ADDIS ABABA UNIVERSITY

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GENETIC DIVERSITY STUDY IN NAPIER GRASS (Cenchrus purpureus) COLLECTIONS AND PROGENIES

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List of Abbreviations

AFLP	Amplified Fragment Length Polymorphism	
DArT	Diversity Array Technology	
EMBRAPA	BRAPA Brazilian Agricultural Research Corporation	
EST	Expressed Sequence Tag	
ICRISAT	International Crop Research Institute for the Semi_Arid Tropics	
GenAlEx	Genetic Analysis in Excel	
NGS	Next Generation Sequencing	
NTSYSpc	(SYSpc) Numerical Taxonomy System for personal computer	
PCA	CA Principal Component Analysis	
RAPD	Random Amplified Polymorphic DNA	
RFLP	Restriction Fragment Length Polymorphism	
SNP	Single Nucleotide Polymorphism	
SSR	Simple Sequence Repeat	
UPGMA	Unweighted Paired Group Method Using Arithmetic Mean	
UPGMC	Unweighted Paired Group Method Using Centroid	
II DI		

ILRI International Livestock Research Institute

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Title: Genetic diversity study in Napier grass (Cenchrus purpureus) collections and progenies

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Abstract

Application of currently available advanced molecular tools and characterization of Napier grass forage crop is limited. Furthermore, ILRI genebank held collection of this forage grass from different part of the world, but with little information on the genetic diversity of the collection. Therefore, the objective of this study was to evaluate the extent of the genetic diversity of Napier grass accessions collected from different parts of the world and progeny plants raised from naturally produced seeds. The genomic DNA was extracted from leaves of 347 Napier grass genotypes and genotyped by the DArTseq platform. A total of 96,454 Silico DArT and 96,321 SNP markers were generated, of which highly informative 1001 SNP markers were selected for diversity analysis after screening. The average polymorphic information content (PIC) values of Silico DArT and SNP markers were 0.21 and 0.15 and average heterozygosity of 0.26 and 0.18 respectively. Two major groups and ten sub-clusters were identified by population stratification and diversity analysis using STRUCTURE and hierarchal clustering. Discriminant analysis of principal component (DAPC) further confirmed the sub-clusters. Analysis of molecular variance (AMOVA) showed significant (P < 0.00) variation among the populations. The mean values of fixation index (Fst) per cluster ranged from 0.34 in cluster VI, that consisted of progeny plants, to 0.76 in cluster VII, that consisted mostly of the ILRI collections, and the largest divergence (0.38) was also between sub-cluster IV and VII. All these parameters showed the presence of high diversity and genetic differentiation among the assayed Napier grass genotypes.

Key words: Cenchrus purpureus, DArTseq markers, Genetic diversity, Napier grass, progeny plants

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1 Introduction

Napier grass (Cenchrus purpureus (Schumach) Morrone syn. Pennisetum purpureum Schumach.) or elephant grass is a monocot perennial grass that belongs to the family Poaceae (grass family) and genus *Cenchrus*. The genus *Cenchrus* is very diverse, consisting of a heterogeneous group of species with different basic chromosome number and ploidy level. Among these species Napier grass is an allotetraploid species with a basic chromosome number of 7 (2n = 4x = 28) (Singh and Obeng, 2013). Napier grass is widely cultivated as a forage crop in the tropical and subtropical regions of the world. It is mainly used as animal feed due to its ease of establishment and management, high palatability and high dry matter production that can provide up to 78 tons of dry matter/ ha/ year (Negawo et al., 2017; Oliveira et al., 2017; Maria et al., 2014). In addition, Napier grass has the potential to produce biofuels such as alcohol, ethanol and butanol, and methane since it has high cellulose content that can be used as carbon (energy) source (Romero et al., 2019; Roslan et al., 2020; Yasuda et al., 2013). It is also, considered as one of the exceptional potential phytoremediator plants to tackle heavy metal and chemical environmental pollutions due to its rapid growth rate, ease of establishment, low cost of management system and high level of biomass that can withstand and enable it to accumulate large amounts of pollutants (contaminants) in their system; ability to grow on different environmental situations even on poor soil conditions (Lotfy and Mostafa, 2014; Roslan et al., 2020; Tananonchai and Sampanpanish, 2018). According to Yang et al. (2020) through the use of liquid extraction method that directly remove heavy metals from plants; the content of heavy metal in Napier grass was lowered (detoxified) while the crude protein was retained to be used safely as animal feed or energy production materials. Moreover, some Napier grass cultivars were identified as potential plants in the "push-pull insect management strategy" and used to trap African stem borer, Busseola fusca Fuller (Lepidoptera: Noctuidae), an insect that causes production loss in maize and sorghum (Khan et al., 2007).

The international livestock research institute (ILRI) forage genebank holds more than 130 Napier grass accessions collected from different parts of the world. These collections consist from different sources where 60 accessions are ILRI's 'in trust' collection which represents a diverse set of genotypes assembled from a range of environments and origins, of which eight accessions are hybrids of *C. purpureus* \times *C. americanus* crosses, and the remaining 62 accessions were introduced to ILRI from different institutions, including the Brazilian Agricultural Research Corporation (EMBRAPA), the Crop Genetics and Breeding Research Unit, USDA–ARS, Tifton, Georgia and the International Crop

Research Institute for the Semi-Arid Tropics (ICRISAT), India. These collections are maintained insitu at the Bishoftu and Ziway (Batu) sites in Ethiopia (Negawo et al., 2017). The collection is characterized by a high amount of genetic diversity and is highly variable in terms of agronomic and morphological characters (Muktar et al., 2019). Though the collections represent diverse sets of genotypes that are variable in genetic and phenotypic traits, still the diversity and population size is very limiting in selecting different traits for East Africa's different agroecological zones. Furthermore, the study by Muktar et al. (2019) indicated the presence of long haplotype blocks and less linkage disequilibrium (LD) decay in the ILRI in trust collections which were maintained via vegetative propagation than the EMBRAPA materials, that had passed through an active breeding program, suggesting breeding the ILRI collection and developing progeny population would increase the available genetic variability. Thus, developing a breeding population would be utilized for molecular genetic studies such as quantitative trait loci (QTL) mapping and genetic diversity analysis. Introducing new materials from international institutes would be good and the easiest way to increase the diversity, even though the use of the introduced materials may be restricted by research compliances and copy rights. The other way to increase the diversity is by crossing the most genetically distant genotypes and analysing the progenies to identify unique genotypes.

The application of currently available advanced molecular tools and characterization of this forage crop is limited. Utilization of molecular technologies is essential to assess and identify the variability found in this forage crop accurately in a short time and can support improvement and conservation efforts. But so far, a handful of publications that are based on the use of low density molecular markers mostly for diversity analysis (Azevedo *et al.*, 2012; Paudel *et al.*, 2018; Wanjala *et al.*, 2013) and a couple of publications that are based on the advanced sequencing technologies (Muktar *et al.*, 2019; Zhou *et al.*, 2018), have been produced on Napier grass genotyping. Zhou *et al.* (2018) used Illumina-based sequencing technology to develop about 50 EST-SSR markers and 6 SNP markers through transcriptome analysis that were used to facilitate the genetic diversity study in Napier grass. A recent report (Muktar *et al.*, 2019) used the GBS method of the DArTseq platform for the development of genome-wide sequence-based molecular markers (dominant (SilicoDArT) and co-dominant (SNP) markers) for 105 Napier grass accessions.

The current study evaluated the level and patterns of genetic diversity in Napier grass accessions and progeny plants maintained in the ILRI forage gene-bank using genome-wide markers from the DArT-seq platform to generate baseline information for breeding, conservation and its future utilization.

1.2. Objectives of the study

General objective

To evaluate the level and patterns of genetic diversity in Napier grass accessions and progeny plants maintained in the ILRI forage gene-bank using genome-wide markers from the DArT-seq platform to generate baseline information for breeding, conservation and its wise use.

Specific objectives

- To assess genetic variation among and within accessions of Napier grass (*Cenchrus purpureus*) collections and progeny plants.
- To identify potential genotypes that could be used in the future Napier grass breeding program
- To identify potential duplicate and unique genotypes

2 Literature review

2.1 Origin and distribution of Napier grass

Napier grass (*Cenchrus purpureus* (Schumach.) Morrone syn. *Pennisetum purpureum* Schumach.), commonly called elephant grass originated from the tropical region of sub-Saharan Africa (Clayton *et al.*, 2013) and has been distributed as a forage crop into most tropical and subtropical regions all over the world (Negawo *et al.*, 2017). It was introduced into the USA in 1913 (Burton, 1990), into Brazil from Cuba in around 1920 (Daher *et al.*, 2002), into Central and South America and the West Indies in the 1950s and into Australia in the 1960s. Currently, it is naturalized to these areas and sometimes becomes invasive (CABI, 2014). It is often considered as a weed in crops, growing along roadsides, waterways, wetlands, floodplains, swamps, forest edges, disturbed areas and wastelands (Francis, 2004). Since it withstands drought, it is a pioneer species in arid lands such as the Galapagos Islands (CABI, 2014).

2.2 Taxonomy and botanical descriptions of Napier grass

2.2.1 Taxonomy

Elephant grass or Napier grass (*C. purpureus*), originally described and classified as *Pennisetum purpureum* Schumach (Stapf and Hubbard, 1934) and the taxon *Cenchrus purpureus* (Schumach.) Morrone was proposed in 2010 as a replacement for *Pennisetum purpureum* Schumach (Chemisquy *et al.*, 2010).

Cenchrus is the richest genus that consists about 140 species, including important cultivated species such as Napier grass, Pearl millet and Kikuyu grass (Kikuyu and Mithen, 1987). The species belonging to this genus constitute a heterogeneous assemblage with different basic chromosome numbers of 5, 7, 8 and 9, varying ploidy levels from diploid to octoploid with sexual or apomictic reproductive behavior and annual, biennial or perennial life cycles (Negawo *et al.*, 2017).

2.2.2 Botanical descriptions and ecology of Napier grass

Elephant grass (*C. purpureum*) is a perennial and one of the highest yielding monocot C4 major tropical grasses. It is a very versatile species that can be grown under a wide range of ecological conditions and systems: dry or wet conditions, smallholder or large-scale agriculture. It is a valuable forage and very popular throughout the tropics, as cut-and-carry animal feed (Negawo *et al.*, 2017).

Napier grass is a summer-growing grass that grows from sea level up to an altitude of 2500 m. It is more productive in places where temperatures range from 25 °C to 40 °C, annual rainfall is over 1500 mm and on rich, deep soils, such as friable loams (FAO, 2015; Skerman and Riveros 1990). It is not able to grow below 15 °C and is sensitive to frost, though it can regrow from the stolons if the soil is not frozen (Duke, 1983). It has some level of tolerance to drought and can grow in areas where the rainfall range is 200-4000 mm (Singh *et al.*, 2013), but it undergoes some morphological changes including leaf rolling, reduced stomatal conductance and enhanced water use efficiency to withstand water stress conditions (Negawo *et al.*, 2017). On the other hand, this grass doesn't tolerate flooding; it prefers well-drained soils. When there is a poor drainage system, it is best to grow it on raised beds (Göhl, 1982). However, it is able to grow on poorly drained clays, with a fairly heavy texture, or excessively drained sandy soils with a pH ranging from 4.5 to 8.2 (FAO, 2015). Elephant grass is a perennial full day sunlight species that can still grow under partial shade but does not withstand complete shade under a dense tree canopy (Francis, 2004). Morphologically it is described as a robust, tall, perennial grass with a vigorous root system that penetrates deep into the soil, developing from the nodes of its rhizomes; and it forms dense thick clumps, up to 1 m across. The culms are coarse, perennial, and up to 4-7 m in height, branched overhead. The leaves are flat, linear, and hairy at the base and up to 100-120 cm in length and 1-5 cm width, with a bluish-green color (some genotypes have purple coloured leaves), leaf margin is finely toothed and leaf blade has a prominent midrib (Singh *et al.*, 2013). The inflorescence is a stiff terminal bristly spike, up to 15-20 cm in length, yellow-brown to purplish in color. The spikelets are 4-6 mm long, surrounded by 2 cm long plumose bristles and arranged around a hairy axis that falls at maturity. There is little or no seed formation due to the grass's self-incompatibility and exogamous nature. When seeds are present, they are very small (3,000,000 seed/kg) (Francis, 2004; Mannetje, 1992).

2.3 Mode of propagation

2.3.1 Sexual propagation of Napier grass

The sexual propagation of Napier grass is limited due to their vegetative propagation nature, however some extent of sexual propagation through cross pollination have been reported (Negawo *et al.*, 2017). The sexual propagation through cross pollination often results in seeds of a mixed lot, thus the plants produced from them are not uniform and their performance is also unpredictable. In addition, the seeds germinate poorly, and seedlings are weak even when they grow fully, poor seed-setting and shattering make seed availability a problem. Still, some of the ILRI Napier grass accessions growing in the Bishoftu and Ziway sites produce seeds, which need further investigation whether the seeds have been produced through apomixis or sexual reproduction. A preliminary genetic diversity analysis among progeny plants raised from seeds suggested that sexual reproduction by cross pollination is possibly the major mechanism for the seeds production. In addition, Napier grass is naturally cross-compatible with pearl millet (*Cenchrus americanus*, 2n = 2x = 14) and their crosses result in the production of triploid hybrids, which are sterile and can be propagated vegetatively by means of stem cuttings (Negawo *et al.*, 2017).

According to Souza *et al.* (2019), about 95% of Napier grass seeds are predominantly derived from cross-fertilization and the outcrossing rate in their study reveals it is an allogamous grass with multilocus (tm) and unilocus outcrossing rates (ts) of 0.953 and 0.895, respectively. Their result is also consistent with the previous reports by Azevedo *et al.* (2012); Harris *et al.* (2010) which described

Napier grass as an allogamous species with high heterozygosity since outcrossing guarantees genetic variability, and thus creates new combinations of alleles within a species.

Napier grass, being a strictly out crosser (95%), is mostly self-incompatible and plants produced by self-fertilized (which is very rare) produce few viable seeds and the seeds show low germination index and the resulting plants have low vigour (Pereira *et al.*, 2010; Souza *et al.*, 2019; Hanna *et al.*, 2004). Self-incompatibility varies greatly among accessions and is directly affected by environmental factors such as temperature, humidity, different environments and artificial pollinating techniques (Souza *et al.*, 2019).

In addition, protogyny, which affects the production of sexual seeds, has already been described in this species. According to Pereira *et al.* (2010), stigmas are receptive between 3 to 5 days prior to the pollen grains release which makes fertilization within the same inflorescence unfeasible. Difficulties in self-fertilization limit the development of hybrids which offer opportunities for greater uniformity, higher selection intensities, absolute parental control and maximum exploitation of heterosis in Napier grass (Canto *et al.*, 2016).

2.3.2 Asexual propagation of Napier grass

Asexual or vegetative propagation of Napier grass is the most common method of propagation. Asexual propagation of Napier grass through stem chopping consists of at least 3 nodes in which two of these nodes are buried in the soil. The planting row width ranges from 50 to 200 cm and distance within rows ranges from 50 to 100 cm (Mannetje, 1992). After planting, elephant grass grows vigorously and can be up to 4 m high in three months (Skerman *et al.*, 1990). It is fast-growing and has a high annual productivity that depends on the climate, especially temperature and rainfall (Aroeira *et al.*, 1999; Artus-Poliakoff *et al.*, 1991). Elephant grass requires high levels of fertilizer and regular water supply (Mannetje, 1992) and the yields range from 20 to 80 t DM/ha/year under high fertilizer inputs (Francis, 2004; Skerman *et al.*, 1990), while 2-10 t DM/ha/year when there is no, or inadequate, fertilizer input (Bogdan, 1977). Cuttings can be made at 45-90-day intervals, depending on location (FAO, 2015).

2.4 Economic value of Napier grass

Napier grass has a lot of economic importance, it is primarily used as a fodder crop for small scale farmers since it grows with little nutrient supply and ease of management. The perennial nature of the grass and year-round availability makes it the first choice as a forage crop. It is also used as an

alternative bioenergy source, as a phytoremediation plant and as pest management in the push - pull management system (Negawo *et al.*, 2017).

A number of traits, including high dry matter production, ease of establishment and regeneration, persistence, and enhanced water use efficiency make Napier grass the primary forage of choice by small-scale dairy farmers (Nymbati *et al.*, 2010). Napier grass follows C4 photosynthetic pathway and is considered to have a competitive advantage over C3 grass species when grown in tropical and sub-tropical regions (Taylor *et al.*, 2011). It also has the capacity to reduce shoot dry matter and maximize carbon assimilation during times of water stress, making it a desirable forage crop in areas prone to droughts (Cardoso *et al.*, 2015). It is often fed fresh in cut and carry systems. It can also be chopped into pieces prior to feeding to reduce coarseness of leaves and stems for the animal. Moreover, wilting after chopping in the sun for several hours reduces moisture, facilitates rumination, stimulates appetite and thus improves forage utilization (Moran, 2011).

Napier grass (*C. purpureus*) genotypes were also evaluated by (de Morais *et al.*, 2009) for bioenergy production and they indicated that genotypes such as Cameroon, CNPGL F 06-3 and Bag 02 were promising for bioenergy production purposes. Other studies in Brazil's semiarid area using gypsum by (dos Santos *et al.*, 2015) also state that elephant grass, mainly Cameroon and Gramafante varieties, is a great potential energy crop. According to (Favare *et al.*, 2019), Elephant grass is an excellent alternative for bioenergy production mainly, due to its high percentage of stem and dry matter yield.

On other hand, eight Napier grass varieties were evaluated for their potential role as trap plants in the management of the African stemborer, *Busseola fusca* Füller (Lepidoptera: Noctuidae) in a push–pull strategy (Khan *et al.*, 2007). They concluded that the Napier grass cultivar called Bana had potential for use as a trap plant in the management of *B. fusca* in a 'push–pull' strategy, but the effectiveness of such a strategy would strictly depend on proper establishment and management of these companion plants.

In addition, Napier grass is also considered as one of the exceptional phytoremediator plants to tackle environmental pollutions due to its fast growth rate, high level of biomass yield, ease of establishment, low management cost and tolerance to pollutions; that enable it to accumulate large amount of pollutants (contaminants) in their system and able to degrade it (Lotfy and Mostafa, 2014; Roslan *et al.*, 2020; Tananonchai and Sampanpanish, 2018).

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2.4.1 Napier grass cultivation and its limitations

Napier grass as a major fodder for a dairy farm in East and central Africa, grown by more than 70 percent of smallholder dairy farmers in Kenya (Mulaa *et al.*, 2013); Uganda (Kabirizi *et al.*, 2007) and Tanzania (Pallangayo *et al.*, 2008). It constitutes between 40 to 80% of the forage for small holder dairy farmers (Staal *et al.*, 1997). In general, ease of establishment, management and regeneration, and its enhanced water-use efficiency make it primary forage of choice in the regions of tropics and subtropics for smallholder dairy farmers (Negawo *et al.*, 2017) however, the production of this forage crop is currently facing challenges from abiotic and biotic conditions such as climate change, Napier grass stunt and head smut diseases, respectively caused by a phytoplasma and a fungus *Ustilago kameruniensis*, which spread by wind, farm tools, infected plants, water, and animal manure respectively, have caused forage yield reduction of up to 90% (Mulaa *et al.*, 2013), and these biotic factors are currently the biggest threats to forage production and hence dairy sector in the East and Central Africa region.

Thus, information on the range of diversity of the available genetic resources of this important forage crop will help in the development of varieties that can overcome this production and productivity constraints.

2.5 Genetic diversity study of Napier grass

Precise assessment of the level of genetic diversity can be invaluable in plant breeding for diverse applications including: analysis of genetic variability in cultivars, identifying diverse parental combinations to create progenies with maximum genetic variability for further selection, and introgression of desirable genes from diverse germplasm into the available genetic base (Mohammadi and Prasanna, 2003). Therefore, a major focus of research in molecular genetics has been to determine the amount of genetic variation in populations and describe the possible mechanisms of maintaining such variability in meeting new environmental challenges (Weir, 1996).

Specifically, diversity information that can be derived from molecular analyses of Napier grass germplasm will help to determine the degree of relationships of the different germplasm within and among breeding populations and the germplasm held at the ILRI genebank, used in future association study to identify outstanding morphological and agronomic characters present in the germplasm. I will also provide the basic data for the effective utilization of the germplasm in the grass breeding programs for its improvement and conservation.

2.6 Genetic markers and their applications for genetic diversity analysis in Napier grass

Analysis of genetic diversity within and among populations involves the use of different genetic markers. Now, genetic markers are used in both basic plant research and plant breeding programs to characterize plant germplasm for gene isolation, marker-assisted introgression of favourable alleles, to produce improved varieties (Henry, 2001), and to obtain information about the genetic variation of populations for conservation and management purposes. There are three main classes of genetic markers for genetic diversity analysis: morphological markers, biochemical or protein markers, and DNA based molecular markers.

2.6.1 Morphological markers

Qualitative traits are usually controlled by a single locus and their expression is the same over a range of environmental conditions. These traits can be used as markers as they are usually visually described. These traits include seed shape, flower color, and seed colour (Bagali *et al.*, 2010). The advantage of using this marker for genetic diversity analysis within and among the population is that it does not require any sophisticated equipment and it is the most direct method. However, they are limited in number and subject to changes in environmental factors and may vary at different developmental stages (Mondini *et al.*, 2009). In Napier grass morphological markers were used by Van De Wouw *et al.* (1999) to assess variation among 53 accessions and these researchers indicated that some accessions could not be distinguished from each other by any of the characters observed. In general, morphological markers are not as such a reliable method to measure genetic differences related to productivity since most of the yield related traits are quantitate traits that are controlled by multi-loci and gene expression is influenced by different factors, including environmental factors.

2.6.2 Biochemical or protein based molecular markers

To overcome the limitation of morphological markers, protein-based biochemical markers and DNA based markers have been developed (De Vicente and Fulton, 2003). Biochemical marker-based analysis of genetic diversity is the separation of proteins into specific banding patterns. It is a fast method that requires only small amounts of isozymes or protein. Napier grass polymorphism based on isozymes and total proteins (TP) was assessed by Bhandari *et al.* (2006) and could distinguish variations among the assessed accessions. Since only a limited number of enzymes are available and

not evenly distributed throughout the genome of the organism, they do not clearly show diversity at the genome level (Mondini *et al.*, 2009).

2.6.3 DNA based molecular markers

Molecular markers use naturally occurring polymorphisms in DNA sequences due to mutation, recombination, or in some cases due to errors during DNA replication (Gupta *et al.*, 1999). Molecular markers are highly polymorphic and heritable, relatively simple to detect, distributed throughout the genome and completely independent of environmental conditions. They enable the detection of genetic variability at any stage of plant development. Main disadvantage of this technique is it requires more complex equipment and procedures. DNA based molecular markers are classified into two as hybridization-based marker and PCR based markers.

2.6.3.1 Restriction Fragment Length Polymorphism (RFLP)

Restriction Fragment Length Polymorphism (RFLP) is the first DNA based molecular marker that works by hybridization and has been an important tool for plant genome mapping (Jiang, 2013). It involves the digestion of genomic DNA into fragments with specific restriction enzymes and utilizes the variation in fragment length to assess differences. Variations in fragment length arise due to change (mutation) at an enzyme recognition site, single base pair substitution, insertion of genetic materials, such as transposable element, or by tandem duplications, deletions, translocations, or other rearrangements (Gupta *et al.*, 2002).

Smith *et al.* (1993) used RFLP and RAPD to study genetic diversity of Napier grass. RFLP is codominant locus-specific marker and is highly reproducible which can be used for different related organisms since they allow synteny (conserved order of genes between related organisms). The main shortcoming of RFLP markers is its tedious and time-consuming procedures.

2.6.3.2 PCR based markers

I. Random Amplified Polymorphic DNA (RAPD)

RAPD marker usually uses 10 nucleotides long single arbitrary primer to amplify the genomic DNA of different amplicon sizes. The difference in fragment length is due to some changes in the sequences found in primer annealing sites and between the two priming sites (Williams *et al.*, 1990). As stated by (Negawo *et al.*, 2017), Smith and co-workers were the first to use RAPD and AFLP markers to analyze diversity in Napier grass and able to clearly distinguish Napier grass accessions from its hybrid and

pearl millet. The main limitation of RAPD marker is low reproducibility and its dominant nature (Williams *et al.*, 1990).

I. Amplified Fragment Length Polymorphism (AFLP)

AFLP marker was originally developed for universal DNA fingerprinting analysis (Vos *et al.*, 1995). It is robust and relatively insensitive to PCR reaction conditions and highly reproducible. AFLP is used for the rapid screening of genetic diversity and intraspecific variation (Russell *et al*, 1997). Wanjala *et al.* (2013) used AFLP to study genetic diversity of Napier grass collections from east African region maintained by the ILRI forage genebank and they were able to discriminate and group the collections efficiently. The limitations of AFLP include difficulties in interpretation of band profiles in terms of loci, allele, and dominance.

II. Simple Sequence Repeat (SSR)

Simple sequence repeat (SSR) markers also called microsatellite markers are very short DNA (mono, di-, tri-, tetra-, penta-, and hexanucleotide) motifs usually characterized by a high degree of repetition even in thousands. They are evenly distributed throughout the genome of the organism (Singh *et al.,* 2010) and their polymorphism reflects differences in simple repetitive sequences of defined regions of the genome. Products of different sizes or lengths can be amplified with a pair of primers flanking different microsatellite regions. Typically, only a single locus is amplified resulting in single or double bands depending on the homozygosity or heterozygosity of that specific locus (Singh *et al.,* 2010). In general, SSR markers are locus-specific and co-dominant, simple to operate, abundant throughout the genome and highly reproducible. It has high rates of transferability from species to species. The cross-amplification rate of microsatellite markers between species of the same genus can vary from 50 to 100% (Azevedo *et al.,* 2012).

This transferability property of SSR markers was assessed between pearl millet (*C. americanus* syn. *Pennisetum glaucum*) and Napier grass (*C. purpureus* syn. *Pennisetum purpureum*) by Azevedo *et al.* (2012) to evaluate the genetic diversity and phylogenetic relationships among Napier grass accessions by using SSR markers that were originally developed for pearl millet. According to their study, about 55.5% of the primers showed successful cross-amplification to study the diversity of Napier grass and enabled them to cluster 107 accessions into three main groups. Kandel *et al.* (2016) also studied the genetic diversity of Napier grass using microsatellite, Single-Nucleotide Polymorphism (SNP) and Insertion-Deletion markers, which were originally developed for pearl millet. Those studies showed

the possibility of cross-amplification of the markers within the genus. The main limitation of this marker is the requirement for prior sequence information for the primer flanking regions to develop a pair of primers.

III. Inter Simple Sequence Repeat (ISSR)

Inter Simple Sequence Repeat (ISSR) marker was introduced in 1994 to asses genetic variation below the species level; mainly for studying population structure and differentiation of cultivated plants (Gupta *et al.*, 1994).This marker technique involves the amplification of DNA segment present in between two identical microsatellite repeat regions oriented in opposite direction (Joshi *et al.*, 2000). In Napier grass, Babu *et al.* (2009) used ISSR, together with RAPD, markers in the analysis of genetic diversity among thirty Napier grass genotypes from a wide geographical distribution.

IV. Single-Nucleotide Polymorphism (SNP)

SNP (Single Nucleotide Polymorphism) markers occur at high frequencies and are distributed across the genome. SNP marker differences in DNA sequence among genotypes are identified through an expensive and laborious DNA sequencing process. At first, SNPs revolutionize biomedicine, but since the technology depends on intensive genomic sequencing and a high cost of analysis that cannot be covered in agriculture or basic research. However, the development of a solid-state, open-platform method for DNA polymorphism analysis, for example the application of genotyping-by-sequencing (GBS) method, offers a low-cost high-throughput, robust system with minimal DNA sample requirement and is capable and has been providing high quality genome-wide coverage information even in organisms without any DNA sequence information such as Napier grass (Muktar *et al.*, 2019).

2.6.4 DArTseq markers

The DArTseq technology is one of the GBS methods, it uses a combination of genome complexity reduction using restriction enzymes and next generation sequencing (NGS) and produces high-density genome-wide dominant (SilicoDArT) and co-dominant (SNP) markers. This technology is an excellent approach and has been developed for the whole-genome profiling of Napier grass breeding programs and genetic resources conservation at the ILRI forage gene bank (Muktar *et al.*, 2019).

2.7 Parameters for genetic diversity analysis

Genetic diversity can be measured using different statistical methods. It can be in terms of the Coefficient of Correlation, genetic distance, and heterozygosity.

Coefficient of Correlation is defined as the probability that alleles of two individuals are identical by descent or by state. The value of coefficient of correlation ranges from zero, when the genotypes are completely unrelated; to one, when the two individuals have many alleles in common or are almost identical (Martin *et al.*, 1991). This Coefficient of Correlation is used to cluster genotypes into similar groups (Bered *et al.*, 2002). Coefficient of correlation (rxy) can be computed for all pairwise combinations of genotypes from pedigree information using the formula given by Falconer and Mackay (1996):

$$rxy = 2f xy | \sqrt{(1+Fx)(1+Fy)}$$

Where fxy= a coefficient of co-ancestry, Fx and Fy = inbreeding coefficients of X and Y, respectively.

Genetic distance is used to measure the genetic divergence between species or populations within a species, whether the distance measures degree of differentiation or time of divergence from a common ancestor. Many types of estimation of genetic distance are available and the appropriate choice of a genetic distance measure, on the basis of the type of the variable and the scale of measurement, is an important component in analysis of genetic diversity among genotypes. For the first time Nei defined genetic distance as the difference between two entities that is explained by allelic variation (Nei, 1972). Later on, it was defined in a more comprehensive way by Beumont et al. (1998) as any quantitative measure of genetic difference at either sequence or allele frequency level calculated between individual genotypes or populations. Currently there are different methods that are used to measure genetic diversity from molecular data. Some of them include Euclidean distance, Roger's distance, Fixation index (Fst) and analysis of molecular variance (AMOVA). Allelic diversity is used when genetic marker data or molecular marker data are interpreted in terms of locus/allele model. This allelic diversity may be expressed as the: a) percentage of polymorphic loci (p); b) mean number of alleles per locus (n), and; c) total gene diversity or expected heterozygosity (He), and polymorphic information content (PIC) (Bhanu, 2017). Percentage of polymorphic loci (p) gives an estimate of number of polymorphic loci with respect to total loci including polymorphic and monomorphic loci and is calculated as:

$$p = \frac{Np}{Nt} * 100$$

 N_p is the number of polymorphic loci and N_t is the number of total loci

Heterozygosity (He) and polymorphic information content (PIC), which is an indirect estimate of the number of alleles per locus, can be calculated as:

$$He = 1 - \sum_{i=1}^{n} (Pi)^2$$
 and PIC = $1 - \sum p_i^2 - \sum \sum p_i^2 p_j^2$

Where, P_i and p_j are the frequency of i^{th} and j^{th} allele at a particular locus

The PIC is a good index for genetic diversity evaluation and used to evaluate the level of gene variation, with values ranging from zero to one. A PIC value > 0.5, indicates a locus with high diversity, a PIC value < 0.25 a locus with low diversity and a PIC value between 0.25 and 0.5 for a locus with intermediate diversity (David *et al.*, 1980).

Hardy-Weinberg Equilibrium (HWE): Hardy-Weinberg equilibrium explains that both gene and genotype frequencies will be constant from generation to subsequent next generations under the assumptions that no genetic drift, mutation, and natural selection happened in the population and the population is closed (no gene flow) and has a random mating pattern (Labate, 2000).

F-statistic (Fst): Fixation indices are the measures of standardized variances in allele frequencies that detect departure from HWE caused by biased inbreeding, out breeding, or population subdivision and genetic drift (Wright, 1950). Hence, the F statistic quantifies the mean heterozygosity difference between populations and subpopulations. Fst is considered to be the most informative statistic for examining the overall level of genetic divergence among subpopulations and can be calculated as:

$$Fst = \frac{(HT - HS)}{HT}$$

Where Fst is reduction in heterozygosity, H_T is average heterozygosity in a population and H_S average heterozygosity in a subpopulation.

The Fst value ranges between zero and one. When it is equal to zero, it means complete sharing of genetic material and when it is one, no sharing (the populations are fixed).

Analysis of Molecular Variance (AMOVA): is a method of estimating variance components within and among populations directly from molecular data and testing hypotheses about differentiation (Excoffier *et al.*, 1992). AMOVA treats molecular data as a vector q_i which is a matrix of 1s and 0s, 1 indicating the presence of a marker and 0 its absence.

Euclidean distance: is one of the genetic distance measures based on allele frequency distribution between pairs of vectors calculated by subtracting the vector of one haplotype from another, according to the formula $(\mathbf{q}_j - \mathbf{q}_k)$. If \mathbf{q}_j and \mathbf{q}_k are visualized as points in *n*-dimensional space indicated by the intersections of the values in each vector, with *n* being equal to the length of the vector, then the Euclidean distance is simply equal to the shortest distance between those two points. The distance between points p and q is the length of the line segment connecting them (pq) (Aremu, 2012).

2.7.1 Multivariate statistics to estimate genetic diversity

Genetic diversity among different varieties and/or species can be assessed using multivariate statistics. This provides reliable information on the real genetic distances between genotypes hence it is a tool for assessment of genetic diversity (Bhanu, 2017). Some of the multivariate techniques include:

I. Cluster analysis

These techniques depict a pattern of similarity/relatedness between genotypes based on their evolutionary relationships and group the similar ones in the same group while differentiating the others. This method is mainly based on the unweighted paired group method using arithmetic mean (UPGMA) to provide precise grouping information on breeding materials used in accordance with pedigrees and calculated results found in agreement with known heterotic groups than the other clusters (Aremu *et al.*, 2007).

II. Principal component analysis (PCA)

Principal component analysis (PCA) can be described as a quantitative type of data reduction technique. This technique transforms multi-correlated variables into different sets of uncorrelated variables for further study (Bhanu, 2017). The new variables are in linear combinations with the original variables. It is based on the development of characteristic values and mutually independent principal components arranged in a decreasing order of variance. The technique is most suitable when different variables have the same unit and is difficult for different scales. This difficulty is avoided by standardizing all the variables and to do this each variable is divided by its estimated standard deviation. This technique is a means for further analysis, not an end (Bhanu, 2017).

2.8 Some statistical tools for genetic diversity analysis

I. Numerical Taxonomy System for personal computer (NTSYSpc)

NTSYSpc is used to analyze genetic diversity from different molecular marker data and works based on similarity indices as a 0, 1 matrix of genotypic data. It is used for cluster analysis, principal component, and/or principal coordinate analysis (Rohlf, 1998).

II. Genetic Analysis in Excel (GenAlEx)

GenALEx is an Excel add-in easy and user-friendly program designed for use with SSR, SNP, and AFLP, allozyme, multi locus markers and sequencing DNA data in genetic diversity analysis. It works with three data type's codominant data, dominant, and geographic data. Analyses performed include: observed and expected heterozygosity, marker index, fixation index, allelic patterns, haploid diversity by population, haploid diversity by locus, haploid disequilibrium and, Nei's genetic distance, principal component analysis and Shannon index (Bhanu, 2017).

III. Popgene

Popgene software is a user-friendly package developed for the analysis of genetic diversity among and within natural populations. Codominant data, dominant and quantitative trait data are the data types it works on and performs population genetic structure analysis using markers or phenotype/trait data based on gene frequency, number of allele, polymorphic loci, gene diversity, Shannon index, homozygosity, gene flow, genetic distance (based on Nei coefficient) and produces a dendrogram based on UPGMA and neighbor-joining methods (Bhanu, 2017).

IV. R statistical software

R is a free, open source, user friendly software based on a programming language developed in 1995 at the University of Auckland as an environment for statistical computing and graphics (Ikaha and Gentleman, 1996). Currently, several scientific disciplines, including medicine, agriculture, soil science, and ecology preferentially use R software due to its graphical capabilities and its free availability with different packages that are designed for analyzing different data types.

The free software R (https://www.r-project.org/) is a standard and preferable for the analysis of genetic data, offering packages that are dedicated to population genetics (Paradis, 2010), phylogenetics (Schliep, 2011) or genome-wide association studies (Clayton and Leung, 2007). When the R software is known as a standard for genetic data analysis, classical population genetics tools are being challenged by the increasing availability of genomic sequences since dedicated tools are needed for harnessing the large amount of information generated by currently available sequencing technologies.

The R package Adegenet for multivariate analysis: this R package is contributed and used to implement classes and functions to facilitate the multivariate analysis using genetic markers. The package allowed the use of big marker data and compresses and defines new formal classes of

genotypes (genind) or groups of genotypes (genpop), which can be used as an input to multivariate analysis. A lot of functions are also implemented to manipulate and analyze these objects, including recent developments in spatial genetics and data simulations. By assuring a good interoperability of data, adegenet contributes to making the R software a unifying platform for the analysis of genetic markers (Jombart *et al.*, 2008).

The R package Poppr for population genetic analysis: the R package Poppr gives significant, accessible tools for the analysis of clonal, partially clonal, and sexual populations available in one environment on all major operating systems. The capacity to analyze data for multiple populations across a user-defined hierarchy and provide novel functionality in R, combined with R's graphing abilities of publication-ready figures is thus obtained conveniently (Kamvar *et al.*, 2014).

The R package Ape for Phylogenetics and Evolution Analysis: Analysis of Phylogenetics and Evolution (APE) is a package written in the R language for use in molecular evolution and phylogenetics. APE enables the manipulation of phylogenetic trees, as well as several advanced methods for phylogenetic and evolutionary analysis such as comparative and population genetic methods (Paradis *et al.*, 2004, 2019).

The R package synbreed: the package synbreed was developed within the synbreed project for synergistic plant and animal breeding (<u>www.synbreed.tum.de</u>) and genetic data analysis. This package executes data processing, data analysis, and visualization since it contains a collection of functions that are embedded within the framework of a single, unified dataset that is required for genomic analysis (Wimmer *et al.*, 2012). Moreover, its implementation is flexible with respect to different ranges of data formats.

Pegas R package for AMOVA analysis: the package Pegas provides functions for data reading, writing, plotting, analyzing and manipulating allelic and haplotypic data. It is used for analysis of linkage disequilibrium (LD), population structure (fixation index (Fst)), analysis of molecular variance (AMOVA) and Hardy-Weinberg Equilibrium (HWE) (Knaus and Winter, 2020).

3 Materials and Methods

3.1 Plant material

A total of 363 Napier grass accessions that comprises 60 accessions from the ILRI genebank collection; 45 (25 Centro Nacional de Pesquisa de Gado de Leite (CNPGL), EMBRAPA's elite lines and 20 Brazilian Active Germplasm Bank of Napier Grass (BAGCE)) accessions from the Brazilian Agricultural Research Corporation (EMBRAPA), Brazil, 22 accessions from the United States Department of Agriculture (USDA), Tift, Georgia, USA, 31 accessions from the ICRISAT genebank, Patancheru, India (Supplementary Table S1) and a total of 205 progeny plants raised from seeds of 13 ILRI accessions (Table 1), were included in this genetic diversity study.

Napier grass seeds collected from 13 seed-bearing accessions and maintained in the ILRI forage genebank, approximately 20 to 30 seeds per accession were obtained and pre-germinated on agar medium containing potassium nitrate. The germinated seeds were transplanted into soil-filled pots and maintained in the screen house until they produce 3 to 4 leaves for sample collection. On average, about 16 progeny plants from each of the 13 accessions were sampled to determine the level of genetic diversity within and among genotypes. On the other hand, the recently introduced accessions from USDA and ICRISAT were propagated through stem cuttings. As indicated by Mannetje (1992), Napier grass stems were chopped into fragments consisting of at least 3 nodes in which two of the nodes were buried in the soil at a 45⁰ angle during planting and were watered two times a day.

Acc. No.	Species	Progenies	No. of progenies
ILRI_1026	Cenchrus purpureus	ILRI_NS_1	15
ILRI_16789	Cenchrus purpureus	ILRI_NS_2	7
ILRI_16839	Cenchrus purpureus	ILRI_NS_3	20
ILRI_16783	Cenchrus purpureus	ILRI_NS_4	17
ILRI_14983	Cenchrus purpureus	ILRI_NS_5	17
ILRI_16835	Cenchrus purpureus x C. glaucum	ILRI_NS_6	19
ILRI_16837	Cenchrus purpureus x C. glaucum	ILRI_NS_7	18
ILRI_16803	Cenchrus purpureus	ILRI_NS_8	14
ILRI_16821	Cenchrus purpureus	ILRI_NS_9	10
ILRI_16818	Cenchrus purpureus	ILRI_NS_10	13
ILRI_16810	Cenchrus purpureus	ILRI_NS_11	19
ILRI_14984	Cenchrus purpureus	ILRI_NS_12	18
ILRI_16790	Cenchrus purpureus	ILRI_NS_13	18

Table 1: Napier grass progeny plants raised from the 13 accessions. The detail of each progeny and accessions used in the study is shown in Supplementary Table S1.

3.2 DNA extraction and genotyping

Young leaf samples were collected from each individual plant into 2ml Eppendorf tubes, using ice in an ice-box and transferred to a -80 freezer as quickly as possible. The leaf samples freeze-dried for about 48 hours and ground into fine powder using a tissue grinder (tissue lyser). Genomic DNA was extracted from the leaf powder using a DNeasy[®] Plant Mini Kit (250) (Qiagen Inc.,Valencia, CA) following the manufacturer's procedures. The genomic DNA concentration and quality was checked using a Nano-drop spectrometer (DeNovix DS-11 FX spectrophotometer) and by using a 0.8% agarose gel electrophoresis. The concentration of the DNA was adjusted to 50 to 100 ng/µl and sent for DArTseq sequence genotyping under Integrated Genotyping Service and Support (IGSS) platform at BecA-ILRI, Nairobi, Kenya and genotyped as described in Muktar *et al.* (2019).

3.3 Data analysis

The genotype data was analyzed using different statistical tools; the missing percentage of data, expected heterozygosity, and polymorphic information content of the markers were calculated in Excel (Microsoft Excel for Office 365). The distributions of these markers were also analyzed and visualized using the R package synbreed (version 0.12-12) (Wimmer et al., 2012). From the genome wide SNP markers, a subset of robust markers was selected based on their expected heterozygosity (He), polymorphic information content (PIC), missing value percentage, minor allele frequency (MAF), and genome wide distribution. The markers were further filtered based on their best contribution to the genetic differentiation and diversity by employing discriminant analysis of principal components (DAPC) (Jombart and Ahmed, 2011). The selected robust markers were used for genetic diversity and population structure analysis by using Euclidean genetic distance and neighbor-joining (NJ), Unweighted paired group method using arithmetic mean (UPGMA). Hierarchal clustering were calculated using the R functions dist () and NJ () and dendrogram or phylogenetic trees were generated using the Adegenet (Jombart et al., 2008) and Poppr (Kamvar et al., 2014) packages in R statistical software. The degree of genetic similarity between genotypes was analyzed using Nei's genetic distance (Nei, 1972; Pagnotta, 2018). Genetic diversity and population stratification were analyzed by the Bayesian algorithm implemented in the STRUCTURE software (Corander et al., 2013; Pritchard, 2009) and DAPC (Jombart and Ahmed, 2011), using the filtered robust SNP markers. In addition, major clusters and sub clusters were detected using hierarchical clustering with complete linkage analysis (Kamvar et al., 2014; Mohammadi and Prasanna, 2003) and Bayesian model based Population

Structure analysis was carried out using STRUCTURE software and the true value of K was determined using the method of Evano *et al.* (2005). AMOVA (Excoffier *et al.*, 1992) and fixation index (Meirmans and Hedrick, 2011) were employed to determine the extent of population differentiation among different groups and sub groups that were obtained from population structure analyses using the R package Pegas (Knaus and Winter, 2020). Diversity among and within accessions and progeny plants was analyzed using statistics on R software.

4 **Results**

4.1 Genome-wide distribution and polymorphism of the DArTseq markers

A total of 363 Napier grass genotypes from different collections and progeny plants were genotyped using the DArTseq platform and 96,454 SilicoDArT and 96,321 SNP markers were generated. Prior to further diversity analysis, genotype missing percentage was checked and 16 genotypes (6 progenies, 2 EMBRAPA collections, 1 EMBRAPA elite line, 3 ICRISAT, 3 USDA and 1 ILRI accession), which had high missing value (\geq 50%) were excluded from further analysis hence a total of 347 Napier grass genotypes were used for diversity analysis.

The percentages of missing values of the markers ranged from 0.0 to 29.0 % for SilicoDArT markers and 0 to 83.7 % for SNP markers. The expected heterozygosity (He) and polymorphic information content (PIC) of the markers ranged from 0 to 0.5 and 0 to 0.38 respectively for both SilicoDArT and SNP markers while the average He was 0.26 and 0.18, and the average PIC was 0.21 and 0.15 for the SilicoDArT and SNP markers, respectively (Fig. 1).

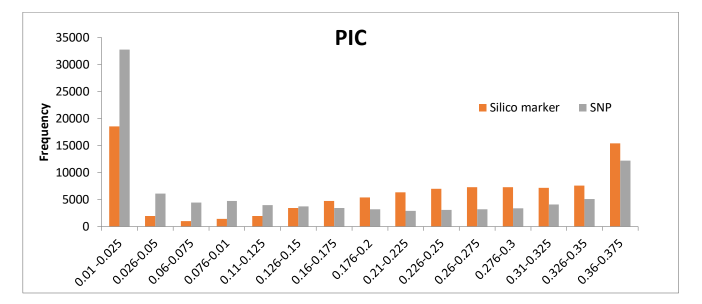
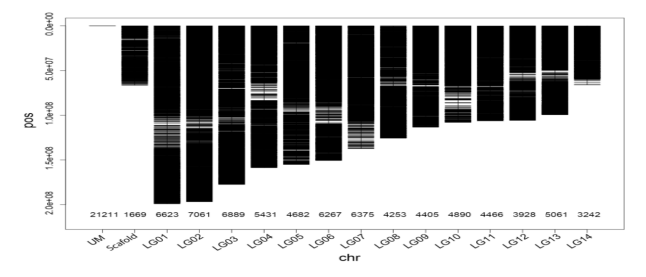


Figure 1: Frequency distribution of polymorphic information content (PIC) values of SilicoDArT and SNP markers

In line with the distribution of SilicoDArT markers throughout the genome of Napier grass, out of 96,453 markers, 76.28 % (73,573) mapped on to the fourteen linkage groups of the Napier grass genome (Yan *et al.*, 2020), while 1.73 % (1,669) mapped onto scaffolds and 21.99% (21,211) were found to be unmapped (unknown location). The highest number of markers were mapped to linkage group two (LG2) (7.32% (7,061) and (LG3) 7.14% (6,889) followed by LG1, LG7 and LG6, in which 6.87% (6,623), 6.61% (6,375) and 6.50% (6,267) were mapped respectively, while LG14 contained the least number of markers (3.36% (3,242)) (Fig. 2a).

From the 96,400 SNP makers, 88.82% (85,619) were aligned on to the fourteen linkage groups while 1.87% (1,802) mapped on to different scaffolds and 9.31% (8,979) were not able to be mapped on to the draft genome. Out of the aligned 88.82% markers, 8.74% were mapped on LG2 and 8.51% on LG3. The percentage of mapping for LG1, LG7, LG6, LG4 and LG13 were 7.69% (7,416), 7.40% (7,129), 7.32% (7,058), 6.50% (6,264) and 6.15% (5,925), respectively. The lowest numbers of markers 3.79% (3,655) were mapped on LG14 (Fig.2b).



a)

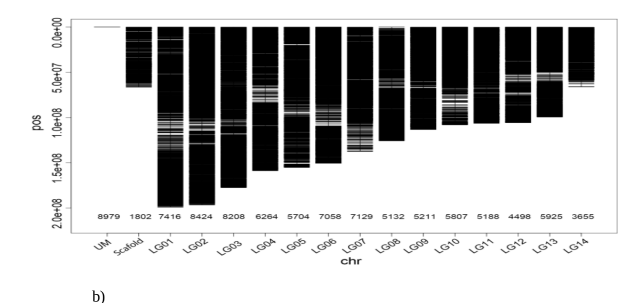


Figure 2: The Distribution of SilicoDArT (a) and SNP (b) markers across the fourteen linkage groups (LG) of the Napier grass (*C. purpureus*) draft genome.

4.1.1 Marker selection for diversity study

From the 96,321 SNP markers generated, a subset of robust markers was selected based on the following criteria:

- I. Markers with missing data percentage less than or equal to ten percent (NA% ≤ 10 %)
- II. Minor allele frequency greater than or equal to five percent (MAF \geq 5 %)
- III. Polymorphic information content (PIC) greater than or equal to zero point two (PIC ≥ 0.2)
- IV. Expected heterozygosity greater than or equal to zero point two (He ≥ 0.2)
- V. Distribution of the markers across the linkage groups (LG) (genome-wide distribution)
- VI. Markers contribution to diversity using the loading function in discriminant analysis of principal components (DAPC).

Stepwise marker selection was conducted in such a way that each criterion is taken into account after the previous criteria are considered. Hence, initially markers were selected based on missing value percentage (NA %) in which markers with missing value percentage of less than or equal to ten were retained while the rest were removed so that 79,831SNP markers were retained. Based on the second criterion of minor allele frequency (MAF \geq 5 %), 3,629 SNP makers were retained. Further the third and fourth criteria of PIC and He reduced the markers to 2,357 SNP markers. Finally, 1001 robust and genome-wide distributed markers were selected based on the markers contribution to genetic diversity and genetic differentiation according to the DAPC analysis by using the R package Adegenet. These selected 1001 SNP markers, with an average PIC value of 0.31, were used in genetic diversity and population structure analysis. The distribution of the selected markers was visualized using the R package synbreed and out of these selected markers, 1.8% (18) were unmapped, 0.9% (9) mapped to scaffolds, and the rest were distributed across the linkage groups with the highest distribution (map) number on LG2, (9.2% or 92 markers) and lowest number on LG14, which was 4.6% or 46 markers (Fig. 3).

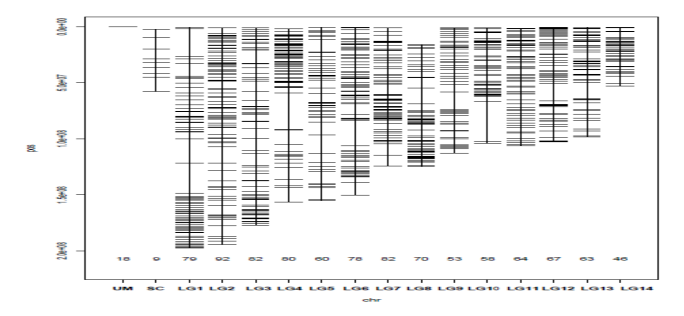


Figure 3: Genome wide distribution of 1001 SNP markers selected based on different criteria, including the Discriminant Analysis of Principal Components (DAPC).

4.1.2 Genetic diversity in Napier grass populations

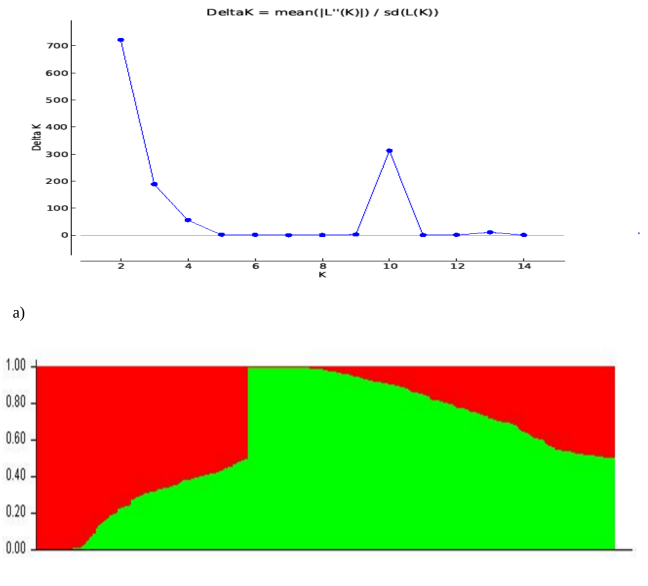
Genetic diversity and population stratification were detected by using DAPC, Structure software, and hierarchal clustering approaches and a significant amount of genetic diversity and population stratifications in the Napier grass collections and progeny plants with a high degree of consistency were obtained. The degree of genetic similarity between genotypes was identified using Nei's genetic distance (Nei, 1972) (Table 6). Furthermore, a high amount of genetic diversity and the extent of variation among clusters and sub clusters were confirmed by AMOVA (Table 7), and the fixation index (Fst) (Table 6).

4.1.3 Genetic diversity revealed by the Bayesian algorithm in STRUCTURE software

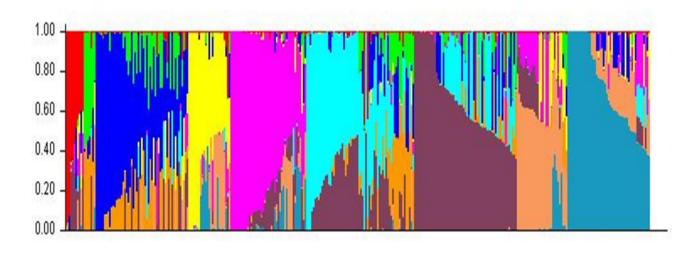
Population structure analysis using the Bayesian algorithm in structure software revealed two major clusters with ten sub clusters as suggested by the delta k (Dent, 2012) (Fig. 4a). Cluster I comprises almost half of the progeny plants (91 out of 199) and most of the EMBRAPA collection (BAGCEs, 15 out of 19), some accessions from the ICRISAT collection (9 out of 25), some from the ILRI collection (20 out of 59), two of the EMBRAPA elite lines (CNPGLs), and just one from the USDA collection. Cluster II is composed mainly of genotypes from the ILRI collection, progeny plants, most of the EMBRAPA elite lines (CNPGLs) and the USDA collection.

The collections were further grouped into ten sub-clusters, in which cluster I was further divided into six sub-clusters, while cluster II was divided into four sub-clusters (Supplementary Table S2). Eighty-one individual genotypes were found to be admixed (membership probability < 50 %) among the sub-clusters (Supplementary Table S3). The first sub-cluster from the population structure analysis consisted of 47 progeny plants (13 of the ILRI_NS_11, 16 of the ILRI_NS_7 and ILRI_NS_9, 14 of the ILRI_NS_3 and ILRI_NS_6, 3 of the ILRI_NS_13 and 1 of the ILRI_NS_4). The second sub cluster consisted of 6 genotypes that were entirely ILRI accessions. The third sub cluster consisted of 7 genotypes, all progeny plants from ILRI_NS_6. The fourth sub-cluster consisted of 23 genotypes (11 of the ILRI collection, 9 genotypes of the ICRISAT collection and 3 genotypes of the EMBRAPA collection). The fifth sub-cluster consists of 18 genotypes, out of these 12 were from the EMBRAPA collection, 3 were from the ILRI collection, 2 were EMBRAPA elite lines and one USDA genotype. The sixth sub-cluster consisted of 37 genotypes, all of which were progeny plants (10 of ILRI_NS_12 and ILRI_NS_13, 6 of ILRI_NS_10, 7 of ILRI_NS_7, 4 of ILRI_NS_4 and 2 of ILRI_NS_6). The

seventh sub-cluster consisted of 29 genotypes, of which 21 (72.41%) were from the ILRI collection, 4 (23.79%) were EMBRAPA elite lines, two were from the USDA and one from each of the ICRISAT and EMBRAPA collections. The eighth sub-cluster consisted of 27 genotypes, mostly genotypes from the USDA collection and EMBRAPA elite lines. The ninth sub-cluster consisted of 7 genotypes, of which six were from the ILRI collection and one was a USDA genotype. The last sub-cluster (sub-cluster X) consists of 66 genotypes, mostly represented by progeny plants (ILRI_NS_1, ILRI_NS_2, ILRI_NS_3, and ILRI_NS_5 and ILRI_NS_8) with only two genotypes from the ILRI collection and one from the ICRISAT collection.



b)



c)

Figure 4. Population structures identified by structure analysis in the Napier grass collections and progenies: a) Delta k that shows picks at K = 2 and K = 10, suggesting possible number of populations/clusters; b) The two major clusters of the populations/structures; c) The ten populations/sub-clusters and the admixtures.

4.1.4 Genetic diversity revealed by the Discriminant Analysis of Principal Components (DAPC)

DAPC clustered the population into ten clusters (K groups) (Fig. 5). The first cluster (K1) consisted of a total of 52 progeny plants, of which 38 (73.1%) were from ILRI_NS_12, ILRI_NS_13 and ILRI_NS_7. Generally, cluster I represents progeny plants while the second cluster consisted of 52 genotypes, out of these 50 genotypes were progeny plants and the other two were USDA and ICRISAT accessions. The third cluster consisted of 33 genotypes and of these 32 were progenies and one ILRI accession. The fourth cluster consisted of 44 genotypes and of these 19 were progeny plants while the remaining were contributed by BAGCE (1), CNPGL (8), USDA (9), ICRISAT (4) and ILRI (3) collections. The fifth cluster consisted of 28 genotypes, most were ILRI and CNPGL genotypes (11 and 9, respectively). The remaining four were ICRISAT and USDA genotypes. The sixth cluster consisted of six genotypes which were entirely from ILRI. The seventh cluster consisted of 30 genotypes, while half (15) of these were ILRI genotypes, 11 were ICRISAT genotypes, 3 were BAGCE and 1 was a USDA genotype. The eighth cluster consisted of 48 genotypes, while 46 were progeny plants and the remaining 2 were ILRI accessions. The ninth cluster consisted of 34 genotypes from BAGCE (13), ILRI (8), ICRISAT (5), USDA (4) and CNPGL (4). The last cluster (cluster 10) consisted of 20 genotypes of these 16 were ILRI accessions, 3 were EMBRAPA elite lines and 1 was from the EMBRAPA collection.

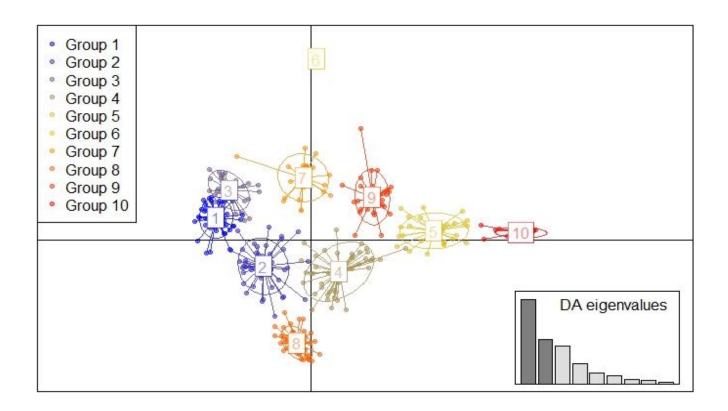


Figure 5: The ten clusters detected by the Discriminant Analysis of Principal Components (DAPC) of the 347 Napier grass genotypes. The axes represent the first two Discriminants. Each circle represents a cluster, and each dot (point) represents an individual.

k1	k2	k3	k4	k5	k6	k7	k8	k9	k10
ILRI_NS_4.2	ILRI_NS_1.4	ILRI_NS_3.2	ILRI_NS_1.9	CNPGL_93.08.1	ILRI_16808	BAGCE_24	ILRI_Ns_1.1	BAGCE_16	BAGCE_63
ILRI_NS_4.10	ILRI_NS_1.16	ILRI_NS_6.5	ILRI_NS_3.11	CNPGL_93.32.2	ILRI_16809	BAGCE_80	ILRI_NS_1.2	BAGCE_56	CNPGL_92.38.2
ILRI_NS_4.11	ILRI_NS_1.18	ILRI_NS_6.8	ILRI_NS_5.5	CNPGL_94.07.2	ILRI_16810	Tift_N130	ILRI_NS_1.3	BAGCE_7	ILRI_14984
ILRI_NS_4_15	ILRI_NS_3.1	ILRI_NS_6.10	ILRI_NS_5.8	Tift_N147	ILRI_16818	India_149_21785	ILRI_NS_1.5	BAGCE_75	ILRI_16785
ILRI_NS_7.1	ILRI_NS_3.3	ILRI_NS_6.12	ILRI_NS_5.12	Tift_N109	ILRI_16821	India_126_22231	ILRI_NS_1.6	BAGCE_94	ILRI_16786
ILRI_NS_7.2	ILRI_NS_3.4	ILRI_NS_6.14	ILRI_NS_5.19	Tift_N131	ILRI_16822	India_118_22241	ILRI_NS_1.10	Tift_N37	ILRI_16787
ILRI_NS_7.4	ILRI_NS_3.6	ILRI_NS_6.15	ILRI_NS_6.4	Tift_N138		India_124_22233	ILRI_NS_1.12	Tift_N210	ILRI_16789
ILRI_NS_7.5	ILRI_NS_3.7	ILRI_NS_6.19	ILRI_NS_6.11	India_150_21784		India_125_22232	ILRI_NS_1.13	Tift_N71	ILRI_16792
ILRI_NS_7_6	ILRI_NS_3.8	ILRI_NS_7.3	ILRI_NS_6.13	India_144_21964		India_115_22243	ILRI_NS_1.14	Tift_N137	ILRI_16795
ILRI_NS_7.12	ILRI_NS_3.9	ILRI_NS_7.7	ILRI_NS_6.18	India_141_21967		India_132_22225	ILRI_NS_1.15	India_127_22230	ILRI_16798
ILRI_NS_7.13	ILRI_NS_3.10	ILRI_NS_7.10	ILRI_NS_6.20	India_147_21787		India_121_22237	ILRI_NS_2.1	India_146_21788	ILRI_16800
ILRI_NS_7.14	ILRI_NS_3.13	ILRI_NS_7.11	ILRI_NS_8.2	ILRI_14355		India_123_22234	ILRI_NS_2.2	India_142_21966	ILRI_16801
ILRI_NS_7.15	ILRI_NS_3.14	ILRI_NS_9.1	ILRI_NS_8.4	ILRI_14389		India_89_22236	ILRI_NS_2.3	India_131_22226	ILRI_16803
ILRI_NS_7.17	ILRI_NS_3.15	ILRI_NS_9.2	ILRI_NS_8.9	ILRI_14982		India_116_22242	ILRI_NS_2.6	India_151_21783	ILRI_16804
ILRI_NS_7.18	ILRI_NS_3.16	ILRI_NS_9.8	ILRI_NS_8.14	ILRI_15357		ILRI_1026	ILRI_NS_2.7	ILRI_15743.MOTT.	ILRI_16806
ILRI_NS_7.20	ILRI_NS_3.17	ILRI_NS_9.9	ILRI_NS_8.16	ILRI_16793		ILRI_16782	ILRI_NS_2.8	ILRI_16783	ILRI_16836
ILRI_NS_10.2	ILRI_NS_3.18	ILRI_NS_9.10	ILRI_NS_8.17	ILRI_16799		ILRI_16790	ILRI_NS_2.9	ILRI_16788	ILRI_18438
ILRI_NS_10.4	ILRI_NS_3.20	ILRI_NS_9.12	ILRI_NS_12.10	ILRI_16819		ILRI_16794	ILRI_NS_3.5	ILRI_16791	CNPGL_9279.2
ILRI_NS_10.5	ILRI_NS_4.1	ILRI_NS_9.13	ILRI_NS_12.19	ILRI_16837		ILRI_16796	ILRI_NS_3.12	ILRI_16802	CNPGL_92.66.3
ILRI_NS_10.6	ILRI_NS_4.3	ILRI_NS_9.15	BAGCE_1	ILRI_16902		ILRI_16797	ILRI_NS_3.19	ILRI_16812	Napier_Addis
ILRI_NS_10.7	ILRI_NS_4.4	ILRI_NS_9.18	CNPGL_91.06.2	ILRI_16840		ILRI_16805	ILRI_NS_4.14	ILRI_16813	
ILRI_NS_10_8	ILRI_NS_4.5	ILRI_NS_11.4	CNPGL_91.11.2	CNPGL_93.18.2		ILRI_16807	ILRI_NS_5.1	ILRI_16815	
ILRI_NS_10.9	ILRI_NS_4.6	ILRI_NS_11.5	CNPGL_91.25.1	CNPGL_93.01.1		ILRI_16814	ILRI_NS_5.3	BAGCE_97	
ILRI_NS_10.11	ILRI_NS_4.7	ILRI_NS_11.8	CNPGL_92.190.01	CNPGL_96.27.3		ILRI_16816	ILRI_NS_5.2	CNPGL_00.1.1	
ILRI_NS_10.12	ILRI_NS_4.9	ILRI_NS_11.12	CNPGL_93.06.1	CNPGL_92.133.3		ILRI_16817	ILRI_NS_5.4	BAGCE_30	
ILRI_NS_11_6	ILRI_NS_4.12	ILRI_NS_11.13	Tift_N200	CNPGL_92.56.2		ILRI_16834	ILRI_NS_5.6	BAGCE_53	
ILRI_NS_12.1	ILRI_NS_4.13	ILRI_NS_11.14	Tift_N172	CNPGL_94.13.1		ILRI_16838	ILRI_NS_5.9	BAGCE_90	
ILRI_NS_12.3	ILRI_NS_4.18	ILRI_NS_11.15	Tift_N43	Maralfalfa.1		ILRI_18448	ILRI_NS_5.10	BAGCE_81	
ILRI_NS_12.4	ILRI_NS_4.19	ILRI_NS_11.16	Tift_N23			ILRI_18662	ILRI_NS_5.13	BAGCE_34	
ILRI_NS_12.5	ILRI_NS_4.20	ILRI_NS_11.17	Tift_N225			BAGCE_17	ILRI_NS_5.14	BAGCE_86	
ILRI_NS_12.6	ILRI_NS_5.16	ILRI_NS_11.18	Tift_N75				ILRI_NS_5.15	CNPGL_96.23.1	
ILRI_NS_12.8	ILRI_NS_6.1	ILRI_NS_11.20	Tift_N8				ILRI_NS_5.17	CNPGL_92.198.7	
ILRI_NS_12.11	ILRI_NS_6.3	ILRI_16835	Tift_N223				ILRI_NS_5.18	PIONEIRO	
ILRI_NS_12.12	ILRI_NS_6.6		Tift_N68				ILRI_NS_6_2	BAGCE_100	
ILRI_NS_12.13	ILRI_NS_6.7		India_145.121965				ILRI_NS_8.3		
ILRI_NS_12.14	ILRI_NS_6.9		India_120_22238				ILRI_NS_8.7		
ILRI_NS_12.17	ILRI_NS_6.16		India_119_22239				ILRI_NS_8.8		
ILRI_NS_12.18	ILRI_NS_7.9		India_129_22228				ILRI_NS_8.10		
ILRI_NS_12.20	 ILRI_NS_9.7		ILRI_16784				ILRI_NS_8.11		
ILRI_NS_13.1	ILRI_NS_10.1						ILRI_NS_8.15		
ILRI_NS_13.2	ILRI_NS_10.3		 CNPGL_96.21.1				ILRI_NS_8.19		
ILRI_NS_13.3	 ILRI_NS_10.10		 CNPGL_93.37.5				ILRI_NS_8.20		1
 ILRI_NS_13.5	ILRI_NS_10.13		 CNPGL_93.04.2				ILRI_NS_11.3		
ILRI_NS_13.7	ILRI_NS_11.1		mott_new2				ILRI NS 11.9		1

 Table 2: List of genotypes under different K groups from Discriminant Analysis of Principal Components (DAPC)

ILRI_NS_13.8	ILRI_NS_11.2			ILRI_NS_12.7	
ILRI_NS_13.9	ILRI_NS_11.10			ILRI_NS_12.15	
ILRI_NS_13.11	ILRI_NS_11.11			ILRI_14983	
ILRI_NS_13.12	ILRI_NS_11.19			ILRI_16839	
ILRI_NS_13.13	ILRI_NS_12.2				
ILRI_NS_13.15	ILRI_NS_13.17				
ILRI_NS_13.16	Tift_N128				
ILRI_NS_13.19	India_128_22229				

4.1.5 Genetic diversity revealed by the hierarchal cluster analysis

Hierarchical clustering with complete linkage (hclust) analysis resulted into two major clusters and up to ten sub-clusters (Fig. 6; Table 3), which is highly similar to the STRUCTURE clustering except that the genotypes identified as admixed by STRUCTURE are distributed across the different sub-clusters in this case.

Under the hierarchal cluster analysis, sub-cluster I represented 51 genotypes of these 50 were progenies (9 progenies from each ILRI NS7 and ILRI NS9, 10 progenies from each ILRI NS12 and ILRI_NS13, 4 progenies from each ILRI_NS_10 and ILRI_NS_11, 3 from ILRI_NS_4 and 1 ILRI_NS_6) and the rest were ILRI genotypes. The second sub-cluster consisted of 34 genotypes in which all of them are progeny plants (ILRI_NS_1, 10, 11, 12, 13, 3, 4, 6 and 7 with respective number of 1, 8, 2, 4, 4, 3, 6, 2, and 4). The third sub-cluster consisted of 27 of progenies from ILRI_NS_11, ILRI_NS_3, ILRI_NS_4, ILRI_NS_6 and ILRI_NS_7 in proportion of 8, 7, 2, 6 and 4 respectively. The fourth sub-cluster consisted of 25 genotypes and of these 17 were ILRI, 5 were ICRISAT and 3 BAGCE genotypes. The fifth sub-cluster consisted of 23 genotypes out of these 22 were progenies (6 ILRI_NS_3, 4 for each ILRI_NS_4 and ILRI_NS_6, 1 for each ILRI_NS_5, ILRI_NS_9 and ILRI_NS_10, 2 and 3 ILRI_NS_1 and ILRI_NS_11 respectively, and the last 1 was an ICRISAT genotype. The sixth sub-cluster consisted of 46 genotypes from BAGCE (13), ILRI (11), ICRISAT (13), USDA (6), CNPGL (2) and ILRI_NS_4 (1). This sub-cluster generally represents most of the Napier grass collections. The seventh sub-cluster consisted of 48 genotypes of these 46 were progeny plants (10 from ILRI NS 1, 2 from each ILRI NS 11 and ILRI NS 12, 7 were ILRI NS 2, 3 were ILRI_NS_3, 1 from each ILRI_NS_4, ILRI_NS_12, ILRI_NS_5, and 8 ILRI_NS_8). The remaining two were from the ILRI collection. The eighth sub-cluster consisted of 26 genotypes, of these 7 and 2 were from USDA and ICRISAT, respectively. The remaining 17 genotypes were progeny plants of ILRI_NS_5, ILRI_NS_6 and ILRI_NS_8. Sub-cluster nine consisted of 36 genotypes out of these 27 were ILRI, 6 were CNPGL, 1 was BAGCE and 2 were USDA genotypes. The last sub-cluster (subcluster ten) consisted of 31 genotypes most of these were from EMBRAPA (16 CNPGL and 1 BAGCE). The rest, four from each of the ILRI, ICRISAT and USDA collections and two from the progeny plants.

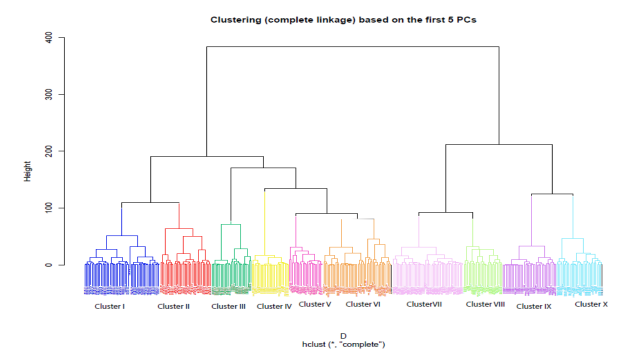


Figure 6. A dendrogram according to the hierarchal clustering based on complete linkage analysis, showing clusters and sub-clusters of the Napier grass genotypes

			Cluster I			Cluster II				
sub-cluster I	sub-cluster II	sub-cluster III	sub-cluster IV	sub-cluster V	sub-cluster VI	sub-cluster VII	sub-cluster VIII	sub-cluster IX	sub-cluster X	
ILRI_NS_9.9	ILRI_NS_1.16	ILRI_NS_11.12	BAGCE_24	ILRI_NS_5.16	BAGCE_100	ILRI_14983	ILRI_NS_1.9	BAGCE_63	BAGCE_1	
ILRI_NS_9.8	ILRI_NS_10.1	ILRI_NS_11.14	BAGCE_17	ILRI_NS_1.18	BAGCE_16	ILRI_16839	ILRI_NS_3.11	CNPGL_92.133.3	CNPGL_00.1.1	
ILRI NS 9.2	ILRI_NS_10.10	ILRI NS 11.16	BAGCE 80	ILRI NS 1.4	BAGCE 30	ILRI_Ns_1.1	ILRI NS 5.12	CNPGL 92.38.2	CNPGL 91.06.2	
ILRI_NS_9.18	ILRI_NS_10.11	ILRI_NS_11.17	 ILRI_16782	ILRI_NS_10.3	BAGCE_34	ILRI_NS_1.10	ILRI_NS_5.19	 CNPGL_9279.2	 CNPGL_91.11.2	
ILRI_NS_9.15	ILRI_NS_10.12	ILRI_NS_11.18	ILRI_16794	ILRI_NS_11.11	BAGCE_53	ILRI_NS_1.12	ILRI_NS_5.5	CNPGL_93.01.1	CNPGL_91.25.1	
ILRI_NS_9.13	ILRI_NS_10.13	ILRI_NS_11.4	ILRI_16796	ILRI_NS_11.19	BAGCE_56	ILRI_NS_1.13	ILRI_NS_5.8	CNPGL_93.32.2	CNPGL_92.190.01	
ILRI_NS_9.12	ILRI_NS_10.5	ILRI_NS_11.5	ILRI_16797	ILRI_NS_11.2	BAGCE_7	ILRI_NS_1.14	ILRI_NS_6.11	CNPGL_94.13.1	CNPGL_92.56.2	
ILRI_NS_9.10	ILRI_NS_10.6	ILRI_NS_11.8	ILRI_16805	ILRI_NS_3.13	BAGCE_75	ILRI_NS_1.15	ILRI_NS_6.13	ILRI_14355	CNPGL_92.66.3	
ILRI_NS_9.1	ILRI_NS_10_8	ILRI_NS_3.1	ILRI_16807	ILRI_NS_3.15	BAGCE_81	ILRI_NS_1.2	ILRI_NS_6.18	ILRI_14389	CNPGL_93.04.2	
ILRI_NS_7_6	ILRI_NS_11.1	ILRI_NS_3.10	ILRI_16808	ILRI_NS_3.16	BAGCE_86	ILRI_NS_1.3	ILRI_NS_6.20	ILRI_14982	CNPGL_93.06.1	
ILRI_NS_7.5	ILRI_NS_11.10	ILRI_NS_3.18	ILRI_16809	ILRI_NS_3.20	BAGCE_90	ILRI_NS_1.5	ILRI_NS_6.4	ILRI_14984	CNPGL_93.08.1	
ILRI_NS_7.4	ILRI_NS_12.11	ILRI_NS_3.2	ILRI_16810	ILRI_NS_3.4	BAGCE_94	ILRI_NS_1.6	ILRI_NS_8.14	ILRI_15357	CNPGL_93.18.2	
ILRI_NS_7.20	ILRI_NS_12.13	ILRI_NS_3.3	ILRI_16816	ILRI_NS_3.9	BAGCE_97	ILRI_NS_11.3	ILRI_NS_8.16	ILRI_16785	CNPGL_93.37.5	
ILRI_NS_7.2	ILRI_NS_12.2	ILRI_NS_3.6	ILRI_16818	ILRI_NS_4.1	CNPGL_92.198.7	ILRI_NS_11.9	ILRI_NS_8.17	ILRI_16786	CNPGL_94.07.2	
ILRI_NS_7.18	ILRI_NS_12.20	ILRI_NS_3.8	ILRI_16821	ILRI_NS_4.18	ILRI_1026	ILRI_NS_12.15	ILRI_NS_8.2	ILRI_16787	CNPGL_96.21.1	
ILRI_NS_7.15	ILRI_NS_13.11	ILRI_NS_4.13	ILRI_16822	ILRI_NS_4.6	ILRI_15743.MOTT.	ILRI_NS_12.7	ILRI_NS_8.4	ILRI_16789	CNPGL_96.23.1	
ILRI_NS_7.14	ILRI_NS_13.12	ILRI_NS_4.4	ILRI_16834	ILRI_NS_4.9	ILRI_16783	ILRI_NS_2.1	ILRI_NS_8.9	ILRI_16792	CNPGL_96.27.3	
ILRI_NS_7.1	ILRI_NS_13.17	ILRI_NS_6.10	ILRI_16838	ILRI_NS_6.1	ILRI_16788	ILRI_NS_2.2	India_129_22228	ILRI_16793	ILRI_16784	
ILRI_NS_6.19	ILRI_NS_13.7	ILRI_NS_6.12	ILRI_18448	ILRI_NS_6.16	ILRI_16791	ILRI_NS_2.3	India_145.121965	ILRI_16795	ILRI_16811	
ILRI_NS_4_15	ILRI_NS_3.14	ILRI_NS_6.14	ILRI_18662	ILRI_NS_6.6	ILRI_16802	ILRI_NS_2.6	Tift_N172	ILRI_16798	ILRI_16813	
ILRI_NS_4.11	ILRI_NS_3.17	ILRI_NS_6.15	India_123_22234	ILRI_NS_6.7	ILRI_16812	ILRI_NS_2.7	Tift_N200	ILRI_16799	ILRI_NS_12.10	
ILRI_NS_4.10	ILRI_NS_3.7	ILRI_NS_6.5	India_124_22233	ILRI_NS_9.7	ILRI_16814	ILRI_NS_2.8	Tift_N223	ILRI_16800	ILRI_NS_12.19	
ILRI_NS_13.9	ILRI_NS_4.12	ILRI_NS_6.8	India_125_22232	India_128_22229	ILRI_16815	ILRI_NS_2.9	Tift_N225	ILRI_16801	India_141_21967	
ILRI_NS_13.8	ILRI_NS_4.19	ILRI_NS_7.10	India_126_22231		ILRI_16817	ILRI_NS_3.12	Tift_N68	ILRI_16803	India_144_21964	
ILRI_NS_13.5	ILRI_NS_4.2	ILRI_NS_7.11	India_149_21785		ILRI_16835	ILRI_NS_3.19	Tift_N75	ILRI_16804	India_147_21787	
ILRI_NS_13.3	ILRI_NS_4.3	ILRI_NS_7.3			ILRI_NS_4.20	ILRI_NS_3.5	Tift_N8	ILRI_16806	India_150_21784	
ILRI_NS_13.2	ILRI_NS_4.5	ILRI_NS_7.7			India_115_22243	ILRI_NS_4.14		ILRI_16819	mott_new2	
ILRI_NS_13.19	ILRI_NS_4.7				India_116_22242	ILRI_NS_5.1		ILRI_16836	Tift_N109	
ILRI_NS_13.16	ILRI_NS_6.3				India_118_22241	ILRI_NS_5.10		ILRI_16837	Tift_N138	
ILRI_NS_13.15	ILRI_NS_6.9				India_119_22239	ILRI_NS_5.13		ILRI_16840	Tift_N23	
ILRI_NS_13.13	ILRI_NS_7.12				India_120_22238	ILRI_NS_5.14		ILRI_16902	Tift_N43	
ILRI_NS_13.1	ILRI_NS_7.13				India_121_22237	ILRI_NS_5.15		ILRI_18438		
ILRI_NS_12.8	ILRI_NS_7.17				India_127_22230	ILRI_NS_5.17		Maralfalfa.1		
ILRI_NS_12.6	ILRI_NS_7.9				India_131_22226	ILRI_NS_5.18		Napier_Addis		
ILRI_NS_12.5					India_132_22225	ILRI_NS_5.2		Tift_N131		
ILRI_NS_12.4					India_142_21966	ILRI_NS_5.3		Tift_N147		
ILRI_NS_12.3					India_146_21788	ILRI_NS_5.4				
ILRI_NS_12.18					India_151_21783	ILRI_NS_5.6				
ILRI_NS_12.17					India_89_22236	ILRI_NS_5.9				
ILRI_NS_12.14					PIONEIRO	ILRI_NS_6_2				
ILRI_NS_12.12					Tift_N128	ILRI_NS_8.10				
ILRI_NS_12.1					Tift_N130	ILRI_NS_8.11				
ILRI_NS_11_6					Tift_N137	ILRI_NS_8.15				

Table 3. The list of genotypes under different clusters and sub-clusters detected by hierarchal clustering analysis

ILRI_NS_11.20			Tift_N210	ILRI_NS_8.19		
ILRI_NS_11.15			Tift_N37	ILRI_NS_8.20		
ILRI_NS_11.13			Tift_N71	ILRI_NS_8.3		
ILRI_NS_10.9				ILRI_NS_8.7		
ILRI_NS_10.7				ILRI_NS_8.8		
ILRI_NS_10.4						
ILRI_NS_10.2						
ILRI_16790						

4.2 Population differentiation and divergence analysis

The Nei's genetic distance reflected the diversity among the Napier grass collections and progeny plants and among clusters and sub-clusters. Based on the overall dataset, the pairwise Nei's genetic distance ranged from 0.07071 for sub-clusters I and VI to 0.5118 for sub-clusters II and IX (Table 4). The six ILRI genotypes in sub-cluster II showed high similarity among themselves, with 0.005 to 0.02 range of Nei's genetic distance, probably representing potential duplicates. Analysis of molecular variance (AMOVA) was used to partition the existing genetic variation into different components. In the current study, analysis of one level of molecular variance was carried out using the R package Pegas and the number of subpopulations which were determined with STRUCTURE software were used for AMOVA analysis. Variance components obtained by AMOVA were highly significant (P < 0.00) among populations (Table 5).

Estimated Mean value of fixation index (Fst) per cluster ranged from 0.3398 in cluster VI that contain progeny plants to 0.7621 in cluster VII, which mainly contains the ILRI collection (Table 6). Divergence among populations was estimated based on allele-frequency or Net nucleotide distance and the largest divergence (0.3777) was between sub-cluster IV and VII (Table 7).

 Table 4. The Nei's genetic distance among the ten sub-clusters of Napier grass collections and progeny plants

	C1	C2	C3	C4	C5	C6	C7	C8	С9	C10
C1	0	0.20691	0.16473	0.11548	0.1695	0.07071	0.43496	0.24225	0.46183	0.18314
C2	0.20691	0	0.29624	0.11065	0.21632	0.24717	0.47953	0.29654	0.5118	0.29953
C3	0.16473	0.29624	0	0.20794	0.18506	0.15965	0.30173	0.14491	0.29771	0.11329
C4	0.11548	0.11065	0.20794	0	0.13455	0.15303	0.41527	0.21879	0.44283	0.22442
C5	0.1695	0.21632	0.18506	0.13455	0	0.17097	0.20149	0.10591	0.23945	0.16033
C6	0.07071	0.24717	0.15965	0.15303	0.17097	0	0.36253	0.19599	0.36941	0.13015
C7	0.43496	0.47953	0.30173	0.41527	0.20149	0.36253	0	0.13362	0.08839	0.21433
C8	0.24225	0.29654	0.14491	0.21879	0.10591	0.19599	0.13362	0	0.14961	0.11054
C9	0.46183	0.5118	0.29771	0.44283	0.23945	0.36941	0.08839	0.14961	0	0.19898
C10	0.18314	0.29953	0.11329	0.22442	0.16033	0.13015	0.21433	0.11054	0.19898	0

Table 5. AMOVA showing the genetic variation among the ten sub clusters (sub populations) of Napier grass accessions and progeny plants, *at 1000 number of permutations for test of hypothesis

Source of	SSD	MSD	df	Variance	Phi statistics	p.value*	CV
variation				components			
pops	13.757165	1.52857390	9	0.05969	0.70824	0.00	
Error	6.295303	0.02459103	256	0.02459			25.195
Total	20.052468	0.07566969	265				

Table 6. Mean values of Fst for the ten sub clusters detected by STRUCTURE analysis

Sub-	Sub-	Sub-	Sub-	Sub-	Sub-	Sub-	Sub-	Sub-	Sub-
cluster I	cluster II	cluster III	cluster IV	cluster V	cluster VI	cluster VII	clusterVIII	cluster IX	cluster X
0.442	0.5041	0.3951	0.6142	0.4168	0.3398	0.7621	0.4052	0.5096	0.5854

Table 7. Divergence among the ten sub-clusters based on allele-frequency or nucleotide distance computed using the STRUCTURE software

	Sub-cluster I	2	3	4	5	6	7	8	9	10
1	-	0.1481	0.144	0.1736	0.1366	0.1395	0.2651	0.1093	0.0927	0.2204
2	0.1481	-	0.1545	0.1277	0.1747	0.1228	0.3126	0.1756	0.1455	0.2624
3	0.144	0.1545	-	0.2128	0.1005	0.1015	0.1887	0.097	0.1712	0.1493
4	0.1736	0.1277	0.2128	-	0.1992	0.168	0.3777	0.2168	0.0957	0.3355
5	0.1366	0.1747	0.1005	0.1992	-	0.1354	0.2257	0.0891	0.1647	0.1851
6	0.1395	0.1228	0.1015	0.168	0.1354	-	0.1645	0.1182	0.1518	0.1244
7	0.2651	0.3126	0.1887	0.3777	0.2257	0.1645	-	0.1797	0.3146	0.1042
8	0.1093	0.1756	0.097	0.2168	0.0891	0.1182	0.1797	-	0.1528	0.1333
9	0.0927	0.1455	0.1712	0.0957	0.1647	0.1518	0.3146	0.1528	-	0.2736
10	0.2204	0.2624	0.1493	0.3355	0.1851	0.1244	0.1042	0.1333	0.2736	-

5 Discussion

5.1 Enhancement of diversity in ILRI Napier grass collection

Napier (Elephant) grass originated from the tropical region of sub-Saharan Africa and has been distributed as forage crop into most tropical and subtropical regions of the world (Clayton *et al.*, 2013). Characterization of genetic diversity in Napier grass is a prerequisite for wise and effective germplasm conservation and utilization as well as developing efficient breeding programs (Negawo *et al.*, 2018; Negawo *et al.*, 2017). The ILRI forage genebank collected and conserved a diverse set of genotypes that are very variable in genetic and phenotypic traits. However, the diversity and population size is very limiting in selecting different desirable traits for different agro-ecological conditions. Therefore, the strategies followed to increase diversity by developing progeny plants raised from seeds produced by open pollination. This will increase the chance of developing new genetic makeup through recombination and unique genotypes can be identified and incorporated into the existing collection (Nielsen *et al.*, 2014; Zhou *et al.*, 2018). Furthermore, it is crucial to assess genetic diversity in order to ensure that the most diverse populations are identified and selected to widen the genetic base of this grass.

Marker-assisted breeding in Napier grass is usually hindered due to low genetic information as this forage crop is under researched. Therefore, development and implementation of large-scale informative markers like SNPs, assist breeders in differentiating the Napier grass germplasm at a genome level (Muktar *et al.*, 2019; Zhou *et al.*, 2018). Today, different molecular markers have been utilized for genetic diversity analysis in Napier grass (Bhandari *et al.*, 2006; Muktar *et al.*, 2019; Van De Wouw *et al.*, 1999; Wanjala *et al.*, 2013; Zhou *et al.*, 2018). DArTseq marker technology is a rapid, low-cost, and efficient method for genotyping, providing a broad genome coverage, and as a result, has been increasingly used in different plant species for different purpose's (Baloch *et al.*, 2017; Mace *et al.*, 2008; Wenzl *et al.*, 2004), as well as in Napier grass (Muktar *et al.*, 2019). In this study, the genetic diversity in different Napier grass collections and progeny plants was estimated using the DArTseq SNP markers, and considerable genetic variation among populations as well as genotypes was identified. Furthermore, potential duplicates and divergent groups were detected.

5.2 Marker diversity and genome-wide distribution

In this study, 347 Napier grass genotypes and progeny plants were used to assess genetic diversity. Initially, 363 genotypes were incorporated in the study but, later 16 genotypes were excluded from

further investigation due to their high missing value percentages (≥ 50 %). The GBS method of the DArT-seq markers was utilized to investigate genetic variability and differentiation within and among the 347 Napier grass collection and progeny plants. A total of 96,454 SilicoDArT and 96,321 SNP markers were generated, of which 1001 highly informative SNPs were selected for the diversity study. Previously, since Napier grass reference genome sequence was not generated, the closely related pearl millet (*Cenchrus americanus*) genome was used to identify the genomic position and genome-wide distribution of the SilicoDArT and SNP markers (Muktar *et al.*, 2019). In this study, the markers were mapped onto the new reference genome sequence of Napier grass, in which most of the markers were able to be mapped with a higher precision.

Expected heterozygosity and PIC of the DArTseq generated markers ranged from 0 to 0.5 and 0 to 0.38 for both silicoDArT and SNP markers, with the average He =0.26 and 0.18 and PIC = 0.21 and 0.15, respectively. The results of the markers quality parameters were comparable with that of other species. The average PIC values of the generated SilicoDArT and SNP markers was similar with Lesquerella and related species (0.21) (Cruz *et al.*, 2013) and lower than that of values identified in DArT markers of sorghum (0.41) (Emma *et al.*, 2008) and wheat (0.44) (Mona *et al.*, 2006) also, similar with PIC value (0.212) of SSR markers that were originally developed for pearl millet and used to asses genetic diversity in Napier grass due to the cross-species transferability of microsatellite markers (Kandel *et al.*, 2016) and slightly greater than the previous report made on Napier grass using a similar marker platform (Muktar *et al.*, 2019).

According to Botstein *et al.* (1980), the PIC values can be classified into highly informative (PIC value > 0.5), moderately informative (PIC value $0.25 \le 0.5$) and slightly informative (PIC value ≤ 0.25) for multi allelic markers, such as SSR. However, the PIC value of SNP markers is restricted to the extreme PIC values of 0.5 (when the two alleles have the same frequencies) due to the bi-allelic nature of the SNP markers. The advantage of SNP markers is their abundance in the genome and their distribution across the genome, which has been observed in the Napier grass draft genome of the markers used in this study. The obtained result was also consistent with the previous report by Muktar *et al.* (2019) while a slightly higher average in He and PIC values were obtained in the current study indicating high marker polymorphism and distribution that could be attributed to the sample size difference of the population in this study.

5.3 Genetic diversity among Napier grass

The revealed patterns of genetic diversity were interesting in which STRUCTURE software identified two major classes and ten sub-clusters of the assayed genotypes (Fig. 4a and 4b). Moreover, the result from hierarchal clustering in complete linkage was in accordance with result from structure analysis (Fig. 6). The DAPC analysis resulted in ten K groups (clusters) indicating also highly consistent results with STRUCTURE and Hierarchal clustering for genetic diversity analysis. In addition, AMOVA showed that the two major groups and the ten subgroups detected were significantly different from each other. The high level of diversity was observed and this high variation, and population stratification could be attributed to the outcrossing and self-incompatibility nature of Napier grass (Souza *et al.*, 2019). Variation in geographic origin of the accession might have also contributed to the high genetic diversity, since the accessions were collected from different parts of the world. Selection and breeding system were additional factors with the possibility of their contribution for the genetic variation. However, in Napier grass genetic contribution of genetic drift and gene flow on genetic variation is expected to be low since this grass is mostly propagated through stem cutting, and due to its low seed germination rate (Wanjala *et al.*, 2013; Xie *et al.*, 2009).

In this study, half of the progenies and most of the BAGCE accessions with some accessions from ICRISAT and ILRI were clustered under the main cluster one while most of ILRI collections, CNPGL (EMBRAPA elite lines), USDA collections and half of the progenies also clustered under the second main cluster. Most of the progenies clustered under sub-cluster I, III, VI, and X while most of the ILRI collections were clustered under sub-cluster II, IV, VII, and IX. The majority of the CNPGL and USDA collections were clustered under sub-cluster VIII while most of the BAGCE collections clustered under sub-cluster V. In some cases, progenies with the same maternal plants and clustered under more than one sub-cluster, implying they had different pollen sources. However, mostly the clustering of the genotypes did not seem to be based on the geographical origin which is also consistent with the findings by Negawo *et al.* (2018); Kandel *et al.* (2016); and Muktar *et al.* (2019); while it is in contrast with reports made by Harris *et al.* (2009) and Lowe *et al.* (2003).

5.3.1 Population differentiation and genetic divergence

According to Weising *et al.* (2005), the extent of genetic variation in a species and its distribution among and within populations is determined by the interactions of various factors, including the evolutionary history of the species, genetic drift, mating system, gene flow, mutation, and selection. In general, outcrossing species retain most of their variation within populations whereas selfing species allocate more variation among populations. For DNA based markers, among population genetic differentiation is often estimated according to Nei's (1972) genetic distance. However, AMOVA is nowadays even more widely used for the partitioning of genetic variation among populations and subpopulations (Excoffier et al. 1992). In accordance with this, the one level AMOVA of the current study estimated the variance among the sub populations and high levels of genetic differentiation among the sub populations were revealed. Fixation index (Fst) is also another important parameter for differentiation of populations and subpopulations and measures the degree of differentiation among populations in terms of allele frequencies. The values of Fst can be grouped into four categories with respect to genetic differentiation; very high (Fst > 0.25), high (0.15 - 0.25), intermediate (0.05 - 0.15), and low (0.0 - 0.05) (De Vicente and Fulton, 2003). In this study, the estimate of population differentiation in Napier grass collections and progeny plants using fixation index (Fst) revealed a relatively high level of differentiation among the populations. Mean fixation index (Fst) per cluster of the current study ranged from 0.3398 in cluster VI, that contains entirely progeny plants, to 0.7621 in cluster VII that consisted of most of the ILRI accessions, some EMBRAPA elite lines and the USDA collections. More differentiation was observed among progeny plants and the accessions and this could be attributed to the outcrossing nature of Napier grass (Souza et al., 2019).

From allele-frequency or Net nucleotide distance, the highest divergent result (0.3777) was between populations of sub-cluster IV (which consisted mostly of ILRI accessions with a few ICRISAT and EMBRAPA's accessions) and VII (which consisted of two ILRI accessions and 46 progeny plants), indicating the divergence between the accessions and progenies and showing the potential of progeny plants, that were produced from open pollination, for enhancement of diversity of the ILRI Napier grass collection with potential for future utilization in breeding programs. Most of the introduced accessions clustered away from the ILRI collections (Table. 3) hence increase the genetic base for future use.

Six potential duplicate genotypes are also identified in the ILRI collection and the accessions showed high genetic similarity among themselves, with 0.005 to 0.02 range of Nei's genetic distance. Identification of duplicate or redundant genotypes is important especially for the genebank to reduce the cost of conservation.

6 Conclusions and Recommendations

6.1 Conclusions

In this study, a large number of SNP and SilicoDArT markers with high distribution throughout the genome and high polymorphism were generated using the GBS method of the DArTseq genotyping platform. The markers have valuable information for genome wide identification of genetic diversity in Napier grass. The DArTseq-SNP markers were employed for genetic diversity and population structure analysis in Napier grass collections from different institutes EMBRAPA, USDA, ICRISAT, and ILRI gene-bank and progenies raised from 13 ILRI accessions. The genetic diversity using the molecular markers identified high genetic variation and detected potential duplicates and unique genotypes to enhance Napier grass diversity in the ILRI collection. Therefore, this genetic diversity information is valuable for various purposes, including germplasm conservation, wise utilization, and implementation of effective breeding programs.

Moreover, Results from the diversity parameters showed that the presence of up to ten sub populations with considerable variation among the sub populations. Sub-population IV and subpopulation VII were also found as the most divergent and genotypes from these sub-populations could be used in a Napier grass heterotic breeding program as potential parental plants. In general, the results of this study indicate the suitability of the populations in the future molecular genetic studies in Napier grass.

6.2 Recommendations

Based on the findings obtained in this study, the following recommendations were forwarded.

- Napier grass breeding programs should focus on the divergent genotypes from progenies and accessions and also from distantly related populations of ILRI and USDA accessions. The unique genotypes from the progeny and other collections should be incorporated into the existing collection.
- The ILRI accession from sub-cluster II showed high similarity/low genetic diversity and hence, attention should be given to characterize these genotypes and to avoid redundant genotypes
- Characterization of more populations is also very important and an in-depth study should be made in the future including quantitative trait loci (QTL) mapping and genome wide association mapping.

7 References

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Appendices

Appendix 1: (Supplementary Table S1) List of Napier grass genotypes that were used in the study. The 13 ILRI accessions that were used to raise progenies and the progenies raised from them indicated by the same highlighted color with respect to their progenies. The 16 genotypes that were excluded from further analysis are highlighted in yellow.

						year
No.	Acc. No.	Genus	Species/ cross	Origin	Collection	acquired
1	ILRI_1026	Cenchrus	purpureus	Burundi	ILRI	1986
2	ILRI_14355	Cenchrus	purpureus	Ethiopia	ILRI	1985
3	ILRI_14389	Cenchrus	purpureus	Nigeria	ILRI	1985
4	ILRI_14982	Cenchrus	purpureus x glaucum	USA	ILRI	1986
5	ILRI_14983	Cenchrus	purpureus	USA	ILRI	1986
6	ILRI_14984	Cenchrus	purpureus	USA	ILRI	1986
7	ILRI_15357	Cenchrus	purpureus x glaucum	NA	ILRI	1986
8	ILRI_15743	Cenchrus	purpureus	USA	ILRI	1988
9	ILRI_16621	Cenchrus	purpureus	Namibia	ILRI	1991
10	ILRI_16782	Cenchrus	purpureus	Tanzania	ILRI	1992
11	ILRI_16783	Cenchrus	purpureus	Tanzania	ILRI	1992
12	ILRI_16784	Cenchrus	purpureus	Tanzania	ILRI	1992
13	ILRI_16785	Cenchrus	purpureus	Tanzania	ILRI	1992
14	ILRI_16786	Cenchrus	purpureus	Swaziland	ILRI	1992
15	ILRI_16787	Cenchrus	purpureus	Swaziland	ILRI	1992
16	ILRI_16788	Cenchrus	purpureus	Swaziland	ILRI	1992
17	ILRI_16789	Cenchrus	purpureus	Swaziland	ILRI	1992
18	ILRI_16791	Cenchrus	purpureus	Swaziland	ILRI	1992
19	ILRI_16792	Cenchrus	purpureus	Mozambique	ILRI	1992
20	ILRI_16793	Cenchrus	purpureus	Cuba	ILRI	1992

21	ILRI_16794	Cenchrus	purpureus	Mozambique	ILRI	1992
22	ILRI_16795	Cenchrus	purpureus	Zimbabwe	ILRI	1992
23	ILRI_16796	Cenchrus	purpureus	Zimbabwe	ILRI	1992
24	ILRI_16797	Cenchrus	purpureus	Zimbabwe	ILRI	1992
25	ILRI_16798	Cenchrus	purpureus	Zimbabwe	ILRI	1992
26	ILRI_16799	Cenchrus	purpureus	Zimbabwe	ILRI	1992
27	ILRI_16800	Cenchrus	purpureus	Zimbabwe	ILRI	1992
28	ILRI_16801	Cenchrus	purpureus	Zimbabwe	ILRI	1992
29	ILRI_16802	Cenchrus	purpureus	Zimbabwe	ILRI	1992
30	ILRI_16803	Cenchrus	purpureus	Zimbabwe	ILRI	1992
31	ILRI_16804	Cenchrus	purpureus	USA	ILRI	1992
32	ILRI_16805	Cenchrus	purpureus	USA	ILRI	1992
33	ILRI_16806	Cenchrus	purpureus	USA	ILRI	1992
34	ILRI_16807	Cenchrus	purpureus	USA	ILRI	1992
35	ILRI_16808	Cenchrus	purpureus	USA	ILRI	1992
36	ILRI_16809	Cenchrus	purpureus	USA	ILRI	1992
37	ILRI_16810	Cenchrus	purpureus	USA	ILRI	1992
38	ILRI_16811	Cenchrus	purpureus	USA	ILRI	1992
39	ILRI_16812	Cenchrus	purpureus	USA	ILRI	1992
40	ILRI_16813	Cenchrus	purpureus	USA	ILRI	1992
41	ILRI_16814	Cenchrus	purpureus	USA	ILRI	1992
42	ILRI_16815	Cenchrus	purpureus	USA	ILRI	1992
43	ILRI_16816	Cenchrus	purpureus	USA	ILRI	1992
44	ILRI_16817	Cenchrus	purpureus	USA	ILRI	1992
45	ILRI_16818	Cenchrus	purpureus	USA	ILRI	1992
46	ILRI_16819	Cenchrus	purpureus	USA	ILRI	1992
47	ILRI_16821	Cenchrus	purpureus	Zimbabwe	ILRI	1992
48	ILRI_16822	Cenchrus	purpureus	Malawi	ILRI	1992
49	ILRI_16834	Cenchrus	purpureus x glaucum	Zimbabwe	ILRI	1992
50	ILRI_16835	Cenchrus	purpureus x glaucum	Zimbabwe	ILRI	1992
51	ILRI_16836	Cenchrus	purpureus	Zimbabwe	ILRI	1992

52	ILRI_16837	Cenchrus	purpureus x glaucum	Zimbabwe	ILRI	1992
53	ILRI_16838	Cenchrus	purpureus x glaucum	Zimbabwe	ILRI	1992
54	ILRI_16839	Cenchrus	purpureus	Zimbabwe	ILRI	1992
55	ILRI_16840	Cenchrus	purpureus x glaucum	Zimbabwe	ILRI	1992
56	ILRI_16902	Cenchrus	purpureus	Zimbabwe	ILRI	1992
57	ILRI_16790	Cenchrus	purpureus	Swaziland	ILRI	1992
58	ILRI_18438	Cenchrus	purpureus	Tanzania	ILRI	1995
59	ILRI_18448	Cenchrus	purpureus	Tanzania	ILRI	1995
60	ILRI_18662	Cenchrus	purpureus x glaucum	South_Africa	ILRI	2006
61	BAGCE-1	Cenchrus	purpureus	Colombia	EMBRAPA_collection	1976
62	BAGCE-100	Cenchrus	purpureus	Brazil	EMBRAPA_collection	1993
63	BAGCE-16	Cenchrus	purpureus	Brazil	EMBRAPA_collection	1976
64	BAGCE-17	Cenchrus	purpureus	Costa Rica	EMBRAPA_collection	1976
65	BAGCE-22	Cenchrus	purpureus	NA	EMBRAPA_collection	1976
66	BAGCE-24	Cenchrus	purpureus	NA	EMBRAPA_collection	1976
67	BAGCE-25	Cenchrus	purpureus	India	EMBRAPA_collection	1976
68	BAGCE-30	Cenchrus	purpureus	Brazil	EMBRAPA_collection	1976
69	BAGCE-343	Cenchrus	purpureus	Brazil	EMBRAPA_collection	1976
70	BAGCE-53	Cenchrus	purpureus	Brazil	EMBRAPA_collection	1976
71	BAGCE-56	Cenchrus	purpureus	Brazil	EMBRAPA_collection	1989
72	BAGCE-63	Cenchrus	purpureus	Cuba	EMBRAPA_collection	1991
73	BAGCE-7	Cenchrus	purpureus	Brazil	EMBRAPA_collection	1976
74	BAGCE-75	Cenchrus	purpureus	Brazil	EMBRAPA_collection	1992
75	BAGCE-80	Cenchrus	purpureus	Brazil	EMBRAPA_collection	1992
76	BAGCE-81	Cenchrus	purpureus	Brazil	EMBRAPA_collection	1992
77	BAGCE-86	Cenchrus	purpureus	NA	EMBRAPA_collection	1992
78	BAGCE-90	Cenchrus	purpureus	NA	EMBRAPA_collection	NA
79	BAGCE-94	Cenchrus	purpureus	NA	EMBRAPA_collection	1993
80	BAGCE-97	Cenchrus	purpureus	NA	EMBRAPA_collection	1993
81	CNPGL_00-1-1	Cenchrus	purpureus	NA	EMBRAPA_elite_lines	NA
82	CNPGL_91-06-2	Cenchrus	purpureus	NA	EMBRAPA_elite_lines	NA

83	CNPGL_91-112	Cenchrus	purpureus	NA	EMBRAPA_elite_lines	NA
84	CNPGL_91-25-1	Cenchrus	purpureus	NA	EMBRAPA_elite_lines	NA
85	CNPGL_92-133-3	Cenchrus	purpureus	NA	EMBRAPA_elite_lines	NA
86	CNPGL_92-198-7	Cenchrus	purpureus	NA	EMBRAPA_elite_lines	NA
87	CNPGL_92-190-1	Cenchrus	purpureus	NA	EMBRAPA_elite_lines	NA
88	CNPGL_92-38-2	Cenchrus	purpureus	NA	EMBRAPA_elite_lines	NA
89	CNPGL_92-56-2	Cenchrus	purpureus	NA	EMBRAPA_elite_lines	NA
90	CNPGL_92-66-3	Cenchrus	purpureus	NA	EMBRAPA_elite_lines	NA
91	CNPGL_9279-2	Cenchrus	purpureus	NA	EMBRAPA_elite_lines	NA
92	CNPGL_93-01-1	Cenchrus	purpureus	NA	EMBRAPA_elite_lines	NA
93	CNPGL_93-04-2	Cenchrus	purpureus	NA	EMBRAPA_elite_lines	NA
94	CNPGL_93-06-1	Cenchrus	purpureus	NA	EMBRAPA_elite_lines	NA
95	CNPGL_93-08-1	Cenchrus	purpureus	NA	EMBRAPA_elite_lines	NA
96	CNPGL_93-18-2	Cenchrus	purpureus	NA	EMBRAPA_elite_lines	NA
97	CNPGL_93-32-2	Cenchrus	purpureus	NA	EMBRAPA_elite_lines	NA
98	CNPGL_9337-5	Cenchrus	purpureus	NA	EMBRAPA_elite_lines	NA
99	CNPGL_94-07-2	Cenchrus	purpureus	NA	EMBRAPA_elite_lines	NA
100	CNPGL_94-13-1	Cenchrus	purpureus	NA	EMBRAPA_elite_lines	NA
101	CNPGL_96-21-1	Cenchrus	purpureus	NA	EMBRAPA_elite_lines	NA
102	CNPGL_96-23-1	Cenchrus	purpureus	NA	EMBRAPA_elite_lines	NA
103	CNPGL_96-24-1	Cenchrus	purpureus	NA	EMBRAPA_elite_lines	NA
104	CNPGL_96-27-3	Cenchrus	purpureus	NA	EMBRAPA_elite_lines	NA
105	PIONEIRO	Cenchrus	purpureus	NA	EMBRAPA_elite_lines	NA
106	N 8	Cenchrus	purpureus K-12	PUERTO RICO	USDA	1975
107	N 19	Cenchrus	purpureus	PUERTO RICO	USDA	NA
108	N 23	Cenchrus	purpureus	SWAZILAND	USDA	NA
109	N 36	Cenchrus	purpureus	SCHANK	USDA	NA
110	N 37	Cenchrus	purpureus	SCHANK	USDA	NA
111	N 43	Cenchrus	purpureus	NA	USDA	NA
112	N 68	Cenchrus	purpureus	SCHANK	USDA	NA
113	N 71	Cenchrus	purpureus	SCHANK	USDA	NA

114	N 75	Cenchrus	purpureus	NA	USDA	1977
115	N109	Cenchrus	Purpureus (SPAIN NAPIER)	NA	USDA	1979
116	N128	Cenchrus	81-D62-1 (Dwarf Napier plant)	NA	USDA	NA
117	N130	Cenchrus	Purpureus (NB21 Museum Plot)		USDA	NA
118	N131	Cenchrus	purpureus	NA	USDA	NA
119	N137	Cenchrus	purpureus	BATORE	USDA	NA
120	N138	Cenchrus	purpureus	BATORE	USDA	NA
121	N147	Cenchrus	purpureus	NA	USDA	1983
122	N172	Cenchrus	purpureus	NA	USDA	NA
123	N200	Cenchrus	purpureus	NA	USDA	F2_1986
124	N210	Cenchrus	purpureus	NA	USDA	F2_1986
125	N223	Cenchrus	purpureus	NA	USDA	1988
126	N225	Cenchrus	purpureus	NA	USDA	1988
127	N228	Cenchrus	purpureus	NA	USDA	1988
128	145_1_21965	Cenchrus	purpureus	NA	ICRISAT	2019
129	120_22238	Cenchrus	purpureus	NA	ICRISAT	2019
130	127_22230	Cenchrus	purpureus	NA	ICRISAT	2019
131	117_22240	Cenchrus	purpureus	NA	ICRISAT	2019
132	149_21785	Cenchrus	purpureus	NA	ICRISAT	2019
133	126_22231	Cenchrus	purpureus	NA	ICRISAT	2019
134	150_21784	Cenchrus	purpureus	NA	ICRISAT	2019
135	118_22241	Cenchrus	purpureus	NA	ICRISAT	2019
136	128_22229	Cenchrus	purpureus	NA	ICRISAT	2019
137	124_22233	Cenchrus	purpureus	NA	ICRISAT	2019
138	125_22232	Cenchrus	purpureus	NA	ICRISAT	2019
139	115_22243	Cenchrus	purpureus	NA	ICRISAT	2019
140	119_22239	Cenchrus	purpureus	NA	ICRISAT	2019
141	146_21788	Cenchrus	purpureus	NA	ICRISAT	2019
142	132_22225	Cenchrus	purpureus	NA	ICRISAT	2019
143	144_21964	Cenchrus	purpureus	NA	ICRISAT	2019
144	141_21967	Cenchrus	purpureus	NA	ICRISAT	2019

145	142_21966	Cenchrus	purpureus	NA	ICRISAT	2019
146	131_22226	Cenchrus	purpureus	NA	ICRISAT	2019
147	147_21787	Cenchrus	purpureus	NA	ICRISAT	2019
148	151_21783	Cenchrus	purpureus	NA	ICRISAT	2019
149	121_22237	Cenchrus	purpureus	NA	ICRISAT	2019
150	129_2228	Cenchrus	purpureus	NA	ICRISAT	2019
151	123_22234	Cenchrus	purpureus	NA	ICRISAT	2019
152	89_22236	Cenchrus	purpureus	NA	ICRISAT	2019
153	136_21968	Cenchrus	purpureus	NA	ICRISAT	2019
154	116_22242	Cenchrus	purpureus	NA	ICRISAT	2019
155	G1 (Gaint Napier)	Cenchrus	purpureus	NA	ICRISAT	2019
156	Ns 1-1	Cenchrus	purpureus	ILRI_collection_collection	ILRI	2019
157	NS 1_2	Cenchrus	purpureus	ILRI_collection	ILRI	2019
158	NS 1_3	Cenchrus	purpureus	ILRI_collection	ILRI	2019
159	NS 1_4	Cenchrus	purpureus	ILRI_collection	ILRI	2019
160	NS 1_5	Cenchrus	purpureus	ILRI_collection	ILRI	2019
161	NS 1_6	Cenchrus	purpureus	ILRI_collection	ILRI	2019
162	NS 1_9	Cenchrus	purpureus	ILRI_collection	ILRI	2019
163	NS 1_10	Cenchrus	purpureus	ILRI_collection	ILRI	2019
164	NS 1_11	Cenchrus	purpureus	ILRI_collection	ILRI	2019
165	NS 1_12	Cenchrus	purpureus	ILRI_collection	ILRI	2019
166	NS 1_13	Cenchrus	purpureus	ILRI_collection	ILRI	2019
167	NS 1_14	Cenchrus	purpureus	ILRI_collection	ILRI	2019
168	NS 1_15	Cenchrus	purpureus	ILRI_collection	ILRI	2019
169	NS 1_16	Cenchrus	purpureus	ILRI_collection	ILRI	2019
170	NS 1_18	Cenchrus	purpureus	ILRI_collection	ILRI	2019
171	NS 2_1	Cenchrus	purpureus	ILRI_collection	ILRI	2019
172	NS 2_2	Cenchrus	purpureus	ILRI_collection	ILRI	2019
173	NS 2_3	Cenchrus	purpureus	ILRI_collection	ILRI	2019
174	NS 2_6	Cenchrus	purpureus	ILRI_collection	ILRI	2019
175	NS 2_7	Cenchrus	purpureus	ILRI_collection	ILRI	2019

176	NS 2_8	Cenchrus	purpureus	ILRI_collection	ILRI	2019
177	NS 2_9	Cenchrus	purpureus	ILRI_collection	ILRI	2019
178	NS 3_1	Cenchrus	purpureus	ILRI_collection	ILRI	2019
179	NS 3_2	Cenchrus	purpureus	ILRI_collection	ILRI	2019
180	NS 3-3	Cenchrus	purpureus	ILRI_collection	ILRI	2019
181	NS 3-4	Cenchrus	purpureus	ILRI_collection	ILRI	2019
182	NS 3-5	Cenchrus	purpureus	ILRI_collection	ILRI	2019
183	NS 3-6	Cenchrus	purpureus	ILRI_collection	ILRI	2019
184	NS 3-7	Cenchrus	purpureus	ILRI_collection	ILRI	2019
185	NS 3-8	Cenchrus	purpureus	ILRI_collection	ILRI	2019
186	NS 3-9	Cenchrus	purpureus	ILRI_collection	ILRI	2019
187	NS 3-10	Cenchrus	purpureus	ILRI_collection	ILRI	2019
188	NS 3-11	Cenchrus	purpureus	ILRI_collection	ILRI	2019
189	NS 3-12	Cenchrus	purpureus	ILRI_collection	ILRI	2019
190	NS 3-13	Cenchrus	purpureus	ILRI_collection	ILRI	2019
	NS 3-14	Cenchrus	purpureus	ILRI_collection	ILRI	2019
192		Cenchrus	purpureus	ILRI_collection	ILRI	2019
193	NS3-16	Cenchrus	purpureus	ILRI_collection	ILRI	2019
	NS 3-17	Cenchrus	purpureus	ILRI_collection	ILRI	2019
195	NS 3-18	Cenchrus	purpureus	ILRI_collection	ILRI	2019
196	NS 3-19	Cenchrus	purpureus	ILRI_collection	ILRI	2019
197	NS 3-20	Cenchrus	purpureus	ILRI_collection	ILRI	2019
198	NS 4-1	Cenchrus	purpureus	ILRI_collection	ILRI	2019
199	NS 4-2	Cenchrus	purpureus	ILRI_collection	ILRI	2019
200	NS 4-3	Cenchrus	purpureus	ILRI_collection	ILRI	2019
201	NS 4-4	Cenchrus	purpureus	ILRI_collection	ILRI	2019
202	NS 4-5	Cenchrus	purpureus	ILRI_collection	ILRI	2019
203	NS 4-6	Cenchrus	purpureus	ILRI_collection	ILRI	2019
204	NS 4-7	Cenchrus	purpureus	ILRI_collection	ILRI	2019
205	NS 4-9	Cenchrus	purpureus	ILRI_collection	ILRI	2019
206	NS 4-10	Cenchrus	purpureus	ILRI_collection	ILRI	2019

207	NS 4-11	Cenchrus	purpureus	ILRI_collection	ILRI	2019
208	NS 4-12	Cenchrus	purpureus	ILRI_collection	ILRI	2019
209	NS 4-13	Cenchrus	purpureus	ILRI_collection	ILRI	2019
210	NS 4-14	Cenchrus	purpureus	ILRI_collection	ILRI	2019
211	NS 4_15	Cenchrus	purpureus	ILRI_collection	ILRI	2019
212	NS 4-18	Cenchrus	purpureus	ILRI_collection	ILRI	2019
213	NS 4- 19	Cenchrus	purpureus	ILRI_collection	ILRI	2019
214	NS 4-20	Cenchrus	purpureus	ILRI_collection	ILRI	2019
215	NS 5-1	Cenchrus	purpureus	ILRI_collection	ILRI	2019
216	NS 5-2	Cenchrus	purpureus	ILRI_collection	ILRI	2019
217	NS 5-3	Cenchrus	purpureus	ILRI_collection	ILRI	2019
218	NS 5 -4	Cenchrus	purpureus	ILRI_collection	ILRI	2019
219	NS 5-5	Cenchrus	purpureus	ILRI_collection	ILRI	2019
220	NS 5-6	Cenchrus	purpureus	ILRI_collection	ILRI	2019
221	NS 5-8	Cenchrus	purpureus	ILRI_collection	ILRI	2019
222	NS 5-9	Cenchrus	purpureus	ILRI_collection	ILRI	2019
223	NS 5-10	Cenchrus	purpureus	ILRI_collection	ILRI	2019
224	NS 5-12	Cenchrus	purpureus	ILRI_collection	ILRI	2019
225	NS 5-13	Cenchrus	purpureus	ILRI_collection	ILRI	2019
226	NS 5-14	Cenchrus	purpureus	ILRI_collection	ILRI	2019
227	NS 5-15	Cenchrus	purpureus	ILRI_collection	ILRI	2019
228	NS 5-16	Cenchrus	purpureus	ILRI_collection	ILRI	2019
229	NS 5-17	Cenchrus	purpureus	ILRI_collection	ILRI	2019
230	NS 5-18	Cenchrus	purpureus	ILRI_collection	ILRI	2019
231	NS 5-19	Cenchrus	purpureus	ILRI_collection	ILRI	2019
232	NS 6-1	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
233	NS 6_2	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
234	NS 6-3	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
235	NS 6-4	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
236	NS 6- 5	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
237	NS 6-6	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019

238	NS 6-7	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
239	NS 6-8	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
240	NS 6-9	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
241	NS 6-10	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
242	NS 6-11	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
243	NS 6-12	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
244	NS 6-13	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
245	NS 6-14	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
246	NS 6-15	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
247	NS 6-16	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
248	NS 6-18	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
249	NS 6-19	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
250	NS 6-20	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
251	NS 7-1	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
252	NS 7-2	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
253	NS 7-3	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
254	NS 7-4	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
255	NS 7-5	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
256	NS 7_6	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
257	NS 7-7	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
258	NS 7-8	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
259	NS 7-9	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
260	NS 7-10	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
261	NS 7-11	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
262	NS 7-12	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
263	NS 7-13	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
264	NS 7-14	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
265	NS 7-15	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
266	NS 7-17	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
267	NS 7-18	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019
268	NS 7-20	Cenchrus	purpureus x glaucum	ILRI_collection	ILRI	2019

269	NS 8_2	Cenchrus	purpureus	ILRI_collection	ILRI	2019
270	NS 8_3	Cenchrus	purpureus	ILRI_collection	ILRI	2019
271	NS 8_4	Cenchrus	purpureus	ILRI_collection	ILRI	2019
272	NS 8_7	Cenchrus	purpureus	ILRI_collection	ILRI	2019
273	NS 8_8	Cenchrus	purpureus	ILRI_collection	ILRI	2019
274	NS 8_9	Cenchrus	purpureus	ILRI_collection	ILRI	2019
275	NS 8_10	Cenchrus	purpureus	ILRI_collection	ILRI	2019
276	NS 8_11	Cenchrus	purpureus	ILRI_collection	ILRI	2019
277	NS 8_14	Cenchrus	purpureus	ILRI_collection	ILRI	2019
278	NS 8_15	Cenchrus	purpureus	ILRI_collection	ILRI	2019
279	NS 8-16	Cenchrus	purpureus	ILRI_collection	ILRI	2019
280	NS 8-17	Cenchrus	purpureus	ILRI_collection	ILRI	2019
281	NS 8-19	Cenchrus	purpureus	ILRI_collection	ILRI	2019
282	NS 8-20	Cenchrus	purpureus	ILRI_collection	ILRI	2019
283	NS 9-1	Cenchrus	purpureus	ILRI_collection	ILRI	2019
284	NS 9-2	Cenchrus	purpureus	ILRI_collection	ILRI	2019
285	NS 9-7	Cenchrus	purpureus	ILRI_collection	ILRI	2019
286	NS 9-8	Cenchrus	purpureus	ILRI_collection	ILRI	2019
287	NS 9-9	Cenchrus	purpureus	ILRI_collection	ILRI	2019
288	NS9-10	Cenchrus	purpureus	ILRI_collection	ILRI	2019
289	NS 9-12	Cenchrus	purpureus	ILRI_collection	ILRI	2019
290	NS 9-13	Cenchrus	purpureus	ILRI_collection	ILRI	2019
291	NS 9-15	Cenchrus	purpureus	ILRI_collection	ILRI	2019
292	NS 9-18	Cenchrus	purpureus	ILRI_collection	ILRI	2019
293	NS 10-1	Cenchrus	purpureus	ILRI_collection	ILRI	2019
294	NS 10-2	Cenchrus	purpureus	ILRI_collection	ILRI	2019
295	NS 10-3	Cenchrus	purpureus	ILRI_collection	ILRI	2019
296	NS 10-4	Cenchrus	purpureus	ILRI_collection	ILRI	2019
297	NS 10-5	Cenchrus	purpureus	ILRI_collection	ILRI	2019
298	NS 10-6	Cenchrus	purpureus	ILRI_collection	ILRI	2019
299	NS 10-7	Cenchrus	purpureus	ILRI_collection	ILRI	2019

300	NS 10_8	Cenchrus	purpureus	ILRI_collection	ILRI	2019
301	NS 10-9	Cenchrus	purpureus	ILRI_collection	ILRI	2019
302	NS 10-10	Cenchrus	purpureus	ILRI_collection	ILRI	2019
303	NS 10-11	Cenchrus	purpureus	ILRI_collection	ILRI	2019
304	NS 10_12	Cenchrus	purpureus	ILRI_collection	ILRI	2019
305	NS 10-13	Cenchrus	purpureus	ILRI_collection	ILRI	2019
306	NS 11- 1	Cenchrus	purpureus	ILRI_collection	ILRI	2019
307	NS 11-2	Cenchrus	purpureus	ILRI_collection	ILRI	2019
308	NS 11-3	Cenchrus	purpureus	ILRI_collection	ILRI	2019
309	NS 11_4	Cenchrus	purpureus	ILRI_collection	ILRI	2019
310	NS 11-5	Cenchrus	purpureus	ILRI_collection	ILRI	2019
311	NS 11_6	Cenchrus	purpureus	ILRI_collection	ILRI	2019
312	NS 11-8	Cenchrus	purpureus	ILRI_collection	ILRI	2019
313	NS 11-9	Cenchrus	purpureus	ILRI_collection	ILRI	2019
314	NS 11_10	Cenchrus	purpureus	ILRI_collection	ILRI	2019
315	NS 11-11	Cenchrus	purpureus	ILRI_collection	ILRI	2019
316	NS 11_12	Cenchrus	purpureus	ILRI_collection	ILRI	2019
317	NS 11-13	Cenchrus	purpureus	ILRI_collection	ILRI	2019
318	NS 11_14	Cenchrus	purpureus	ILRI_collection	ILRI	2019
319	NS 11-15	Cenchrus	purpureus	ILRI_collection	ILRI	2019
320	NS 11-16	Cenchrus	purpureus	ILRI_collection	ILRI	2019
321	NS 11_17	Cenchrus	purpureus	ILRI_collection	ILRI	2019
322	NS 11-18	Cenchrus	purpureus	ILRI_collection	ILRI	2019
323	NS 11-19	Cenchrus	purpureus	ILRI_collection	ILRI	2019
324	NS 11-20	Cenchrus	purpureus	ILRI_collection	ILRI	2019
325	NS 12-1				ILRI	2019
326					ILRI	2019
327	NS 12-3				ILRI	2019
328	NS 12-4				ILRI	2019
329	NS 12-5				ILRI	2019
330	NS 12-6				ILRI	2019

331	NS 12-7	Cenchrus	purpureus	ILRI_collection	ILRI	2019
332					ILRI	2019
333					ILRI	2019
334					ILRI	2019
335					ILRI	2019
336					ILRI	2019
337					ILRI	2019
338					ILRI	2019
339					ILRI	2019
340					ILRI	2019
341					ILRI	2019
342					ILRI	2019
343	NS 13-1	Cenchrus	purpureus	ILRI_collection	ILRI	2019
344	NS 13-2	Cenchrus	purpureus	ILRI_collection	ILRI	2019
345	NS13-3	Cenchrus	purpureus	ILRI_collection	ILRI	2019
346	NS 13-4	Cenchrus	purpureus	ILRI_collection	ILRI	2019
347	NS 13-5	Cenchrus	purpureus	ILRI_collection	ILRI	2019
348	NS 13_6	Cenchrus	purpureus	ILRI_collection	ILRI	2019
349	NS 13-7	Cenchrus	purpureus	ILRI_collection	ILRI	2019
350	NS 13-8	Cenchrus	purpureus	ILRI_collection	ILRI	2019
351	NS 13-9	Cenchrus	purpureus	ILRI_collection	ILRI	2019
352	NS 13-10	Cenchrus	purpureus	ILRI_collection	ILRI	2019
353	NS 13-11	Cenchrus	purpureus	ILRI_collection	ILRI	2019
354	NS 13-12	Cenchrus	purpureus	ILRI_collection	ILRI	2019
355	NS 13-13	Cenchrus	purpureus	ILRI_collection	ILRI	2019
356	NS 13-15	Cenchrus	purpureus	ILRI_collection	ILRI	2019
357	NS 13-16	Cenchrus	purpureus	ILRI_collection	ILRI	2019
358	NS 13-17	Cenchrus	purpureus	ILRI_collection	ILRI	2019
359	NS 13-18	Cenchrus	purpureus	ILRI_collection	ILRI	2019
360	NS 13-19	Cenchrus	purpureus	ILRI_collection	ILRI	2019
361	Napier_Addis	Cenchrus	purpureus	NA	ILRI	NA

362	Maralfalfa_1	Cenchrus	purpureus	NA	ILRI	NA
363	mott_new2	Cenchrus	purpureus	NA	ILRI	NA

Appendix 2: (Supplementary Table S2) List of individual plants under the different clusters and sub-clusters according to the population structure analysis in STRUCTURE software

	Cluster I						Cluster II			
sub-cluster I	sub-cluster II	sub-cluster III	sub-cluster IV	sub-cluster V	sub-cluster VI	sub-cluster VII	sub-cluster VIII	sub-cluster IX	sub-cluster X	
ILRI_NS_6-10	ILRI_16808	ILRI_NS_6-11	BAGCE_17	BAGCE_100	ILRI_NS_7-20	BAGCE_63	CNPGL_91-06-2	ILRI_14355	ILRI_14983	
ILRI_NS_11-14	ILRI_16809	ILRI_NS_6-13	BAGCE_24	BAGCE_16	ILRI_NS_7-18	CNPGL_92-38-2	CNPGL_91-11-2	ILRI_15357	ILRI_16839	
ILRI_NS_11-17	ILRI_16821	ILRI_NS_6-4	BAGCE_80	BAGCE_30	ILRI_NS_7-15	CNPGL_92-66-3	CNPGL_91-25-1	ILRI_16799	ILRI_NS_10-1	

ILRI NS 11-16	ILRI 16822	ILRI NS 6-6	ILRI 16782	BAGCE 34	ILRI NS 7-13	CNPGL 9279-2	CNPGL 92-190-01	ILRI 16837	ILRI_NS_10-13
ILRI NS 11-8		ILRI_NS_6-16	ILRI 16794	BAGCE_53	ILRI_NS_7_6	CNPGL 93-32-2	 CNPGL_93-04-2		ILRI Ns 1-1
ILRI NS 11-5	ILRI 16810	ILRI NS 6-1		BAGCE_56	ILRI_NS_6-9	 ILRI 14389	 CNPGL 93-06-1	 Maralfalfa-1	ILRI NS 1-10
ILRI NS 6-8	_	ILRI_NS_6-7	ILRI 16797	BAGCE 75	ILRI NS 6-3	ILRI 14982		Tift N147	ILRI NS 11-11
ILRI NS 6-5			ILRI 16805	BAGCE 81	ILRI NS 4-2				ILRI NS 11-2
 ILRI NS 6-14				BAGCE 86	ILRI_NS_4-11		 ILRI 16784		ILRI_NS_1-12
ILRI NS 11-18			ILRI_16816	BAGCE 90	ILRI NS 4-10	ILRI 16786	ILRI 16811		ILRI NS 11-3
ILRI NS 6-15			ILRI 16834	BAGCE 94	ILRI_NS_4_15	ILRI 16787	ILRI_NS_12-10		ILRI_NS_1-13
ILRI NS 11-4			ILRI 16838	CNPGL 92-198-7	ILRI NS 13-8	ILRI 16789	ILRI NS 12-19		ILRI_NS_1-14
ILRI NS 11-12			ILRI 18448	CNPGL 96-23-1	ILRI_NS_13-7	ILRI 16792	ILRI NS 8-14		ILRI NS 1-15
ILRI NS 7-11			ILRI 18662	ILRI 16812	ILRI NS 13-5	ILRI 16793	India 119 22239		ILRI NS 1-16
ILRI NS 7-7			India_116_22242	ILRI 16813	ILRI NS 13-3	ILRI 16795	India 129 22228		ILRI NS 11-9
ILRI NS 7-3			India 121 22237	ILRI 16815	ILRI NS 13-2	ILRI 16798	India 145(1) 21965		ILRI NS 1-2
ILRI_NS_9-2			India_123_22234	PIONEIRO	ILRI NS 13-19	ILRI 16800	mott new2		ILRI_NS_12-15
ILRI NS 3-2			India 124 22233	Tift N71	ILRI NS 13-16	ILRI 16801	Tift N172		ILRI NS 12-7
ILRI NS 9-12			India 125 22232		ILRI_NS_13-15	ILRI 16803	Tift N200		ILRI_NS_1-3
ILRI NS 9-15			India 126 22231		ILRI NS 13-13	ILRI_16804	Tift N223		ILRI NS 1-5
 ILRI_NS_9-8			India_132_22225		ILRI_NS_13-11		Tift N225		ILRI_NS_1-6
 ILRI NS 11-13			India 149 21785		ILRI NS 12-6		Tift N23		ILRI NS 2-1
ILRI_NS_9-13			India_89_22236		ILRI_NS_12-5	ILRI 16836	Tift_N43		ILRI_NS_2-2
 LRI NS 11-15					ILRI NS 12-4		 Tift_N68		ILRI NS 2-3
LRI NS 6-19					ILRI NS 12-3	ILRI 18438	Tift N75		ILRI NS 2-6
ILRI_NS_9-9					ILRI NS 12-20	India 144 21964	Tift N8		ILRI NS 2-7
 ILRI NS 9-18					ILRI_NS_12-18	Napier Addis			ILRI_NS_2-8
ILRI NS 11-20					ILRI NS 12-17	Tift N109			ILRI NS 2-9
ILRI NS 4-4					ILRI NS 12-13	Tift N131			ILRI NS 3-1
 ILRI NS 9-1					ILRI NS 12-11	_			ILRI_NS_3-12
ILRI NS 7-10					ILRI NS 12-1				ILRI NS 3-13
ILRI NS 13-12					ILRI NS 10-9				ILRI NS 3-16
ILRI NS 3-3					ILRI NS 10-5				ILRI NS 3-19
 ILRI NS 7-14					ILRI NS 10-4				ILRI_NS_3-20
LRI NS 3-8					ILRI NS 10-2				ILRI NS 3-5
LRI NS 13-1					ILRI_NS_10-12				ILRI_NS_3-9
LRI NS 3-18					ILRI NS 10 8				ILRI NS 4-1
LRI NS 6-12									ILRI NS 4-13
 LRI NS 7-4									ILRI_NS_4-14
LRI NS 3-10									ILRI NS 4-6
 LRI_NS_13-9						1	1		ILRI_NS_5-1
 LRI_NS_11-19						1	1		ILRI_NS_5-10
LRI_NS_11_6									ILRI_NS_5-12
 ILRI_NS_7-1						1	1		ILRI_NS_5-13
 ILRI_NS_3-1						1	1		ILRI_NS_5-14
LRI_NS_3-6						1	1		ILRI_NS_5-15
ILRI_NS_7-5									ILRI_NS_5-17
						1			ILRI_NS_5-18
			1						ILRI NS 5-2
	1			1	1	1	1		ILRI NS 5-3

				ILRI_NS_5-4
				ILRI_NS_5-6
				ILRI_NS_5-9
				ILRI_NS_6_2
				ILRI_NS_8-10
				ILRI_NS_8-11
				ILRI_NS_8-15
				ILRI_NS_8-17
				ILRI_NS_8-19
				ILRI_NS_8-2
				ILRI_NS_8-20
				ILRI_NS_8-3
				ILRI_NS_8-7
				ILRI_NS_8-8
				ILRI_NS_8-9
				India_128_22229

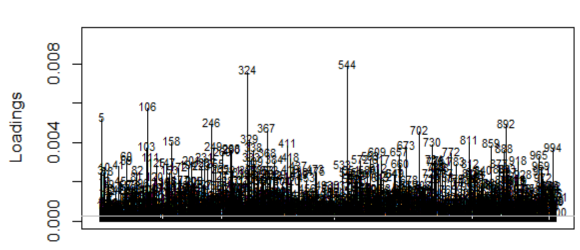
Appendix 3: (Supplementary Table S3): Membership probability of individual genotypes under different clusters and sub clusters identified by STRUCTURE analysis

	Membership probability									
	sub-	sub-cluster								
Genotype	cluster I	П	III	IV	v	VI	VII	VIII	IX	х
BAGCE_1	0.00	0.00	0.00	0.00	0.29	0.00	0.17	0.43	0.09	0.00
BAGCE_7	0.00	0.00	0.00	0.35	0.47	0.00	0.17	0.01	0.00	0.00
BAGCE_97	0.00	0.00	0.00	0.40	0.41	0.00	0.13	0.00	0.06	0.00
CNPGL_00-1-1	0.01	0.01	0.00	0.01	0.48	0.00	0.17	0.32	0.00	0.00
CNPGL_92-133-3	0.00	0.00	0.00	0.00	0.03	0.00	0.47	0.24	0.26	0.00
CNPGL_92-56-2	0.00	0.00	0.00	0.06	0.21	0.00	0.28	0.24	0.21	0.00
CNPGL_93-01-1	0.00	0.00	0.00	0.11	0.06	0.00	0.45	0.00	0.38	0.00
CNPGL_93-08-1	0.00	0.00	0.00	0.00	0.22	0.00	0.38	0.20	0.20	0.00
CNPGL_93-18-2	0.00	0.00	0.00	0.00	0.16	0.00	0.39	0.29	0.16	0.00
CNPGL_94-07-2	0.00	0.00	0.00	0.00	0.12	0.00	0.44	0.38	0.05	0.00
CNPGL_94-13-1	0.00	0.00	0.00	0.00	0.12	0.00	0.45	0.11	0.32	0.00
CNPGL_96-27-3	0.00	0.00	0.00	0.00	0.44	0.00	0.34	0.00	0.22	0.00
ILRI_1026	0.01	0.20	0.00	0.44	0.00	0.01	0.16	0.02	0.15	0.00
ILRI_15743(MOTT)	0.11	0.01	0.00	0.27	0.00	0.17	0.44	0.00	0.00	0.00
ILRI_16783	0.12	0.01	0.00	0.26	0.00	0.15	0.45	0.00	0.00	0.00
ILRI_16788	0.13	0.00	0.00	0.26	0.00	0.16	0.44	0.00	0.00	0.00
ILRI_16790	0.28	0.01	0.00	0.48	0.00	0.23	0.00	0.00	0.00	0.00

U.D. 46704	0.10	0.01	0.00	0.26	0.01	0.40	0.42	0.00	0.00	0.00
ILRI_16791	0.10	0.01	0.00	0.26	0.01	0.19	0.43	0.00	0.00	0.00
ILRI_16802	0.11	0.01	0.00	0.26	0.01	0.21	0.41	0.00	0.00	0.00
ILRI_16814	0.27	0.01	0.44	0.22	0.00	0.00	0.00	0.06	0.00	0.00
ILRI_16817	0.28	0.00	0.46	0.23	0.00	0.00	0.00	0.04	0.00	0.00
ILRI_16835	0.43	0.06	0.29	0.21	0.00	0.00	0.00	0.00	0.00	0.00
ILRI_NS_10-10	0.18	0.00	0.00	0.00	0.00	0.49	0.00	0.00	0.00	0.33
ILRI_NS_10-11	0.28	0.00	0.01	0.00	0.00	0.41	0.00	0.08	0.00	0.21
ILRI_NS_10-3	0.26	0.00	0.03	0.01	0.00	0.29	0.00	0.00	0.00	0.42
ILRI_NS_10-7	0.40	0.00	0.00	0.00	0.00	0.46	0.00	0.00	0.00	0.13
ILRI_NS_11-1	0.15	0.05	0.00	0.01	0.00	0.37	0.00	0.00	0.00	0.42
ILRI_NS_11-10	0.12	0.03	0.00	0.01	0.00	0.34	0.09	0.10	0.00	0.31
ILRI_NS_1-18	0.45	0.00	0.00	0.00	0.00	0.07	0.00	0.00	0.00	0.48
ILRI_NS_12-12	0.43	0.00	0.00	0.00	0.00	0.47	0.00	0.00	0.00	0.10
ILRI_NS_12-14	0.38	0.00	0.02	0.00	0.00	0.47	0.00	0.00	0.00	0.13
ILRI_NS_12-2	0.13	0.00	0.00	0.00	0.00	0.43	0.00	0.00	0.00	0.44
ILRI_NS_12-8	0.45	0.00	0.01	0.00	0.00	0.49	0.00	0.00	0.00	0.05
ILRI_NS_13-17	0.04	0.01	0.01	0.00	0.00	0.47	0.00	0.00	0.00	0.47
ILRI_NS_1-4	0.43	0.01	0.00	0.01	0.00	0.10	0.00	0.00	0.00	0.45
ILRI_NS_1-9	0.00	0.00	0.00	0.00	0.06	0.00	0.16	0.34	0.00	0.44
ILRI_NS_3-11	0.00	0.00	0.00	0.00	0.00	0.00	0.12	0.37	0.00	0.49
ILRI_NS_3-14	0.09	0.02	0.02	0.00	0.00	0.47	0.01	0.07	0.01	0.31
ILRI_NS_3-15	0.39	0.00	0.00	0.00	0.00	0.00	0.00	0.07	0.07	0.47
ILRI_NS_3-17	0.10	0.07	0.00	0.03	0.00	0.35	0.00	0.00	0.00	0.45
ILRI_NS_3-4	0.35	0.00	0.02	0.00	0.00	0.15	0.00	0.00	0.00	0.48
ILRI_NS_3-7	0.29	0.00	0.00	0.00	0.00	0.24	0.00	0.00	0.00	0.46
ILRI_NS_4-12	0.18	0.00	0.00	0.00	0.00	0.49	0.00	0.00	0.00	0.33
ILRI_NS_4-19	0.37	0.00	0.00	0.00	0.00	0.38	0.00	0.00	0.00	0.24
ILRI_NS_4-20	0.22	0.00	0.00	0.17	0.01	0.00	0.06	0.19	0.01	0.34
ILRI_NS_4-5	0.22	0.00	0.00	0.00	0.00	0.48	0.00	0.00	0.00	0.29
ILRI_NS_4-7	0.21	0.01	0.00	0.04	0.00	0.36	0.00	0.00	0.00	0.38
ILRI_NS_4-9	0.39	0.00	0.00	0.00	0.00	0.23	0.00	0.00	0.00	0.38
ILRI_NS_5-16	0.00	0.00	0.10	0.44	0.00	0.00	0.00	0.00	0.00	0.46
ILRI_NS_5-19	0.00	0.00	0.00	0.00	0.03	0.00	0.09	0.43	0.01	0.45
ILRI_NS_5-5	0.00	0.00	0.00	0.00	0.00	0.00	0.19	0.38	0.01	0.43
ILRI_NS_5-8	0.00	0.00	0.06	0.00	0.07	0.00	0.02	0.38	0.00	0.46
ILRI_NS_6-18	0.06	0.00	0.44	0.09	0.03	0.06	0.01	0.21	0.10	0.00
ILRI_NS_6-20	0.00	0.00	0.47	0.00	0.00	0.00	0.14	0.29	0.05	0.04
ILRI_NS_7-12	0.31	0.00	0.00	0.00	0.00	0.43	0.00	0.00	0.00	0.25
ILRI_NS_7-17	0.32	0.01	0.00	0.03	0.01	0.39	0.00	0.00	0.00	0.24
ILRI_NS_7-2	0.49	0.00	0.00	0.04	0.01	0.46	0.00	0.00	0.00	0.00
ILRI_NS_7-9	0.42	0.00	0.00	0.00	0.02	0.30	0.00	0.03	0.02	0.20
ILRI_NS_8-16	0.00	0.00	0.00	0.00	0.01	0.00	0.14	0.43	0.00	0.41
ILRI_NS_9-10	0.48	0.01	0.00	0.19	0.00	0.31	0.00	0.00	0.00	0.00
ILRI_NS_9-7	0.29	0.03	0.08	0.14	0.00	0.04	0.00	0.00	0.00	0.42
India_115_22243	0.10	0.18	0.00	0.48	0.08	0.00	0.03	0.00	0.13	0.00
India_118_22241	0.14	0.12	0.02	0.36	0.01	0.01	0.12	0.19	0.04	0.00
India_120_22238	0.01	0.04	0.10	0.27	0.00	0.01	0.09	0.44	0.04	0.01

India_127_22230	0.00	0.13	0.00	0.43	0.04	0.00	0.17	0.01	0.21	0.01
India_131_22226	0.03	0.09	0.00	0.47	0.00	0.00	0.17	0.00	0.22	0.00
India_141_21967	0.00	0.00	0.00	0.06	0.01	0.00	0.37	0.23	0.33	0.00
India_142_21966	0.01	0.07	0.00	0.35	0.05	0.00	0.22	0.01	0.29	0.01
India_146_21788	0.02	0.09	0.00	0.06	0.42	0.00	0.26	0.02	0.14	0.00
India_147_21787	0.00	0.06	0.00	0.07	0.02	0.00	0.49	0.01	0.35	0.00
India_150_21784	0.01	0.00	0.01	0.16	0.02	0.00	0.38	0.08	0.33	0.02
India_151_21783	0.08	0.00	0.00	0.45	0.00	0.00	0.14	0.00	0.32	0.00
Tift_N128	0.19	0.06	0.36	0.03	0.00	0.04	0.00	0.32	0.00	0.00
Tift_N130	0.21	0.00	0.00	0.37	0.05	0.01	0.00	0.35	0.00	0.00
Tift_N137	0.00	0.00	0.02	0.35	0.08	0.00	0.21	0.02	0.29	0.02
Tift_N138	0.01	0.00	0.00	0.30	0.04	0.00	0.26	0.03	0.36	0.00
Tift_N210	0.17	0.00	0.00	0.47	0.03	0.01	0.32	0.00	0.00	0.00
Tift_N37	0.00	0.00	0.00	0.40	0.43	0.00	0.15	0.00	0.02	0.00
ILRI_NS_10-6	0.28	0.00	0.00	0.00	0.00	0.39	0.00	0.00	0.00	0.32
ILRI_NS_4-3	0.25	0.00	0.01	0.00	0.00	0.37	0.00	0.00	0.00	0.36
ILRI_NS_4-18	0.22	0.00	0.00	0.00	0.00	0.39	0.00	0.00	0.00	0.38

Appendix 4: loading plot for 1001 SNP markers that were selected from DPAC analysis



Loading plot

Variables

Appendix 5: Genotype composition plot (compoplot) that shows membership probabilities of individual genotypes from DPAC analysis



Individuals