Adaptability and Phenotypic Stability of Resistance to Two Viral Diseases and Yield Traits in Cassava

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Abstract
Cassava productivity is hampered by pests and diseases including cassava mosaic disease (CMD) and cassava brown streak disease (CBSD). The main objective of this study was to identify stable superior genotypes that combine disease resistance and high yield. Sixteen cassava genotypes were planted in a randomized complete block design with three replications for six planting seasons (years) at five sites in Tanzania. The genotypes were assessed using the additive main effect and multiplicative interaction (AMMI) analysis, and highly significant (P < 0.001) effects of genotype, environment, and genotype-by-environment (G*E) interactions were observed for all traits studied. Percent sum of squares (SS) due to environment (12.66% - 85.23%) was the highest followed by G*E (14.12% - 39.56%) for CMD foliar symptoms, root weight and dry matter. On the other hand, % SS due to genotype (52.14 % - 69.14%) was highest followed by G*E (26.14% - 35.91%) for CBSD foliar and root symptoms indicating that the environment and G*E greatly influenced trait expression. The most stable genotypes which combined disease resistance and high yield were NDL 2003/31 and NDL 2003/111. The findings of this study will give impetus for the release of new cassava varieties that are not only high yielding but are also dually resistant to both CMD and CBSD in different locations and sites.

Keywords
Cassava Brown Streak Disease, Cassava Mosaic Disease, Disease Resistance, Genotype Environment Interaction, High Yield
1. Introduction

Cassava (*Manihot esculenta* Crantz) is a vital food staple in sub-Saharan Africa (SSA), ranked as the number one root crop, followed by sweet potato and yam [1]. With over 300 million MT of annual root production [1], cassava is a major source of carbohydrates in the diet of millions of people in SSA and is grown as a famine reserve crop owing to its tolerance of harsh environmental conditions [2] [3]. The crop also has industrial application as it is used to produce high-quality flour, starch, beverages, animal feeds, alcohol, biofuel, detergents, textiles, plastics and pharmaceutical products [4] [5] [6] [7].

Although Tanzania has the largest area (885,091 ha) under cassava production in East Africa, its average yield is low at 5.7 t/ha [1], which is far below the estimated yield potential of cassava (50 - 60 t/ha) [8]. This is due to many biotic and abiotic factors including the two viral diseases: cassava mosaic disease (CMD), and cassava brown streak disease (CBSD) [9] [10] [11]. Cassava roots affected by CBSD have a brown necrotic rot and are unfit for consumption. By contrast, storage roots of cassava plants severely affected by CMD fail to bulk because their leaves become chlorotic and mottled, thus having impeded photosynthesis and leading to stunted growth [12]. Dual infections of CMD and CBSD are common and a serious threat to cassava production and food security as losses more than 80% have been reported in susceptible varieties [13].

Deployment of cassava varieties with dual resistance to both diseases is currently being pursued as the most effective and sustainable way to manage the devastating effects of the viral diseases in Eastern and Southern Africa [14]. CMD, CBSD and yield traits expression in cassava can be influenced by the environment leading to varied phenotypes in different environments [15] [16] [17]. This is defined as genotype-by-environment (*G*E) interaction [18] and it can result from differences in the sensitivities of genotypes to the conditions in the target environment [19]. This leads to inconsistent performances across different environments; therefore, limiting the efficiency of selection of superior genotypes.

The objective of most cassava improvement programmes is to identify and select diseases free, high yielding and stable genotypes across several environments and seasons. The efficiency and success of such selections depend on the consistency of the performances of genotypes in varying environments [20] [21]. For this reason, genotypes are tested in diverse environments to assess their adaptability and stability. Genotypes whose *G*E effects are not significant are said to be stable [22]. Several methods have been used to assess the *G*E effect and stability in crop performances including the additive main effect and multiplicative interaction (AMMI) model [23] [24].

The AMMI model fits the sum of several multiplicative terms rather than only one multiplicative term in assessing the performance of genotypes in different environments [25]. AMMI analysis can be used to determine the stability of the genotypes across locations using the PCA (principal component axis) scores and AMMI stability value (ASV) [26]. The ASV is based on the AMMI model’s IPCA1
and IPCA2 (interaction principal components axes 1 and 2), respectively scores for each genotype [27]. Genotypes having the least ASV are considered as widely adapted genotypes. Similarly, IPCA2 score near zero indicates more stable genotypes whilst large values represent more responsive and less stable genotypes.

However, the stability parameter alone does not give much information about the yield or performance of a genotype and cannot be used as the only selection parameter since most stable genotypes would not necessarily be the best with regards to desirable traits. Therefore, Jiwuba et al. [15], Nduwumuremyi et al. [28] and Tumuhimbise et al. [29] used yield stability index (YSI) and genotype stability index (GSI) which incorporate high yield or performance with stability. Both the YSI and the GSI are based on the sum of the ranking due to ASV scores and yield or performance ranking. Low GSI value indicates desirable genotypes with high mean yield or performance and stability.

The main aim of this research was to analyze the effects of G*E interaction on resistance to CMD, CBSD and yield traits on 16 cassava genotypes using the AMMI model. The specific objectives were to 1) Identify superior genotypes that exhibit high stability which combine CMD and CBSD resistance and high yield; 2) Identify environments that best represent the target environment for high expression of the traits.

2. Materials and Methods
2.1. Study Location and Germplasm

The study was done in five sites: Chambezi, Mtopwa, Nachingwea, Naliendele and Mtopwa for six planting seasons (2013, 2014, 2015, 2016, 2017 and 2018) (Table 1). Advanced breeding lines including released improved varieties and local landraces were evaluated in the study (Table 2). The advanced breeding lines and improved varieties were obtained from the TARI-Naliendele or the

| Descriptions       | Sites                                      |
|-------------------|-------------------------------------------|
| Site location     | Chambezi [42] [43] Mtopwa [44] Nachingwea [44] Naliendele [44] Ségera [45] [46] |
| Co-ordinates      | 06°55’S, 38°91’E 10°41’S, 39°23’E 10°20’S, 38°46’E 10°22’S, 40°10’E 05°31’S and 38°54’E |
| Altitude          | 46 m 760 m 465 m 111 m 290 m               |
| Soil Type         | Ferralic Cambisol Ferralic Cambisol Veti-acric Ferrasols - Xanthic Ferrasols - Rhodic Veti-acric Ferrasols - Xanthic Rhodic Ferrasols (Orthic, Xanthic) |
| Soil texture      | Sandy soils Deep, highly weathered, well drained sandy clay loam Deep, highly weathered, red sandy clay loam Deep, highly weathered, well drained sandy clay loam Well drained, moderately deep or deep reddish and yellowish sandy clay loam to clay. Well-structured soil with low fertility |
| Soil pH           | 5.0 - 7.0 4.5 - 6.5 4.5 - 8.2 4.5 - 6.5 5.5 - 7.5 |
### Table 2. Pedigree and status of advanced breeding lines and local cassava cultivars.

| Genotype                | Female parent | Male parent | Remarks              | Status                      |
|-------------------------|---------------|-------------|----------------------|-----------------------------|
| Albert                  | Unknown       | Unknown     | Putative fullsib     | Local landrace              |
| KBH 2002/26 (Mkuranga 1)| KBH 95/082    | Unknown     | Halfsib              | Released                    |
| KBH 2002/477            | Kiroba        | Unknown     | Halfsib              | Not released                |
| KBH 2002/482 (Kizimbani)| Kiroba        | Unknown     | Halfsib              | Released                    |
| KBH 2002/494            | Kiroba        | Unknown     | Halfsib              | Not released                |
| KBH 2002/66 (Kipusa)   | 196/1632      | Unknown     | Halfsib              | Released                    |
| KBH 96/1056             | Kiroba        | Unknown     | Halfsib              | Candidate line for official release |
| Kiroba                  | Unknown       | Unknown     | Halfsib              | Released                    |
| Mahiza                  | Unknown       | Unknown     | Halfsib              | Local landrace              |
| Mkumba                  | Namikonga     | AR42-4      | Fullsib              | Released                    |
| Naliendele 034          | Kibaha        | Unknown     | Halfsib              | Released but CBSD resistance broken |
| NDL 2003/111            | Namikonga     | Kalulu      | Fullsib              | Candidate line for official release |
| NDL 2003/031            | Nachinyaya    | Kiroba      | Fullsib              | Candidate line for official release |
| NDL 2005/1471           | Nachinyaya    | Unknown     | Halfsib              | Candidate line for official release |
| NDL 2005/1472           | Nachinyaya    | Unknown     | Halfsib              | Candidate line for official release |
| Pwani                   | Namikonga     | AR42-4      | Fullsib              | Released                    |

Fullsib—genotypes with known male and female parents; Halfsib—genotypes with a known mother that was open pollinated hence male parent is unknown.

International Institute for Tropical Agriculture’s breeding programmes, while the local landraces were obtained from farmers’ fields.

#### 2.2. Experimental Design

A randomized complete block design with three replicates was used for this study. Cassava cuttings (about 25 cm long with 4 to 5 nodes and viable buds) from each of the genotypes were planted in 4 rows with 10 cuttings each at a spacing of 1.0 m × 1.0 m, resulting in a total of 40 plants/plot/replicate. To increase disease inoculum pressure, susceptible cassava varieties Albert and Limbanga were planted as spreader rows for CBSD and CMD, respectively [30]. Albert and Limbanga cutting were planted alternately after every 8 plots and as a border row around each replicate. Released varieties and landraces including Albert, Kiroba, Pwani, Mahiza, Mkumba and Naliendele 134 were planted as controls in the experiment. Neither fertilizer nor irrigation was applied; the field was rain-fed throughout the growing period but was kept weed-free.
2.3. Data Collection

Data on several parameters were collected including CMD and CBSD foliar severity at 3, 6, and 9 MAP; root necrosis; root weight (t/ha), and dry matter content during harvest at 12 MAP. CMD foliar severity was scored on a 1 - 5 scale where: 1 = no visible symptoms; 2 = mild distortion only at the base of leaflets with the rest of leaflets appearing green and healthy/mild chlorotic pattern over entire leaflets; 3 = conspicuous mosaic pattern throughout the leaf, narrowing and distortion of lower 1/3 of leaflets; 4 = severe mosaic, distortion of two-thirds of leaflets and general reduction of leaf size; and 5 = severe mosaic, distortion of ¾ of leaflets, twisted and malformed leaves [31].

CBSD foliar severity was scored on a 1 - 5 scale where: 1 = no visible symptoms; 2 = mild foliar mosaic on some leaves and no stem lesions; 3 = foliar mosaic with mild stem lesions and no die back; 4 = foliar mosaic and pronounced stem lesions and no dieback; and 5 = defoliation with pronounced stem lesions and dieback [32]. At 12 MAP, plants were harvested, and roots were examined for CBSD root symptoms. Roots from each plant were chopped longitudinally and transversely to identify the presence of necrotic patches on the starch bearing tissues. Scoring for root necrosis severity was also done based on a 1 - 5 where: 1 = no clear symptoms; 2 = <5% of root necrotic; 3 = 5% - 25% of root necrotic; 4 = 25% - 50% root necrotic and mild root constriction; and 5 = >50% of root necrotic [32] [33] [34]. Roots from each plant were harvested and chopped longitudinally and transversely to check for root necrosis on the starch bearing tissues. Root weight in tonnes per hectare (t/ha) was estimated according to Masinde et al. [16] while root dry matter content using the specific gravity method [35].

\[
\text{Root weight (t/ha)} = \frac{\text{Root weight (kg/m}^2\text{)} \times 10000}{1000}
\]

\[
\text{Dry matter content} = 158.3 \times \left[ \frac{\text{Weight of roots in air}}{\text{Weight of roots in air} - \text{Weight of roots in water}} \right] - 142
\]

2.4. Data Analysis

The AMMI model was used to determine the stability of the genotypes across environments. The AMMI model first fits the additive effects for the genotypes and the growing environments (five growing sites and six seasons) and multiplicative term for G*E interactions. The AMMI model according to Gauch [36] and Farshadfar et al. [37] is presented as

\[
Y_{ij} = \mu + g_i + e_j + \sum_{k=1}^{n} \lambda_k a_k r_{ik} + e_{ij}
\]

where

\( Y_{ij} \) = Phenotypic trait e.g. yield of the \( i^{th} \) genotype in the \( j^{th} \) environment
\( \mu \) = Grand mean
\( g_i \) and \( e_j \) = Genotype and environment deviations from the grand mean,
respectively

\[ k = \text{The number of principal components retained in the model} \]
\[ \lambda_k = \text{The eigenvalue of the PCA axis } k \]
\[ \alpha_{ik} \text{ and } \gamma_{jk} = \text{the principal component scores for PCA axis } k \text{ of the } i^{th} \text{ genotype and the } j^{th} \text{ environment, respectively} \]
\[ e_{ij} = \text{Residual} \]

The ASV was calculated for each genotype according to the relative contributions of IPCA1 and IPCA2 to the interaction sum of squares. The ASV has been defined as the distance from the coordinate point to the origin in a two-dimensional scatterplot of the first IPCA1 scores against the second IPCA2 [27] [38]. The IPCA1 accounts for most of the \( G^E \) variation. The IPCA1 scores are weighted by the ratio of IPCA1 SS (from the AMMI ANOVA) to IPCA2 SS in the ASV formula as

\[
\text{ASV} = \sqrt{\left( \frac{\text{IPCA1 sum of squares}}{\text{IPCA2 sum of squares}} \right) (\text{IPCA1 score})^2 + (\text{IPCA2 score})^2}
\]

(4)

The larger the IPCA score is, either negative or positive, the more adapted a genotype is to a certain environment. Smaller ASV scores indicate a more stable genotype across environments [39]. Genotype stability index (GSI) was also calculated using the sum of the ranking based on trait and ranking based on the AMMI stability value. GSI incorporates both the mean and stability of the trait being studied in a single criterion. Low values of both parameters show suitable genotypes for example those with high mean yield and stability [29] [40]. Both AMMI and biplot analysis were computed using the R package Agricolae [41].

\[
\text{GSI} = \text{RASV} + \text{RY},
\]

where

\[
\text{RASV} = \text{Rank of the genotypes based on the AMMI stability value},
\]
\[
\text{RY} = \text{Rank of the genotypes based on yield across environments.}
\]

3. Results

3.1. CMD Foliar Symptoms

The results of the combined AMMI analysis of variance revealed highly significant (\( P \leq 0.001 \)) effects of genotype, environment and \( G^E \) for CMD foliar symptoms at 3, 6, and 9 MAP (Table 3). Percent sum of squares (SS) due to environment (41.04% - 42.61%) was higher than % SS due to \( G^E \) (30.19% - 36.35%) and genotype (22.61% - 27.24%) indicating that environment greatly influenced the expression of CMD symptoms. \( G^E \) interaction SS was partitioned into four significant Interactive Principal Components Axes (IPCA) for CMD 3 MAP while CMD 6 and 9 MAP had three significant IPCAs. IPCA1 and IPCA2 accounted for a total SS of 67.88%, 80.5% and 73.19% of the \( G^E \) variation for CMD at 3, 6 and 9 MAP, respectively. This justified the use of AMMI2 (IPCA2 vs IPCA1) biplot model for CMD foliar symptoms. The mean CMD foliar symptoms were 1.17, 1.24 and 1.21 at 3, 6, and 9 MAP, respectively (Table S1).
### Table 3. Combined AMMI ANOVA for CMD and CBSD foliar symptoms at 3, 6, and 9 MAP of 16 cassava genotypes evaluated across 30 environments (6 planting seasons × 5 sites).

| Source          | df    | CMD 3 MAP | CMD 6 MAP | CMD 9 MAP | CBSD 3 MAP | CBSD 6 MAP | CBSD 9 MAP |
|-----------------|-------|-----------|-----------|-----------|------------|------------|------------|
| Treatment       | 479   | 0.19***   | 0.36***   | 0.26***   | 0.88***    | 1.26***    | 1.02***    |
| Genotype (G)    | 15    | 1.37***   | 2.92***   | 2.23***   | 17.15***   | 25.70***   | 22.43***   |
| Environment (E) | 29    | 1.28***   | 2.54***   | 1.80***   | 0.61***    | 1.30***    | 0.79***    |
| Block           | 60    | 0.08***   | 0.10*     | 0.08***   | 0.11       | 0.16*      | 0.09       |
| Interaction (G'E) | 435   | 0.08***   | 0.13***   | 0.09***   | 0.34***    | 0.42***    | 0.29***    |
| IPCA1           | 43    | 0.33***   | 0.58***   | 0.38***   | 1.43***    | 1.73***    | 1.09***    |
| IPCA2           | 41    | 0.20***   | 0.48***   | 0.27***   | 0.83***    | 1.36***    | 0.84***    |
| IPCA3           | 39    | 0.13***   | 0.11*     | 0.11***   | 0.39***    | 0.47*      | 0.27***    |
| IPCA4           | 37    | 0.05*     | 0.06      | 0.05      | 0.35*      | 0.34       | 0.28*      |
| IPCA5           | 35    | 0.03      | 0.04      | 0.04      | 0.22       | 0.25       | 0.24       |
| Error           | 899   | 0.03      | 0.07      | 0.04      | 0.13       | 0.11       | 0.09       |

#### Sum of squares

| Source          | df    | Sum of squares |
|-----------------|-------|----------------|
| Treatment       | 479   | 90.70          |
| Genotype (G)    | 15    | 20.51          |
| Environment (E) | 29    | 37.22          |
| Block           | 60    | 4.47           |
| Interaction (G'E) | 435   | 32.97          |
| IPCA1           | 43    | 14.08          |
| IPCA2           | 41    | 8.30           |
| IPCA3           | 39    | 5.11           |
| IPCA4           | 37    | 1.98           |
| IPCA5           | 35    | 1.20           |
| Error           | 899   | 31.2           |

% treatment SS due to G: 22.61, 25.41, 27.24, 60.90, 63.90, 69.14
% treatment SS due to E: 41.04, 42.61, 42.56, 4.18, 6.25, 4.72
% treatment SS due to G'E: 36.35, 31.98, 30.19, 34.91, 29.84, 26.14
% G'ESS due to IPCA1: 42.71, 45.20, 43.66, 41.76, 41.35, 36.89
% G'ESS due to IPCA2: 25.17, 35.31, 29.53, 22.93, 31.07, 26.94
% G'ESS due to IPCA3: 15.50, 7.55, 11.51, 10.27, 10.13, 8.32
% G'ESS due to IPCA4: 6.00, 3.84, 5.40, 8.75, 7.04, 8.17
% G'ESS due to IPCA5: 3.64, 2.75, 3.57, 5.24, 4.88, 6.64

Higher mean and % SS due to IPCA1 and IPCA2 were observed at 6 MAP in comparison to at 3 and 9 MAP. This indicated that there were fewer interactions, therefore, more stable symptoms expression at 6 MAP.

ASV ranked the genotypes based on the least scores where low scores represented the most stable genotypes. Low ASV coupled with low disease severity resulted
in the selection of stable genotypes with minimal CMD symptoms. All the genotypes had low CMD severity of $\leq 1.8$ (Table S1). Based on CMD foliar symptoms at 6 MAP, the most stable genotypes with regards to low ASV values and their position relative to the biplot origin (0,0) were Albert, NDL 2003/111, KBH 2002/66 and NDL 2003/31 with a means $\leq 1.27$ (Figure 1, Table S1). The GSI ranking combines both stability and higher scores of a trait. Accordingly, site Chambezi’s environments had moderate stability with the highest mean CMD

Figure 1. AMM2 biplot for CMD foliar symptoms. Environments. Chambezi 2013-2018 (c - c6), Nachingwea 2013-2018 (nc1 - nc6), Mtopwa 2013-2018 (m1 - m6), Naliendele 2013-2018 (nl - nl6), Segera 2013-2018 (s1 - s6). Genotypes: Albert (1), KBH 2002/26 (2), KBH 2002/477 (3), KBH 2002/482 (4), KBH 2002/494 (5), KBH 2002/66 (6), KBH 96/1056 (7), Kiroba (8), Mahiza (9), Mkumba (10), Naliendele 134 (11), NDL 2003/111 (12), NDL 2003/31 (13), NDL 2005/1471 (14), NDL 2005/1472 (15), Pwani (16).
foliar severity of 1.24 (Figure 2, Table S1).

3.2. CBSD Foliar Symptoms

There was a highly significant (P ≤ 0.001) effect of genotype, environment and G’E interaction for CBSD foliar symptoms at 3, 6, and 9 MAP (Table 3). Percent SS due to genotype (60.90% - 69.14%) was higher than due to environment.

Figure 2. AMM1 biplot for CMD foliar symptoms. Environments. Chambezi 2013-2018 (c - c6), Nachingwea 2013-2018 (nc1 - nc6), Mtopwa 2013-2018 (m1 - m6), Naliende 2013-2018 (nl - nl6), Segera 2013-2018 (s1 - s6). Genotypes: Albert (1), KBH 2002/26 (2), KBH 2002/477 (3), KBH 2002/482 (4), KBH 2002/494 (5), KBH 2002/66 (6), KBH 96/1056 (7), Kiroba (8), Mahiza (9), Mkumba (10), Naliende 134 (11), NDL 2003/111 (12), NDL 2003/31 (13), NDL 2005/1471 (14), NDL 2005/1472 (15), Pwani (16).
(4.18% - 6.25%) and $GE$ (29.84% - 34.91%) indicating that most of the variations observed were due to genetic make-up. Four IPCAs were significant ($P \leq 0.05$) for CBSD 3 and 9 MAP while 6 MAP had three significant IPCAs. IPCA1 and IPCA2 accounted for a total SS of 64.69%, 72.42% and 63.83% of the $GE$ variation for CBSD 3, 6 and 9 MAP, respectively. The mean CBSD foliar severity was 1.46, 1.61 and 1.57 at 3, 6, and 9 MAP. Similar to CMD symptoms, a higher mean and % SS due to IPCA1 and IPCA2 were observed at 6 MAP, indicating more stable symptoms expression at this time point.

Based on foliar symptoms at 6 MAP, all the genotypes had low CBSD foliar severity $\leq 1.7$ apart from Naliendele 134, Mkumba, Mahiza and Albert. The most stable genotypes with low ASV included KBH 2002/66, NDL 2005/1471, KBH 96/1056 and NDL 2003/111 with mean of $\leq 1.27$ (Figure 3, Table S2). Although Mahiza too had a low ASV of 0.49, it had a higher severity of 2.27. Similar to CMD symptoms, site Chambezi's environments had the highest mean CBSD foliar severity of 1.77 with moderate GSI ranking indicating moderate stability (Figure 4, Table S2).

### 3.3. CBSD Root Necrosis

The effect of genotype, environment and $GE$ interaction was highly significant ($P \leq 0.001$) for root necrosis (Table 4). Percent SS due to genotype was highest at 52.14% followed by $GE$ (35.19%) and environment (12.66%). The findings indicated that genetic make-up greatly influenced the expression of root symptoms. Five IPCAs had significant ($P \leq 0.05$) mean squares and IPCA1 and IPCA2 accounted for a total SS of 60.51% of the $GE$ variation. The most stable genotypes with low ASV included NDL 2003/11, Pwani, NDL 2005/1471, KBH 2002/482 and NDL 2003/31 (0.42) with a mean of $\leq 1.72$ (Figure 5, Table S3). All the genotypes had low root necrosis severity ($\leq 1.9$) below the grand mean.
Figure 3. AMMI2 biplot for CBSD foliar symptoms. Environments: Chambezi 2013-2018 (c - c6), Nachingwea 2013-2018 (nc1 - nc6), Mtopwa 2013-2018 (m1 - m6), Naliendele 2013-2018 (nl - nl6), Segera 2013-2018 (s1 - s6). Genotypes: Albert (1), KBH 2002/26 (2), KBH 2002/477 (3), KBH 2002/494 (4), KBH 2002/66 (5), KBH 96/1056 (7), Kiroba (8), Mahiza (9), Mkumba (10), Naliendele 134 (11), NDL 2003/111 (12), NDL 2003/31 (13), NDL 2005/1471 (14), NDL 2005/1472 (15), Pwani (16).

Table 4. Combined AMMI ANOVA for root necrosis, root weight and dry matter of 16 cassava genotypes evaluated across 30 environments (6 planting seasons × 5 sites).

| Source      | df | Root necrosis | Root weight | Dry matter |
|-------------|----|---------------|-------------|------------|
| Treatment   | 479| 3.60***       | 1778.05***  | 159.99**   |
| Genotype (G)| 15 | 59.97***      | 10,009.40***| 33.38*     |
| Environment (E)| 29 | 7.53***       | 12,572.60***| 2252.12*** |
| Block       | 60 | 0.95**        | 1220.80***  | 67.46***   |
| Interaction (G*E)| 435| 1.40***       | 774.60***   | 24.88***   |
| IPCA1       | 43 | 4.90***       | 2830.10***  | 94.28***   |
| IPCA2       | 41 | 3.82***       | 2010.01***  | 73.80***   |
| IPCA3       | 39 | 1.41***       | 886.32***   | 34.06***   |
| IPCA4       | 37 | 1.17***       | 722.53***   | 17.55***   |
| IPCA5       | 35 | 1.15*         | 507.32***   | 14.20      |
| Error       | 899| 0.56          | 251.5       | 19.31      |

Sum of squares

| Source      | df | Root necrosis | Root weight | Dry matter |
|-------------|----|---------------|-------------|------------|
| Treatment   | 479| 1725.14       | 851,688.00  | 76,633.00  |
| Genotype (G)| 15 | 899.54        | 150,149.00  | 501.00     |
| Environment (E)| 29 | 218.45        | 364,607.00  | 65,311.00  |
| Block       | 60 | 56.87         | 73,249.00   | 4048.00    |
| Interaction (G*E)| 435| 607.15        | 336,932.00  | 10,821.00  |
| IPCA1       | 43 | 210.79        | 121,694.39  | 4054.13    |
| IPCA2       | 41 | 156.57        | 82,410.25   | 3025.58    |
| IPCA3       | 39 | 54.81         | 34,566.54   | 1328.29    |
| IPCA4       | 37 | 43.13         | 26,733.41   | 649.33     |
| IPCA5       | 35 | 40.39         | 17,756.27   | 497.10     |
Continued

|                | 899  | 502.29 | 226,307.0 | 17,380.00 |
|----------------|------|--------|-----------|-----------|
| % treatment SS due to $G$ | 52.14 | 17.62  | 0.65      |
| % treatment SS due to $E$   | 12.66 | 42.81  | 85.23     |
| % treatment SS due to $GE$  | 35.19 | 39.56  | 14.12     |
| $G^2SS$ due to IPCA1       | 34.72 | 36.12  | 37.47     |
| $G^2SS$ due to IPCA2       | 25.79 | 24.46  | 27.96     |
| $G^2SS$ due to IPCA3       | 9.03  | 10.26  | 12.28     |
| $G^2SS$ due to IPCA4       | 7.10  | 7.93   | 6.00      |
| $G^2SS$ due to IPCA5       | 6.65  | 5.27   | 4.59      |

**Figure 4.** AMM2 biplot for CBSD foliar symptoms. Environments: Chambezi 2013-2018 (c - c6), Nachingwea 2013-2018 (nc1 - nc6), Mtopwa 2013-2018 (m1 - m6), Naliendele 2013-2018 (nl - nl6), Segera 2013-2018 (s1 - s6). Genotypes: Albert (1), KBH 2002/26 (2), KBH 2002/477 (3), KBH 2002/482 (4), KBH 2002/494 (5), KBH 2002/66 (6), KBH 96/1056 (7), Kiroba (8), Mahiza (9), Mkumba (10), Naliendele 134 (11), NDL 2003/111 (12), NDL 2003/31 (13), NDL 2005/1471 (14), NDL 2005/1472 (15), Pwani (16).
of 2.29 apart from Mkumba (2.29), Naliendele (3.52), Mahiza (3.08), KBH 2002/477 (2.22) and KBH 2002/66 (2.04). The genotypes with the highest root necrosis severity also had high ASV, therefore, unstable. Among the environments, Chambezi 2013 to 2018 has the highest mean root necrosis severity of 2.33 with moderate GSI ranking indicating moderate stability (Figure 6, Table S3).

3.4. Root Weight

The effect of genotype, environment and $G\times E$ interaction was highly significant

![Figure 5](image-url)
Figure 6. AMM1 biplot for root necrosis, root weight and dry matter content. Environments. Chambezi 2013-2018 (c - c6), Nachingwea 2013-2018 (nc1 - nc6), Mtopwa 2013-2018 (m1 - m6), Naliendele 2013-2018 (nl - nl6), Segera 2013-2018 (s1 - s6). Genotypes: Albert (1), KBH 2002/26 (2), KBH 2002/477 (3), KBH 2002/482 (4), KBH 2002/494 (5), KBH 2002/66 (6), KBH 96/1056 (7), Kiroba (8), Mahiza (9), Mkumba (10), Naliendele 134 (11), NDL 2003/111 (12), NDL 2003/31 (13), NDL 2005/1471 (14), NDL 2005/1472 (15), Pwani (16).

(P ≤ 0.001) for root weight (Table 4). Percent SS due to environment was highest at 42.81% followed closely by $G^E$ (35.19%) and genotype (17.62%). The findings indicated that both the environment and $G^E$ greatly influenced the expression of root weight. Five IPCAs had significant (P ≤ 0.05) mean squares and IPCA1 and IPCA2 accounted for a total SS of 60.58% of the $G^E$ variation. The most stable genotypes with low ASV included KBH 2002/477, NDL 2003/31, Mahiza and NDL 2005/477 (Figure 5, Table S3). Since ASV measure alone is
not sufficient for the selection of superior genotypes, GSI ranking was used as it combines both genotype stability and high yield. Accordingly, the most stable and high yielding genotypes included NDL 2003/31 (51.55 t/ha), NDL 2003/111 (51.86 t/ha), KBH 2002/477 (42.86 t/ha) and NDL 2005/1472 (35.59 t/ha) (Table S3). Chambezi 2013 to 2018 environments had the highest mean root weight (45.21 t/ha) with moderate to high stability based on GSI ranking (Figure 6, Table S3). Higher yields were observed in favourable environments for example Chambezi which had higher rainfall than other sites (Figure 7, Table 1, Table S3). Site Segera having received the least rainfall was one of the sites with a lower combined mean root weight of 33.47 t/ha.

3.5. Dry Matter Content

The effect of genotype, environment and GE interaction was significant (P ≤
0.001) for dry matter content (Table 4). Percent SS due to environment was very high at 85.23% followed by $G\times E$ (14.12%) and very low SS due to genotype (0.68%).

Four IPCAs had significant ($P \leq 0.05$) mean squares and IPCA1 and IPCA2 accounted for a total SS of 65.43% of the $G\times E$ variation. The mean dry matter contents for genotypes were close ranging from 26.95% - 28.77% (Figure 6, Table S3). The genotypes with high stability and dry matter content included KBH 2002/494 (28.77%), KBH 2002/477 (28.69%), KBH 2002/66 (28.55%) Kiroba (28.37%) and KBH 2002/482 (28.18%) (Figure 5, Table S3). Segera which received the least rainfall had the highest combined mean dry matter content of 30.43%. Root necrosis may have affected dry matter content since environments with higher root necrosis had corresponding low dry matter content and vice versa. For example, among the environments in Chambezi, Chambezi 2016 had the highest root necrosis severity (3.06) and the lowest dry matter content (16.96%). Similar observations were made in other sites.

4. Discussion

The performance of cassava is subject to the strong influence of genotype, environment and $G\times E$ interactions [15] [29] [47]. TARI-Naliendele has been developing improved genotypes, however, only a few varieties have been released. The newly developed breeding lines are in their final stages of breeding. Therefore, evaluating them in diverse environments and providing recommendations for suitable ones will contribute to increasing cassava production and improved food and nutrition security.

The AMMI model was used in this study and the effects of genotype, environment, and $G\times E$ interactions were significant. Percent SS due to environment was the highest followed by $G\times E$ interaction in CMD foliar symptoms severity, root weight and dry matter content showing that environment and $G\times E$ interaction greatly influenced the variations observed. On the other hand, % SS due to genotype was the highest followed by $G\times E$ interaction in CBSD foliar symptoms and root necrosis. A considerable percentage of $G\times E$ interaction was explained by IPCA1 (34.725% - 45.20%), followed by IPCA2 (22.93% - 35.31%) and lastly IPCA3 (7.55% - 15.50%). Several studies have shown similar findings where a significant and greater percentage of $G\times E$ interaction was explained by IPCA1 and IPCA2 [15] [29] [47] [48].

Mean CMD and CBSD foliar symptoms severity increased from 3 to 6 MAP then dropped at 9 MAP. The total % SS due to both IPCA1 and IPCA2 was the highest at 6 MAP for both CMD and CBSD. A possible explanation for this is that at 3 MAP, some plants may still have low viral titre [49] and may not express symptoms thus causing significant variations in the replications and environments. This may result in the representation of substantial % SS by other IPCAs apart from IPCA 1 and 2. CBSD foliar symptoms are more difficult to recognize in older plants as the lower leaves with prominent symptoms senesce and fall off, causing variation in symptoms expression among the plants partic-
ularly at 9 MAP [50]. Additionally, younger leaves are more susceptible to CMD resulting in a decrease in CMD symptoms in some plants with increasing plant age [51]. In our earlier study we reported a higher heritability at 6 MAP for CMD and CBSD foliar symptoms thus emphasising the importance of assessment at this time point [20].

Stability analysis methods are often used by breeders to identify genotypes that have stable performance and respond positively to improvements in environmental conditions [39] [40]. With regards to CMD and CBSD, suitable genotypes would have low ASV and low disease severity. Further, genotypes with CMD foliar severity scores (<2.0) are classified as resistant while those with (≥2.0) as susceptible [52]. In this study, all genotypes had low foliar severity (>1.35) apart from Mahiza which was slightly higher at 1.77. The most stable genotypes with low CMD foliar severity (≤1.28) were Albert, NDL 2003/111, KBH 2002/66, KBH 2002/26, NDL 2005/1472, NDL 2003/31 and KBH 96/1056.

CBSD-resistant varieties exhibit minimal symptoms with a severity of (<2.0) both on leaves and roots, tolerant once have more severe symptoms on leaves (≥2.0) coupled with minimal symptoms on roots (<2.0) while susceptible ones develop severe symptoms on both leaves and roots (>2.0) [16] [53] [54]. All genotypes apart from Naliendele 134, Mkumba, Mahiza, Albert, KBH 2002/26 and KBH 2002/66 had minimal symptoms (<2.0) both on leaves and roots. Stable genotypes with low ASV on CBSD foliar severity did not necessarily have stable root necrosis severity. This could be due to the different QTLs affecting CBSD foliar symptoms and root necrosis leading to varied expression of symptoms on leaves and roots [53]. Stable resistant genotypes with minimal symptoms on both leaves and roots (<2.0) were NDL 2005/1471 NDL 2003/111, KBH 2002/482 and NDL 2003/31. The environments in site Chambezi had the highest combined means for CMD (1.24), CBSD (1.77), and root necrosis (2.06). Masmuma et al. [53] reported higher CMD and CBSD severity suggesting the suitability of this site for disease resistance evaluation. The environments, however, had higher GSI ranking portraying moderate stability. Virus transmission and disease spread are determined by inoculum pressure and their variation from season to season may have contributed to the lower stability observed [55].

Based on GSI ranking, the most stable high yielding genotypes included NDL 2003/31 (51.55 t/ha), NDL 2003/111 (51.86 t/ha), KBH 2002/477 (42.86 t/ha) and NDL 2005/1472 (35.59 t/ha). Higher yields were recorded in favourable environments indicating that genotypes can exploit their full potential to yield well under good environmental conditions. Accordingly, the environments in site Chambezi had the highest combined root weight means of 45.21 t/ha. Chambezi had higher rainfall particularly during the first six months, a critical period for root initiation and development [56] [57]. Most of the genotypes had moderately high dry matter content ranging from 26.95% - 28.77%. Low rainfall results in high dry matter content as was observed in the environments in site Segera which had the highest combined dry matter content mean of 30.43 [16] [58].
Additionally, the environments with the least CBSD root necrosis symptoms had the highest dry matter content and vice versa, indicating that presence of root symptoms can affect key agronomic traits leading to loss of farmer preferred traits [16] [58].

5. Conclusion

$G^*E$ was significant ($P \leq 0.05$) for CMD foliar symptoms, CBSD foliar and root symptoms, root weight and dry matter content. This emphasised the importance of testing genotypes in multiple environments before an effective selection is made. Besides, variations were also significant among the test environments. Site Chambezi had the highest mean CMD and CBSD severity; therefore, it has been empirically confirmed as the most suitable environment for evaluation for disease resistance. Cassava produces high yield under favourable conditions such as adequate rainfall and soil fertility as was observed in Chambezi. The most stable genotypes which combined CMD and CBSD resistance, high yield were NDL 2003/31 and NDL 2003/111. These genotypes outperformed the checks Albert, Kiroba, Pwani, Mahiza, Mkumba and Naliendele 134 indicating that they have the potential to increase cassava productivity and should therefore be recommended for release to cassava farmers or further breeding prospects.

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Conflicts of Interest

The authors declare no conflicts of interest regarding the publication of this paper.

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### Appendix

#### Table S1. Mean, AMMI stability value (ASV), Genotype stability index (GSI) and rank (r) of CMD foliar symptoms for genotypes and sites.

| Genotype       | CMD 3 MAP |       |       |       | CMD 6 MAP |       |       |       | CMD 9 MAP |       |       |       |
|----------------|-----------|-------|-------|-------|-----------|-------|-------|-------|-----------|-------|-------|-------|
|                | Mean ASV  | GSI rASV | rGSI | Mean ASV  | GSI rASV | rGSI | Mean ASV  | GSI rASV | rGSI | Mean ASV  | GSI rASV | rGSI |
| Albert         | 1.16      | 0.35 | 12.0 | 4 | 3 | 1.28 | 0.07 | 6.0 | 1 | 1 | 1.20 | 0.06 | 10.0 | 1 | 3 |
| KBH 2002/26    | 1.02      | 0.72 | 28.0 | 12 | 15 | 1.02 | 0.61 | 28.0 | 6 | 15 | 1.03 | 0.69 | 28.0 | 12 | 15 |
| KBH 2002/477   | 1.16      | 0.47 | 15.0 | 6 | 8 | 1.22 | 0.43 | 16.0 | 7 | 7 | 1.20 | 0.67 | 19.0 | 11 | 11 |
| KBH 2002/482   | 1.13      | 0.30 | 14.0 | 2 | 7 | 1.34 | 0.60 | 14.0 | 11 | 5 | 1.25 | 0.09 | 7.0 | 2 | 1 |
| KBH 2002/494   | 1.20      | 0.32 | 9.0 | 3 | 2 | 1.25 | 0.57 | 16.0 | 9 | 7 | 1.24 | 0.48 | 14.0 | 8 | 6 |
| KBH 2002/66    | 1.18      | 0.82 | 20.0 | 13 | 11 | 1.14 | 0.23 | 15.0 | 3 | 6 | 1.16 | 0.33 | 17.0 | 6 | 7 |
| KBH 96/1056    | 1.13      | 0.30 | 12.0 | 1 | 3 | 1.15 | 0.41 | 16.0 | 5 | 7 | 1.14 | 0.31 | 17.0 | 5 | 7 |
| Kiroba         | 1.20      | 1.34 | -     | - | - | 1.15 | 0.41 | -     | - | - | 1.14 | 0.31 | -     | - | - |
| Mahiza         | 1.51      | 0.90 | 15.0 | 14 | 8 | 1.77 | 1.79 | 17.0 | 16 | 11 | 1.68 | 1.24 | 17.0 | 16 | 7 |
| Mkumba         | 1.05      | 1.09 | 28.0 | 15 | 15 | 1.03 | 0.78 | 30.0 | 15 | 16 | 1.05 | 0.96 | 29.0 | 14 | 16 |
| Naliendele 134 | 1.33      | 0.72 | 13.0 | 11 | 5 | 1.41 | 0.58 | 12.0 | 10 | 3 | 1.39 | 0.56 | 11.0 | 9 | 4 |
| NDL 2003/111   | 1.20      | 0.66 | 13.0 | 9 | 5 | 1.19 | 0.20 | 12.0 | 2 | 3 | 1.20 | 0.29 | 13.0 | 3 | 5 |
| NDL 2003/31    | 1.24      | 0.39 | 8.0 | 5 | 1 | 1.27 | 0.30 | 10.0 | 4 | 2 | 1.25 | 0.30 | 8.0 | 4 | 2 |
| NDL 2005/1471  | 1.15      | 0.57 | 17.0 | 7 | 10 | 1.29 | 0.77 | 18.0 | 14 | 12 | 1.26 | 1.17 | 18.0 | 15 | 10 |
| NDL 2005/1472  | 1.03      | 0.70 | 25.0 | 10 | 14 | 1.08 | 0.48 | 21.0 | 7 | 13 | 1.06 | 0.57 | 24.0 | 10 | 14 |
| Pwani          | 1.03      | 0.59 | 22.0 | 8 | 13 | 1.22 | 0.42 | 16.0 | 12 | 7 | 1.06 | 0.46 | 20.0 | 7 | 12 |
| **Grand mean** | 1.17      |       |       |       | 1.24 |       |       |       |       | 1.21 |       |       |
| **CV**         | 16.1      |       |       |       | 15.2 |       |       |       |       | 17.1 |       |       |

#### Environments

| Chambezi mean | 1.45 | 1.62 | 1.54 |
|---------------|------|------|------|
| Mtopwa mean   | 1.04 | 1.07 | 1.09 |

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| Genotype     | CBSD 3 MAP |   |   |   | CBSD 6 MAP |   |   |   | CBSD 9 MAP |   |   |   |
|--------------|------------|---|---|---|------------|---|---|---|------------|---|---|---|
|              | Mean | ASV | GSI | rASV | rGSI | Mean | ASV | GSI | rASV | rGSI | Mean | ASV | GSI | rASV | rGSI |
| Albert       | 2.23 | 3.04 | 18.0 | 16 | 8 | 2.66 | 1.33 | 16.0 | 15 | 4 | 2.54 | 1.22 | 16.0 | 15 | 4 |
| KBH 2002/26  | 1.13 | 0.15 | 14.0 | 2 | 3 | 1.42 | 0.82 | 19.0 | 10 | 10 | 1.28 | 0.44 | 15.0 | 5 | 3 |
| KBH 2002/477 | 1.57 | 1.52 | 18.0 | 13 | 8 | 1.73 | 1.27 | 20.0 | 14 | 13 | 1.69 | 1.15 | 19.0 | 13 | 13 |
| KBH 2002/482 | 1.25 | 0.38 | 14.0 | 5 | 3 | 1.34 | 0.80 | 18.0 | 8 | 8 | 1.33 | 0.64 | 17.0 | 8 | 5 |
| KBH 2002/494 | 1.51 | 1.77 | 20.0 | 14 | 11 | 1.69 | 0.56 | 11.0 | 4 | 4 | 1.64 | 0.93 | 18.0 | 11 | 10 |
| KBH 2002/66  | 1.26 | 0.64 | 16.0 | 8 | 7 | 1.27 | 0.35 | 12.0 | 1 | 3 | 1.26 | 0.37 | 14.0 | 3 | 2 |
| KBH 96/1056  | 1.07 | 0.58 | 21.0 | 7 | 13 | 1.02 | 0.56 | 21.0 | 5 | 14 | 1.06 | 0.34 | 17.0 | 1 | 5 |
| Kiroba       | 1.15 | 0.20 | 14.0 | 3 | 3 | 1.52 | 0.80 | 17.0 | 9 | 7 | 1.39 | 0.60 | 15.0 | 7 | 3 |
| Mahiza       | 2.13 | 2.01 | 18.0 | 15 | 8 | 2.49 | 0.49 | 5.0 | 3 | 1 | 2.35 | 1.02 | 15.0 | 12 | 3 |
| Mkumba       | 2.31 | 0.98 | 13.0 | 12 | 2 | 2.27 | 1.72 | 19.0 | 16 | 10 | 2.36 | 1.37 | 18.0 | 16 | 10 |
| Naliendele 134 | 1.48 | 0.04 | 8.0 | 1 | 1 | 2.09 | 0.92 | 16.0 | 12 | 4 | 1.83 | 1.17 | 19.0 | 14 | 13 |
| NDL 2003/111 | 1.06 | 0.43 | 21.0 | 6 | 13 | 1.08 | 0.57 | 19.0 | 6 | 10 | 1.08 | 0.37 | 18.0 | 4 | 10 |
| NDL 2003/31  | 1.06 | 0.28 | 20.0 | 4 | 11 | 1.03 | 0.58 | 22.0 | 7 | 15 | 1.06 | 0.35 | 17.0 | 2 | 5 |
| NDL 2005/1471 | 1.12 | 0.66 | 22.0 | 9 | 16 | 1.05 | 0.49 | 16.0 | 2 | 4 | 1.11 | 0.68 | 22.0 | 9 | 15 |

Table S2. Mean, AMMI stability value (ASV), Genotype stability index (GSI) and rank (r) of CBSD foliar symptoms for genotypes and sites

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| Environment | Year | Mean | CV  |
|-------------|------|------|-----|
| NDL 2005/1472 | 1.20 | 0.88 | 21.0 |
| Pwani | 1.78 | 0.84 | 14.0 |

**Grand mean**

| Environment | Mean | CV  |
|-------------|------|-----|
| Chambezi | 1.46 | 1.61 |
| CV | 25.1 | 20.8 |

**Environments**

| Environment | Year | Mean | CV  |
|-------------|------|------|-----|
| Chambezi 2013 | 1.40 | 0.35 | 29.0 |
| Chambezi 2014 | 1.40 | 0.31 | 21.0 |
| Chambezi 2015 | 1.41 | 0.34 | 22.0 |
| Chambezi 2016 | 1.41 | 0.35 | 24.0 |
| Chambezi 2017 | 1.28 | 0.86 | 47.0 |
| Chambezi 2018 | 1.40 | 0.34 | 26.5 |

**Chambezi mean**

| Environment | Year | Mean | CV  |
|-------------|------|------|-----|
| Mtopwa 2013 | 1.48 | 1.00 | 32.0 |
| Mtopwa 2014 | 1.42 | 0.88 | 34.5 |
| Mtopwa 2015 | 1.42 | 0.88 | 33.5 |
| Mtopwa 2016 | 1.49 | 1.11 | 30.0 |
| Mtopwa 2017 | 1.50 | 1.14 | 32.0 |
| Mtopwa 2018 | 1.67 | 1.26 | 32.0 |

**Mtopwa mean**

| Environment | Year | Mean | CV  |
|-------------|------|------|-----|
| Nachingwea 2013 | 1.57 | 0.77 | 19.0 |
| Nachingwea 2014 | 1.63 | 0.48 | 14.0 |
| Nachingwea 2015 | 1.58 | 0.74 | 17.0 |
| Nachingwea 2016 | 1.56 | 0.27 | 8.0 |
| Nachingwea 2017 | 1.51 | 0.81 | 23.0 |
| Nachingwea 2018 | 1.67 | 0.33 | 5.0 |

**Nachingwea mean**

| Environment | Year | Mean | CV  |
|-------------|------|------|-----|
| Naliendele 2013 | 1.47 | 0.66 | 24.0 |
| Naliendele 2014 | 1.40 | 0.36 | 30.5 |
| Naliendele 2015 | 1.40 | 0.35 | 28.0 |
| Naliendele 2016 | 1.49 | 0.78 | 25.0 |
| Naliendele 2017 | 1.43 | 0.84 | 30.0 |
| Naliendele 2018 | 1.70 | 1.61 | 31.0 |

**Naliendele mean**

| Environment | Year | Mean | CV  |
|-------------|------|------|-----|
| Segera 2013 | 1.30 | 1.17 | 54.4 |
| Segera 2014 | 1.38 | 1.15 | 48.5 |
| Segera 2015 | 1.04 | 1.14 | 52.5 |
| Segera 2016 | 1.64 | 1.20 | 54.5 |
| Segera 2017 | 1.36 | 1.17 | 54.5 |
| Segera 2018 | 1.32 | 1.15 | 47.5 |

**Segera mean**

| Environment | Year | Mean | CV  |
|-------------|------|------|-----|
| DOI: 10.4236/ajps.2021.124046 | 703 | 1.34 | 1.68 |

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### Table S3. Mean, AMMI stability value (ASV), Genotype stability index (GSI) and rank (r) of root necrosis, root weight and dry matter content for genotypes and sites.

| Genotype       | Root necrosis | Root weight | Dry matter |
|----------------|---------------|-------------|------------|
|                | Mean | ASV | GSI | rGSI | Mean | ASV | GSI | rGSI | Mean | ASV | GSI | rASV | rGSI |
| Albert         | 4.11 | 1.34 | 13.0 | 12   | 17.36 | 4.92 | 27.0 | 11   | 27.18 | 2.65 | 25   | 10   | 14   |
| KBH 2002/26    | 2.22 | 1.36 | 18.0 | 13   | 36.56 | 1.66 | 10.0 | 1    | 27.88 | 1.67 | 19   | 8    | 11   |
| KBH 2002/477   | 1.58 | 0.56 | 17.0 | 7    | 42.86 | 3.60 | 12.0 | 6    | 28.69 | 0.46 | 5    | 2    | 2    |
| KBH 2002/482   | 1.72 | 0.41 | 12.0 | 4    | 40.20 | 4.74 | 18.0 | 10   | 28.18 | 1.14 | 16   | 6    | 5    |
| KBH 2002/494   | 1.58 | 1.23 | 22.0 | 11   | 43.23 | 5.54 | 16.0 | 12   | 28.77 | 0.13 | 2    | 1    | 1    |
| KBH 2002/66    | 2.04 | 2.23 | 21.0 | 15   | 30.71 | 4.62 | 20.0 | 9    | 28.55 | 0.46 | 8    | 3    | 3    |
| KBH 96/1056    | 1.46 | 0.75 | 24.0 | 9    | 44.18 | 17.59 | 19.0 | 16   | 28.40 | 3.14 | 17   | 11   | 6    |
| Kiroba         | 1.53 | 0.50 | 19.0 | 6    | 40.83 | 4.10 | 15.0 | 8    | 28.37 | 1.39 | 14   | 7    | 4    |
| Mahiza         | 3.08 | 1.11 | 13.0 | 10   | 20.80 | 2.86 | 18.0 | 3    | 28.62 | 3.53 | 17   | 13   | 6    |
| Mkumba         | 2.29 | 1.54 | 18.0 | 14   | 22.60 | 7.54 | 28.0 | 14   | 27.35 | 3.64 | 27   | 14   | 15   |
| Nalindele 134  | 3.52 | 2.57 | 18.0 | 16   | 27.50 | 8.10 | 28.0 | 15   | 28.21 | 1.92 | 18   | 9    | 9    |
| NDL 2003/111   | 1.50 | 0.16 | 15.0 | 1    | 51.86 | 3.15 | 6.0  | 5    | 28.30 | 3.43 | 20   | 12   | 12   |
| NDL 2003/31    | 1.36 | 0.42 | 21.0 | 5    | 51.55 | 1.67 | 4.0  | 2    | 26.95 | 1.02 | 21   | 5    | 13   |
| NDL 2005/1471  | 1.53 | 0.36 | 15.0 | 3    | 42.89 | 5.89 | 18.0 | 13   | 27.27 | 0.84 | 18   | 4    | 9    |
| NDL 2005/1472  | 1.92 | 0.63 | 15.0 | 8    | 35.59 | 3.07 | 14.0 | 4    | 27.63 | 4.19 | 28   | 16   | 16   |
| Pwani          | 1.59 | 0.33 | 11.0 | 2    | 29.87 | 3.66 | 19.0 | 7    | 28.71 | 3.99 | 17   | 15   | 6    |
| Grand mean     | 2.06 |       |      |      |       |      |      |      | 36.1  |       |      |      |      |
| CV             | 36.2 |       |      |      |       |      |      |      | 43.8  |       |      |      |      |

### Environments

| Environments | Mean | ASV | GSI | rASV | Mean | ASV | GSI | rASV | Mean | ASV | GSI | rASV | rGSI |
|--------------|------|-----|-----|------|------|-----|-----|------|------|-----|-----|------|------|
| Chambezi 2013 | 2.31 | 0.92 | 28.0 | 19   | 45.45 | 2.01 | 19.0 | 10   | 24.29 | 0.55 | 34.0 | 13   | 20   |
| Chambezi 2014 | 1.56 | 0.17 | 30.0 | 2    | 25.57 | 1.38 | 26.0 | 4    | 27.62 | 0.92 | 35.0 | 18   | 22   |
| Chambezi 2015 | 2.42 | 1.20 | 32.0 | 25   | 43.79 | 4.50 | 34.0 | 24   | 23.29 | 1.35 | 41.0 | 19   | 26   |
| Chambezi 2016 | 3.06 | 1.51 | 31.0 | 30   | 60.27 | 7.06 | 29.0 | 26   | 16.96 | 0.31 | 38.0 | 8    | 24   |
| Chambezi 2017 | 2.58 | 0.92 | 20.0 | 17   | 42.10 | 2.83 | 26.0 | 15   | 28.71 | 3.99 | 17   | 15   | 6    |
| Chambezi 2018 | 2.10 | 1.06 | 33.0 | 22   | 54.05 | 7.24 | 33.0 | 27   | 36.35 | 0.82 | 20.0 | 17   | 3    |
| Chambezi mean | 2.33 |       |      |      |      |      |      |      | 45.21 |       |      |      |      |
| Mtopwa 2013   | 1.79 | 0.12 | 25.0 | 1    | 29.70 | 1.72 | 23.0 | 7    | 27.38 | 1.66 | 42.0 | 24   | 28   |
| Mtopwa 2014   | 1.68 | 0.42 | 34.0 | 8    | 24.47 | 2.08 | 36.0 | 12   | 29.15 | 1.94 | 41.0 | 26   | 26   |
| Mtopwa 2015   | 1.92 | 0.96 | 38.0 | 20   | 36.02 | 1.59 | 21.0 | 6    | 18.34 | 0.10 | 29.0 | 2    | 11   |
| Mtopwa 2016   | 2.54 | 0.51 | 16.0 | 11   | 67.39 | 3.98 | 24.0 | 22   | 17.45 | 0.18 | 32.0 | 3    | 16   |
| Mtopwa 2017   | 1.90 | 0.65 | 33.0 | 14   | 14.03 | 1.40 | 32.0 | 5    | 32.92 | 1.65 | 31.0 | 23   | 13   |
| Mtopwa 2018   | 1.21 | 1.30 | 57.0 | 27   | 25.06 | 2.06 | 34.0 | 11   | 36.12 | 0.18 | 8.0  | 4    | 1    |
| Mtopwa mean   | 1.84 |       |      |      |      |      |      |      | 32.79 |       |      |      |      |
| Nachingwea 2013 | 2.25 | 0.54 | 22.0 | 12   | 20.32 | 3.41 | 46.0 | 21   | 26.89 | 0.19 | 25.0 | 6    | 6    |
| Nachingwea 2014 | 1.96 | 0.18 | 20.0 | 3    | 45.65 | 5.77 | 33.0 | 25   | 39.81 | 6.93 | 31.0 | 30   | 13   |
Continued

| Year       | Nachingwea | Naliendele | Segera  |
|------------|------------|------------|---------|
| 2015       | 2.06       | 2.06       | 1.87    |
| 2016       | 2.50       | 2.56       | 1.90    |
| 2017       | 2.08       | 1.85       | 1.87    |
| 2018       | 1.49       | 2.00       | 1.75    |
| Mean       | 2.06       | 1.98       | 2.10    |

| Year       | Nachingwea | Naliendele | Segera  |
|------------|------------|------------|---------|
| 2015       | 19.5       | 39.5       | 31.5    |
| 2016       | 6          | 26         | 5       |
| 2017       | 3          | 18         | 9       |
| 2018       | 29         | 24         | 7       |
| Mean       | 39.17      | 29.95      | 33.47   |

| Year       | Nachingwea | Naliendele | Segera  |
|------------|------------|------------|---------|
| 2015       | 68.36      | 27.60      | 13.56   |
| 2016       | 12.72      | 18.97      | 39.83   |
| 2017       | 31.0       | 34.0       | 3.04    |
| 2018       | 30         | 18.0       | 3.14    |
| Mean       | 39.17      | 29.95      | 33.47   |

| Year       | Nachingwea | Naliendele | Segera  |
|------------|------------|------------|---------|
| 2015       | 0.05       | 1.75       | 3.14    |
| 2016       | 18.42      | 3.40       | 46.5    |
| 2017       | 14.0       | 18.0       | 43.0    |
| 2018       | 0.05       | 18.0       | 9.0     |
| Mean       | 27.88      | 27.90      | 30.43   |

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