PCV were recorded for the characters similar to the findings of Poudel et al. (2021) and among all the characters, the highest values having been registered for yield per plant corroborated with the earlier reports (Arya et al. 2017a). Coefficient of variation for tillers per square meter and grains per spike was higher compared to the other characters which agreed well to earlier reports (Kumar et al. 2013; Yadav et al. 2014). Revelation of moderate variability for yield per plant, tillers per square meter and grains per spike suggested that there was still chance of improving these traits by selection while low variability for the other characters indicated that there was hardly any opportunity for genetic enhancement of these characters through selection (Mitra et al. 2020) and induction of variability by means of hybridization or mutation followed by selection can improve the traits (Poudel et al. 2021).

Heritability depicts the percentage of variability that is transmitted from parents to offspring. However, heritability alone cannot provide a reliable picture for genetic gain (Arya et al. 2017a; Sejake et al. 2020) as evident for characters like plant height and days to maturity which registered high heritability (broad sense) but low genetic advance as percent of mean. Combination of these genetic variability parameters indicated that these characters might not be governed by additive gene action hence, direct selection will not be effective. High heritability combined with high genetic advance reflects governing of the characters by additive genes thus, direct selection for those characters could be rewarding (Poudel et al. 2021). High heritability coupled with high genetic advance as percent of mean recorded for tillers per square meter, grains per spike and yield per plant suggested direct and early generation selection for these characters (Bhanu et al. 2018; Al-Nager et al. 2020). Days to 50 % heading and days to 50 % flowering registered very high heritability with moderate genetic advance as percent of mean indicating selection for these traits might be beneficial through their phenotypic performance (Hailu 2020; Poudel et al. 2021), while moderate heritability and low genetic advance as percent of mean for test weight suggested direct selection for this trait would be non-rewarding.

Selection based on yield alone, a complex character is generally not very effective. Correlation studies provide important information to identify and verify whether the selection for a certain character influences another one, to quantify indirect gains due to selection of correlated traits and to evaluate the complexity of the traits (Tiwari and Upadhyay 2011). Correlation coefficients for most of the characters at genotypic level were higher than the corresponding coefficients at phenotypic level indicating the presence of inherent genetic relationship among the characters as suggested earlier (Tripathi et al. 2015). Yield per plant registering significant negative correlation with AUDPC at both genotypic and phenotypic levels suggested that spot blotch disease was a major biotic constraint for realizing high yield in wheat as reported earlier (Meena et al. 2014; Ayana et al. 2018). Days to 50 % heading, days to 50 % flowering, tillers per square meter, spike length and grains per spike revealed significant positive correlation with yield per plant at both genotypic and phenotypic level. Days to maturity showed significant positive correlation at genotypic level but it was not significant at phenotypic level. This might be possible since the genetic advance as percent of mean for days to maturity was low indicating the character to be governed by non-additive genes and hence influenced by environment (Addisu and Shumet 2015; Hossain et al. 2021). The characters, days to 50 % heading, days to 50 % flowering, tillers per square meter and grains per spike per square meter registered significant negative correlation with AUDPC at both genotypic and phenotypic level. The negative association between AUDPC and 50 % days to heading and 50 % days to flowering suggested a negative relation between the disease severity and duration of the crop (Mahto 2001; Sharma et al. 2006). Earlier reports also suggested negative correlation between disease severity and yield attributing parameters like, number of grains per spike and tillers per square meter (Gilchrist et al. 1991; Sharma et al. 1997; Sharma and Duveiller 2003, Singh et al. 2008). Non-significant correlation between AUDPC and plant height, spike length and test weight indicated a scope of simultaneous improvement of these traits and disease resistance. Selection of the characters having strong positive correlation with yield and simultaneous negative correlation with AUDPC would be rewarding in future wheat breeding programs (Sharma and Duveiller 2003). Correlation alone does not provide a clear picture of character association, since two characters might show correlation as a result of their correlation with a common third one (Poudel et al. 2021). Path coefficient analysis provides the actual information by splitting the correlation coefficients into measures of direct and indirect effects of the set of quantitative characters on yield per plant (Mecha et al. 2017). Grains per spike registered maximum direct positive effect on yield per plant at both genotypic and phenotypic level and its correlation with yield per plant was also very high. Days to 50 % heading and days to 50 % flowering also revealed positive direct effects although the values were very low. Other traits which had significant positive correlation with yield had negative direct effects. The low value of residual effect at both genotypic and phenotypic level suggested the inclusion of most of the responsible factors for grain yield per plant. AUDPC being complex and dependent on several factors, is difficult and often misleading to estimate empirically and data mining techniques in the form of MLR has been reported to be more accurate and is
becoming a new trend in understanding the sensitivity of several characters towards complex traits like AUDPC. (Nourani and Fard 2012). MLR extends linear modelling ideas to a wider class of response types, such as count data or binary responses (Sengupta et al. 2021) and is able to figure out the interrelationship among the input (Quantitative characters) and the output data (AUDPC) and predict each output with its corresponding output. MLR analysis revealed days to 50% heading to be most sensitive towards AUDPC in a negative direction and the MLR equation suggested that the character contributed 66.1% towards AUDPC. Sensitivity analysis using MLR was in accordance with the findings of correlation study which indicated days to 50% heading to be negatively correlated with AUDPC. Correlation study along with MLR validated the character days to 50% heading to be most important for spot blotch resistance screening which might be due to the fact that in South East Asian countries the prevalence of spot blotch coincides with the heading and post-heading stage of wheat (Chowdhury et al. 2013). In this context it may be pointed out that breeding for short duration crops will be rewarding since reduction in days to 50% heading can alleviate the impact of spot blotch by escaping the critical stage (flowering).

The genotypes could be grouped into 10 clusters indicating presence of divergence among them. The clustering pattern suggested no parallelism between genetic diversity and geographical origin as recorded earlier in soybean, Glycine max L. (Malik et al. 2011) and tomato, Solanum lycopersicon L. (Debnath et al. 2020). Grouping of the genotypes of same geographical origin into different clusters might be due to change in certain characters as a result of natural or artificial selection (Narayan et al. 2018). The intra and inter cluster distance among the genotypes indicates the distance among the genotypes in a single cluster and between the genotypes of different cluster respectively. The intra and inter cluster D² distance depicted the diversity present within the genotypes in a particular cluster and among the clusters, respectively. Maximum intra cluster distance was recorded for cluster IV, while minimum was recorded for cluster III. High intra cluster distance suggested the genotypes within the cluster had high degree of divergence and would produce desirable breeding materials for attaining maximum genetic advance (Dobariya et al. 2006; Chandramohan et al. 2016) whereas low intra cluster distance suggests presence of homogeneity among the genotypes within the cluster and selection of genotypes within the cluster would be ineffective. High inter cluster distance was recorded between cluster VII and cluster VIII, cluster III and cluster V, cluster V and cluster VII and cluster VIII and cluster X. Higher inter cluster distance suggested that the genotypes grouped in these clusters revealed broad spectrum of genetic diversity and can be utilised in future wheat breeding program to isolate desirable transgressive segregates for developing potential high yielding wheat varieties (Singh et al. 2010; Arya et al. 2017b). Low inter cluster distance between cluster II and cluster IV and cluster II and cluster VII depicted close relationship between the genotypes present in the clusters. Yield per plant, days to 50 % flowering, grains per spike and days to 50 % heading contributed majorly towards total divergence. The characters contributing maximum towards divergence might offer good scope of improvement through selection (Anuradha et al. 2020; Kumar et al. 2020). From this context, the cluster mean analysis of the major contributing characters revealed maximum cluster mean for the days to 50 % heading and days to 50 % flowering were in in cluster VIII, while the minimum cluster mean for these characters were recorded in cluster IV. Maximum cluster mean for grains per spike and yield per plant were registered in cluster IV and minimum cluster mean for these characters were in cluster VIII and cluster V respectively. The contrasting value of cluster mean for the characters were evident from the high inter cluster distances between the clusters and inter crossing among the genotypes from these clusters might be rewarding.

The disease reaction of wheat genotypes was characterized by their response measured as area under disease progress curve (AUDPC) as suggested earlier by Duveller et al. (1998) which significantly differed among the genotypes. Frequency distribution for AUDPC revealed more or less normal distribution suggesting that the variable was clustered more near the mean. This implied the presence of more number of moderately resistant/ moderately susceptible genotypes than that of highly susceptible or highly resistant ones. No genotypes showed complete resistance for the disease similar to the earlier reports on lack of resistance in the south Asian wheat cultivars by Siddique et al. (2006) and Sharma et al. (2006). AUDPC score of the susceptible check Sonalika was higher which was in accordance with earlier reports (Sharma et al. 2004; Sharma et al. 2006; Singh et al. 2007; Kumar et al. 2019). The genotypes 29872 and 30001 showing higher AUDPC value than Sonalika could be utilized as susceptible check variety only after screening their performance under high disease pressure in in-vitro and in-vivo conditions. Low AUDPC score along with acceptable level of yield performance with respect to the yield checks revealed by the genotypes 29882, 29610, 29473, 29940, 29748, 29748 and 30081 provided an opportunity to be used as a variety as such or as promising parent in hybridization programme.

The present investigation screened the genotypes having low AUDPC score for spot blotch disease along with high yield under hot spot region which presented a reliable picture on the disease reaction of the genotypes. Moreover, the present study aimed at understanding the interaction of the genotypes with the quantitative characters through heat map analysis and also developing a selection criterion for screening better genotypes through identification of important yield attributing characters with negative correlation with AUDPC with further validation using multiple linear regression model. This kind of selection approach for screening high yielding resistant lines ensures great deal of novelty.

**Conclusion**

Breeding of high yielding and spot blotch resistance is of utmost importance in wheat especially in the South Asian countries like India. In the present investigation, the characters 50% days to flowering, 50% days to heading and grains per spike came out as the major yield attributing characters since they showed high heritability coupled with high genetic advance as percent of mean, significant positive correlation with yield per plant and simultaneous significant negative correlation with AUDPC and also conferring positive direct effect to yield per plant in path analysis. Multiple linear regression (MLR) identified days to 50% heading to be most sensitive towards AUDPC in a negative direction indicating breeding for short duration to be rewarding. D² analysis revealed presence of wide divergence among the genotypes by grouping them into ten clusters. Presence of superior
genotypes in different clusters of varied diversity broadens the scope of using them in combinational breeding programme to exploit transgressive segregates in positive direction. Among the 50 genotypes, 29882, 29610, 29473, 29940, 29477, 29748 and 30081 revealed acceptable yield performances with low AUDPC value. Identification of high yielding and less susceptible genotypes for spot blotch disease in the present investigation offered an opportunity for wheat improvement through selective breeding.

**Declarations**

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The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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**Authors’ contributions**

All authors contributed significantly towards the final make-up of the paper. Conceptualisation (Anirban Maji and Soham Hazra); Data curation (Soham Hazra, Shouvik Gorai and Sudip Bhattacharya); Formal analysis (Pritam Roy, Shouvik Gorai and Sudip Bhattacharya); Investigation and methodology (Pritam Roy, Mousumi Murmu and Shouvik Gorai); Supervision (Anirban Maji, Dhiman Mukherjee, and Subhra Mukherjee); Writing-original draft (Soham Hazra and Shouvik Gorai); Writing-reviewing and editing (Anirban Maji).

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Research involving Human Participants and/or Animals:

This article does not contain any studies with human participants or animals performed by any of the authors.

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**Figures**

Heat map clustering analysis of 50 genotypes across 9 characters. The rows of a microarray heat map represent characters, and the columns represent the genotypes. Each cell is colorized based on the level of expression of that character in that sample. Here, G represents genotype followed by numerical as described in Table 1 and Ch represents characters as plant height (Ch1), days to 50% heading (Ch2), days to 50% flowering (Ch3), Tillers/sq.m (Ch4), Days To maturity (Ch5), Spike length (Ch6), Grains per spike (Ch7), Test weight (Ch8), and yield/plant (Ch9)
Figure 2

GCV, PCV, Heritability (broad sense), Genetic advance as percent of mean of the nine quantitative characters (pooled data of two years)

Figure 3

GCV, PCV, Heritability (broad sense), Genetic advance as percent of mean of the nine quantitative characters (pooled data of two years)
a. Genotypic correlation of yield, and yield contributing traits including AUDPC (pooled data of two years). Significance levels are provided by * (5%) and ** (1%) respectively. b. Phenotypic correlation of yield, and yield contributing traits including AUDPC (pooled data of two years). Significance levels are provided by * (5%) and ** (1%) respectively.

**Figure 4**

a. Genotypic path coefficient analysis indicating direct and indirect effects of the independent variables over yield (solid line represents direct effect, while dotted line refers to indirect effects, and the arrow signifies the relatable variables; as pooled data of two years). b. Phenotypic path coefficient analysis indicating direct and indirect effects of the independent variables over yield (solid line represents direct effect, while dotted line refers to indirect effects, and the arrow signifies the relatable variables; as pooled data of two years).
Figure 5

Inter cluster and intra cluster distance of the genotypes (not to scale); (pooled data of two years)

- Plant height (cm)
- Days to 50% heading
- Days to 50% flowering
- Tills/m²
- Days To maturity
- Spike length (cm)
- Grains per spike
- Test weight (g)
- Yield/plant (g)

Figure 6

Percentage contribution of the characters towards divergence(pooled data of two years)
Figure 7

Frequency distribution of AUDPC among the genotypes(pooled data of two years)

Figure 8

Yield per plot and AUDPC score of the selected genotypes as compared to the check varieties (pooled data of two years).