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# Multi Environment Analysis of Soybean Genotypes to Delineate Stability and Adaptability for Yield and Quality Parameters

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#### Authors' contributions

This work was carried out in collaboration among all authors. All authors read and approved the final manuscript.

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#### ABSTRACT

The present investigation was carried out at three locations of Gujarat under the jurisdiction of Anand Agricultural University (AAU) during *kharif*-2022. Three diverse environments were selected for conducting an experiment. The experimental material comprised of 40 genotypes of soybean sown in a randomized block design with three replications at all environments. Based on the AMMI analysis for seed yield per plant, E<sub>3</sub> was considered as the most suitable environment for seed yield

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per plant while,  $E_2$  was considered as the poor environment. The genotypes G4, G16 and G40 were considered as the most stable genotypes and had general adaptation for seed yield per plant among all the genotypes since they were least affected by the environment. The genotypes G15, G22, G24, G25, G27, G29, G32 and G39 performed well in the poor environment ( $E_2$ ). Genotypes G10, G20 and G33 had specific adaptability in potential environment ( $E_3$ ) Based on the AMMI stability value G24, G25 and G22 were considered as stable genotypes over environment while, G33, G20 and G35 were considered as unstable genotypes. According to the AMMI 2 biplot G15, G16, G22, G24, G25, G27, G32 and G39 were considered as high yielding with general adaptability. The vertex genotypes for  $E_3$  were G10, G12, G20 and G33. G31 was for  $E_2$  while, vertex genotype for  $E_1$  was G26.

Keywords: AMMI analysis; AMMI stability value; GGE biplot and stability.

#### 1. INTRODUCTION

Soybean [*Glycine max* (L.) Merrill, 2n=2x=40] is an important oilseed crop known as "Golden bean" which belongs to the family *Leguminosae* along with many essential food legume crops such as chickpea, peanut, alfalfa, faba bean, lentil and pea.

Soybean yield is a complex guantitative trait that is influenced by interactions between genes and the environment [1]. Environmental factors, particularly day length at different latitudes, have a significant impact on soybean yield [2,3,4]. According to Camara et al. [5] the primary climatic element affecting soybean adaptation to various ecoregions is photoperiod. The growing each soybean cultivar region of was constrained to a relatively small range of latitudes due to photoperiodic sensitivity in order to achieve its highest yield [6]. When grown 2°N usual cultivation latitudes. of their some soybean cultivars had markedlv lower yields [7]. However, soybean is grown around the world at latitudes ranging from 50°N to 35°S [8].

Plant breeders frequently tried to increase yield by focusing on yield-related traits [9]. The genotypic correlation coefficients could be effectively divided into direct and indirect impacts and their relationships with grain yield could be clearly understood with the use of correlation studies and path coefficient analyses [10]. The features that have a substantial impact on soybean grain yield had been identified using path analysis [11] Yahaya and Ankrumah, [12] Al-Ballat and Al-Araby, [13]. The interpretation of genetic tests and predictions was hampered by the genotype-environment (G X E) interaction, which is a significant issue in the study of quantitative traits like yield and yield component [14,15].

biplot is an efficient method GGE for recommending specific genotypes in specific mega-environments, evaluating the mean performance and stability of genotypes, and analysing the power of target environments to genotypes [16,17]. distinguish lt visuallv examines the relationships between genotypes, test environments and genotype-by-environment interactions. When genotypes are tested in a variety of environments where the interaction between genotype and environment was significant (as in the advanced stages of testing), as well as when a large number of hybrids are evaluated in fewer locations (as in the early stages of testing), where the main goal is to discard inferior genotypes, the use of GGE biplot is more prevalent. In conclusion, GGE biplot tool has become increasingly popular in cultivar evaluation and mega-environment investigation for plant breeders and agricultural researchers [18] and many studies on  $G \times E$  interaction effect on crop growth, yield and other agronomic traits have been conducted in various plants using GGE biplot.

Keeping all these in view present study was conducted to delineate the adaptability and stability of genotypes of soybean.

#### 2. MATERIALS AND METHODS

Experimental materials were comprised of 40 genotypes (Table 1) of soybean sown in three different locations of Gujarat, under the jurisdiction of Anand Agricultural University during kharif-2022. The diverse (AAU) environments selected for conducting an Experimental Farm, experiment were Department of Genetics & Plant Breeding, B. A. College of Agriculture, AAU, Anand considered as E<sub>1</sub>, Agricultural Research Station, AAU, Derol as E<sub>2</sub> and Tribal Research cum Training Centre, AAU, Devgadh Baria considered as E<sub>3</sub>. The

Sr. No.	Name	Source	Sr. No.	Name	Source
1	Gourav	AAU, Devgadh Baria	21	GS 2	JAU, Amreli
2	Doko	AAU, Devgadh Baria	22	GS 3	JAU, Amreli
3	Birsa Soya	AAU, Devgadh Baria	23	PS 1670	JAU, Amreli
4	Lee	AAU, Devgadh Baria	24	BAUS 31-17	JAU, Amreli
5	JS 335	AAU, Devgadh Baria	25	MACS 1701	JAU, Amreli
6	JS 20-34	AAU, Devgadh Baria	26	RSC 11-35	JAU, Amreli
7	JS 20-69	AAU, Devgadh Baria	27	SL 1212	JAU, Amreli
8	NRC 131	AAU, Devgadh Baria	28	PS 1661	JAU, Amreli
9	EC 37939	AAU, Devgadh Baria	29	PS 1660	JAU, Amreli
10	EC 39045	AAU, Devgadh Baria	30	KDS 114	JAU, Amreli
11	EC 39044	AAU, Devgadh Baria	31	JS 22-14	JAU, Amreli
12	EC 100804	AAU, Devgadh Baria	32	JS 22-11	JAU, Amreli
13	NRC 193	AAU, Devgadh Baria	33	AS-15	JAU, Amreli
14	EC 109543	AAU, Devgadh Baria	34	NRC 128	JAU, Amreli
15	NRC 192	AAU, Devgadh Baria	35	PS 1659	JAU, Amreli
16	RVS 2011-4	JAU, Amreli	36	MACS NRC 1711	JAU, Amreli
17	BAUS 96-17	JAU, Amreli	37	IVT-E	JAU, Amreli
18	AUKS 203	JAU, Amreli	38	CAUM 52	JAU, Amreli
19	GS 1	JAU, Amreli	39	RVSM 2012-11	JAU, Amreli
20	DS 3152	JAU, Amreli	40	NRC 37	JAU, Amreli

Table 1. List of soybean genotypes and their source

experimental material comprised of 40 diverse genotypes of soybean sown in a randomized block design with three replications at all environments. Each genotype was grown in a single row plot of 3.0 m length spaced at  $45 \times 10$ cm. The recommended cultural practices including plant protection-measures were followed uniformly to raise a good crop and the observations were recorded and analyzed using AMMI model of stability analysis.

AMMI model was used in the analysis of data obtained from three different locations. The model for the analysis id given below

ANOVA: 
$$Y_{ij} = \mu \square \alpha_g \square \square_e \square \alpha \square_{ge} \square \rho_{ij} \square \epsilon_{ijk}$$
  
PCA:  $Y_{ij} \square \mu \square n \square n \square_{gn} \square_{en} \square \rho_{ij} \square \epsilon_{ijk}$   
AMMI:  $Y_{ij} \square \mu \square \alpha_g \square \square_e \square n \square n \square_{gn} \square_{en} \square \rho_{ij}$   
 $\square \epsilon_{ijk}$ 

Where,

The additive parameters

μ = Grand mean

 $\alpha_g$  = Deviation of genotype 'g' from grand mean

 $\beta_{a}$  = Deviation of environment 'e'

The multiplicative parameters

 $\lambda_n$  = Singular value for Interaction Principal Component Axis n (IPCA) 
$$\begin{split} \gamma_{gn} &= \text{Genotype eigenvector for axis 'n'} \\ \delta_{en} &= \text{Environment eigenvector} \\ \rho_{ij} &= \text{Residual} \\ \epsilon_{ijk} &= \text{Error term} \end{split}$$

AMMI stability value Purchase et al., [19] was calculated to correlate the mean value and stability based on IPCA scores. It was calculated using following formula.

 $\mathsf{ASVI} = \sqrt{\{\frac{SSIPCA1}{SSIPCA2}[IPCA1 \ score]\}2 + [ICPA2score]}2$ 

#### 3. RESULTS AND DISCUSSION

#### 3.1 Analysis of Variance

The analysis of variance for all the individual location (Table 2) revealed highly significant mean sum of squares due to genotypes for all the characters *viz.*, Days to 50% flowering, days to maturity, plant height, primary branches per plant, pods per plant, pod length, seeds per pod, seed yield per plant, 100 seed weight, protein and oil content indicating presence of high genetic variability in the experimental material.

The broad range of variability was revealed from the pooled analysis of variance (Table 3) among the tested genotypes as it was found significant at both 5% and 1% level of significance for all the characters studied. The selected environments were also found significantly different for all the characters. This indicated the environment selected for the present study were different. For all characters except days to maturity  $G \times E$  interaction were found highly significant which indicated the role of environment for expression of all these characters (Table 3). AMMI stability analysis was performed for all the characters showing significant G X E interaction (Table 4).

#### 3.2 Combined Analysis of Variance for Yield and its Attributing Traits

The effect of G x E on expression of characters in soybean can be best explained with the use of AMMI model ANOVA of AMMI showed significant variations for genotypes, environments and G x E at both level of significance for all the characters studied except days to maturity. Hence the GGE biplot analysis was not possible for that character.

The findings of the pooled analysis of variance as per the AMMI model are shown in Table 4. From the ANOVA of AMMI model, the maximum per cent of variation was contributed by environment for days to flowering (64.02%), plant height (79.11%), primary branches per plant (49.26%), pods per plant (41.34%), protein content (33.80%) and grain yield per plant (48.62%) and genotypes contributed highest variation to pod length (61.47%), seeds per pod (52.71%) and oil content (70.44%). The GEI effect was significant for all the characters studied. Which explains that most of the characters were largely influenced by the environment for complete expression.

#### 3.3 Mean Value Comparison among Genotypes Across Environments (Table 5)

The mean value for all the characters across environments was significant for all the characters. The mean value for 50% flowering ranged from 52.00 to 46.78 with an average of 49.39 days. The genotypes, G6 and G5 required less days for 50% flowering. However, the mean values for seed yield per plant was ranged from 37.99g/plant to 7.79g/plant. The genotype G10 had higher seed yield per plant whereas, G6 had lower seed yield per plant. The average oil content in these genotypes was 17.61 with a range of 11.8%-23.43%. The highest oil content was found in the G17 while, G28 had lowest oil content.

#### 3.4 AMMI Model Analysis for Days to 50% Flowering, Seed Yield Per Plant and Oil Content

According to the model of stability analysis, the genotypes, environment and G x E had significant effect on the expression of days to 50% flowering, seed yield per plant and oil content.

For 50% flowering the significantly (p<0.05) higher variation was contributed by the environments (64.02%) followed by genotypes (21.84%) and G x E (14.14%). It also reported that first two IPCAs explained 100% of GEI with 59.40% of variation due to PC1 and 40.60% of variation due to PC2. Amare and Tamado (2014) indicated the most accurate model for AMMI can be forecasted by using the first two IPCA. The PC1 accounted about 59.4% and PC 2 accounted 40.6% of the variation. Similar results of G x E variation was observed by Dabessa et al. [20] and Silva et al. [21].

In case of seed yield per plant, the significantly higher variation was explained by environment (48.62%) followed by GEI (27.40%) and genotypes (23.98%). Further the PC1 explained 85.3% and PC2 explained 14.7% of variation combined making 100% of variation of GEI. Both the above characters are majorly explained by the environmental sum of squares which indicates that the selected environments were diverse and the influence of environment on expression of these characters. Similar results were also reported by the Atnaf et al. [22] Mukuze et al. [23] Mwiinga et al. [24] and Razmi et al. [25].

Soybean being the oilseed crop, oil content is one of the most important quality parameter to be considered. For oil content, the genotypes (70.44%) contributed highest variation followed by GEI (28.51%) and environment (1.04%). This explains that environment had a least significant role in the expression of this character. Further GEI was divided into only first two IPCAs, PC1 explained 96.6% of the variation and PC2 explained 3.4% of the variation with significant differences (p<0.05). Similar result was reported by Dabessa et al. [20] Kocaturk et al. [26] and Mukuze et al. [23].

Sources	df	Days to 50% flowering		ing	Days to maturity			Plant height (cm)			Primary branches per plant		
		E1	E <sub>2</sub>	E <sub>3</sub>	E1	E2	E <sub>3</sub>	E1	E <sub>2</sub>	E <sub>3</sub>	E1	E2	E3
Replication	2	9.56	11.43	8.57	2.04*	0.88*	1.79*	10.20*	24.31	78.74	2.04*	0.88*	1.79*
Treatment	39	23.44**	11.51**	19.51**	6.04**	4.00**	11.22**	56.96**	226.76**	660.17**	6.04**	4.00**	11.22**
Error Total	78 119	6.18	4.68	3.46	0.57	0.21	0.53	2.19	11.13	38.38	0.57	0.21	0.53

Sources	df	df Pods per plant			Pod len	Pod length (cm) Se			er pod		Seed yield per plant (g)		
		E <sub>1</sub>	E <sub>2</sub>	E <sub>3</sub>	E1	E2	E <sub>3</sub>	E₁	E <sub>2</sub>	E <sub>3</sub>	E1	E <sub>2</sub>	E <sub>3</sub>
Replication	2	252.66	120.40	450.43	0.12	0.01	0.06	0.07*	0.09*	0.12*	35.61*	28.31*	40.32
Treatment	39	2250.73**	1953.83**	5096.36**	0.30**	0.27**	0.61**	0.13**	0.30**	0.47**	238.54**	117.82**	538.15
Error	78	83.38	64.48	112.76	0.08	0.08	0.10	0.02	0.02	0.03	7.42	5.87	18.50
Total	119												

Sources	df	100 seed	l weight (g)		Oil conten	il content (%) Protein content (%				%)		
		E1	E2	E3	E1	E2	E₃	E1	E <sub>2</sub>	E₃		
Replication	2	0.25	0.29	0.58	0.47	1.78*	2.88*	2.19*	1.62	4.08*		
Treatment	39	8.13**	6.13**	9.19**	42.66**	33.03**	25.46**	50.87**	53.09**	64.55**		
Error	78	0.38	0.22	0.26	0.50	0.48	0.66	0.63	0.83	1.04		
Total	119											

#### Table 3. Pooled RBD analysis of all environments for all characters

Sources	df	DF	DM	PH (cm)	РВ	PP	PL (cm)	SPP	SYP (g)	100 SW (g)	OC (%)	PC (%)
Rep/Env	2	2	2	2	2	2	2	2	2	2	2	2
Environment	2	1889.90**	2202.68**	69691.60**	402.60**	127849.70**	1.94**	0.01**	16506.40**	233.49**	20.81**	517.27**
Genotypes	39	33.07**	68.70**	520.89**	9.64**	5227.12**	0.79**	0.48**	417.43**	12.83**	72.00**	64.62**
GXE	78	10.70**	20.95	211.51**	5.81**	2036.89**	0.20**	0.21**	238.54**	5.31**	14.57**	51.95**
Pooled error	234	4.78	17.55	17.24	0.44	86.87	0.09	0.02	10.60	0.28	0.55	0.83
Total	359											
DF-Days to 50%	flowering	PP- Pa	ods per plant		100SW-1	00 Seed Weight						

DF-Days to 50% flowering

DM- Days to maturity PL-Pod length PH-Plant height

SPP-Seeds per pod

100SW-100 Seed Weight OC-Oil Content

PC-Protein Content

PB-Primary branches per plant SYP-Seed Yield per Plant

\*, \*\* Significant at 5% and 1% respectively

			Character	3 of Soybean		
Source	of	df	Days to 5	0% flowering	Plant h	eight (cm)
variation			MSS	% Explained	MSS	% Explained
			1		2	
Trials		119	49.61		1480.63	
Environments		2	1889.91**	64.02	69691.61**	79.11
Genotypes		39	33.07**	21.84	520.89**	11.53
G×E		78	10.70**	14.14	211.51**	9.36
IPCA I		40	12.38**	59.40	352.24**	85.4
IPCA II		38	8.92**	40.60	63.37**	14.6
Pooled Error		234	4.78		17.23	
Source	of	df		nches per plant		per plant
variation			MSS	% Explained	MSS	% Explained
			3		4	
Trials		119	13.73**		5196.93	
Environments		2	402.60**	49.26	127849.70**	41.34
Genotypes		39	9.64**	23.00	5227.12**	32.96
G×E		78	5.81**	27.74	2036.89**	25.69
IPCA I		40	8.51**	75.1	3369.32**	84.8
IPCA II		38	2.97**	24.9	634.34**	15.2
Pooled Error		234	0.44		86.87	
Source of varia	ation	df	Pod le	ength (cm)	Seed	s per pod
			MSS	% Explained	MSS	% Explained
			5		6	-
Trials		119	0.42		0.30	
Environments		2	1.94**	7.74	0.01**	0.05
Genotypes		39	0.79**	61.47	0.48**	52.71
G×E		78	0.20**	30.79	0.21**	47.23
IPCA I		40	0.31**	79.2	0.29**	70.3
IPCA II		38	0.08**	20.8	0.13**	29.7
Pooled Error		234	0.09		0.02	
Source	of	df		d per plant (g)		d weight (g)
variation			MSS	% Explained	MSS	% Explained
			7		8	
Trials		119	570.80		11.61	
Environments		2	16506.39**	48.62	233.49**	33.80
Genotypes		39	417.43**	23.98	12.83**	36.21
G×E		78	238.54**	27.40	5.31**	29.99
IPCA I		40	396.70**	85.3	5.77**	55.7
		~~			1.00***	

## Table 4. Pooled analysis of variance over environments as per AMMI model for variouscharacters of soybean

Source of variation	df	Oil	content (%)	Protei	n content (%)	
		MSS	% Explained	MSS	% Explained	
		9	-	10	-	
Trials	119	33.50		63.92		
Environments	2	20.81**	1.04	517.27**	13.60	
Genotypes	39	72.00**	70.44	64.62**	33.13	
G×E	78	14.57**	28.51	51.95**	53.27	
IPCA I	40	27.44**	96.6	52.78**	52.1	
IPCA II	38	1.02**	3.4	51.07**	47.9	
Pooled Error	234	0.55		0.83		

14.7

4.83\*\*

0.28

44.3

72.07\*\*

10.60

38

234

IPCA II

Pooled Error

\*, \*\* Significant at 5% and 1% levels, respectively

Sr. No.	Genotype	DF	PH	NPB	NPP	PL	SPP	SYP	100 SW	00	PC
1	Gourav	46.11	38.88	7.23	108.08	3.81	3.00	27.98	8.73	12.79	36.30
2	Doko	50.44	37.17	6.82	98.19	4.11	3.24	31.20	7.98	15.97	43.11
3	Birsa Soya	45.22	34.33	6.91	102.90	3.92	2.61	24.30	10.18	18.64	34.54
4	Lee	48.33	28.89	5.98	123.33	4.07	2.61	33.04	9.87	17.78	36.00
5	JS-335	42.11	25.12	7.89	103.23	3.85	2.86	28.37	10.02	16.38	38.38
6	JS-20-34	46.78	22.44	3.92	30.88	4.00	2.67	7.79	9.08	19.36	38.54
7	JS-20-69	47.44	35.97	9.11	97.51	3.84	2.78	24.36	8.58	13.34	35.82
8	NRC-131	47.89	43.09	7.57	110.29	3.92	3.12	28.51	8.88	12.36	38.82
9	EC-37939	44.56	43.16	8.33	122.67	3.49	3.13	31.72	8.53	18.03	37.81
10	EC 39045	48.33	50.36	9.80	144.98	3.59	2.94	37.99	7.37	15.51	36.78
11	EC 39044	52.00	32.66	8.51	91.70	3.62	2.86	21.39	9.30	13.06	41.93
12	EC 100804	50.11	51.67	8.96	119.91	3.59	3.02	26.01	6.72	13.43	35.10
13	NRC 193	50.44	44.08	7.68	114.57	3.53	2.50	27.08	8.19	12.78	32.99
14	EC109543	48.77	31.54	7.38	106.68	3.72	3.04	33.98	9.89	15.81	36.15
15	NRC 192	45.00	45.83	6.38	74.23	3.96	2.93	26.09	11.03	16.40	39.98
16	RVS-2011-4	47.11	39.35	6.07	81.98	3.95	3.07	26.78	10.92	21.12	40.06
17	BAUS-96-17	49.11	31.69	7.63	120.68	3.76	2.92	36.62	10.97	23.43	37.74
18	AUKS-203	46.33	35.08	6.80	43.06	4.29	3.16	10.87	10.13	15.18	37.26
19	GS 1	48.44	39.84	7.96	124.66	3.29	2.69	29.28	8.04	14.39	35.24
20	DS-3152	47.44	33.97	7.69	105.84	3.89	2.91	36.46	10.40	18.12	37.21
21	GS 2	50.33	61.26	8.71	130.92	4.02	3.03	34.31	10.02	17.62	37.24
22	GS 3	47.77	33.90	7.72	64.56	3.80	2.93	18.67	8.32	17.20	31.94
23	PS-1670	48.00	28.64	7.04	91.91	4.06	2.69	27.19	9.92	15.56	38.13
24	BAUS 31-17	50.00	32.92	7.06	83.81	3.58	2.66	21.64	9.62	15.50	34.56
25	MACS 1701	45.55	28.55	6.93	57.39	4.58	3.60	20.02	10.03	16.11	36.45
26	RSC-11-35	46.11	38.48	7.28	97.84	3.88	3.16	31.01	11.20	16.46	38.62
27	SL-1212	46.66	29.75	7.44	80.61	3.42	2.54	17.85	7.56	17.48	35.50
28	PS-1661	48.44	42.17	6.91	89.24	3.46	3.13	20.74	8.40	11.80	37.20
29	PS-1660	47.78	34.82	7.38	67.63	3.36	2.84	18.61	7.74	14.08	41.68
30	KDS-114	46.77	31.23	7.61	82.72	4.10	3.11	23.95	8.80	21.36	36.31
31	JS-22-14	48.89	34.94	7.30	81.64	4.26	3.09	24.38	11.69	15.80	40.19
32	JS-22-11	46.89	40.17	7.73	85.14	3.65	3.23	21.59	8.24	20.87	41.42
33	AS-15	47.33	40.24	9.24	85.19	3.69	3.04	29.39	8.63	17.71	34.46
34	NRC-128	49.22	36.17	7.20	69.82	3.97	2.82	17.05	9.54	18.73	35.40

#### Table 5. Mean values comparison among genotypes across the locations

Sr. No.	Genotype	DF	PH	NPB	NPP	PL	SPP	SYP	100 SW	00	PC
35	PS-1659	49.11	28.70	8.12	103.57	3.57	3.03	28.32	8.77	14.79	36.12
36	MACS NRC 1711	49.00	30.64	6.24	101.66	4.19	3.20	32.91	9.31	17.67	37.25
37	IVT-E	46.22	31.37	7.12	64.08	3.79	3.12	21.13	11.27	15.41	36.82
38	CAUM-52	45.44	39.68	7.06	91.28	3.88	3.21	29.55	9.83	13.04	44.15
39	RVSM-2012-11	48.78	38.37	6.21	76.76	4.36	3.08	25.52	10.84	12.79	41.34
40	NRC-37	48.44	27.46	6.90	118.21	4.36	3.19	35.82	9.22	21.63	36.91
	SEm	1.25	2.12	0.37	5.34	0.16	0.08	1.81	0.30	0.43	0.53
	C. V. (%)	4.60	10.22	8.75	10.32	7.63	4.79	12.50	5.69	4.50	2.42
	C. D. (5%)	3.53	5.96	1.05	15.05	0.48	0.23	5.12	0.86	1.20	1.47
	Grand mean	47.72	36.36	7.39	93.73	3.85	2.97	26.23	9.34	16.38	37.54
	Max.	52.00	61.26	9.8	144.98	4.58	3.60	37.79	11.69	23.43	44.15
	Min.	46.78	22.44	3.92	30.88	3.29	2.50	7.79	6.72	11.80	31.94

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Footnote: CV- Coefficient of variation, CD- Critical Difference, SEm- Standard Error of Mean

Sr. No.	Genotype	IPCA I	IPCA II	ASV	Rank of ASV
1	Gourav	-0.31	0.29	0.54	9
2	Doko	-0.53	-0.69	1.04	28
3	Birsa Soya	-1.57	0.90	2.48	40
4	Lee	-0.63	0.39	1.00	27
5	JS-335	-0.81	0.29	1.22	33
6	JS-20-34	-0.04	0.84	0.84	22
7	JS-20-69	-0.36	-0.23	0.58	10
8	NRC-131	-0.59	0.68	1.10	29
9	EC-37939	-0.48	0.41	0.81	19
10	EC 39045	0.14	-0.23	0.30	3
11	EC 39044	-0.33	-1.10	1.20	32
12	EC 100804	0.05	-0.82	0.83	21
13	NRC 193	-0.32	-0.51	0.70	16
14	EC109543	0.12	-0.44	0.48	7
15	NRC 192	-0.07	-0.40	0.42	4
16	RVS-2011-4	0.02	-0.54	0.54	8
17	BAUS-96-17	-0.32	-0.51	0.70	15
18	AUKS-203	0.15	-0.37	0.43	5
19	GS 1	0.35	0.53	0.74	17
20	DS-3152	0.77	-0.06	1.12	31
21	GS 2	0.27	0.52	0.66	13
22	GS 3	0.66	0.80	1.25	34
23	PS-1670	0.93	-0.18	1.38	37
24	BAUS 31-17	0.60	-0.30	0.94	24
25	MACS 1701	0.39	-0.11	0.58	11
26	RSC-11-35	-0.89	-0.53	1.42	38
27	SL-1212	0.44	0.03	0.66	12
28	PS-1661	0.03	0.26	0.27	1
29	PS-1660	0.64	-0.004	0.94	25
30	KDS-114	1.04	0.90	1.60	39
31	JS-22-14	-0.35	-0.59	0.79	18
32	JS-22-11	0.32	-0.49	0.68	14
33	AS-15	0.50	0.56	0.92	23
34	NRC-128	-0.62	-0.34	0.96	26
35	PS-1659	-0.54	0.25	0.83	20
36	MACS NRC 1711	-0.09	-0.26	0.29	2
37	IVT-E	-0.45	1.07	1.26	35
38	CAUM-52	0.77	-0.06	1.12	30
39	RVSM-2012-11	0.83	0.32	1.26	36
40	NRC-37	0.28	0.15	0.44	6
E1	Anand (E1)	-1.11	-2.47		
E <sub>2</sub>	Derol (E <sub>2</sub> )	-1.79	2.10		
E3	Devgadh Baria (E <sub>3)</sub>	2.90	0.36		

### Table 6. IPCA scores and ASV of different soybean genotypes and environments for days to 50 % flowering

#### 3.5 AMMI Biplot Analysis and ASV Rankings

The GGE refers to the genotype main effect (G) and the genotype x environment interaction (GE), which are the two most important sources of variation for cultivar evaluation in a multi environment trials [27]. A GGE biplot displays the genotypic main effect (G) and genotype by

environment interaction (GE) of a genotype byenvironment dataset [18].

Genotypes having a zero IPCA 1 score are less influenced by the environment and better adapted to all environments. According to AMMI biplot (Fig.1a) analysis for 50% flowering, G16, G18, G27 and G32 genotypes had low mean and near zero IPCA value indicated the adaptation to all environments. Genotypes G3, G5, G6 and G26 were positioned far away from the origin with lower mean value indicated that they were unstable genotypes and adapted only to favourable environment. Based on ASV values

the genotypes, G28, G36 and G10 had lowest ASV values hence considered as highly stable genotypes and G3, G30 and G26 had lowest ASV values hence they were identified as highly unstable genotypes.

Table 7. IPCA scores and ASV of different soybean genotypes and environments for seed yield
per plant (g)

Sr. No.	Genotype	IPCA I	IPCA II	ASV	Rank of ASV
1	Gourav	-1.00	1.62	6.00	22
2	Doko	1.80	-0.24	10.47	31
3	Birsa Soya	-0.71	-1.27	4.31	18
4	Lee	0.25	1.24	1.89	9
5	JS-335	-0.60	0.66	3.54	16
6	JS-20-34	-1.96	-0.54	11.40	34
7	JS-20-69	-1.15	-0.53	6.67	24
8	NRC-131	1.75	0.27	10.18	30
9	EC-37939	1.96	-0.28	11.38	33
10	EC 39045	1.90	0.66	11.05	32
11	EC 39044	-1.05	0.73	6.12	23
12	EC 100804	2.04	-1.26	11.92	35
13	NRC 193	-0.43	-1.55	2.94	13
14	EC109543	-0.68	0.69	3.98	17
15	NRC 192	0.26	0.04	1.52	6
16	RVS-2011-4	-0.15	-0.77	1.15	5
17	BAUS-96-17	-2.16	-0.97	12.56	36
18	AUKS-203	-1.67	-0.99	9.73	29
19	GS 1	-2.16	0.61	12.57	37
20	DS-3152	2.79	0.26	16.19	39
21	GS 2	0.36	0.99	2.32	12
22	GS 3	0.14	-0.07	0.82	3
23	PS-1670	1.29	0.29	7.47	27
24	BAUS 31-17	0.00	0.17	0.17	1
25	MACS 1701	0.02	-0.14	0.18	2
26	RSC-11-35	-1.09	2.20	6.69	25
27	SL-1212	-0.33	-0.46	1.99	10
28	PS-1661	0.93	-0.79	5.45	20
29	PS-1660	-0.28	0.50	1.69	8
30	KDS-114	1.03	-0.17	5.97	21
31	JS-22-14	-1.55	-1.86	9.21	28
32	JS-22-11	-0.27	0.36	1.60	7
33	AS-15	2.94	0.03	17.04	40
34	NRC-128	-0.56	-0.57	3.28	14
35	PS-1659	-2.60	0.62	15.10	38
36	MACS NRC 1711	1.25	-0.01	7.26	26
37	IVT-E	0.59	-0.58	3.47	15
38	CAUM-52	-0.82	-0.11	4.76	19
39	RVSM-2012-11	0.16	-0.49	1.06	4
40	NRC-37	-0.26	1.70	2.28	11
E1	Anand (E1)	-3.80	3.76		
E2	Derol (E <sub>2</sub> )	-3.15	-4.00		
E <sub>3</sub>	Devgadh Baria (E <sub>3)</sub>	6.95	0.24		

Sr. No.	Genotype	Mean (%)	Rank	IPCA I	IPCA II	ASV	Rank of ASV
1	Gourav	12.79	36	-0.34	0.12	9.65	14
2	Doko	15.97	21	-0.50	-0.46	14.10	22
3	Birsa Soya	18.64	8	-1.06	-0.09	30.03	35
4	Lee	17.78	11	0.96	0.06	27.38	33
5	JS-335	16.38	19	1.47	0.87	41.83	39
6	JS-20-34	19.36	6	-0.11	0.30	3.08	3
7	JS-20-69	13.34	33	-0.34	0.48	9.72	15
8	NRC-131	12.36	39	-0.53	0.35	15.00	26
9	EC-37939	18.03	10	-0.88	0.17	25.13	32
10	EC 39045	15.51	25	0.26	0.04	7.39	11
11	EC 39044	13.06	34	-0.76	0.12	21.63	30
12	EC 100804	13.43	32	0.38	-0.16	10.72	18
13	NRC 193	12.78	38	1.00	-0.26	28.36	34
14	EC109543	15.81	22	1.25	-0.30	35.51	37
15	NRC 192	16.40	18	1.73	0.52	49.26	40
16	RVS-2011-4	21.12	4	-0.60	-0.11	17.09	27
17	BAUS-96-17	23.43	1	0.22	-0.33	6.29	8
18	AUKS-203	15.18	28	0.24	0.07	6.89	10
19	GS 1	14.39	30	-0.23	0.55	6.55	9
20	DS-3152	18.12	9	1.33	-0.29	37.83	38
21	GS 2	17.62	14	1.14	-0.28	32.52	36
22	GS 3	17.20	16	-0.84	0.20	23.77	31
23	PS-1670	15.56	24	-0.36	0.05	10.26	17
24	BAUS 31-17	15.50	26	0.51	-0.14	14.48	24
25	MACS 1701	16.11	20	0.49	-0.12	13.87	21
26	RSC-11-35	16.46	17	-0.14	0.01	3.92	4
27	SL-1212	17.48	15	-0.18	-0.03	5.20	6
28	PS-1661	11.80	40	-0.50	0.08	14.31	23
29	PS-1660	14.08	31	-0.04	-0.09	1.01	2
30	KDS-114	21.36	3	-0.29	-0.37	8.33	_ 12
31	JS-22-14	15.80	23	-0.63	0.47	17.89	29
32	JS-22-11	20.87	5	-0.43	-0.32	12.35	20
33	AS-15	17.71	12	-0.53	0.07	14.94	25
34	NRC-128	18.73	7	-0.34	-0.24	9.60	13
35	PS-1659	14.79	29	-0.34	-0.21	9.75	16
36	MACS NRC 1711	17.67	13	-0.61	0.18	17.40	28
37	IVT-E	15.41	27	-0.39	-0.20	10.98	19
38	CAUM-52	13.04	35	-0.20	-0.05	5.76	7
39	RVSM-2012-11	12.79	37	0.20	-0.05	0.29	1
40	NRC-37	21.62	2	0.17	-0.59	4.79	5
Environ		21.02	-	0.17	0.00		5
Environ E1	Anand (E <sub>1</sub> )	16.79	1	3.56	-0.13		
$E_1$ $E_2$	Derol (E <sub>2</sub> )	15.96	3	-2.05	-0.13		
E2 E3	Devgadh Baria (E <sub>3)</sub>	16.41	2	-2.05 -1.51	-1.27 1.40		
<b>L</b> 3	Grand mean	16.38	2	-1.51	1.40		

## Table 8. IPCA scores and ASV of different soybean genotypes and environments for oil content (%)

Whereas, biplot (Fig. 2a) for seed yield per plant genotypes G4, G16 and G40 were considered as the most stable genotypes among all the genotypes since they were least affected by the environment. The genotypes G15, G22, G24, G25, G27, G29, G32 and G39 performed well in the poor environment. Genotypes G10, G20 and G33 were considered as specifically adapted to the favourable environment (E<sub>3</sub>). Based on ASV ranking genotype G24, G25 and G22 considered as high stable genotypes. G33, G20 and G35 got highest ranking 40, 39 and 38 respectively and considered as a highly unstable genotypes.

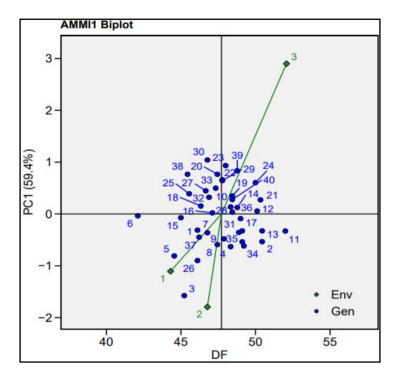


Fig. 1a. AMMI 1 biplot display of all genotypes for days to 50% flowering

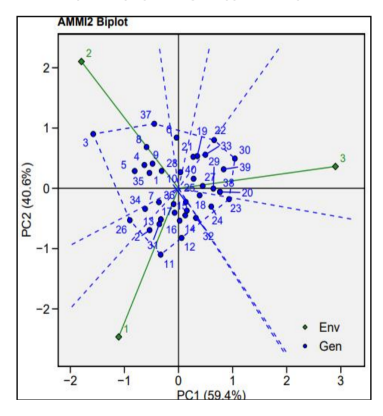


Fig. 1b. AMMI 2 biplot display of all genotypes for days to 50% flowering

Based on the biplot (Fig. 3a) for oil content, the genotypes G6, G17, G26 and G27 having general adaptability as it was found at the right-hand side of the grand mean level and close to

IPCA=0 line. G29, G39, G10, G18, G19 and G38 genotypes had low mean and near zero IPCA value indicated their adaptation to poor environment. Genotypes G4, G5, G15, G20 and

G21 were positioned far away from the origin with higher IPCA value indicated that unstable genotypes and adapted to only favourable environment (E<sub>1</sub>). According to the ASV parameter the genotypes, G39 G29 and G6 were considered as highly stable genotypes with lowest ASV values whereas, genotypes G15, G5 and G20 were considered as unstable genotypes with highest ASV values.

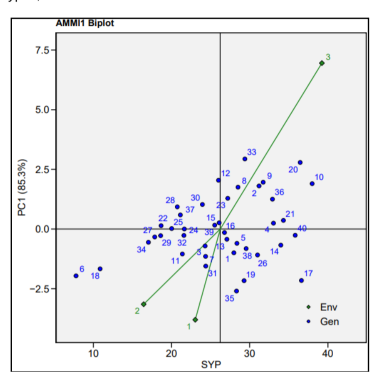


Fig. 2a. AMMI 1 biplot display of all genotypes for seed yield per plant

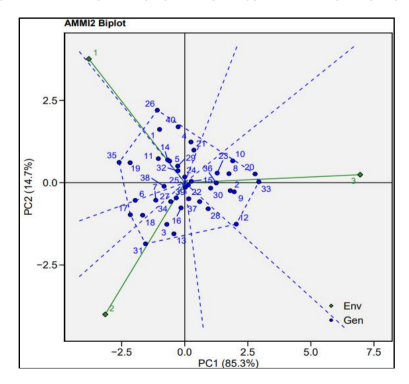


Fig. 2b AMMI 2 biplot display of all genotypes for seed yield per plant

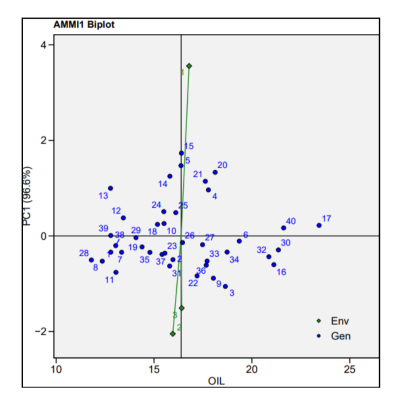


Fig. 3a. AMMI 1 biplot display of all genotypes for oil content p

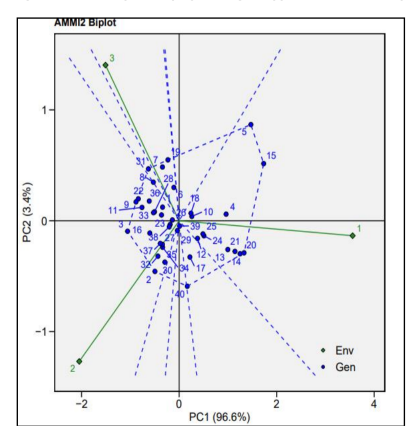


Fig. 3b. AMMI 2 biplot display of all genotypes for oil content

#### 3.6 Which won where Polygon View of GGE Biplot

GGE biplot produces best polygons to view or visualize the genotype x environment interaction pattern [16] Visualization of the 'Which-wonwhere' pattern in the polygon view is helpful to estimate possible existence of different megaenvironments in the target environment [28] Yan et al., [18] Yan and Tinker, [29]. Fig. 1b, 2b and 3b presents a polygon view of thirty two soybean genotypes tested at three environments. With this biplot, a polygon was constructed by connecting the vertex genotypes (located farthest away from the biplot origin in various directions) with straight lines and as a result, the rest of the genotypes placed inside the polygon. The polygon is formed by connecting the signs of the genotypes that are farthest away from the biplot origin, such that all other genotypes are contained in the polygon and perpendicular lines divide the polygon into sectors. Sectors help to visualize the mega-environments. This means that winning genotypes for each sector are placed at the vertex [30].

The AMMI model analysis of variance revealed a significant IPCA II value. This aided in the creation of the AMMI II biplot (Fig. 1b). According to Fig. 2b whole biplot was divided in to seven sectors. The genotypes in a sector were similar in performance compared to the genotypes in other sectors. First sector contained Devgadh Baria  $(E_3)$  with a vertex genotype G30 which indicated that this genotype showed highest days for 50% flowering at Devgadh Baria (E<sub>3</sub>). Anand environment was found in the 3rd sector with a vertex genotypes of G11 and G12 which indicated that these genotypes took more days for flowering. The 5<sup>th</sup> sector contained Derol (E2) with a vertex genotypes of G3 and G37 which indicated that these genotypes showed highest days for flowering at Derol [31].

According to the biplot 2 (Fig. 2b) of seed yield per plant, the polygon was formed with 7 sectors. The E3 was found in the 2<sup>nd</sup> sector with a vertex genotypes of G10, G12, G20 and G33. The 4<sup>th</sup> sector contained the E2 with a vertex genotype G31 and the 7<sup>th</sup> sector contained E1 with vertex genotype G26. However, genotypes within the polygon, particularly those located near the biplot origin were less responsive than the genotypes on the vertices and the ideal genotype would be one closest to the origin. The genotypes G15, G16, G22, G24, G25, G27, G32 and G39 were found near the origin which indicated that they were insensitive to the environmental conditions. Genotype G33 and G36 found on the PC2 line which indicated that these were high yielding genotypes.

Biplot 2 for oil content (Fig. 3b) was drawn by considering both PC1 and PC2 values which resulted in the formation of eight sectors.1<sup>st</sup> sector contained E1 environment with a vertex genotype G5, G14, G15 and G20. E2 environment was found in 5<sup>th</sup> sector with a vertex genotypes G3, G9 and G22. The 7<sup>th</sup> sector contained E3 environment with a vertex genotype G19. In this study, genotypes G6, G26, G27, G38 and G39 were found very close to the origin and hence were unaffected by environmental interaction forces, resulting in general adaptation to all the environments.

#### 4. CONCLUSION

The present study was conducted to delineate the stability and adaptability of genotypes of soybean by using combined ANOVA, AMMI analysis and GGE biplot technique. Based on the individual analysis of variance all the parameters viz., days to 50% flowering, days to maturity, plant height, primary branches per plant, pods per plant, pod length, seeds per pod, seed yield per plant, 100 seed weight, oil content and protein content showed significant differences among the genotypes across all the locations. In pooled analysis of variance all parameters except days to maturity showed significant G X E interaction which resulted in the analysis of stability by AMMI model. Analysis of variance based on AMMI model indicated that variation due to environment and G X E played a significant role in the expression of most of the characters studied. G X E further partitioned into interaction principal component axis (IPCA I and IPCA II). As per the per cent contribution from IPCA I and IPCA II, AMMI 1 and AMMI 2 biplots were constructed. For days to flowering the genotypes G1, G7, G15, G16 and G18 took less days to flower with higher stability hence, these genotypes can be exploited for earliness character with stability in soybean.

According to the AMMI 1 model for seed yield per plant,  $E_3$  was considered as the most suitable environment for seed yield per plant while,  $E_2$  was considered as the poor environment. The genotypes G4, G16 and G40 were considered as the most stable genotypes and had general adaptation for seed yield per plant among all the genotypes since they were least affected by the environment. The genotypes G15, G22, G24, G25, G27, G29, G32 and G39 performed well in the poor environment (E<sub>2</sub>), Genotypes G10, G20 and G33 had specific adaptability in potential environment (E<sub>3</sub>) because of the similar sign of environment with their IPCA I scores. Based on the AMMI stability value G24, G25 and G22 were considered as stable genotypes over environment while, G33, G20 and G35 were considered as unstable genotypes. According to the AMMI 2 biplot G15, G16, G22, G24, G25, G27, G32 and G39 were considered as high yielding with general adaptability. The vertex genotypes for E<sub>3</sub> were G10, G12, G20 and G33. G31 was for E<sub>2</sub> while, vertex genotype for E1 was G26. Genotypes G4, G16 and G40 were considered as best genotypes with high average seed yield per plant and general adaptability to all the locations which can be used for further breeding activities.

#### **COMPETING INTERESTS**

Authors have declared that no competing interests exist.

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