



# AMMI and GGE Biplot Analysis of Yield and Related Traits among Selected Mini-core Pigeonpea (*Cajanus cajan* L. Millsp.) Accessions

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## Authors' contributions

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

## Article Information

DOI: 10.9734/AJBGMB/2023/v14i1307

## Open Peer Review History:

This journal follows the Advanced Open Peer Review policy. Identity of the Reviewers, Editor(s) and additional Reviewers, peer review comments, different versions of the manuscript, comments of the editors, etc are available here: <https://www.sdiarticle5.com/review-history/99808>

Original Research Article

Received: 08/03/2023

Accepted: 12/05/2023

Published: 22/05/2023

## ABSTRACT

Pigeon pea [*Canjanus cajan* (L.) Millsp.] is an orphan crop that has remained greatly unimproved in quality and quantity in Africa even though it has great potential of supplying food to the growing population. Its yield is affected by various biotic and abiotic factors and a lack of varieties with broad adaptability to a range of environments. It is therefore important to select accessions that are stable in different environments for yield and yield components. A randomized complete block design layout was used to carry out a multi-locational trial using 107 pigeon pea accessions. Data was taken for the total number of pods per plant at maturity, pod length, seed weight, seeds per pod, and seed yield per plant. Additive main effects and multiplicative interaction (AMMI) with genotype and genotype by environment interaction (GGE) biplot were used to interpret the data. The mean squares for the PCA explained by the first two principal components account for 100% of

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the G x E interaction. The results revealed that the differences in the yield and yield components across the three locations were mostly due to the genotypes of the pigeon pea and to some extent the genotype by environment interaction. Six accessions were observed to be high-yielding and stable across all three environments.

**Keywords:** Pigeonpea; GE interaction; GGE biplot; AMMI; stability; yield components.

## 1. INTRODUCTION

Pigeon pea [*Canjanus cajan* (L.) Millsp.] has been reported to be an orphan crop that has remained greatly unimproved in quality and quantity in Africa even though it has great potential [1]. Its production in Nigeria has been at a subsistence level among small-scale farmers [2] but has recently been considered a cash crop for national income earnings [3]. This is not peculiar to only Nigeria or Africa, as pigeon pea production is distinguished from other major pulses in the world for being almost entirely produced by smallholder farmers [4].

This pulse is a rich source of 20-30% protein, especially sulfur-containing amino acids such as methionine and cysteine [5]. It also contains cajanin, cahanones 2-2 methyl cajanon, 2-hydroxy genistein, and isoflavones that confer antioxidant properties. It has been used generally for treating dysentery, jaundice, diabetes, skin irritations, sores, hepatitis, constipation, the expulsion of bladder stones, inducing lactation, and for stabilizing the menstrual period [6-9]. It is used for the production of noodles and confectionaries [10], improvement of soils, and as a national income earner, as India offered a hundred billion dollars to Nigeria to export pigeon pea and other legumes to India [11].

Pigeon pea cultivation in Nigeria is predominantly in the agroecological zones of the Guinea Savanna [2]. However, this region is presently the largest and most threatened agroecological zone [12] due to desert encroachment and its proximity to the Sudano-Sahelian region [13]. Similarly, there is a southward shift of the rainforest, creating a derived savanna, with a loss of forest biodiversity, and a change in agricultural practices [14]. The implication of this is that the crops being cultivated in the Guinea Savanna can now be cultivated in the derived savanna and the depleting rainforest region of Nigeria.

Currently, pigeon pea production in Nigeria is 3.52% of the world's 5.4 million hectare production area [15]. Its yield is affected by various biotic and abiotic factors ranging from moisture, altitude, temperature, photoperiodism, insect pests, mineral stresses, diseases, and a lack of varieties with broad adaptability to a range of environments [16,17]. In India, crop yield was reported to be 652 kg ha<sup>-1</sup>, 1,268 kg ha<sup>-1</sup> in Malawi, 921 kg ha<sup>-1</sup> in Myanmar, 300-400 kg ha<sup>-1</sup> in Mozambique, and 1,345 kg ha<sup>-1</sup> in the guinea savanna of Nigeria, while global productivity is at 774 kg ha<sup>-1</sup> [4,18,19]. To mitigate against this gap in pigeon pea yield, varieties that are resistant to abiotic and biotic stresses need to be developed [20]. However, without multi-environment trials, the effects of genotypes by environment interaction (GEI) will result in inconsistencies in the performance of these varieties. It is therefore important to select genotypes that are stable in different environments for yield and yield components.

There are different components used for determining yield in pigeon pea hybrids. Seed yield per hectare, seeds per pod, pod length, 100 seed weight, seed yield per plant, and pods per plant are reported to be significantly associated with pigeon pea yield [21,22]. Since yield is not a monogenic trait, stability in these yield and yield components is therefore important when selecting pigeon pea for yield. Using multi-environment trials could become problematic to manipulate due to the large data generated. However, the additive main effects and multiplicative interaction (AMMI) analysis, with genotype and genotype by environment interaction (GGE) biplot models are some of the powerful means of interpreting this kind of data [23]. The AMMI model makes use of variance and principal component analysis to represent the GEI in several dimensions, while the GGE combines both genotype main effects and those of GEI for analysis [24,25]. AMMI and GGE have been used to identify traits such as high yield performance and stability across rainfed environments in India [23], grain yield and stability in Malawi [26], yield stability of vegetable

pigeon pea in Kenya [27], number of primary branches/plant, pod length, number of grains/pod, 100-seed weight, and seed yield/plant in Manipur, India [28]. However, no such information exists for pigeon pea in the guinea savanna agroecological zone of Nigeria. Hence, the objective of this study was to evaluate the yield stability and adaptability of Nigerian pigeon pea accessions using AMMI and GGE analyses.

## 2. MATERIALS AND METHODS

### 2.1 Description of the Study Locations

This study was conducted at three locations which include Ejule (Kogi State), Ekpoma (Edo State), and the University of Lagos (Lagos State). Ejule is 369.6 m above sea level (m.a.s.l), in the Guinea Savanna agroecological zone where pigeon pea is cultivated [15]. Ekpoma is 256.6 m.a.s.l in the Derived Savanna agroecological zone and pigeon pea is cultivated there as well. Lagos is 7.3 m.a.s.l, in the humid forest, however, pigeon pea is not cultivated there.

Rainfall (mm) was recorded at Ekpoma and Ejule where crops were planted on the field, while temperature (°C) was recorded at all locations where the experiment was conducted. The average rainfall, and minimum and maximum temperatures throughout the research are presented in Table 2. Soil samples were collected from both Ejule and Ekpoma fields before land preparation was carried out.

### 2.2 Description of Plant Samples

A total of 129 pigeon pea accessions were obtained from the regional genebank of the International Crops Research Institute for the Semi-Arid Tropics (ICRISAT), located in Niamey, Niger Republic. The collections contained high, medium, and low-yielding accessions. These accessions were gathered from different parts of the world and kept in the genebank. The White accession is a local high-yielding genotype grown by many farmers in Kogi state. This was used as a local check, thereby, bringing the total accessions to 130.

### 2.3 Experimental Design and Planting

The 130 pigeon pea accessions were planted in a randomized complete Block design (RCBD). However, only 107 viable accessions germinated and were used for the multi-locational trials in the study (Table 1). Multi-locational trials were conducted at Ejule (Kogi State), Ekpoma (Edo State), and the University of Lagos (Lagos State). A 60 cm row-to-row and 20 cm plant-to-plant spacing was adopted for the sowing as described by Navneet [29] on ridges in Ejule and Ekpoma, while polyethene pots in the screen house were used in the University of Lagos. Accessions were thinned to a single plant per pot after three weeks of establishment. Five stands (replicates) per accession were used for the study. Normal agronomic practices were done throughout the duration of the study, including watering plants in the greenhouse every other day.

### 2.4 Collection of Data

Data was as described by International Board for Plant Genetic Resources [30]. Data for yield traits which include pod number, pod length, seeds per pod, hundred seed weight, and seed yield per plant were collected.

### 2.5 Analysis of Data

The AMMI and GGE were analyzed using PB Tools software version 2014. This was carried out for G × E interactions and stability of evaluated yield traits of the genotypes.

The model used for the analysis is given below:

$$Y_{ij} = \mu + G_i + E_j + GE_{ij} + \epsilon_{ij}$$

Where:

$Y_{ij}$  = the observed mean of the  $i^{\text{th}}$  genotype, ( $G_i$ ) in the  $i^{\text{th}}$  environment, ( $E_j$ ),  
 $\mu$  = the overall mean;  
 $G_i$  = effect of the  $i^{\text{th}}$  genotype;  
 $E_j$  = effect of the  $j^{\text{th}}$  environment;  
 $GE_{ij}$  = the interaction effects of the  $i^{\text{th}}$  genotype, and the  $j^{\text{th}}$  environment; and  
 $\epsilon_{ij}$  = the error term.

**Table 1. Pigeon pea accessions used in the study**

S/N	Pigeon pea Accession	Genotype Code	S/N	Pigeon pea Accession	Genotype Code	S/N	Pigeon pea Accession	Genotype Code
1	ISC-1	G1	36	ISC -174	G36	72	ISC -38	G72
2	ISC -10	G2	37	ISC -175	G37	73	ISC -39	G73
3	ISC -100	G3	38	ISC -176	G38	74	ISC -4	G74
4	ISC -104	G4	39	ISC -178	G39	75	ISC -40	G75
5	ISC -109	G5	40	ISC -179	G40	76	ISC -41	G76
6	ISC -11	G6	41	ISC -180	G41	77	ISC -42	G77
7	ISC -111	G7	42	ISC -181	G42	78	ISC -46	G78
8	ISC -115	G8	43	ISC -182	G43	79	ISC -48	G79
9	ISC -118	G9	44	ISC -184	G44	80	ISC -5	G80
10	ISC -120	G10	45	ISC -185	G45	81	ISC -51	G81
11	ISC -122	G11	46	ISC -186	G46	82	ISC -58	G82
12	ISC -123	G12	47	ISC -187	G47	83	ISC -59	G83
13	ISC -124	G13	48	ISC -189	G48	84	ISC -6	G84
14	ISC -129	G14	49	ISC -197	G49	85	ISC -61	G85
15	ISC -13	G15	50	ISC -198	G50	86	ISC -62	G86
16	ISC -134	G16	51	ISC -2	G51	87	ISC -63	G87
17	ISC -135	G17	52	ISC -20	G52	88	ISC -65	G88
18	ISC -137	G18	53	ISC -200	G53	89	ISC -66	G89
19	ISC -14	G19	54	ISC -201	G54	90	ISC -74	G90
20	ISC -140	G20	55	ISC -202	G55	91	ISC -75	G91
21	ISC -141	G21	56	ISC -203	G56	92	ISC -76	G92
22	ISC -147	G22	57	ISC -204	G57	93	ISC -77	G93
23	ISC -150	G23	58	ISC -22	G58	94	ISC -78	G94
24	ISC -153	G24	59	ISC -23	G59	95	ISC -82	G95
25	ISC -155	G25	60	ISC -24	G60	96	ISC -83	G96
26	ISC -157	G26	61	ISC -25	G61	97	ISC -84	G97
27	ISC -158	G27	62	ISC -27	G62	98	ISC -86	G98
28	ISC -16	G28	63	ISC -28	G63	99	ISC -88	G99
29	ISC -166	G29	64	ISC -29	G64	100	ISC -89	G100
30	ISC -167	G30	65	ISC -3	G65	101	ISC -9	G101
31	ISC -168	G31	66	ISC -30	G66	102	ISC -90	G102
32	ISC -169	G32	67	ISC -31	G67	103	ISC -91	G103
33	ISC -170	G33	68	ISC -32	G68	104	ISC -92	G104
34	ISC -171	G34	69	ISC -35	G69	105	ISC -93	G105
35	ISC -172	G35	70	ISC -36	G70	106	ISC -95	G106
36	ISC -174	G36	71	ISC -37	G71	107	White	G107

### 3. RESULTS

#### 3.1 Characteristics of Locations used in the study and Analysis of Variance

Information on the three locations used in the study is presented in Table 2. Ejule (Kogi State) has the highest altitude, followed by Ekpoma (Edo State), while Akoka (Lagos State) has the lowest altitude. The three environments have similar soil texture, but with different mean rainfall. More rainfall was recorded in Lagos, followed by Edo State, while Kogi had the least rainfall in the planting season. Minimum temperatures for the three locations are not so different, but the maximum temperature showed that Kogi had the highest maximum temperature, followed by Edo, and then Lagos. Results for yield and yield components show highly significant differences ( $p < 0.01$ ) within the genotypes, and the test locations, but not significant for genotype by environment interactions for all the yield components except 100 seed weight, where  $G \times E$  showed highly significant differences (0.01) (Table 3).

#### 3.2 Additive Main Effects and Multiplicative Interaction (AMMI 1) Analysis for PC1 and Significant Influence of Traits

The AMMI analysis of variance of 107 pigeon pea genotypes evaluated for yield and yield components across three locations revealed that genotypes accounted for more than 84% of the total variation in all the traits considered, while the environment and genotype by environment each accounted for less than 9% of the total variation (Table 3). The mean squares of the Principal Component Analysis 1 (PCA1) and Principal Component Analysis 2 (PCA2) were significant (0.01), and explained more than 70% and about 27.6% of the total variation, respectively in all the traits (Table 3). PCA3 showed no variation (0%) in the study. Therefore, the PCA1 and PCA2 gave a cumulative contribution of 100.

Results shown in Fig. 1 show the genotype by environment interactions based on the AMMI1 model for total pod number (Fig. 1a), pod length

(Fig. 1b), seeds per pod (Fig. 1c), hundred seed weight (Fig. 1d), and yield per plant (Fig. 1e). The results revealed that genotypes with positive PCA1 scores such as G67, G16, G6, G59, G34, G19, G64 and G82 had positive interactions with Edo (E1) and Kogi (E2) for pod number (Fig. 1a), while for hundred seed weight (Fig. 1d), G101, G83, G67, G82, G2, G45 had positive interactions with Edo (E1) and Lagos (E3). Genotypes with positive PC1 scores such as G36, G100, and G35 for pod length (Fig. 1b), G50, G85, G55 and G88 for seeds per pod (Fig. 1c), had positive interactions with Lagos, and G23, G84, G16, G71, G89, and G29 (Fig. 1e) had positive interactions with E1 (Edo). Conversely, genotypes with negative PCA1 scores such as G86, G106, G1, G90, G103, G24, G104, and G3 (Fig. 1a), had positive interactions with only Lagos (E3). Similarly, genotypes with negative PCA1 scores such as G92, G103, G1, G14, G36, G52, and G100 (Fig. 1d) had positive interactions with only E2. While negative PCA1 scores in G1, G23, G37, G61, G90, G4 (Fig. 1b), and G1, G23, G56, G61, G98, G65 (Fig. 1c) had positive interactions with E1 and E2. E2 and E3 had positive interactions with G33, G1, G91, G25, and G10 (Fig. 1d) which had negative PCA1 scores.

The stable genotypes (Fig. 1) with PCA1 scores close to zero are G88, G97, G32, G57, G77 and G5 (Fig. 1a), G10, G100, G106, G47, G51, G52, G82, and G88 (Fig. 1b), G54, G51, G10, G100, G102, G103 and G106 (Fig. 1c), G101, G78, and G80 (Fig. 1d), G23, G16, G84, G71, G89 and G29 (Fig. 1e). As for high yield, superior genotypes for the total number of pods per plant are G67, G86, G16, G88, G34, and G100, then, G1, G23, G37, G58, and G49 for pod length (Fig. 1b), G23, G56, G61, G49, G54, and G1 for seeds per pod (Fig. 1c), G92, G100, and G88 for hundred seed weight (Fig. 1d), and G23, G16, G84, G33, and G71 for yield per plant (Fig. 1e) across the three environments. Genotypes G88, G82, G102, G101, and G23 are the most stable (Figs. 1a, 1b, 1c, 1d, and 1e respectively), then G86, G1, G23, G92, and G23 (Figs. 1a, 1b, 1c, and 1d) respectively are the superior genotypes, while genotypes G33, G85, and G40 (Figs. 1b, 1c, 1d respectively) and genotypes G1 (Figs. 1a, and 1e) had low yield.

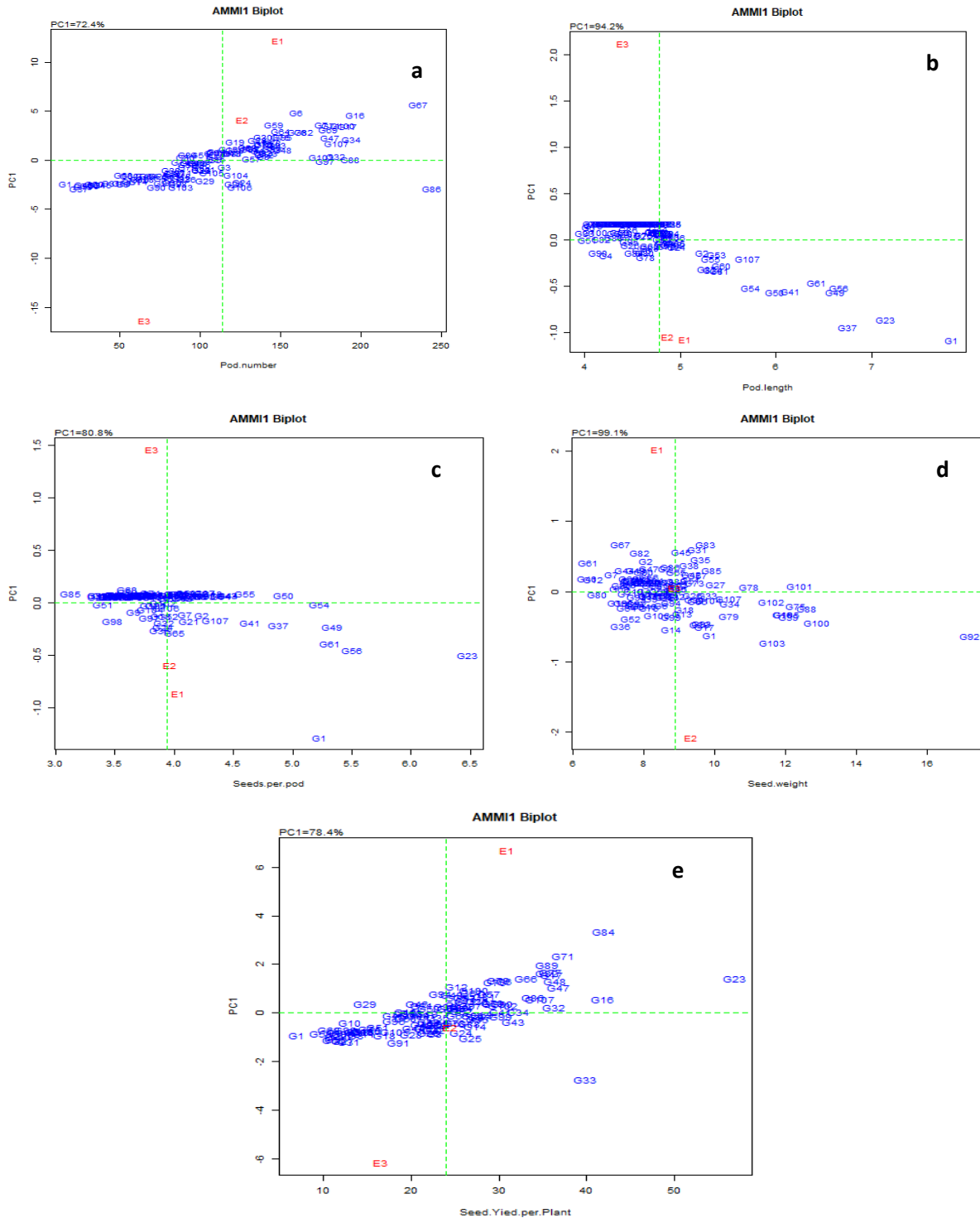
**Table 2. Environmental characteristics of the study locations**

Site	Latitude	Longitude	Altitude (m)	Soil Texture	Mean Rainfall (mm)	Min Temp (°C)	Max Temp (°C)
Edo	6° 79' N	6° 10' E	256.6	Loam	1288.22	23.85	34.44
Kogi	7° 34' N	7° 11' E	369.6	Loam	1127.09	24.47	36.54
Lagos	6° 52' N	3° 39' E	7.3	Loam	2113.00	23.50	30.71

**Table 3. Mean Square from Combined ANOVA for Yield and Yield-contributing Traits of 107 Pigeon pea genotypes tested across three Locations**

Source of Variation	df	PDN		PDL		SPP		HSW		YPP	
		MS	TVE	MS	TVE	MS	TVE	MS	TVE	MS	TVE
<b>Genotype (G)</b>	106	31585.93**	89.35	2.73**	98.21	3.22**	97.38	22.63**	86.92	951.38**	84.06
<b>Environment (E)</b>	2	45745.32**	2.44	7.40**	0.68	0.60**	0.34	73.09**	5.29	4268.71**	7.12
<b>G X E</b>	212	1450.40ns	8.21	0.046ns	1.10	0.04ns	2.28	1.01**	7.78	49.92ns	8.82
<b>Error</b>	642	1455.75	0.00	0.16	0.00	0.09	0.00	0.81	0.00	50.45	0.00
<b>PCA1</b>	107	2326.36**	72.4	0.08ns	94.2	0.07ns	80.8	1.99**	99.07	97.57ns	78.4
<b>PCA2</b>	105	557.75ns	27.6	0.00ns	5.8	0.00ns	19.2	0.02ns	0.93	1.36ns	21.6

\*, \*\* Significant at  $p \leq 0.05$  and  $p \leq 0.01$ , respectively; ns = non-significant; df: degree of freedom; MS: mean square; TVE: total variation explained; PDN: pod number; PDL: pod length; SPP: seeds per pod; HSW: hundred seed weight; YPP: yield per plant



**Fig. 1.** GEI biplot based on AMMI 1 model for the PCA1 scores of 107 pigeon pea genotypes evaluated in three environments in Nigeria for the number of pods (a), Pod length (b), Seeds per pod (c), 100 seed weight (d), and yield per plant (e)

Environment 2 (E2) had a PCA1 score that was nearly zero for pod number and seed yield per plant (Figs. 1a, and 1e), and so contributed most to the stability of the genotypes. While E3 had a high PCA1 score for pod length and seeds per pod, contributing the highest to the GEI component, then E1 and E2 had low PCA1 scores in these two respective traits. A high PCA1 score was recorded in both E1 and E2 for hundred seed weight and low PCA1 in E3. The lowest yield was recorded in E3 for all traits, while the highest yield was recorded in E2 for hundred seed weight and E1 for the other traits. Edo (E1) and Kogi (E2) were correlated with similar PCA1 scores for pod number, pod length, and seeds per pod (Figs. 1a, 1b, and 1c), while Edo (E1) and Lagos (E3) were correlated with similar PCA1 scores for hundred seed weight (Fig. 1d), thus found on the same side of the perpendicular and horizontal lines. For seed yield per plant, Kogi (E2) and Lagos (E3) are seen to be correlated (Fig. 1e).

### 3.3 AMMI 2 Analysis for PC1 and PC2 Scores- Based Graphical Representation

The AMMI biplot demonstrates the magnitude of the G x E interaction by joining environmental scores to the origin by side lines. Short vectors do not show strong interactive forces in terms of discriminating ability, whereas long vectors show a strong ability to discriminate between the genotypes. From Fig. 2, E3 had the longest vector for pod number, pod length, and seeds per pod (Figs. 2a, 2b, and 2c), while E1 has the longest vector for both hundred seed weight and yield per plant (Figs. 2d, and 2e). E2 has the shortest vector for pod number, seeds per pod, and yield per plant (Figs. 2a, 2c, and 2d), while E1 has the shortest vector for pod length and E3 for hundred seed weight. This means that E3 was a more differentiating environment, for pod number, pod length, and seeds per pod, and E1 for hundred seeds weight as well as yield per plant, while E2 is the least discriminating environment in three of these two traits.

The majority of the genotypes were near the origin and so are not sensitive to environmental interaction for all traits considered. Genotypes G6, G30, G7, and G71 are correlated with the vector for E1 and so are best adapted to the environment, while G95, G62, and G1 are correlated with the vector for E2, and G46, and G68 with E3 for pod number (Fig. 2a). For pod

length, genotypes G1, G41, G2, G53, and G24 are correlated with E1, G81, G43, G54 and G37 with E2, while genotypes G91 and G28 were shown to be correlated with E3 (Fig. 2b). For seeds per pod, G56, and G61 are correlated with E1, then G97, G87, and G43 with E2, and G88 with E3 (Fig. 2c), while in hundred seed weight, G83, G67 are correlated with E1, G14, and G16 with E2, and G35, G9 with E3 (Fig. 2d). For yield per plant, genotypes G84, G23, and G89 are correlated with E1, G96, G91, and G81 with E2, and genotypes G33, and G55 are correlated with E3 (E).

### 3.4 Comparisons of Genotypes across the Environments Using GGE Analysis

#### 3.4.1 What-won-where polygon view across the environments

The GGE what-won-where biplot of the 107 pigeon pea genotypes tested in the three environments showed the extent of genotype and environment interaction (Fig. 3). The biplot was divided into 4 sectors for pod number, seeds per pod, seed weight, and seed yield per plant (Figs. 3a, 3c, 3d, and 3e respectively), and 3 sectors for pod length (Fig. 3b), with genotypes present in all sectors of the polygon. Genotypes G67, G6, G30, G1, G3, and G86 located at the vertices of the polygon were considered superior genotypes for pod number (Fig. 3a), genotypes G1, G56, G26, G28, G33, and G3 were superior for pod length (Fig. 3b), genotypes G23, G54, G50, G85, G98, and G1 at the apex for seeds per pod (Fig. 3c), genotypes G92, G83, G67, G61, G40, G80, G36, and G103 for seed weight (Fig. 3d), genotypes G33, G23, G84, G29, and G1 for yield per plant (Fig. 3e).

All three environments were located in one of the sectors for pod length, seeds per pod, and hundred seed weight (Figs. 3b, 3c, and 3d), indicating the presence of a single mega-environment except for pod number where E2 and E3 were located in one sector and E1 in another sector, indicating two mega-environments (Fig. 3a). Similarly, two mega-environments were delineated for yield per plant, where E1 and E2 were located in one sector and E3 in another sector. Genotype G67 was at the apex of the sector with E1, and G86 was at the apex of the sector containing E2 and E3 for pod number (Fig. 3a). G1, G23, and G92 at the apices of the sectors with all three environments in the polygons for pod length, seeds per pod, and hundred seed weight respectively (Figs. 3b,



3c, and 3d), while for yield per plant, G23 was located at the apex of the sector with E1 and E2, and G33 at the apex of the sector with

E3 (Fig. 3e), indicating that these genotypes were the best performers in the three environments

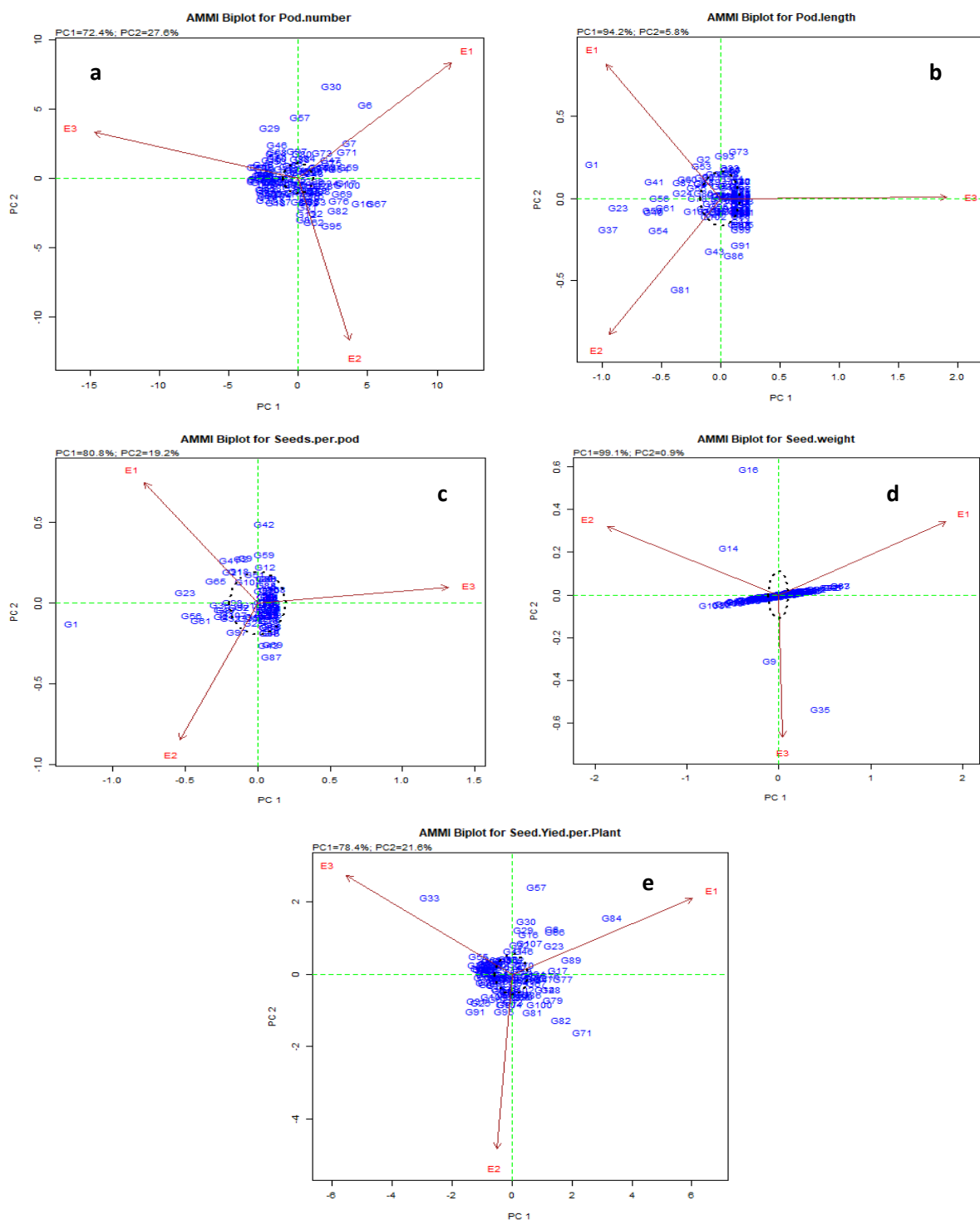
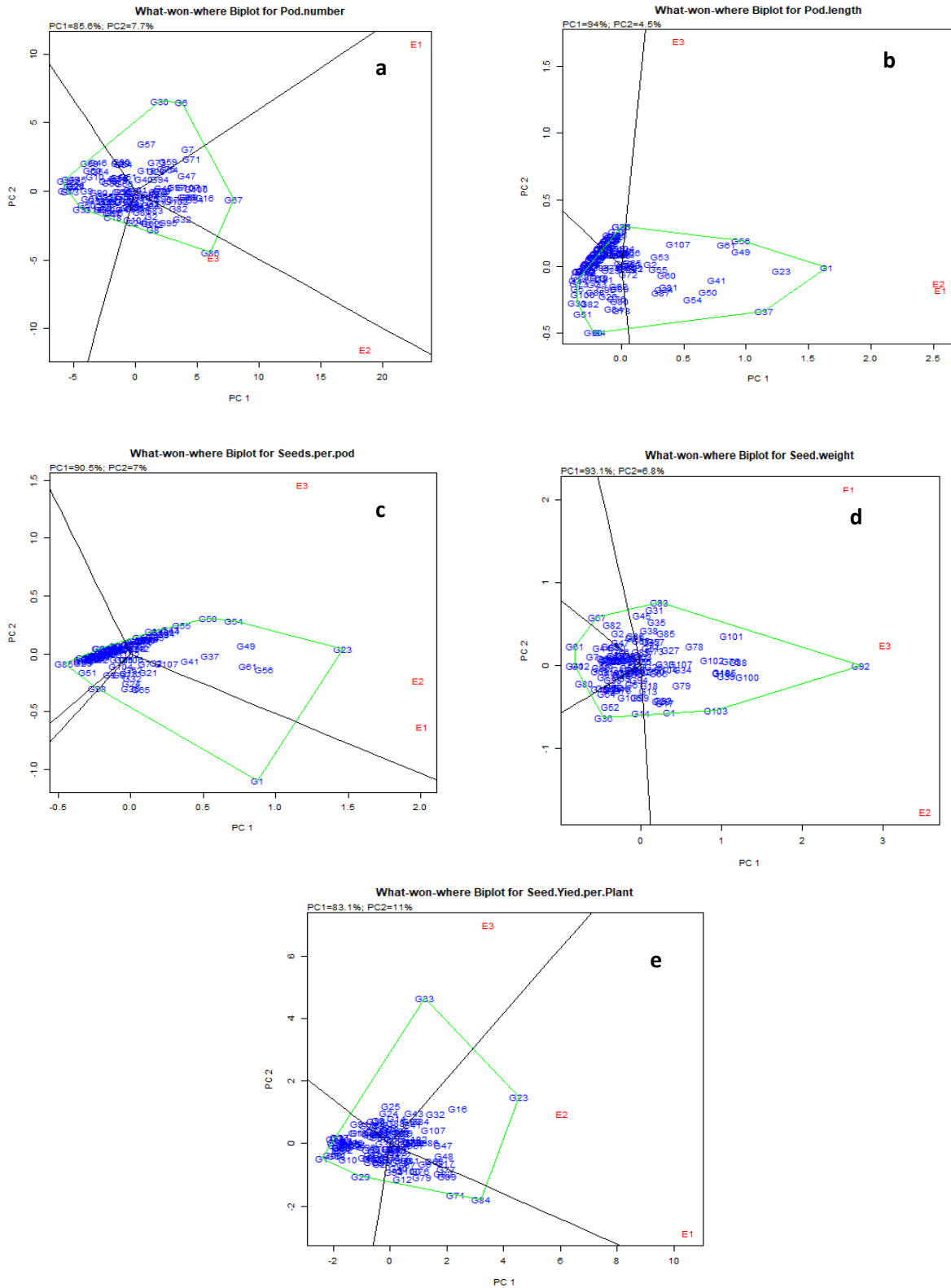


Fig. 2. GEI biplot based on AMMI 1 model for the PCA1 scores and PCA2 scores of 107 pigeon pea genotypes evaluated in three environments in Nigeria for the number of pods (a), Pod length (b), Seeds per pod (c), 100 seed weight (d), and yield per plant (e)



**Fig. 3. What-won-where GGE biplot view displaying the genotype main effect and GE interaction effect of 107 Pigeon pea genotypes in three environments for the number of pods (a), Pod length (b), Seeds per pod (c), 100 seed weight (d), and yield per plant (e)**

### 3.5 GGE Biplot of Mean and Stability Analysis and Ideal Genotype Assessment

What-won-where polygon suggested winning pigeon pea genotypes in the environments. However, there is a need to assess the mean performance and stability of all the pigeon pea genotypes to aid the selection decision. The Average Environment Axis (AEA) makes it possible to visualize the performance and stability of GGE graphically. The blue dotted line is the AEA abscissa and the perpendicular line is the ordinate (Fig. 4). The length of the abscissa to the right of the origin is above average yield, while the length to the left of the origin is below average yield. Length of ordinate on the other hand approximates the GEI. Shorter ordinate signifies lower variability, and higher stability and vice versa.

These results (Fig. 4) revealed that G32, G95, G40, G76 (Fig. 4a), G65, G107 (Fig. 4b), G49, G24, G55, G41, G107, G37 (Fig. 4c), G92, G100, G88, G101 (Fig. 4d), G16, G54, G43, G14 (Fig. 4e) are above average in yield, length, and weight as the case may be, with more stability, whereas G87, G98, G6, G57, G90, G7, G71 (Fig. 4a), G1, G23, G37, G81, G49, G54 (Fig. 4b), G1, G23, G56, G81 (Fig. 4c), G103, G1, G83 (Fig. 4d), and G23, G33, G84, G71 (Fig. 4e) are above average yielding genotypes, but with comparatively lower stability. Genotypes G1, G54, G9, G68 (Fig. 4a), G51 (Fig. 4b), G85, and G51 (Fig. 4c), and G29, G51, and G35 (Fig. 4e) are stable but their yields are below average, with the majority of the genotypes appearing to be relatively stable and clustered around the AEA abscissa. Genotypes G31, G14, G18 (Fig. 4a), G4, G78 (Fig. 4b), G98 (Fig. 4c), G67, G56 (Fig. 4d), and G1, G91 (Fig. 4e) are both below-average yielders and are less stable.

The ideal genotypes have the highest yield and absolute stability in the AEA arrowhead and the concentric circles along the arrowhead rank the inclusion of the genotypes. The closest genotype to the ideal line is G67, followed by G96, and G18 (Fig. 4a), G1 is closest to the ideal line, followed by G23, and G37 (Fig. 4b), G23, followed by G56, G49, G81, and G1 (Fig. 4c), G92, followed by G100 and G88 (Fig. 4d), G23,

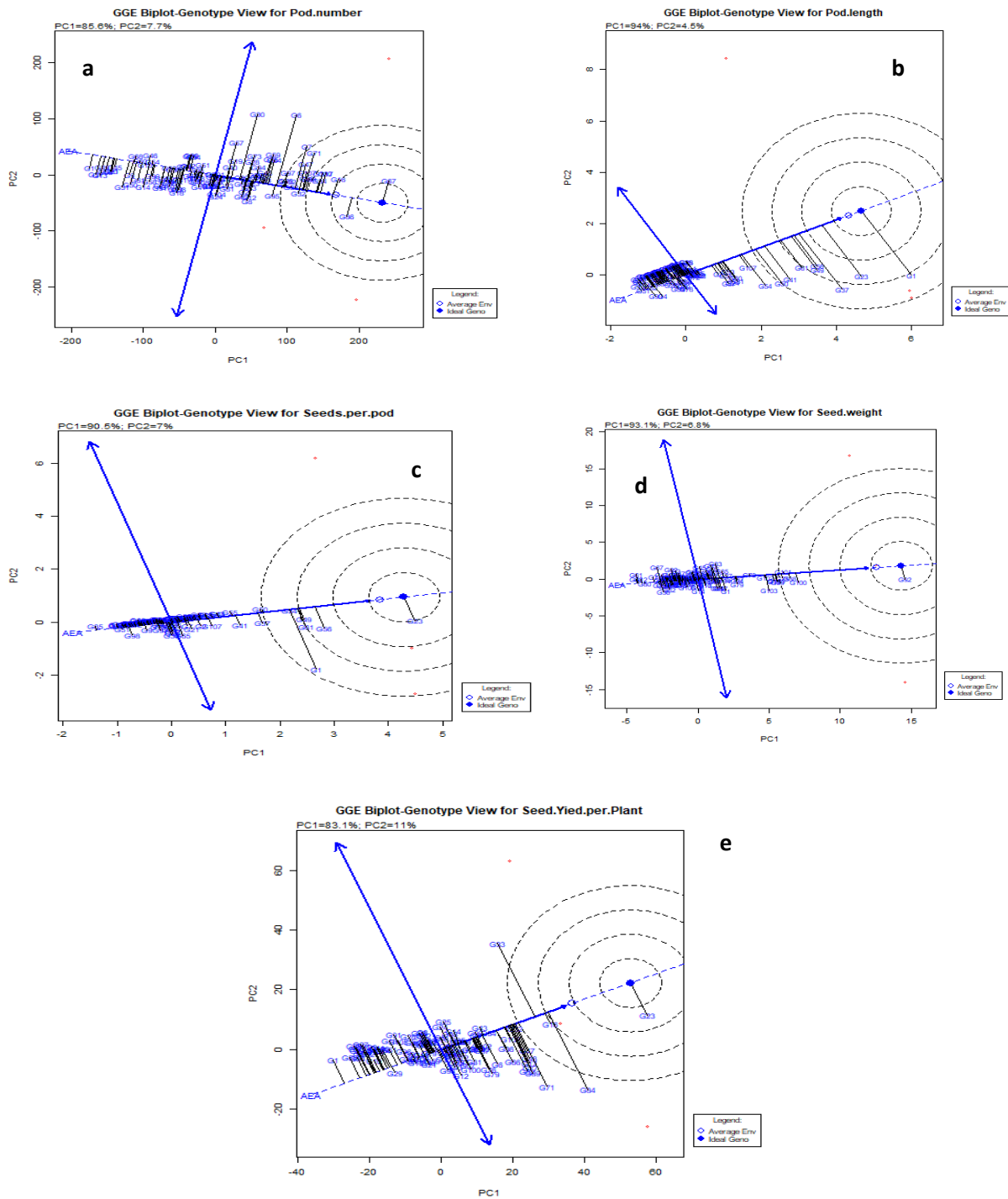
followed by G16 (Fig. 4e). These ideal lines can serve as reference lines in the evaluation of these genotypes.

### 3.6 Relationship among Test Environments

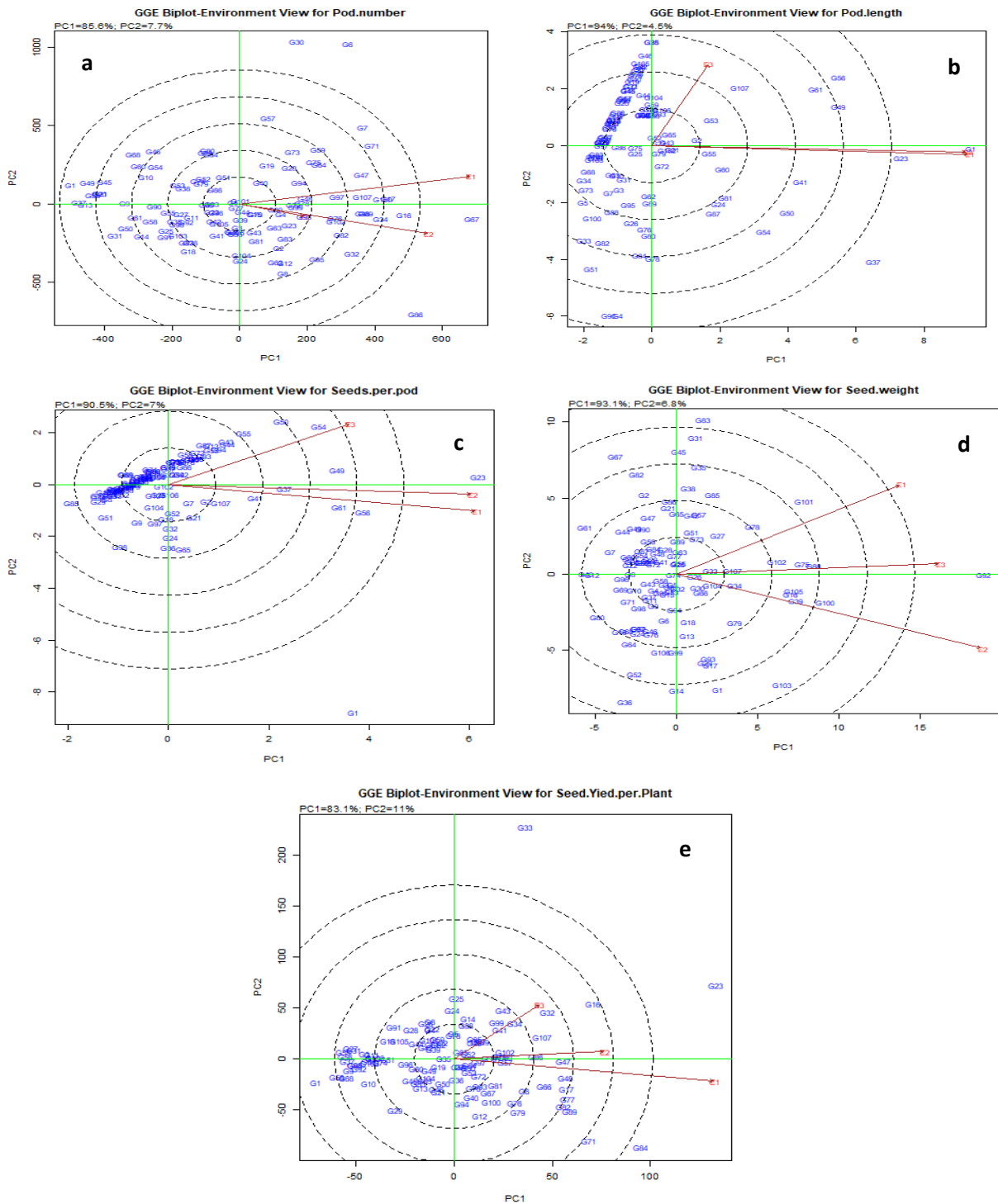
In this study, the three environments were distributed into two groups for all the traits (Fig. 5). For pod length (Fig. 5b), seeds per pod (Fig. 5c), and yield per plant (Fig. 5e), E1 and E2 are the two environments that are closely related, while E2 and E3 are closely related for pod number (Fig. 5a). For hundred seed weight (Fig. 5d), E1 and E3 are two environments that are closely related, while E2 is in a different group. In biplots for all the traits, all three environments are located within an angle of less than 90°, indicating that the three environments share similarities, but E1 and E2 share more.

### 3.7 Discriminateness and Representativeness Pattern of the GGE Biplot

A total of 93.3%, 98.5, 97.5%, 99.9%, and 94.1% (Figs. 6a, 6b, 6c, 6d and 6e respectively) of the variance was explained by the first two principal components (PCs), with PC1 explaining 85.6% and PC2 7.7% for pod number (Fig. 6a), PC1 explaining 94% and PC2 4.5% for pod length (Fig. 6b), PC1 explaining 90.5% and PC2 7% seeds per pod (Fig. 6c), PC1 explaining 93.1% and PC2 6.8% seed weight (Fig. 6d), and PC1 explaining 83.1% and 11% for PC2 seed yield per plant (Fig. 6e). The ideal environment is the environment presented with a blue dot (Fig. 6), while the open red circle represents the average environment. The dotted line is the AEA, and environments with long vectors and small angle with AEA is more informative and representative, while an environment with a short vector gives less information. In this study, E1 has the longest vector in all traits except in hundred seeds weight (Fig. 6d), while E3 has the shortest vector for all traits in the study. The ideal environment lies in between E1 and E2 for pod number (Fig. 6a), and in between E2 and E3 for the other traits (Figs. 6b, 6c, 6d, and 6e), with the average environment located close to E2 for all traits (Figs. 6a, 6b, 6c, and 6e) except for hundred seed weight where the average environment is located near E3 (Fig. 6d).



**Fig. 4. Average Environment Axis (AEA) view to rank genotypes concerning Ideal Genotypes for the number of pods (a), Pod length (b), Seeds per pod (c), 100 seed weight (d), and yield per plant (d)**



**Fig. 5. Environment view of the GGE biplot view displaying the genotype main effect and GE interaction effect of 107 Pigeon pea genotypes in three environments for number pods (a), Pod length (b), Seeds per pod (c), 100 seed weight (d), and yield per plant (e)**

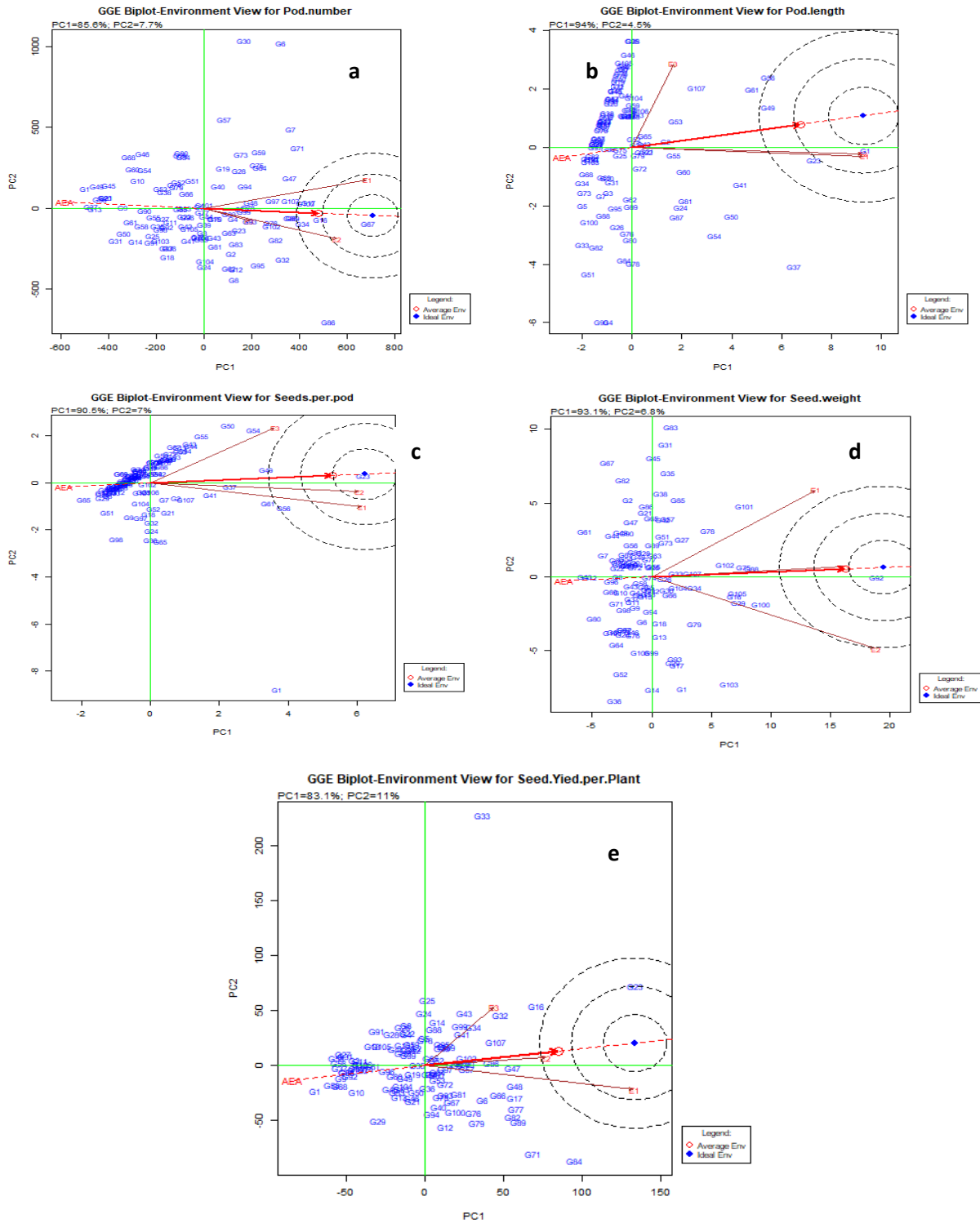


Fig. 6. Discriminativeness and representativeness pattern of the GGE for the number of pods (a), Pod length (b), Seeds per pod (c), 100 seed weight (d), and yield per plant (e)

#### 4. DISCUSSION

The mean squares for the PCA explained by the first two principal components account for 100% of the G x E interaction. Since several authors including Biswas [31] and Rao [32] have confirmed that two PCAs are sufficient for evaluating GEI, these two PCAs were used in analyzing the AMMI biplot. Gebremedhin [33] used significant PCA1 to appropriately explain the GEI, and their adaptation in barley. This revealed that the differences in the yield and yield components across the three locations were mostly due to the genotypes of the pigeon pea and to some extent the genotype by environment. The environment had the least influence on variation except in yield per plant where the environment had a higher influence than the genotype by environment. The minimal influence of the environment on the performances of the pigeon pea genotypes in this study may be because the climatic and edaphic factors of the environments are similar, and so exerted similar selection pressures on the plants. Performances of the genotypes were significant as a result of genotypic variance, and so selection based on the overall mean is dependable since GEI was minimal. These findings agree with that of Singh [28] who reported significant variation in pigeon pea yield and the total number of pods on account of genotype.

This study revealed that E2 contributed most to the stability of the number of pods and seed yield, while E3 contributed most to the stability of the hundred seed weight. These environments provided suitable conditions for the genotypes to reach optimum yield potential compared to other environments for the three traits. Genotypes such as G67, G1, G23, G92, and G23 may be suitable for planting in many environments because they are relatively stable, with an above-average yield as revealed in this study.

The GGE analysis for mean and stability analysis grouped the genotypes into those that are stable and high yielding, low stability and high yielding, those that have high stability, but low yielding, and those with low stability and low yield. The highly stable and high-yielding genotypes are G32, G65, G49, G92, and G16. Those that have low stability and are high yielding are G87, G1, G103, and G23. G1, G51, G85, and G29 have high stability and low yield (although G1 has the longest length of the pod as well as the highest number of seeds per pod, their yield is low in

terms of pod number and yield per plant), while G31, G4, G98, G67, and G1 are both low yielders and less stable. According to Khan [34], genotypes with seemingly undesirable traits are good for breeding programs, as they have yield component compensation criteria. This means that these plants may be low-yielders and less stable, but they may have other beneficial qualities such as the ability to recover quickly from environmental stresses.

The biplot analysis is the most powerful interpretative tool for AMMI analysis [35]. The AMMI model revealed that E1 was the best environment with the highest yield, while the least yield was recorded in E3. Pigeon pea has been cultivated in both Edo and Kogi for years, but the rainfall and lower average temperatures may be some of the reasons why Edo performed comparatively better than Kogi. According to Patil [36], seed yield in pigeon pea decreases with an increase in temperature, and as such, pigeon pea planted in Kogi had different climatic conditions given Kogi's proximity to the encroaching desertification. Similarly, Edo's derived savanna is expected to have better soil since the area was formally a rainforest. Planting was carried out in June in Kogi, and July in Edo and Lagos, and around the time of flowering of medium-duration genotypes (in Kogi), there was a cessation of rainfall for about two weeks in both Edo and Kogi. The crops in both locations were water-stressed during this period. However, crops in Kogi had already started flowering during this stress. This may have affected yield since the crops were at a critical stage of their reproductive development. Sinha [37] cited Lopez to have reported that yields are affected during late flowering and early pod development stages.

The adoption of the polygon to analyze 'what-won-where' is important in GGE. This study placed the three study environments under one or two sector(s) out of the three or four sectors in the biplots. The genotypes at the vertices in the sectors are G1, G23, G92, G33, G67, and G86, so can be recommended for planting in the three locations. These genotypes were the closest to the ideal lines for high yield and stability. Yohane [26] reported that either of the two sites of two mega-environment can be substituted for future pigeon pea genotype trials. Since the study revealed two mega-environments for pod number and yield per plant and the other traits as a single mega-environment, breeding efficiency can be increased by selecting either Edo or Kogi for

evaluation to increase the mega-environment. The relationship among test environment biplots grouped the locations into two. From this study, Edo and Kogi are more alike, and reducing costs will mean dropping either of these locations.

In terms of discrimination and relatedness in this study, E1 has the longest vector for all traits except hundred seed weight, whereas E2 has the longest vector and smaller angle with AEA, and so are the most discriminating environments and can be used to test genotypes for specific adaptability as well as suitable for general adaptation selection. E3 has the shortest vector for all traits considered. E2 is closest to the ideal environment for all traits, while both E2 and E3 appear as the closest to the ideal environment for cultivating pigeon pea in terms of seed weight, unfortunately, the study was carried out in a much more controlled environment in E3, and most likely the reason for its closeness to the ideal environment for the trait. Hence, a field experiment in E3 is needed to determine the certainty of the idealness of this environment for pigeon pea cultivation.

Vector for E3, being the shortest vector for all traits has the least discriminating ability, as it offered little information on the genotype variability. Although the difference in lengths of E1 and E2 are almost indistinguishable in most traits, this is not surprising as the two locations are the agroecological locations where the crop is cultivated. The cosine of the angle between the environments depicts the interrelationship between the environments. The angle between all three locations is less than 90°, showing a positive, but low correlation coefficient between all environments, however, E2 and E1 have a high correlation coefficient than E3 and E1 or E3 and E2.

## 5. CONCLUSIONS

This study evaluated the stability and adaptability of yield and yield components of pigeon pea genotypes using AMMI and GGE biplot analysis. Assessing the adaptability and stability of pigeon pea genotypes in different environmental conditions can be achieved through the use of the Multi-Environmental Trial (MET). In order to determine the suitability of a genotype for commercial cultivation, it is important to evaluate its susceptibility to genotype by environment interaction (GEI). This study revealed that genotypes G67, G86, G1, G23, G92, and G33 are ideal genotypes with high yields and

relatively high stability. The three locations contributed to the morphological traits. Edo contributed the highest to the GEI and thus revealing its suitability for specific adaptation of pigeon pea genotypes, while Kogi contributed to the stability of the genotypes for pod number and seed yield per plant, and Lagos for seed weight. The environment had the least influence on variation. ISC 169, ISC-3, ISC197, ISC-76, and ISC 134 pigeon pea genotypes were highly stable and high yielding across all environments, and Kogi was revealed to be the closest to the ideal environment for pigeon pea production, while Edo is a highly suitable location for specific adaptation of pigeon pea genotypes.

This study grouped the pigeon pea accessions into four major stability and mean yield groups. The first group comprises those that are highly stable and have high yield potentials, and they include G32 (ISC-169), G65 (ISC-3), G49 (ISC-197), G92 (ISC-76), and G16 (ISC-134). The second group is comprised of those that have low stability, but high yield potential and they include G87, G1, G103, and G23. The third group comprises those with high stability but low yield, and they include G1, G51, G85, and G29. The last group comprises genotypes with low stability and low yield, and they include G31, G4, G98, G67, and G1. G88, G16, G23, G71, and G100 performed well across all environments and are relatively stable. Therefore, they are the ideal genotypes for the study and the reference crop for breeding programs across environments. Kogi is the ideal environment for testing genotypes, and since it is an agroecological zone for pigeon pea cultivation, it makes it easier. Lagos has a low altitude and the planting in Lagos was carried out in a screen house where climatic factors were controlled. There is, therefore, a need to replicate this experiment on the field in Lagos to ascertain whether or not Lagos supports the cultivation of pigeon pea.

## ACKNOWLEDGEMENTS

The authors acknowledge the International Crops Research Institute for the Semi-Arid Tropics (ICRISAT) Nigeria and Niger for providing the pigeonpea accessions for the research, and the Department of Botany for providing the screen house for the research.

## COMPETING INTERESTS

Authors have declared that no competing interests exist.



## REFERENCES

1. Odeny DA. The potential of pigeonpea (*Cajanus cajan* (L.) Millsp.) in Africa. Nat. Resour. Forum 2007; 31:297–305. Available: <https://doi.org/10.1111/j.1477-8947.2007.00157.x>.
2. Egbe OM, Kalu BA. Farming System Study: Participatory Rural Appraisal of Pigeonpea Cropping Systems in Southern Guinea Savanna of Nigeria. J. on Environ. (Abia State University Environmental Review). 2006;5(1):37-47.
3. Fatokimi EO, Tanimonure VA. Analysis of the current situation and future outlooks for pigeon pea (*Cajanus Cajan*) production in Oyo State, Nigeria: A Markov Chain model approach. J. Agric. Res. 2021;6:1-8. Available:<https://doi.org/10.1016/j.jafr.2021.100218>.
4. Rawal V, Navarro DK (2019) The Global Economy of Pulses. Food and Agriculture Organization of the United Nations, Rome; 2019. Available: <https://doi.org/10.4060/I7108EN>.
5. Varshney R K, Chen W, Li Y, Bharti AK, Saxena RK, Schlueter JA, Jackson SA. Draft genome sequence of pigeonpea (*Cajanus cajan*), an orphan legume crop of resource-poor farmers. Nat Biotechnol. 2012; 30(1):83. Available: <https://doi.org/10.1038/nbt.2022>
6. Lans C. Comparison of plants used for skin and stomach problems in Trinidad and Tobago with Asian Ethnomedicine. J. Ethnobiol. Ethnomed. 2007;3:3. Available: <https://doi.org/doi:10.1186/1746-4269-3-3>
7. Upadhyay B, Parveen D, Dhaker AK, Kumar A. Ethnomedicinal and Ethnopharmacol-statistical studies of Eastern Rajasthan, India. J. Ethnopharmacol. 2010;129:64-86. Available:<https://doi.org/10.1016/j.jep.2010.02.026>.
8. Yuan-gang Z, Xiao-lei YF, NanWu Y, Michael W. Chemical composition of the SFE-CO<sub>2</sub> extracts from *Cajanus cajan* (L.) Huth and their antimicrobial activity *In vitro* and *In vivo*. J. Phytomed. 2010;17:1095-1101. Available:<https://doi.org/10.1016/j.phymed.2010.04.005>.
9. Pal S, Mishra P, Sachan N, Ghosh AK. Biological Activities and Medicinal Properties of *Cajanus cajan* (L) Millsp. J Adv Pharm Technol Res. 2016;2(4):207-214. Available: <https://doi.org/10.4103/2231-4040.90874>.
10. Fasoyiro SB, Farinde EO, Chete OB, Ajani AO. Knowledge assessment, training and consumer acceptability on cassava-pigeon pea processing at Akufo, Oyo state. Greener J. Agric. Sci. 2019; 9(1):32–36. Available:<https://doi.org/10.15580/GJAS.2019.1.011619016>.
11. National Agricultural Quarantine Service (NAQS), Nigeria. The Export Potential of Pigeon Pea in Nigeria. Nigeria to Export \$100bn Pigeon Pea to India - Punch Newspapers; 2017. Available: <https://punchng.com/nigeria-to-export-100bn-pigeon-pea-to-india/>.
12. CILLS, Comité Permanent Inter-états de Lutte contre la Sécheresse dans le Sahel. Landscapes of West Africa – A Window on a Changing World. U.S. Geological Survey Earth. Resources Observation and Science Centre, Garretson, United States of America; 2016.
13. Macauley BM. Land degradation in Northern Nigeria: the impacts and implications of human-related and climatic factors. Afri. J. Environ. Sci. Technol. 2014;8(5):267-273. Available:<https://doi.org/10.5897/AJEST2013.1584>.
14. Agbelade AD, Fagbemigun OA. Assessment of incentives for forest biodiversity conservation in rain forest and derived savannah vegetation zones of Ekiti State, Nigeria. Forest Research. 2015; 4(3):1-5.
15. Esan V, Ojemola O. Evaluation of production systems, traditional knowledge of pigeon pea (*Cajanus cajan*) and risks of extinction of pigeon pea, jack bean (*Canavalia ensiformis*) and lubia bean (*Lablab purpureus*) in some parts of Southwest Nigeria. J. Exp. Agric. Int. 2018;21(4):1-11. Available:<https://doi.org/10.9734/JEAI/2018/39835>.
16. Chaudhary AK, Sultana R, Pratap A, Nadarajan N, Jha UC. Breeding for abiotic stress in pigeonpea. Journal of Food Legumes. 2011;24(3):165-174.
17. Ezeaku IE, Ajeigbe HA, Okechukwu EC. Evaluation of Introduced Pigeonpea (*Cajanus cajan* (L.) Millsp.) Genotypes for Growth and Yield Performance in Sudano-

- Sahelian Ecology of Nigeria. J. Anim. Plant Sci. 2016;26(1):163-169.
18. Donça MCB, Pedro1 C, Somueque1 SI, Divage BAF, Colial HV, Gimo1 ST, Alexandre DC, Bambo EC, Barbosa IP, Malikouski RG, Muquera AGP. Genotypic Selection for Stability and Yield Adaptability of Short-Duration Pigeon Pea in Mozambique using REML/BLUP Mixed Models. East African Scholars J. Agric. Life Sci. 2021;4(10):201-207. Available:https://doi.org/10.36349/easjals.2021.v04i10.002
  19. Dasbak MAD, Asiegbu JE. Grain yield assessment of six pigeonpea genotypes in production systems and their ratoon ability in a humid tropical agroecology of Nigeria. AJOL. 2012;11(2):38–45. Available:https://doi.org/10.4314/as.v11i2.6.10.4314/as.v11i2.6
  20. Saxena KB, Sawargaonkar SL. First information on heterotic groups in pigeonpea [*Cajanus cajan* (L.) Millsp.]. Euphytica. 2014;200(2):187-196. Available: https://doi.org/10.1007/s10681-014-1142-0
  21. Sawargaonkar SL, Saxena KB, Madrap IA, Rathore A. Stability analysis of yield and related traits in pigeonpea hybrids. Indian J. Gene. 2011;24 (3):184-193.
  22. Mudaraddi B, Saxena KB. Character association between seed yield and component traits among CMS-based pigeonpea hybrids. Electron. J. Plant Breed. 2013;4(1):1086-1089.
  23. Kumar MVN, Ramya V, Kumar CVS, Raju1 T, Kumar NMS, Seshu G, Sathish G, Bhadru D, Ramana MV. Identification of pigeonpea genotypes with wider adaptability to rainfed environments through AMMI and GGE biplot analyses. Indian J. Genet. Plant Breed. 2021; 81(1):63-73. Available:https://doi.org/10.31742/IJGPB.81.1.7
  24. Gauch HG. Statistical analysis of regional trials- AMMI analysis of factorial design. 1st ed. Elsevier: New York; 1992.
  25. Yan W, Hunt LA, Sheng Q, Szlavnic Z. Cultivar evaluation and mega-environment investigation based on the GGE biplot. Crop Sci. 2000;40(3):597-605. Available:https://doi.org/10.2135/cropsci2000.403597x
  26. Yohane EN, Shimelis H, Laing M, Mathew I, and Shayanowako A. Genotype-by-environment interaction and stability analyses of grain yield in pigeonpea [*Cajanus cajan* (L.) Millspaugh]. Acta Agriculturae Scandinavica, Section B-Soil & Plant Science. 2020;71(3):145-155. Available:https://doi.org/10.1080/09064710.2020.1859608
  27. Ojwang JD, Nyankanga R, Ganga Rao NVPRG, Imungi J. Evaluation of vegetable pigeonpea [*Cajanus cajan* (L.) Millsp] genotypes for yield stability. CABI Agric. Biosci. 2021;2(41):1-10. Available: https://doi.org/10.1186/s43170-021-00061-8.
  28. Singh J, Kumar1 A, Fiyaz AR, Singh MK. Stability analysis of pigeon pea genotypes by the deployment of AMMI model under rainfed environment. Legume Res. 2018; 41(2):182-188. Available:https://doi.org/10.18805/lr.v0i0.7851.
  29. Navneet S, Silarwar RS, Sing AK, Kumar RA. Genetic Diversity in Pigeonpea [*Cajanus cajan* L. Millsp.]. Inter. J. Agric. Sci. 2017; 9(18): 4177-4179.
  30. IBPGR and ICRISAT. Descriptors for pigeon pea (*Cajanus cajan* (L) Millsp). International Board for Plant Genetic Resources, Rome, Italy; International Crops Research Institute for the Semi-Arid tropics, Patancheru, India; 1993.
  31. Biswas T, Debasis M, Arpita D, Dinesh PK, Anirban M, Parihar AK, Gupta S. Additive main effects and multiplicative interaction in field pea (*Pisum sativum* L.) genotypes across the major agro-climatic zones in India. Legume Res. 2019;44(8):894-899. Available: https://doi.org/10.18805/LR-4166
  32. Rao PJM, Kishore NS, Sandeep S et al. Evaluation of Performance and Yield Stability Analysis Based on AMMI and GGE-Biplot in Promising Pigeonpea [*Cajanus cajan* (L.) Millspaugh] Genotypes. Legume Res. 2020;5(11): 1414-1420. Available: https://doi.org/10.18805/LR-4299
  33. Gebremedhin W, Firew, M. Tesfye, B. Stability analysis of food barley genotypes in Northern Ethiopia. Afri. Crop Sci. J. 2014;22:145–54.
  34. Khan MMH, Rafii1 MY, Ramlee SI, Jusoh M, Al Mamun M. AMMI and GGE biplot analysis for yield performance and stability assessment of selected Bambara groundnut (*Vigna subterranea* L. Verdc.) genotypes under the

- multi-environmental trials (METs). Sci. Rep. 2021;11:22791.  
Available: <https://doi.org/10.1038/s41598-021-01411-2>
35. Ram K, Munjal R, Kesh H, Kumari S, Kumari A. AMMI and GGE Biplot Analysis for Yield Stability of Wheat Genotypes under Drought and High-Temperature Stress. Int. J. Curr. Microbiol. Appl. sci. 2020;9(5):377-389.  
Available:<https://doi.org/10.20546/ijcmas.2020.905.043>.
36. Patil DD, Pandey V, Gurja R, Patel HP. Effect of Inter-seasonal variation in temperature and rainfall on yield of pigeon pea cultivars using CROPGROW model. J. Agric. Meteorol. 2018;20(4): 286-291.
37. Sinha P, Pazhamala LT, Singh VK, Rachit K, Saxena LK, Azam S, Khan AW, Varshney RK. Identification and Validation of Selected Universal Stress Protein Domain Containing Drought-Responsive Genes in Pigeonpea (*Cajanus cajan* L.). Front. Plant Sci. 2016;6(1065): 1-10.  
Available:<https://doi.org/10.3389/fpls.2015.01065>

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