



## Genetic analysis for various yield components and gluten content in bread wheat (*Triticum aestivum* L.)

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**Abstract:** Genetic analysis was carried out in 55 genotypes (10 parents and 45 F<sub>1</sub>s) through diallel mating design excluding reciprocals in bread wheat (*Triticum aestivum* L.). Analysis of variance showed wide range of variability among the breeding material for all the traits under study. The highest value of phenotypic coefficient of variation (PCV) and genotypic coefficient of variation (GCV) were recorded for grain yield (PCV= 9.07 and GCV= 8.08). Highest heritability with genetic advance was recorded for grain yield ( $h^2=10.60$  and  $GA=14.84$ ), therefore selection will be effective based on grain yield for further study. Grains per spike ( $gr = 0.77$  and  $pr = 0.67$ ) and spikelets per spike ( $gr= 0.63$  and  $pr = 0.52$ ) were found significantly correlated (at <1 % level of significance) with grain yield whereas gluten content showed nonsignificant but positive correlation with grain yield at both genotypic as well as phenotypic level. Similarly, path coefficient analysis estimates for gluten content ( $g= 0.08$  and  $p= 0.03$ ) and grains per spike ( $g=0.36$  and  $p=0.23$ ) showed high positive direct effects on grain yield therefore these traits may be used as an index for selection to high yield in bread wheat genotypes.

**Keywords:** Correlation, Diallel analysis, Yield traits, Gluten content, *Triticum aestivum*

### INTRODUCTION

Bread wheat (*Triticum aestivum* L.) is an important cereal crop in India and play an important role in food security. India holds second position in both area and production after china and also planted to an area of 30.23 million hectares with production amounting to 93.50 million tons (Anonymous 2016) in India. Genetic analysis of wheat yield improvement had shown that grain yield is determined by different component traits, therefore, the knowledge of selection parameters and eye judgment of the breeder will apply after creation of genetic variability. Diallel analysis is most effective for ascertaining the systematic genetic architecture of metric traits within short periods. In the present study, heritability and genetic advance were considered as direct selection parameters, whereas correlation and path coefficient were regarded as indirect selection parameters. The genotypic and phenotypic coefficient of variation is helpful in understanding the clear cut picture of existing variability in the material. Several researchers have also reported earlier the importance of genetic variability and selection parameters for various yield components and quality traits in bread wheat (Singh *et al.*, 2012; Singh *et al.*, 2013, Meena *et al.*, 2014 and Kumar *et al.*, 2016) in the selection of better genotypes of wheat. Hence, in the context of yield improvement, when selecting desirable genotypes it is very important to know the

nature and extent of variation present within a set of breeding material as well as the interrelationship between each yield component and grain yield and the exact contribution of each component to yield *via* direct and indirect effects. It is known that the improvement of the genetic architecture of yield must be based on a more intensive study of individual yield components. Keeping this in view, the present study was undertaken to ascertain the extent of genetic variability, heritability, genetic advance and inter relationship of yield components to establish their implications in the selection of better genotypes of wheat.

### MATERIALS AND METHODS

Ten genotypes of bread wheat (PBW 435, HD 2967, MP 3336, MP 4010, DBW 90, HD 2824, HD 3095, RAJ 4246, NW 5038 and HD 2733) were sown during *rabi* season 2012-2013 for attempting crossing programme in a 10 x 10 diallel fashion excluding reciprocals. In the next crop season *rabi* 2013-2014, experimental material consisted 55 genotypes (10 parents and 45 F<sub>1</sub>s) was sown in a randomized block design with three replications. Each of the parental lines and crosses were sown by dibbling of seeds in two row plot of 2 meter length at spacing of 10 cm between plants within a row and keeping row to row spacing at 23 cm. All the standard agronomical practices (dose of fertilizer and irrigation at all critical stages) were followed

to raise normal crop from sowing to till harvest of crop. Observations were recorded on five randomly selected plants in each replication for days to maturity, plant height (cm), number of spikelets per spike, number of grains per spike, 1000 grain weight (g), grain yield (g) and gluten content (%). The analysis of variance (ANOVA) was based on the model suggested by Panse and Sukhatme, (1967). Heritability (in narrow sense) in  $F_{1s}$  generation was calculated as proposed by Crumpacker and Allard, (1962). The genetic advance was worked out by the formula proposed by Robinson *et al.* (1949). Genotypic and phenotypic coefficient of variation was calculated by the formula given by Burton and De vane, (1953). The genotypic and phenotypic correlation coefficients were calculated as suggested by Al-Jibouri *et al.* (1958). In path analysis; direct, indirect as well as residual effects were calculated. All the analysis work was done in Microsoft Excel as per the procedure given by Singh and Chaudhary (1985).

## RESULTS AND DISCUSSION

**Analysis of variance:** Analysis of variance (Table 2) among all the 55 treatments (parents and their  $F_{1s}$  crosses) showed highly significant differences (at 1 % level of significance) for the traits (days to maturity, plant height, spikelets per spike, grains per spike, 1000 grain weight and grain yield) in parents and in hybrids whereas (plant height, spikelets per spike, grains per spike, 1000 grain weight and grain yield) in parents vs. hybrids, indicating the presence of considerable amount of genetic variability in the present set of material and further genetic analysis would be meaningful. Similar findings on the existence of variability were also reported by Singh *et al.* (2012), Singh *et al.* (2013), Meena *et al.* (2014) and Kumar *et al.* (2016) for different yield and quality traits in wheat crop at different locations in India. The general mean of  $F_1$  crosses (Table 5) were greater than their corresponding

**Table 1.** Pedigree and other details of the parental lines used in crossing programme.

Parents	Species	Parentage/ pedigree	Origin/Source
PBW 435	T. aestivum	HD 2160/CALIDAD	Ludhiana
HD 2967	T. aestivum	ALD/COC//URES/HD2160M/HD2278	New Delhi
MP 3336	T. aestivum	HD2402/4W173	Jabalpur
MP 4010	T. aestivum	ANGOSTURA88	Jabalpur
DBW 90	T. aestivum	HUW468/WH 730	Karnal
HD 2824	T. aestivum	PTO-1/CNO79/PRL/GAA/3/HD 1951	New Delhi
HD 3095	T. aestivum	CPAN300/WR426//HW2007//HD2851	New Delhi
RAJ 4246	T. aestivum	RAJ 3765/WR544	Durgapura
NW 5038	T. aestivum	WAXWING*2/VIVITSI	Kumarganj
HD 2733	T. aestivum	ATTILA/3/TUI/CARC//CHENICHTO/4/ATTILA	New Delhi

**Table 2.** Analysis of variance for yield components and gluten content in bread wheat.

Source of variation	D. F.	Days to maturity	Plant height	Spikelets per spike	Grains per spike	Grain yield	Gluten content	1000 grain weight
Replication	02	0.26	0.27	0.25	0.32	1.26	0.42	3.84
Treatments	54	2.14**	31.28**	1.14**	10.19**	9.85**	0.16	3.58**
Parents	09	4.52**	66.84**	1.89**	13.56**	7.11**	0.23	4.75**
Hybrids	44	1.70**	22.31**	0.88**	8.18**	3.75**	0.15	2.38*
Parents Vs $F_{1s}$	01	0.16	105.46**	5.96**	68.40**	302.73**	0.18	45.96**
Error	108	0.76	0.73	0.10	1.15	0.78	0.15	1.58
Total	164	1.21	10.78	0.44	4.12	3.77	0.16	2.27

\*, \*\* Significant at 5 % and 1 % probability level, respectively.

**Table 3.** Genotypic and phenotypic correlation for yield components and gluten content in bread wheat.

Trait	r	Days to maturity	Plant height	Spikelets per spike	Grains per spike	Gluten content	1000 grain weight	Grain Yield
Days to maturity	G	1.0000	0.6383	0.0967	-0.0329	-0.2757	0.0346	0.0980
	P	1.0000	0.4217**	-0.0393	-0.0189	0.0113	0.0558	0.0118
Plant height	G		1.0000	-0.1899	-0.1548	-0.8577	0.1043	0.0954
	P		1.0000	-0.1533*	-0.1301	-0.1235	0.0694	0.0809
Spikelets per spike	G			1.0000	0.8754	0.5447	-0.3601	0.6355
	P			1.0000	0.6966**	0.1316	-0.1863*	0.5247**
Grains per spike	G				1.0000	0.9719	-0.3769	0.7726
	P				1.0000	0.0819	-0.1739*	0.6721**
Gluten content	G					1.0000	0.0777	1.0000
	P					1.0000	-0.0719	0.0834
1000 grain weight	G						1.0000	-0.3716
	P						1.0000	-0.0540

\*, \*\* Significant at 5 % and 1 % probability level, respectively.

**Table 4.** Direct and indirect effects of component traits on grain yield in bread wheat.

Trait	Path	Days to maturity	Plant height	Spikelets per spike	Grains per spike	Gluten content	1000 grain weight	Grain Yield
Days to maturity	G	0.4508	0.2877	0.0436	-0.0148	-0.1243	0.0156	0.0980
	P	-0.0150	-0.0063	0.0006	0.0003	-0.0002	-0.0008	0.0118
Plant height	G	0.0457	0.0716	-0.0136	-0.0111	-0.0614	0.0075	0.0954
	P	0.0078	0.0186	-0.0029	-0.0024	-0.0023	0.0013	0.0809
Spikelets per spike	G	0.0076	-0.0149	0.0786	0.0689	0.0428	-0.0283	0.6355
	P	0.0000	0.0001	-0.0004	-0.0003	-0.0001	0.0001	0.5247**
Grains per spike	G	0.0119	0.0560	-0.3167	0.3617	-0.3516	0.1363	0.7726
	P	-0.0045	-0.0308	0.1650	0.2369	0.0194	-0.0412	0.6721**
Gluten content	G	-0.0234	-0.0727	0.0462	0.0824	0.0847	0.0066	1.2968
	P	0.0003	-0.0037	0.0039	0.0025	0.0300	-0.0022	0.0834
1000 grain weight	G	-0.0058	-0.0175	0.0604	0.0632	-0.0130	-0.1678	-0.3716
	P	0.0026	0.0033	-0.0088	-0.0082	-0.0034	0.0473	-0.0540

\*, \*\* Significant at 5 % and 1 % probability level, respectively.

**Table 5.** Genetic parameters for yield components and gluten content in bread wheat.

Trait	Grand mean	Range	PCV	GCV	Narrow sense (Hb)	GA as % of means(5%)
Days to maturity	136.07	133.33 - 137.33	0.81	0.50	80.5	0.63
Plant height	90.56	80.20 - 95.57	3.65	3.52	83.5	7.01
Spikelets per spike	17.38	16.13 - 18.73	3.84	3.39	40.3	6.17
Grains per spike	52.53	48.07 - 55.97	3.88	3.31	4.88	5.80
Grain yield	21.51	16.64 - 24.23	9.07	8.08	10.6	14.84
Gluten content	8.53	7.87 - 8.97	4.63	0.61	15.5	0.17
1000 grain weight	42.75	40.90 - 45.25	3.51	1.91	9.3	2.15

\*, \*\* Significant at 5 % and 1 % probability level, respectively.

parents for (days to maturity, plant height, spikelets per spike, grains per spike, grain yield, gluten content and 1000 grain weight). Present results indicated that the variability in progenies which was obtained through hybridization could be properly utilized for selection of suitable genotypes on the basis of different genetic parameters. Similar findings were also reported by Singh *et al.* (2012), Singh *et al.* (2013) and Kumar *et al.* (2016) in wheat crop at different location of India.

**Estimation of variability:** Phenotypic coefficient of variation for all the traits was higher than the genotypic coefficient of variation (Table 5) which indicated the influence of environment in the expression of these metric traits. The highest value of phenotypic coefficient of variation (PCV) and genotypic coefficient of variation (GCV) were recorded for grain yield (PCV=9.07 and GCV=8.08). Whereas remaining traits namely days to maturity, plant height, gluten content, spikelets per spike, 1000 grains weight and grains per spike showed low value of PCV and GCV. Similar findings were also reported by Meena *et al.* (2014) and Kumar *et al.* (2016) for grain yield in wheat crop in India. Similar findings were also reported by Singh *et al.* (2013) for days to maturity, plant height and protein content; Meena *et al.* (2014) for grains per spike; Kumar *et al.* (2016) for 1000 grain weight in wheat crop. Present results revealed that useful variability in progenies may be utilized while making selection of suitable genotypes on the basis of different genetic parameters to improve grain yield in wheat.

**Heritability and genetic advance:** In the present study (Table 5), high heritability in narrow sense

(more than 60 %) was observed for days to maturity (80.5) and plant height (83.5). While for remaining traits like spikelets per spike, grains per spike, 1000 grains weight, gluten content and grain yield were showed moderate to low heritability. Similar findings were also reported by Singh *et al.* (2012) for days to maturity; Singh *et al.* (2013) and Meena *et al.* (2014) for plant height; Emeka *et al.* (2016) for days to maturity and plant height and Kumar *et al.* (2016) for spikelets per spike in wheat crop at different locations in India. Highest genetic advance (more than 10 %) was recorded for grain yield (14.84) indicating that selection based on grain yield would be effective for improvement of wheat crop. Similar findings pertaining to high genetic advance were also reported by Emeka *et al.* (2016) and Kumar *et al.* (2016) for grain yield in wheat crop at different location of India.

Estimates of high heritability coupled with high genetic advance of a character are more effective for making selection due to the fact that expression of these traits is controlled by additive gene action. Highest heritability along with highest genetic advance was recorded for grain yield ( $h^2=10.6$  and  $GA=14.84$ ) revealing the presence of additive and additive x additive type of gene effects in the expression of grain yield. Likewise, high heritability coupled with high genetic advance was also reported by Singh *et al.* (2012) and Kumar *et al.* (2016) in wheat. Therefore, present results indicated that useful variability in progenies developed through hybridization can be properly utilized and selection of suitable genotypes on the basis of different genetic parameters can be done to get high yield and

gluten content in bread wheat.

**Correlation coefficient:** A close relationship between genotypic and phenotypic correlations (Table 3) in most of the traits indicating low environmental effects on the degree of association between various yield components. Genotypic correlation coefficients were, in general, slightly higher than the corresponding phenotypic correlation coefficient for almost all the traits under study. The improvement of yield components and the knowledge of their association with its main component are beneficial in formulating the breeding programme. Correlation coefficients were estimated between grain yield and other component traits revealed that grain yield was highly significant and positively associated with number of grains per spike ( $r_g=0.77$  and  $p_r=0.67$ ) and number of spikelets per spike ( $r_g=0.63$  and  $p_r=0.52$ ) at both genotypic as well as phenotypic level (Table 3) indicating that grain yield and these traits has the same physiological basis for their expression. Similar findings of correlation coefficient were also earlier reported by Meena *et al.* (2014) and Kumar *et al.* (2016) for spikelets per spike and grains per spike in wheat crop at different locations of India. Positive association between spikelets per spike and grains per spike was reported by Meena *et al.* (2014) in wheat crop. It has been observed that number of grains per spike and number of spikelets per spike were positively associated with grain yield. Hence, selection for these traits would also helps in improving grain yield. In the present investigation, the traits which exhibited direct positive correlation with grain yield may be considered as most important traits combinations for yield improvement in bread wheat through selection.

**Path coefficient:** In path coefficient analysis, selection may be done on the basis of direct effects of the components traits towards grain yield for improvement of wheat crop. The results of path coefficient (Table 4) indicated that three yield traits (gluten content, grain per spike and plant height) showed positive phenotypic and genotypic direct effects on grain yield whereas, rest of the traits showed negative direct effect on grain yield. Present finding were similar to the findings of Kumar *et al.* (2016) for days to maturity, grain per spike and plant height at different locations of India in wheat.

## Conclusion

Based on the above results, it may be concluded that grain yield in wheat crop showed highest heritability with highest genetic advance and also have highest PCV and GCV therefore selection will be effective based on grain yield due to the maximum contribution

of additive gene action. Spikelets per spike and grains per spike showed positive significant correlation and also have positive direct effects on grain yield at both genotypic as well as phenotypic level, indicating that we might improve grain yield by improving these traits and thus emphasis should be given on these traits for the selection of elite genotypes from the segregating generations. It is evident that genotypes developed might serve as good source of material for further breeding programme.

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