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# GxE Interactions Analysis of Wheat Genotypes Evaluated Under Peninsular Zone of the Country by AMMI Model

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**To cite this article:**

Ajay Verma, Gyanendra Pratap Singh. GxE Interactions Analysis of Wheat Genotypes Evaluated Under Peninsular Zone of the Country by AMMI Model. *American Journal of Agriculture and Forestry*. Vol. 9, No. 1, 2021, pp. 29-36. doi: 10.11648/j.ajaf.20210901.15

**Received:** December 7, 2020; **Accepted:** December 31, 2020; **Published:** January 30, 2021

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**Abstract:** AMMI analysis of wheat genotypes had highlighted significant effects of environments, interactions and genotypes for the 2017-18 and 2018-19. Number of adaptability measures had been studied as per utilization of number of significant interaction principal components (IPCs). Total of interaction variations exploited by Type-1, 2, 3, 4 & 5 measures were 45.5%, 66.3%, 75.9% & 88.4% respectively. Type-1 measures EV1, D1, ASTAB1 identified (G7, G6, G12) genotypes while SIPC1 selected (G14, G17, G2). EV2, D2, ASTAB2, ASV and ASV1 measures found (G7, G6, G4) as desirable genotypes. Analytic measures based on all significant IPCA's i.e. MASV and MASV1 settled for G6, G7, and G3. Adaptability measures GAI, HM, PRVG & MHPRVG observed G13, G4, and G12 genotypes would be of stable adaptations. Biplot analysis seen largest cluster comprised D3, D5, EV2, EV3, EV5, ASTAB3, ASTAB5, MASV1, MASV and Standard deviation measures. Genotypes were ranked G9, G11, and G6 by values of EV1, D1 & ASTAB1 for second year of study. D2, ASV, ASV1, EV2 & ASTAB2 observed (G9, G6, and G11) as adaptable genotypes. MASV & MASV1 measures also supported G9, G6, G11 genotypes for the considered locations of the zone. Studied measures were clustered in three groups in graphical analysis. Three clusters were observed among studied measures by biplot analysis. Measures EV1, EV2, EV3, D1, D2, D3, ASV, ASV1, MASV, MASV1, ASTAB1, and ASTAB2 & ASTAB3 formed largest cluster.

**Keywords:** AMMI, BLUP, PRVG, MHPRVG, Biplots

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## 1. Introduction

Very well wide adaptation of wheat made it possible to cultivate an important cereal in most of the countries [18]. Multi location trials have been conducted to evaluate the yield performance of several genotypes simultaneously [7]. Field evaluation of genotypes have required an efficient analytic estimation procedure for GxE interactions [1]. Presence of cross over type GxE interactions mask the real potential of deserving genotypes for their specific and general adaptations [17]. Quite large number of analytic approaches have developed especially for adaptation behavior of genotypes [15, 2]. Univariate parametric, non-parametric, multivariate models for additive and multiplicative nature of factors. Over exploited method Additive Main effects and Multiplicative Interaction (AMMI) advocated in agricultural research field experiments [8]. Large portion of the interactions sum of squares had been utilized by AMMI analysis to discriminate environments,

adaptability of the genotypes to specific and general environmental conditions to harvest well yield [5]. This analysis mechanism has proved as an effective analysis with possible opportunities to research workers [6]. The current study was planned with clear objectives (i) Study number of AMMI based measures as per utilization of number of significant interaction principal components (ii) similarity & dissimilarity among adaptability measures.

## 2. Materials and Methods

Peninsular zone comprises mainly of Maharashtra and Karnataka states of our country. Major three species of wheat viz *T. aestivum*, *T. durum*, and *T. dicoccum* are cultivated in this zone. Bread wheat cultivation is concentrated under irrigated environments, whereas, the cultivation of durum and dicoccum wheat is generally confined to rainfed/

restricted irrigation situation. Genotype by environment (GxE) interactions of seventeen advanced wheat genotypes at thirteen major locations during 2017-18 and eleven genotypes at eleven locations in 2018-19 cropping season of the zone by AMMI model. Research field trials were conducted at centers of AICRP by randomized complete block designs with four replications. Recommended agronomic practices were followed to harvest good yield. Details of genotype parentage along with environmental conditions were reflected in tables 1 & 2 for ready reference. AMMI first calculate genotype and environment additive effect using analysis of variance (ANOVA) and then analyse residual from these model using principal components analysis (PCA). AMMI stability value

(ASV) proposed by Purchase [11] to quantify the stability measure by considering relative weight of  $IPCA_1$  and  $IPCA_2$  scores. In certain cases where more than two IPCAs were significant, ASV failed to encompass all the variability explained by GxE interactions. Zali[20] attempted modified version ASV which would cover all available Interaction Principal Components. But in doing so, Zali interpreted the formula of ASV incorrectly compared to the original formula of Purchase [11, 12]. In the present study the original MASV formula of Zali[20] and a revised version of MASV [2] were compared with other AMMI based measures of interaction effects. The description of widely used measures based on AMMI analysis was mentioned for completeness.

Table 1. AMMI analysis based measures

Zobel [21]	1994	EV1	EVF	$EV = \sum_{n=1}^N \lambda_{in}^2 / n$
Sneller et al [16]	1997	SIPC1	SIPCF	$SIPC = \sum_{n=1}^N \lambda_n^{0.5} \gamma_{in}$
Purchase et al [12]	2000	ASV		$ASV = [(\frac{SSIPC1}{SSIPC2} PCI)^2 + (PC2)^2]^{1/2}$
Annicchiarico [4]	1997	D		$D = \sqrt{\sum_{n=1}^N (\lambda_n \gamma_{in})^2}$
Rao and Prabhakaran [13]	2005	ASTAB		$ASTAB = \sum_{n=1}^N \lambda_n \gamma_{ni}^2$
Zali et al [20]	2012	ASV1		$ASV = [(\frac{SSIPC1}{SSIPC2} PCI)^2 + (PC2)^2]^{1/2}$
Zali et al [20]	2012	MASV		$MASV = \sqrt{\sum_{n=1}^{N-1} \frac{SSIPC_n}{SSIPC_{n+1}} (PC_n)^2 + (PC_{n+1})^2}$
Ajay et al [2]	2019	MASV1		$MASV1 = \sqrt{\sum_{n=1}^{N-1} (\frac{SSIPC_n}{SSIPC_{n+1}} PC_n)^2 + (PC_{n+1})^2}$

Table 2. Parentage of wheat genotypes and environmental conditions (2017-18).

Code	Genotype	Parentage	Code	Environments	Latitude	Longitude	Mean sea level
G1	AKAW 4924	DL-9-65-2/AKW1071-1-2	E1	Arabhavi	15° 84 ' N	74° 51' E	625m
G2	GW 491	HD2808/HI1516/PBW573	E2	Dharwad	15° 27' N	75° 0' E	724 m
G3	GW 493	HW2045/HI 1183/PCE2555	E3	Kalloli	16° 26 ' N	74° 86' E	625m
G4	DBW 235	MELON//FILIN/MILAN/3/FILIN/4/TRCH/SRTU//KACHU	E4	Karad	17° 17 ' N	74° 10' E	577 m
G5	HI 1624	GW322/PBW498	E5	Kolhapur	16° 41 ' N	74° 14' E	578 m
G6	GW 495	LOK54/RAJ4083	E6	Mahabaleshwar	17° 55 ' N	73° 39' E	1322 m
G7	MP1338	MILAN/KAUZ//DHARWAR DRY/3/BAV92/4/PAURAO	E7	Mudhol	16° 19 ' N	75° 17' E	546 m
G8	HI 8800	HI8681/HI8663	E8	Nashik	19° 59 ' N	73° 47' E	583 m
G9	MACS 6709	ROLF07/4/BOW/NKT//CBRD/3/CBRD/5/FRET2/TUKURU//FRET2	E9	Niphad	20° 4 ' N	74° 6' E	551 m
G10	HI 1625	GAINT3/HW2045	E10	Nippani	16° 23 ' N	74° 22' E	606 m
G11	PBW 770	PBW585/4/BABAX//IRENA/KAUZ/3/HUITES	E11	Parbhani	19° 15 ' N	76° 46' E	413 m
G12	GW 492	RAJ4040/HD2808	E12	Pune	18° 31 ' N	73° 51' E	562 m
G13	MACS 6222	HD2189*2/MACS2496	E13	Ugar - Khurd	16° 39 ' N	74° 49' E	548 m
G14	DBW 168	SUNSU/CHIBIA					
G15	MACS 3949	STOT//ALTAR84/ALD/3/THB/CEP7780//2*MUSK_4					
G16	MACS 6478	CS/TH.SC//3*VN/3/MIRLO/BUC/4/MILAN/5/TILHI					
G17	UAS 428	GREEN-14/YAN-10/AUK/UAS402					

Table 3. Parentage details of genotypes along with environmental conditions (2018-19).

Code	Genotype	Parentage	Code	Environments	Latitude	Longitude	Mean sea level
G1	PBW 823	T. boeoticum 4992/2*PDW274//2*PBW703	E1	Niphad	20° 4 ' N	74° 6' E	551 m
G2	UAS 428	GREEN-14/YAN-10/AUK/UAS402	E2	Pune	18° 31 ' N	73° 51' E	562 m
G3	DDW 49	PDW314/PDW233	E3	Akola	20° 42' N	77° 0' E	292 m
G4	UAS 3001	UAS259/GW322//HI 977	E4	Parbhani	19° 15 ' N	76° 46' E	413 m
G5	MACS 3949	STOT//ALTAR84/ALD/3/THB/CEP7780//2*MUSK_4	E5	Nasik	19° 59 ' N	73° 47' E	583 m
G6	MACS 6222	HD2189*2/MACS2496	E6	Kolhapur	16° 41 ' N	74° 14' E	578 m
G7	GW 322	PBW173/GW196	E7	Dharwad	15° 27' N	75° 0' E	724 m

Code	Genotype	Parentage	Code	Environments	Latitude	Longitude	Mean sea level
G8	DDW 48	HI8498/PDW233//PDW291	E8	Arbhavi	15° 84 'N 74° 51' E		625m
G9	MACS 6478	CS/TH.SC//3*VN/3/MIRLO/BUC/4/MILAN/5/TILHI	E9	Kalloli	16° 26 'N 74° 86' E		625m
G10	HD 3343M	HI1500/2*GW322 BCRIS/BICUM/LLARETA	E10	Nippani	16° 23 'N 74° 22' E		606 m
G11	WHD 963	INIA/3/DUKEM_12/2*RASCON_21/5/SOMAT_3/GREEN_22/4/ GODRIN/GUTROS/DUKEM/3/THKNEE_11	E11	Ugar-Khurd	16° 39 'N 74° 49' E		548 m

AMMI analysis was performed using AMMISOFT version 1.0, available at <https://scs.cals.cornell.edu/people/hugh-gauch/> and SAS software version 9.3. AMMI based measures were compared with recent analytic measures of adaptability calculated as the relative performance of genetic values (RPGV) and MHGV (Harmonic mean of Genetic Values), based on the harmonic mean of the genotypic values across different environments. Another harmonic mean based measure of the relative performance of the genotypic values (MHRPGV) for the simultaneous analysis of stability, adaptability and yield [14].

$$RPGV_{ij} = GV_{ij} / GV_i$$

$$MHGV_i = \text{Number of environments} / \sum_{i=1}^k \frac{1}{X_i}$$

$$MHRPGV_i = \text{Number of environments} / \sum_{j=1}^k \frac{1}{RPGV_{ij}}$$

$GV_{ij}$  is the genotypic value of the  $i$  genotype, in the  $j$  environment, expressed as a proportion of the average in this environment. Geometric adaptability index (GAI) [10] was calculated as  $\sqrt[n]{\prod_{k=1}^n \bar{X}_k}$ ; in which  $\bar{X}_1, \bar{X}_2, \bar{X}_3, \dots, \bar{X}_m$  are the mean yields of the first, second and  $m$ th genotype across environments and  $n$  is number of environments. Genotypes with higher values of GAI are desirable.

### 3. Results and Discussion

Better understanding of the GxE interaction had been provided by AMMI analysis as this facilitated identification of general and specific adaptations of genotypes and discriminate environments. In fact AMMI exercised family of models with retaining 0, 1, 2, or more significant interaction principal components (IPCs).

#### 3.1. First Year of Study (2017-18)

Estimated sums of squares for GxE signal and GxE noise were 69.9% and 30.2% respectively

Sum of squares for GxE signal is 2.33 times that for genotypes main effects, implied, narrow adaptations are important for trials research dataset. First IPC1 alone is 1.52 times the genotypes main effects whereas GxE noise is 1.01 times the genotypes main effects. Discarding noise improves accuracy, increases repeatability, simplifies conclusions, and accelerates progress [6]. Highly significant environments (49.9), GxE interaction (21.4) and genotypes (6.4) were observed by ANOVA analysis. Diversity of considered locations had justified the selection of environments [3]. Explained variation of GxE interaction accounted by each of highly significant IPCA's, as type-1 measures benefited 45.5%, type-2 measures utilized 66.3%, type 3 measures used up to 75.9%, type-5 measures used up to 88.4% of interaction variations, tough IPCA5, IPCA6 and IPCA7 contributed to the tune of 5.5, 3.9 and 2.9% respectively (Table 4). Use of AMMI derived measures upto first five IPCAs had been justified [9]. Smaller and large values of EV1 observed for (G7, G6, G12) and (G14, G11) while corresponding to D1 were (G7, G6, G12) and (G14, G11), SIPC1 for (G14, G17, G2) and (G11, G15) and for ASTAB1 were (G7, G6, G12) & (G14, G11). EV2 measure pointed towards (G6, G7, G4) as desirable along with undesirable genotypes as (G15, G11) for values of D2 genotypes were (G6, G7, G7) & (G15, G1), whereas as per criterion of SIPC2 were (G2, G14, G5) & (G15, G12) and ASTAB2 values favoured (G6, G7, G4) & (G15, G11) (Tables 6 and 7).

Table 4. AMMI analysis of wheat genotypes evaluated in Peninsular Zone (2017-18).

Source	Degrees of freedom	Mean Sum of Squares	Level of significance	% of Total SS	% of GxE SS	Cumulative% SS by PCA's
Treatments	220	299.70	***	77.70		
Genotypes	16	340.54	***	6.42		
Environments	12	3526.27	***	49.87		
GxE interactions	192	94.64	***	21.41		
IPC1	27	306.00	***		45.47	45.47
IPC2	25	151.14	***		20.79	66.26
IPC3	23	76.03	***		9.62	75.89
IPC4	21	61.18	*		7.07	82.96
IPC5	19	52.36			5.47	88.43
IPC6	17	41.75			3.91	92.34
IPC7	15	35.34			2.92	95.26
Residual	45	19.15				
Error	663	28.54				
Total	883	96.10				

Table 5. AMMI analysis of wheat genotypes evaluated in Peninsular Zone (2018-19).

Source	Degrees of freedom	Mean Sum of Squares	Level of significance	% of Total Sum of Squares	% of GxE Sum of Squares	Cumulative% SS by PCA's
Treatments	120	259.51	***	75.22		
Genotypes	10	445.46	***	10.76		
Environments	10	1823.52	***	44.05		
GxE interactions	100	84.51	***	20.42		
IPC1	19	227.71	***		51.19	51.19
IPC2	17	75.32	***		15.15	66.34
IPC3	15	72.84	*		12.93	79.27
IPC4	13	55.23			8.50	87.77
IPC5	11	44.54			5.80	93.56
IPC6	9	33.63			3.58	97.14
IPC7	7	20.63			1.71	98.85
Residual	9	10.76				
Error	363	28.25				
Total	483	85.71				

Table 6. Principal components analysis of genotypes (2017-18).

17-18	EV1	EV2	EV3	EV5	D1	D2	D3	D5	IPC1	IPC2	IPC3	IPC4	ASV	ASV1	MASV	MASV1
G 1	0.0236	0.0153	0.0102	0.0150	4.53	4.88	4.88	5.76	-1.4656	0.6547	0.0220	-1.1971	2.26	3.27	2.74	3.76
G 2	0.0344	0.0391	0.0271	0.0275	5.46	7.14	7.19	7.87	-1.7694	-1.6394	0.3619	-0.4245	3.09	4.20	3.98	5.55
G 3	0.0099	0.0050	0.0144	0.0180	2.92	2.93	4.19	5.09	-0.9465	0.0541	-1.1786	0.2523	1.40	2.07	2.30	2.89
G 4	0.0051	0.0037	0.0167	0.0228	2.10	2.36	4.14	5.51	0.6787	0.3855	1.3368	-1.3635	1.08	1.53	2.75	3.16
G 5	0.0091	0.0387	0.0328	0.0216	2.81	6.39	6.82	6.95	-0.9094	-2.0497	-0.9331	0.4016	2.45	2.86	4.16	5.52
G 6	0.0002	0.0034	0.0095	0.0090	0.39	1.84	3.04	3.52	0.1274	0.6406	-0.9531	0.4511	0.67	0.70	1.92	2.28
G 7	0.0001	0.0035	0.0116	0.0257	0.23	1.84	3.30	5.41	-0.0748	0.6528	-1.0767	-1.3016	0.66	0.67	2.41	2.73
G 8	0.0142	0.0431	0.0287	0.0262	3.51	6.86	6.86	7.49	1.1360	-2.1040	0.0032	-1.0438	2.69	3.26	4.23	5.69
G 9	0.0085	0.0373	0.0355	0.0247	2.72	6.26	6.91	7.16	-0.8798	2.0155	-1.1524	-0.6118	2.40	2.79	4.25	5.56
G 10	0.0517	0.0369	0.0279	0.0522	6.70	7.45	7.62	9.66	2.1685	-1.1640	0.6426	1.9022	3.41	4.88	4.38	5.91
G 11	0.0830	0.0612	0.0463	0.0335	8.48	9.53	9.77	10.07	2.7459	-1.5576	-0.8306	-0.9904	4.35	6.20	5.17	7.26
G 12	0.0019	0.0283	0.0564	0.0410	1.30	5.29	7.65	8.10	0.4210	1.8329	2.1694	-0.8676	1.94	2.05	4.78	5.84
G 13	0.0044	0.0110	0.0252	0.0380	1.96	3.50	5.17	7.02	0.6353	1.0364	1.4972	1.4594	1.40	1.73	3.42	4.07
G 14	0.0857	0.0456	0.0311	0.0338	8.62	8.77	8.80	9.62	-2.7918	-0.5804	-0.2872	1.2459	4.17	6.13	4.46	6.40
G 15	0.0787	0.0769	0.0737	0.0509	8.26	10.22	11.08	11.39	2.6753	2.1494	-1.6781	1.0840	4.50	6.23	6.17	8.34
G 16	0.0097	0.0096	0.0219	0.0219	2.90	3.60	5.06	5.78	0.9401	-0.7614	1.3962	0.2282	1.59	2.19	2.90	3.62
G 17	0.0797	0.0414	0.0311	0.0383	8.31	8.40	8.56	9.56	-2.6910	0.4344	0.6605	0.7755	4.00	5.90	4.25	6.13

EV = Eigenvector, SIPC = Sum of the value of the IPC Scores, D = Parameter of Annicchiarico; SIPC1 = SIPC for first IPCA, SIPC 2 = SIPC for first two IPCAs., IPCA, principal component of interaction, ASV = AMMI stability value; MASV = Modified AMMI stability value; ASTAB = AMMI stability

Table 7. AMMI based estimates of genotypes (2017-18).

	SIPC1	SIPC2	SIPC3	SIPC5	ASTAB1	ASTAB2	ASTAB3	ASTAB5	MEAN	STDEV	GAI	RPGV	MHRPGV	HM
G 1	-1.46556	-0.81085	-0.78883	-2.35806	20.48	23.84	23.84	33.20	44.30	7.95	43.68	0.9452	0.9388	43.10
G 2	-1.7694	-3.4087	-3.0468	-2.1993	29.85	50.92	51.77	61.93	49.01	10.26	48.02	1.0405	1.0313	47.07
G 3	-0.9465	-0.8923	-2.0709	-3.0099	8.54	8.56	17.55	25.90	46.37	6.39	45.96	0.9935	0.9890	45.54
G 4	0.6787	1.0642	2.4010	0.4187	4.39	5.56	17.11	30.39	49.70	7.60	49.19	1.0634	1.0585	48.69
G 5	-0.9094	-2.9592	-3.8923	-3.8936	7.89	40.83	46.46	48.33	45.93	9.18	45.12	0.9771	0.9693	44.34
G 6	0.1274	0.7680	-0.1851	0.8515	0.15	3.37	9.25	12.39	48.19	7.05	47.73	1.0305	1.0282	47.27
G 7	-0.0748	0.5780	-0.4986	-0.5864	0.05	3.39	10.89	29.31	48.09	8.86	47.35	1.0236	1.0189	46.64
G 8	1.1360	-0.9680	-0.9648	-2.6765	12.30	47.01	47.01	56.04	44.64	8.89	43.85	0.9499	0.9421	43.11
G 9	-0.8798	1.1357	-0.0166	-1.0811	7.38	39.23	47.82	51.21	45.09	7.31	44.55	0.9653	0.9561	44.03
G 10	2.1685	1.0045	1.6471	5.0999	44.83	55.45	58.12	93.29	48.95	9.29	48.12	1.0442	1.0319	47.28
G 11	2.7459	1.1884	0.3578	-0.4409	71.89	90.91	95.37	101.45	49.65	10.75	48.61	1.0555	1.0418	47.63
G 12	0.4210	2.2539	4.4233	4.2427	1.69	28.03	58.46	65.62	49.45	8.85	48.77	1.0560	1.0479	48.14
G 13	0.6353	1.6718	3.1689	3.3110	3.85	12.27	26.77	49.26	49.66	7.07	49.19	1.0645	1.0573	48.71
G 14	-2.7918	-3.3722	-3.6594	-3.4256	74.31	76.95	77.48	92.53	40.50	8.99	39.59	0.8625	0.8452	38.68
G 15	2.6753	4.8247	3.1466	4.3196	68.23	104.46	122.67	129.75	45.83	9.33	44.92	0.9805	0.9568	43.98
G 16	0.9401	0.1787	1.5750	0.6470	8.43	12.97	25.58	33.40	44.72	8.25	44.03	0.9528	0.9463	43.34
G 17	-2.6910	-2.2566	-1.5961	0.7810	69.04	70.52	73.34	91.35	46.70	9.49	45.85	0.9950	0.9829	45.05

ASV and ASV1 observed suitability of (G7, G6, G4) along with unsuitable performance for G15, G11 (Table 6). Considering first two IPCAs in ASV & ASV1 measures utilized 66.3% of GxE interaction sum of squares. The two

IPCAs have different values and meanings and the ASV and ASV1 parameters using the Pythagoras theorem and to get estimated values between IPCA1 and IPCA2 scores to produce a balanced measure between the two IPCA scores [11, 12].

Also, ASV and ASV1 measures used advantages of cross validation due to computation from first two significant IPCAs. Total of 75.9% of Gx $\times$ E interaction utilized by Type-3 measures, values of EV3 preferred G6, G1, G7 and unstable performance of G15, G12 whereas D3 pointed towards G6, G7, G4 and G15, G11 whereas SIPC3 pointed towards G5, G14, G2 & G12, G13; ASTAB3 measure considered G6, G7, G4 & G15, G11 (Tables 5 & 6). Genotypes G6, G1, G4 pointed out by least values EV5 and maximum values found for G10, G15, SIPC5 measure identified G2, G8, G10 and G9, G4 whereas D5 considered G6, G4, G7 as suitable & G15, G11 as unsuitable ones; ASTAB5 selected G6, G3, G7 as suitable & G15, G11 as unsuitable genotypes. Composite measures MASV selected G6, G3, G7 genotypes and G15, G11 however

MASV1 selected G6, G7, G3 for stable performance and G15, G11 would not be recommended for cultivation due to unstable yield behavior. Highly significant differences had been observed among genotypes yield; average yield was considered as an important measure to assess the potential yield of genotypes. G4, G13 and G11 wheat genotypes achieved higher yield as compared to G14, G1. Adaptability index (GAI) pointed towards G13, G4, G12 as of stable adaptations whereas G14 & G1 would be of unstable nature. HM and PRVG measures identified G13, G4, G12 and G14 & G1 for general and specific adaptations. MHPRVG found G4, G13, G12 as genotypes for broad adaptations for studied environmental conditions and G14, G1 for specific conditions.

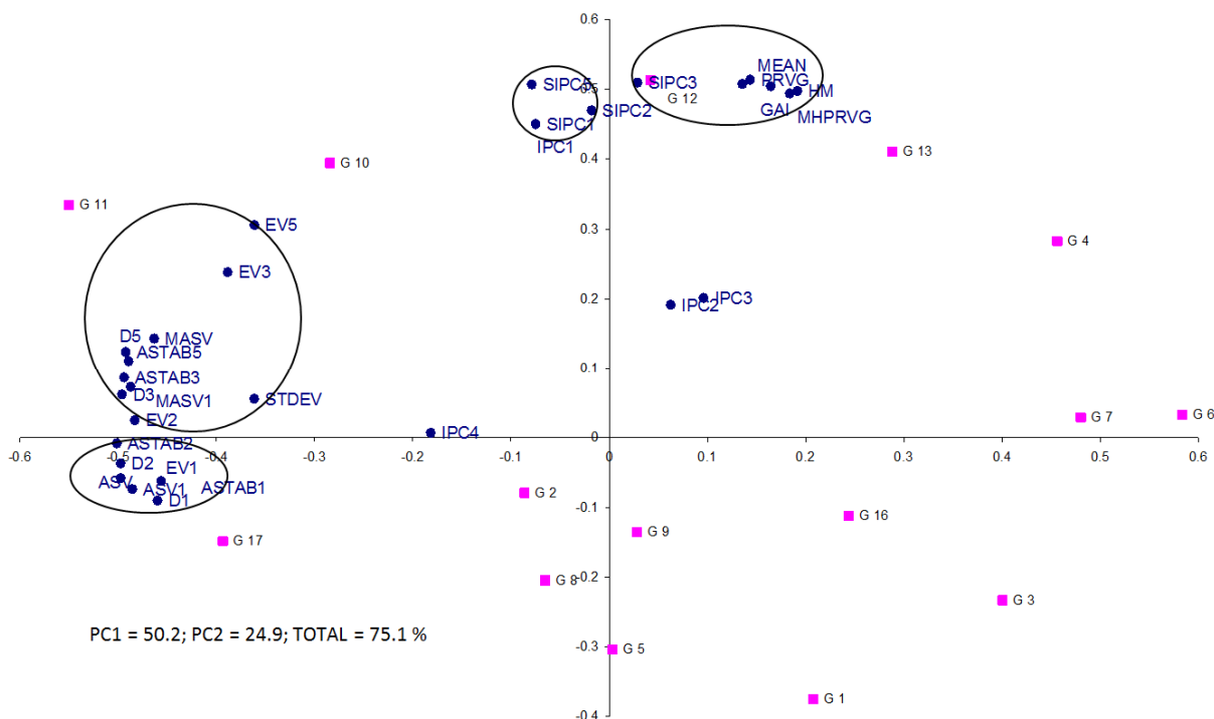


Figure 1. Biplot analysis of AMMI and yield based estimates (2017-18).

Biplot analysis based on first two significant principal components helped to visualize the relationships among AMMI and yield based other measures [19]. In total, PC1 and PC2 accounted for more than 75.1% total variation among measures (Figure 1). Largest cluster consisted of D3, D5, EV2, EV3, EV5, ASTAB3, ASTAB5, MASV1, MASV and Stdev measures. Second nearby cluster consisted of D1, D2, EV1, ASV, ASV1, ASTAB1, ASTAB2, measures. Third close cluster of smaller size clubbed IPC1, SIPC1, SIPC5 measures. Fourth cluster kept all mean based measures, Mean, HM, PRVG, MHPRVG, GAI and SIPC3. Measures IPC2, IPC3 & IPC4 were observed as outliers' measures.

3.2. Second Year of Study (2018-19)

Estimated sums of squares for Gx $\times$ E signal and Gx $\times$ E noise were 66.6% and 33.4% respectively. Note that the SS for Gx $\times$ E signal is 1.26 times that for genotypes main effects.

Hence, narrow adaptations are important for this dataset. Even just IPC1 alone is 0.97 times the genotypes main effects. Also note that Gx $\times$ E noise is 0.63 times the genotypes main effects. Highly significant environments (44.1), Gx $\times$ E interaction (20.4) and genotypes (10.8) were observed by ANOVA analysis. Extent of Gx $\times$ E interaction variation accounted by each of highly significant IPCA's for AMMI based measures, as type-1 benefited 51.2%, type-2 measures utilized 66.3%, type 3 measures used up to 79.3%, contributions of other non significant IPCA4, IPCA5, IPCA6 and IPCA7 were of 8.5, 5.8, 3.6 and 1.7% respectively (Table 4). Ranking of genotypes as per values of EV1 observed (G9, G11, G6) for least and large values by (G1, G2) while D1 corresponds to (G9, G11, G6) and (G1, G2), SIPC1 for (G1, G10, G7) and (G2, G3) and for ASTAB1 were (G9, G11, G6) & (G1, G2). EV2 measure pointed towards (G9, G6, G11) as desirable along with undesirable genotypes as

(G1, G5); values of D2 selected (G9, G11, G6) & (G1, G5), while for SIPC2 were (G1, G7, G5) & (G2, G1) and ASTAB2 values favoured (G6, G6, G11) & (G1, G5) (Tables 7 and 8). ASV observed (G9, G6, G11) and ASV1 recommended (G9, G11, G6) as with stable performance and unsuitable performance for G1, G2 (Table 5). First two IPCAs in ASV & ASV1 measures utilized 66.3% of GxE interaction sum of squares. Type-3 measures had exploited 79.3% of GxE interaction and values of EV3 preferred G9, G8, G4 and unstable performance of G1, G3 whereas D3 pointed towards G9, G4, G11 and G1, G3 whereas SIPC3 pointed towards G1, G7, G5 & G3, G11; ASTAB3 measure considered G9, G4, G11 & G1, G3 (Tables 7 & 8). Composite measures MASV & MASV1 selected G9, G6, G11 genotypes for stable performance and G1, G2 would not be recommended for cultivation due to unstable yield behavior. Average yield was considered as an important measure to assess the genotypes

potential as highly significant yield differences were exhibited. G4, G8, G6 genotypes maintained higher yields as compared to G1 & G11. GAI pointed towards G4, G7, G8 as of stable adaptation and G1 & G11 would be unstable. HM, PRVG and MHPRVG measures identified G4, G7, G8 and G1 & G11 for general and specific adaptations respectively. Lower values of standard error anticipated consistent yield performance for G4, G10, G1 genotypes as far as considered locations were considered. Three clusters were observed among studied measures by biplot analysis by considering first two significant principal components (80.3% total variation) (Figure 2). Larger cluster consisted of EV1, EV2, EV3, D1, D2, D3, ASV, ASV1, MASV, MASV1, ASTAB1, ASTAB2 & ASTAB3. Second cluster comprised of SIPC1, SIPC2, SIPC3, IPC1 & standard deviation. Third cluster clubbed mean based measures, Mean, HM, PRVG, MHPRVG and GAI.

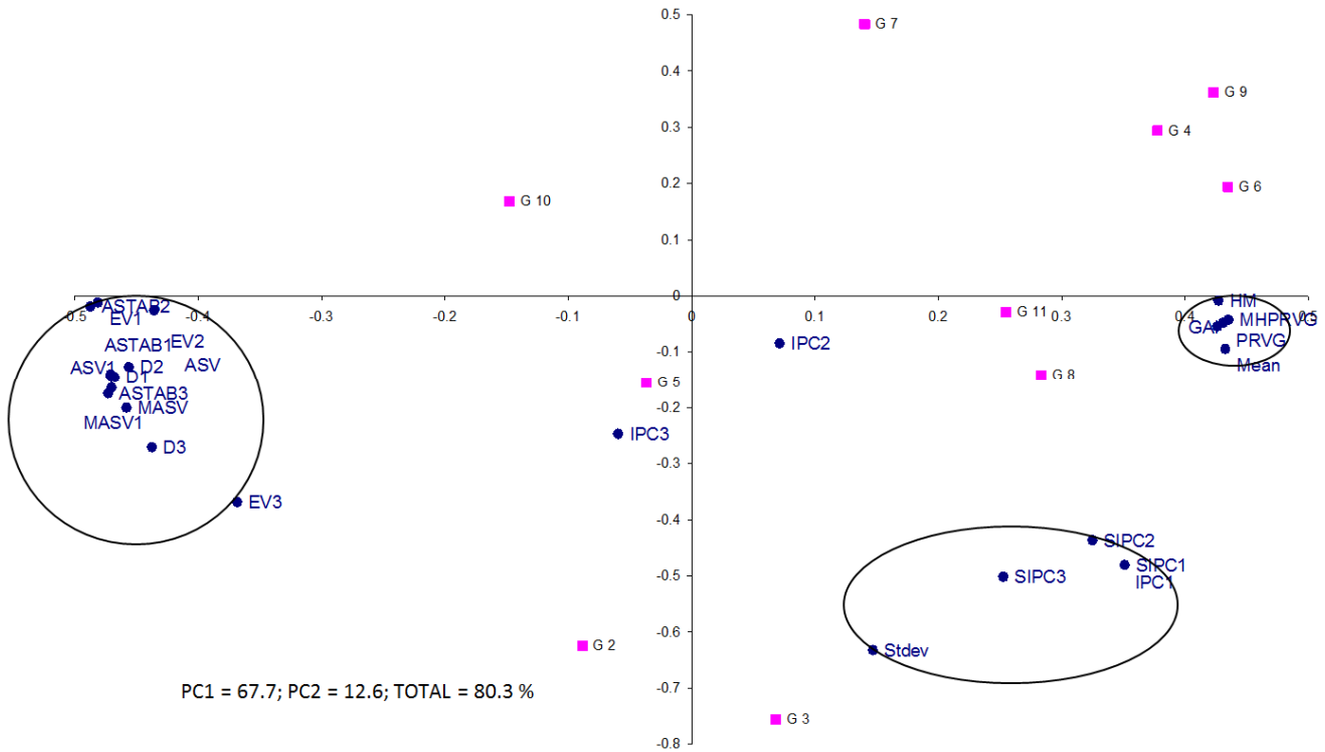


Figure 2. Biplot analysis of AMMI and yield based estimates (2018-19).

Table 8. Principal components analysis of genotypes (2018-19).

	ASV	ASV1	MASV	MASV1	IPC1	IPC2	IPC3	EV1	EV2	EV3	D1	D2	D3
G 1	7.76	14.21	7.84	14.26	-4.1990	-0.7740	0.7300	0.2681	0.1424	0.1003	11.96	12.11	12.23
G 2	4.09	7.20	4.58	7.52	2.0915	1.3958	-1.4129	0.0665	0.0605	0.0604	5.96	6.87	7.66
G 3	3.00	5.51	4.10	6.18	1.6304	-0.0367	2.8037	0.0404	0.0202	0.0928	4.64	4.64	8.17
G 4	1.78	2.68	2.24	3.06	-0.7096	1.2083	0.3737	0.0077	0.0242	0.0176	2.02	3.58	3.69
G 5	3.23	4.51	4.28	5.43	1.1078	-2.5122	-0.6699	0.0187	0.0975	0.0695	3.15	6.91	7.09
G 6	1.13	2.07	1.92	2.59	0.6118	-0.1126	-1.5524	0.0057	0.0030	0.0263	1.74	1.76	4.12
G 7	2.40	3.96	2.96	4.35	-1.1102	-1.2658	-1.0432	0.0187	0.0318	0.0321	3.16	4.42	5.08
G 8	2.69	4.82	2.81	4.89	1.4089	-0.7339	0.1331	0.0302	0.0226	0.0153	4.01	4.40	4.41
G 9	0.59	0.97	0.88	1.18	0.2730	-0.3034	0.5614	0.0011	0.0019	0.0044	0.78	1.07	1.72
G 10	3.58	5.79	4.29	6.32	-1.6050	2.0218	-0.8878	0.0392	0.0767	0.0591	4.57	6.73	7.06
G 11	1.44	2.02	2.11	2.59	0.5003	1.1128	0.9644	0.0038	0.0192	0.0222	1.42	3.07	3.84

Table 9. AMMI based estimates of genotypes (2018-19).

	SIPC1	SIPC2	SIPC3	ASTAB1	ASTAB2	ASTAB3	MEAN	STDEV	GAI	RPGV	MHRPGV	HM
G 1	-4.20	-4.97	-4.24	142.99	146.58	149.64	38.32	5.31	37.98	0.8243	0.8022	37.65
G 2	2.09	3.49	2.07	35.48	47.13	58.61	47.16	9.79	46.33	0.9965	0.9879	45.60
G 3	1.63	1.59	4.40	21.56	21.57	66.76	48.40	10.22	47.55	1.0226	1.0137	46.80
G 4	-0.71	0.50	0.87	4.08	12.82	13.62	49.55	5.28	49.31	1.0584	1.0534	49.09
G 5	1.11	-1.40	-2.07	9.95	47.71	50.29	48.28	9.46	47.56	1.0220	1.0149	46.96
G 6	0.61	0.50	-1.05	3.04	3.11	16.97	48.99	7.86	48.45	1.0392	1.0356	47.94
G 7	-1.11	-2.38	-3.42	10.00	19.58	25.84	48.88	6.10	48.55	1.0416	1.0374	48.23
G 8	1.41	0.68	0.81	16.10	19.32	19.42	49.15	8.78	48.53	1.0414	1.0369	47.99
G 9	0.27	-0.03	0.53	0.60	1.15	2.97	46.29	6.82	45.88	0.9835	0.9815	45.51
G 10	-1.61	0.42	-0.47	20.89	45.34	49.88	47.29	5.29	47.03	1.0113	1.0025	46.76
G 11	0.50	1.61	2.58	2.03	9.44	14.78	45.33	8.43	44.65	0.9592	0.9529	44.01

## 4. Conclusions

AMMI analysis has been proved as an effective tool to explore complex G×E interaction under multi environmental trials. Large number of AMMI based measures had been studied as each measures related to a different concept of stability. Recent analytic measures for adaptability of wheat genotypes exhibited affiliation with AMMI based measures exploiting number of significant IPC scores.

## Conflict of Interest

All the authors do not have any possible conflicts of interest.

## Acknowledgements

The wheat genotypes were evaluated at coordinated centers of AICW&BIP across the country. Authors sincerely acknowledge the hard work of all the staff for field evaluation and data recording.

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