Research Article

Genetic Diversity of Tef [Eragrostis tef (Zucc.) Trotter] as Revealed by Microsatellite Markers

Mahilet Tadesse1,2, Mulugeta Kebede1, and Dejene Girma3

1Adama Science and Technology University, Department of Applied Biology, P. O. Box 1888, Adama, Ethiopia
2Ethiopian Institute of Agricultural Research, Debre Zeit Agricultural Research Center, P. O. Box 32, Debre Zeit, Ethiopia
3Ethiopian Institute of Agricultural Research, National Agricultural Biotechnology Research Center, P. O. Box 249, Holetta, Ethiopia

Correspondence should be addressed to Mahilet Tadesse; mahitad612@gmail.com

Received 13 November 2020; Revised 6 April 2021; Accepted 7 April 2021; Published 23 April 2021

Genetic variability is the fundamental prerequisite of any crop-breeding program to develop superior cultivars. There are about 350 Eragrostis species, of which, tef is the only species cultivated for human consumption. Currently, the Ethiopian Biodiversity Institute (EBI) collected over five thousand tef accessions from different geographical regions, diverse in terms of climate and elevation, which are uncharacterized yet. The objective of this study was to evaluate the genetic diversity among 64 tef accessions using 10 selected polymorphic simple sequence repeats (SSRs) markers. A total of 314 alleles were detected with an average of 14.5 alleles per locus and amplicon size ranged from 90 bp–320 bp. The mean value of polymorphic information content (PIC) was 0.87, appearing polymorphic for all loci. The lowest Fst value (0.05) was recorded among the studied tef populations. The mean value of major allele frequency and the number of effective alleles were 0.33 and 3.32, respectively. The mean value of gene flow (Nm) and Shannon’s information index (I) was 4.74 and 1.65, respectively. The observed (Ho) and expected (He) heterozygosities varied from 0.34 to 0.56 and from 0.58 to 0.76, respectively. The cluster analysis has grouped the 64 tef accessions into three distinct clusters based on their similarity. The PCoA analysis showed that clustering is basing on the geographical origin of accessions. Analysis of molecular variance revealed 56%, 39% and 5% of the total variation due to variation within populations, among individuals and among populations, respectively. Structure bar-plot also inferred three gene pools, but with high level of admixtures. Thus, the present study shows that the identified tef accessions could be of great interest for the initiation of a planned breeding and conservation programs.

1. Introduction

Tef [Eragrostis tef (Zucc.) Trotter] is an indigenous cereal crop of great cultural and economic importance in Ethiopia. It belongs to the grass family Poaceae, subfamily Eragrostidoideae, tribe Eragrostae, and genus Eragrostis. As one of the biggest genus in the grass family, the genus Eragrostis contains over 350 species [1]. However, tef is the only species cultivated for human conception [2]. It is an allotetraploid species with a base chromosome number of 10 (2n = 4x = 40). Cytogenetic studies on tef reported that the genome size of tef is 672 Mbp as described by Cannarozzi et al. [3] and 622 Mbp by Van Buren et al. [4]. Tef is a nutritionally rich cereal crop and contains essential and important nutrients, carbohydrates, protein, fat, fiber, and minerals [5] and has additional health benefits including that the seeds are free from gluten [6]. Some minerals like iron content were significantly higher in tef than in bread wheat [7]. Tef is widely grown across all regions of Ethiopia on a wide range of soil conditions from water-logged vertisols to moisture-stressed, drought-prone soils [8]. Tef is a source of food and income for about 50 million people annually. Over 6 million smallholder farmers in Ethiopia have cultivated it. While its health benefits and nutrition contents, tef is now growing in different countries as food and forage grass including United States, Israel, the Netherlands, Spain, South Africa, India, Australia, and Kenya [8, 9]. Comparatively to other cereal crops, the productivity of tef is significantly low (CSA, 2016). Shattering, the low genetic potential of the local varieties, lodging, and poor agronomic management
practices are the major yield-limiting factors [10]. The wealth of diversity in the species offers ample opportunities for genetic improvement of the crop to develop varieties suitable for different agro-ecologies, cropping systems, and purposes. Because of this fact, efforts have been made in the past to assess and quantify the genetic diversity in the tef germplasm collections using different approaches [11]. Molecular markers are short sections of DNA that differ between varieties and thus can be used for the identification of a germplasm by a specific pattern of polymorphisms, to assess diversity and to determine relationships. DNA-based markers provide a very effective and reliable tool for measuring genetic diversity in crop germplasm and studying evolutionary relationships. The molecular markers are superior to both morphological and biochemical markers because they are relatively abundant throughout the genome, independent of environmental conditions, and can be detected at any developmental stage of the plant [12]. Several studies have been done to examine the diversity in tef germplasm using morphological and agronomic traits. Some of the studies are Assefa et al. [13–17], Adnew et al. [18], Admas and Belay [19] Ayalew et al. [20], Shiferaw et al. [21], and Plaza-Wuthrich et al. [22]. However, few attempts have been undertaken to study the diversity using DNA markers. Ayele [23] by using AFLP markers and Bai et al. [24] by using RAPD markers reported genetic similarity coefficients of 85–90% and 84–96%, which indicated a high level of genetic similarity of the tested genotypes. Previous research [25–31] has shown the usefulness of SSR markers in genetic diversity analysis and in establishing a genetic relationship among tef germplasm. SSR markers are particularly useful since they are codominant, high reproducibility, and polymorphic, have low operational cost and locus-specificity, and revealing of high allelic diversity. To that effect, they have been used extensively in tef and other crops for various breeding and diversity studies [32]. Hence, the objective of this study was to estimate the pattern and level of genetic diversity and relatedness among 64 tef accessions using polymorphic simple sequence repeat (SSR) markers. The study will provide information on the status of genetic variation for further improvement and conservation.

2. Materials and Methods

2.1. Plant Materials. A total of 64 tef accessions representing nine populations (North Shewa, North Gondar, South Wollo, West Gojam, Tigray Central Zone, Tigray West Zone, Kembata Tembaro, East Wellega, and Illubabor) were used in this study as shown in (Table 1). These tef accessions were collected by the Ethiopian Biodiversity Institute from tef growing areas of Ethiopia. The study was carried out at Holetta National Agricultural Biotechnology Research Center (NABRC).

2.2. DNA Extraction. Twenty-day-old plants grown in a greenhouse and fresh leaves were collected from each seedling. The DNA was extracted using a modified cetyl trimethyl ammonium bromide (CTAB) method [33]. The quality of the DNA was determined by running on 1% agarose gel with 1× TAE buffer (Trizma base with EDTA and boric acid; the pH was adjusted to 8.0 with NaOH) at 100 V for 30 minutes. The quantity (concentration) of the DNA was determined using a Nanodrop spectrophotometer (ND-8000, Thermo Scientific). The molecular weight of each amplified product was estimated by comparing the DNA bands with the high-density standard lambda DNA ladder. The patterns of the amplified product were photographed by a UV transilluminator lamp (Bio-Doc Imaging System). The DNA was diluted to 50 ng using TE buffer and stored at 4°C for further PCR work.

2.3. Selection of SSR Markers. A total of 15 SSRs markers were used for initial screening developed by Zeid et al. [34]. Out of which, 10 SSRs primers were selected for final diversity analysis based on their level of polymorphism and specificity to the target loci (Table 2).

2.4. PCR Amplification. Polymerase chain reaction (PCR) based SSR marker system was applied to investigate the genetic relationships among the sixty-four accessions of tef. PCR amplification was carried out in 96-well plates (Eppendorf Master Cycler Pro S) as per the protocol suggested by Williams et al. [35]. Total PCR reaction was optimized to be 12.5 μl, and this included 6.25 μl × One Taq Master Mix (New England Bio Labs), 1 μl of each forward and reverse primers, 0.25 μl Dimethyl-sulfoxide (DMSO, Fisher Scientific), 2 μl nuclease-free water, and 2 μl 50 ng template DNA, respectively. Amplification temperatures were programmed to initial denaturation at 94°C for 3 min followed by 35 cycles of denaturation at 94°C for 1 minute, annealing temperature varies based on the individual primer requirement for 1 minute, primer extension at 72°C for 2 minutes followed by a final extension at 72°C for 10 minutes, and holding temperature at 4°C. The band separation was done by running the PCR products on a 3% agarose gel at 100 V for 2 hours in 1% TAE (Tris-acetate-EDTA) buffer along with a 50 bp DNA ladder. The patterns of the amplified product were photographed by UV transilluminator lamp (Bio-Doc Imaging System), and the data were saved for further analysis.

2.5. Data Analysis. The clear and reproducible alleles amplified by each SSR were scored according to their fragment size (bp) using the PyElph version 1.4 software package [36]. Data from all entries were summarized and converted into a suitable format for various analyses. The number of alleles (Na), number of effective alleles (Ne), Shannon’s Information Index (I), gene flow (Nm), observed heterozygosity (Ho), expected heterozygosity (He), analysis of molecular variance (AMOVA), and Hardy-Weinberg equilibrium (HWE) over the entire populations were determined using GenAlEx software version 6.502 [37]. Locus-based diversity indices including major allele frequency (MAF), gene diversity (H), polymorphic information content (PIC), and fixation index (F) were computed using Power marker version 3.25 software [38]. To examine the genetic relationship between the different accessions, a genetic dissimilarity matrix was calculated using Jaccard’s formula, and the Unweighted Pair Group Method with Arithmetic Mean.
Table 1: Passport data of the tested accessions and their geographical origins.

| Accession | Local name   | Zone                     | Woreda/district | Altitude (m.a.s.l.) |
|-----------|--------------|--------------------------|-----------------|---------------------|
| 15309     | Tef          | North Shewa              | Ataye (Efeson)  | 1443.00             |
| 236957    | Magna        | North Shewa              | Minjarna Shenkora | 1700.00          |
| 236959    | Magna        | North Shewa              | Minjarna Shenkora | 1750.00          |
| 236960    | Tekur tef    | North Shewa              | Minjarna Shenkora | 1780.00          |
| 236961    | Tikur tef    | North Shewa              | Minjarna Shenkora | 2020.00          |
| 243493    | Tikur tef    | South Wollo              | Tenta           | 2060.00             |
| 243502    | Ayiro tef    | South Wello              | Ambasel         | 1750.00             |
| 243507    | Sergenga tef | North Wello              | Habru           | 2020.00             |
| 243508    | Manga tef    | North Wello              | Guba Lafto      | 1835.00             |
| 243509    | Fenkilew     | North Wello              | Guba Lafto      | 1835.00             |
| 243529    | Key tef      | North Gondar             | Addi Arkay      | 1310.00             |
| 243535    | Nech tef     | North Gondar             | Gondar zuria    | 1920.00             |
| 243536    | Sergenga     | North Gondar             | Gondar zuria    | 1920.00             |
| 243537    | Key Fesho tef| North Gondar             | Gondar zuria    | 1920.00             |
| 243538    | Nech tef     | North Gondar             | Gondar zuria    | 1920.00             |
| 243539    | Key tef      | North Gondar             | Gondar zuria    | 2050.00             |
| 243540    | Necht tef    | North Gondar             | Gondar zuria    | 2050.00             |
| 243544    | Key tef      | North Gondar             | Gondar zuria    | 2050.00             |
| 243545    | Deyabo tef   | North Gondar             | Gondar zuria    | 2050.00             |
| 243547    | Nech tef     | West Gojam               | Bahirdar zuria  | 1870.00             |
| 243548    | Nech tef     | West Gojam               | Bahirdar zuria  | 1870.00             |
| 243549    | Key tef/Ansharo | West Gojam            | Bahirdar zuria  | 1870.00             |
| 243550    | Key lafa tef | West Gojam               | Bahirdar zuria  | 1870.00             |
| 24353517  | Sergenga     | West Gojam               | Bahirdar zuria  | 2070.00             |
| 243533    | Key tef      | West Gojam               | Bahirdar zuria  | 1815.00             |
| 243512    | Tseada tef   | Tigray Central Zone      | Kola Temben     | 1990.00             |
| 243517    | Keyih tef    | Tigray Central Zone      | Kola Temben     | 1990.00             |
| 243518    | Goigafe tef  | Tigray Central Zone      | Kola Temben     | 1990.00             |
| 243519    | Tseads Dalga | Tigray Central Zone      | Kola Temben     | 1990.00             |
| 243520    | Murenayi     | Tigray Central Zone      | Kola Temben     | 1990.00             |
| 243521    | Keyih tef    | Tigray Central Zone      | Kola Temben     | 2010.00             |
| 243524    | Tseas tef    | Tigray Central Zone      | Adwa            | 2030.00             |
| 243525    | Sergen tef   | Tigray West Zone         | Tselemti        | 1260.00             |
| 243526    | Tseads tef   | Tigray West Zone         | Tselemti        | 1260.00             |
| 243527    | Tseada tef   | Tigray West Zone         | Tselemti        | 1230.00             |
| 243528    | Sergen tef   | Tigray West Zone         | Tselemti        | 1230.00             |
| 244789    | Dalesha      | Kembata Tembaro          | Alaba           | 1773.00             |
| 244790    | Dorta        | Kembata Tembaro          | Tembaro         | 1764.00             |
| 244791    | Mitazu and Dorta | Kembata Tembaro      | Tembaro         | 1808.00             |
| 244792    | Metazo       | Kembata Tembaro          | Kachabira       | 1717.00             |
| 244793    | Sergenga     | Kembata Tembaro          | Kachabira       | 1714.00             |
| 244794    | Yemushe      | Kembata Tembaro          | Kachabira       | 1714.00             |
| 244796    | Dalesha      | Kembata Tembaro          | Alaba           | 1773.00             |
| 244814    | Xafi         | East Wellega             | Bilaseyo        | 1520.00             |
| 244815    | Taffee       | East Wellega             | Bilaseyo        | 1976.00             |
| 244816    | Taffee       | East Wellega             | Bilaseyo        | 1946.00             |
| 244817    | Tafsi Dima and Adi | East Wellega         | Bilaseyo        | 1946.00             |
| 244818    | Tafsi Adi    | East Wellega             | Bilaseyo        | 1973.00             |
(UPGMA) based Neighbor-Joining tree and hierarchical clustering analysis was carried out based on population and individual accessions using DARwin version 6.0 [39]. The structure software ver. 2.3.4 based on Bayesian algorithm was applied to determine population structure and admixture pattern [40]. To estimate the true number of population cluster ($K$), a burn-in period of 100,000 was used in each run, and data were collected over 200,000 Markov Chain Monte Carlo (MCMC) replications for $K = 1$ to $K = 10$ using 20 iterations for each $K$. The optimum $K$ value was predicted following the simulation system described by Evanno et al. [41] through the web-based STRUCTURE HARVESTER version 0.6.92 [42]. The bar plot for the maximum $K$ was determined using Clumpak beta version [43].

### 3. Results and Discussion

#### 3.1. SSR Polymorphism

The present investigation revealed high genetic variation among the studied tef accessions. These accessions were collected from different agroecology of tef growing areas. The study of genetic diversity in any research work is essential as it serves as baseline information for improvement, conservation approaches, and breeding programs. Applying different modern techniques helps in the development of potential varieties that are suitable and flexible to climate change and emerging issues that are getting global attention [44]. In this study, 15 SSR markers were screened, out of which, 10 SSR markers were found to be polymorphic and suitable for diversity analysis. The use of SSRs for tef diversity study is very crucial as it provides accurate and unbiased assessment and reveals exhaustive information on the genetic divergence of the accessions [31]. A total of 314 alleles were amplified with an average of 14.5 alleles per locus. The PIC values ranged from 0.61 for the CNLTs5 locus to 0.90 for CNLTs6. The number of amplified markers per SSR primers varied from 12 to 17 with maximum number of alleles (17) being amplified by the primer CNLTs6 and CNLTs25.
3.2. Population Genetic Differentiation. Analysis of molecular variance (AMOVA) showed significant differences among and within the populations under assayed. The highest proportion of 56% of the variation was attributed to genetic variability among individuals, while 39% was due to variation among individuals within the same population. In contrast, a smaller portion of 5% of the total variation was among populations (Table 3). Genetic differentiation among the populations was low (FST = 0.05), with a moderate effect of nonrandom mating within the populations (FIS = 0.41). The low (0.05) genetic differentiation among populations was due to the high variability within the populations, which could be due to gene flow. The extensive migration of genes through hybridization of the landraces through developing improved tef varieties could be one of the causes of the current low level of genetic differentiation among populations, and on the other way, the farmers by carelessly handling there could be mixing of the improve varieties with landraces and can be considered as accessions back [45]. Variation of a similar pattern as observed in this study among tef germplasm has been reported in a previous study [28]. 

**Table 3:** Analysis of molecular variance (AMOVA) based on standard permutation across the full data set of tef accessions collected from different geographical origins.

| Source of variation | Df | SS    | MS    | Estimated genetic variance% | P value | F statistics |
|---------------------|----|-------|-------|-------------------------------|---------|--------------|
| Among populations   | 8  | 68.36 | 8.539 | 0.210                         | 0.001   | Fst = 0.05   |
| Among individuals   | 55 | 308.38| 5.607 | 1.632                         | 0.001   | Fis = 0.41   |
| Within population   | 64 | 150.00| 2.344 | 2.344                         | 0.001   | Fit = 0.44   |
| Total               | 127| 526.69| 4.185 | 100                           |         | Nm = 4.742  |

Df: degree of freedom; SS: sum of squares; MS: mean squares.

**Table 4:** Pairwise Nei’s unbiased genetic distance (upper diagonal) and pairwise FST (lower diagonal).

|        | NS  | SW   | NG   | WG   | TCZ  | TWZ  | KT   | EW   | IL   |
|--------|-----|------|------|------|------|------|------|------|------|
| NS     | 1   | 0.198| 0.272| 0.540| 0.208| 0.074| 0.478| 0.316| 0.108|
| SW     | 0.085| 1    | 0.373| 0.341| 0.290| 0.086| 0.117| 0.180| 0.120|
| NG     | 0.059| 0.090| 1    | 0.160| 0.102| 0.270| 0.240| 0.286| 0.126|
| WG     | 0.088| 0.081| 0.072| 1    | 0.308| 0.315| 0.219| 0.270| 0.325|
| CZ     | 0.091| 0.079| 0.088| 0.084| 1    | 0.093| 0.281| 0.241| 0.168|
| TWZ    | 0.095| 0.148| 0.086| 0.108| 0.131| 1    | 0.372| 0.199| 0.090|
| KT     | 0.101| 0.082| 0.078| 0.082| 0.102| 0.129| 1    | 0.157| 0.113|
| EW     | 0.083| 0.062| 0.051| 0.067| 0.088| 0.107| 0.031| 1    | 0.400|
| IL     | 0.076| 0.085| 0.063| 0.099| 0.128| 0.135| 0.091| 0.061| 1    |

NS: North Shewa; SW: South Wollo; NG: North Gondar; WG: West Gojjam; TCZ: Tigray Central Zone; TWZ: Tigray West Zone; KT: Kembata Tembaro; EW: East Wollega; IL: Illubabor.

3.2. Population Genetic Differentiation. Analysis of molecular variance (AMOVA) showed significant differences among and within the populations under assayed. The highest proportion of 56% of the variation was attributed to genetic variability among individuals, while 39% was due to variation among individuals within the same population. In contrast, a smaller portion of 5% of the total variation was among populations (Table 3). Genetic differentiation among the populations was low (FST = 0.05), with a moderate effect of nonrandom mating within the populations (FIS = 0.41). The low (0.05) genetic differentiation among populations was due to the high variability within the populations, which could be due to gene flow. The extensive migration of genes through hybridization of the landraces through developing improved tef varieties could be one of the causes of the current low level of genetic differentiation among populations, and on the other way, the farmers by carelessly handling there could be mixing of the improve varieties with landraces and can be considered as accessions back [45]. Variation of a similar pattern as observed in this study among tef germplasm has been reported in a previous study [28]. In a study, which involved landraces, accessions, and improved varieties collected from different diverse tef growing zones in Ethiopia, 63% of the total variation was attributed to variation within the genotypes. Finding as described by Jifar et al. [31], most genotypes from germplasm accessions had high gene diversity due to more chances of coevolving in nature than those under artificial selection. The breeding lines, however, had the lowest values of all genetic parameters may due to artificial selection towards homogenous populations which drastically reduces the diversity.

**Figure 1:** Principal coordinates analysis (PCoA) biplot showing the clustering pattern of 64 tef from the nine populations.
3.3. Genetic Distance between Populations. The pair-wise population Nei’s unbiased the highest genetic distance ranged from 0.540 between population North Gondar and North Shewa accessions to 0.478 between populations of Kembata Tembaro and North Shewa. The next highest genetic distance was observed between populations of Tigray West zone and Kembata Tembaro (0.372) and between the population of West Gojjam and South Wollo (0.341) followed by Illubabor and West Gojjam (0.325) and East Wellega and North Shewa (0.316). The pair-wise genetic differentiation of low FST value (0.031) was scoured between Kembata Tembaro and East Wellega and followed by 0.051 between East Wellega and North Gondar populations. To explain the properties of subdivided populations, the magnitude of differentiation between and within populations can be quantified using $F$ statistics (Fit, Fis, and Fst) also known as fixation indexes (Fst), which is a means to evaluate the population differentiation due to genetic structure. The low Fst value (smaller genetic differentiation) implies that there is a high frequency of identical alleles among accessions [46]. The pair-wise genetic differentiation of moderate FST value accounts for 94% of the total FST values from 0.061 to 0.135 between the populations. The larger FST value was 0.15, which was recorded between Tigray West Zone and South Wollo as showed (Table 4).

3.4. Principal Coordinate Analysis. Principal coordinate analysis (PCoA) showed that the first three principal coordinates accounted for about 30.72% of the genetic variation present in the SSR molecular data derived from the 64 accessions used in the study. The first, second, and third principal coordinates explained about 13.13%, 9.20%, and 8.39% of the total variation, respectively. The PCoA analysis in the two-dimensional plot displayed in (Figure 1) showed that accessions from different collection sites often clustered together, showing possible gene flow. A similar finding was also reported by Fikre et al. [30]. There was no separate group formed by a single population possibly because of different factors that favor for intermixing of the germplasm including gene flow and hybridization. The result comes up with the NJ dendrogram, in that there was no unique clustering among accessions from a similar population. The PCoA further supports the previous finding by [25, 47] where mixed clustering was observed among accessions from the different origins of

![Figure 2: Unweighted neighbor-joining (NJ) dendrogram showing a genetic relationship of 64 tef accessions using 10 SSR markers.](image-url)
populations. In some cases, accessions of the same population such as East Wollega and North Wollo formed subcluster in the major groups. Even if most of the accessions are forming a subcluster in their specific groups; but there was no separate group formed by a single population. As described on the PCoA biplot, some accession (2, 3, 26, and 38) obtained from populations 1, 5, and 7 were found distantly from the central axis. Therefore, such kinds of accessions are highly recommendable for future. The finding of this study highlighted the need for exploitation of the investigated tef genetic resources to speed up the tef breeding program. Accordingly, the conventional tef varietal refinement efforts should be supported by modern molecular tools and scientific techniques.

3.5. Cluster Analysis and Population Structure. Three major clusters were extracted from all the sixty-four accessions containing 56.25%, 25%, and 18.75% of the total accessions in clusters I, II, and III, respectively. All the clusters were further grouped into two subclusters (Figure 2). The maximum number of populations were included in cluster I having 9 populations and the minimum number in cluster III having 6 populations. Cluster I is composed of accessions from Tigray Central Zone, South Wollo, Illubabor, East Wollega, West Gojjam, Kembata Tembaro, North Shewa, Tigray West Zone, and North Gondar, of which 34. 20% of the total population was from Tigray Central Zone and East Wollega. The clustering pattern indicated the existence of a significant amount of variability among the tef accessions. In the same way, the subclustering of cluster II resulted in two groups composed of six and five populations from Illubabor, South Wollo, North Gondar, North Shewa, Tigray West Zone, East Wollega, and Kembata Tembaro. Likewise, the third cluster contained accessions from East Wollega, West Gojjam, Kembata Tembaro, South Wollo, and Illubabor. The patterns of the grouping of populations across all clusters clearly showed the existence of genetic material independent events of evolutionary forces. Further, the independent events of evolutionary forces such as mutation, selection, genetic drift, and germplasm exchange might separate them into related but different gene pools [48, 49]. The clusters contributing maximum to the divergence were given greater emphasis for deciding the type of cluster for further selection and the choice of the parents of hybridization.

The long-standing concern in population genetics is the identification of genetically homogeneous groups of individuals. In this regard, a recent Bayesian algorithm implemented in the software STRUCTURE allows the identification of such groups [42]. The algorithm allows estimating the true number of clusters (K) in a sample of individuals included in the study. The analysis revealed that maximum delta K (ΔK) (which is a good indicator of the true number of clusters) becomes a peak at delta K. In this study, the delta K value was highest at K = 3 (Figure 3(a)) which suggests that 64 diverse accessions of tef can be divided into three subgroups (Figure 3(b)). The number of sub-groupings was determined from the delta K model described by Evanno et al. [41]. Based on the K value, the Clumpak result (bar plot) showed wide admixtures, as a result, there was no clear geographic origin-based structuring of populations (Figure 3(b)). This indicates that the SSR markers we have employed are very powerful to show the genetic relationship among the studied tef accessions.

4. Conclusions

High genetic diversity was observed among the tested accessions. The current study highlighted the existence of high levels of diversity among 64 tef accessions, which are suitable for the improvement of potential varieties. SSR markers provide unbiased and sufficient information in the identification and confirmation of tef accessions, as a result, the polymorphism detected among the accessions can be used in breeding programs to exploit the use of genetic resources. Most of the studied populations which had high values of polymorphic information content, gene diversity, and Shannon diversity index could be important to capture adequate information on the diversity state of the germplasm accessions in the Gene Bank. It also indicates the need for giving equal emphasis to all tef growing areas of Ethiopia during germplasm collection and experiment investigation. The dendrogram constructed to identify the genetic similarities among these accessions showed that accessions from the same regions were found to cluster mostly together implying a correlation between molecular groupings and their source of collection. The hierarchical clustering and the population structure analysis assembled the tef accessions under three groups. Hence, the selection of parents must be based on the wider intercluster distance and superior mean performance. As a result, the polymorphism detected among the accessions can be used in tef breeding programs for developing superior tef varieties.
Data Availability

The study was conducted at Holeta National Agricultural Biotechnology Research center molecular Laboratory.

Conflicts of Interest

The authors confirm that this article content has no conflict of interest.

Acknowledgments

The authors of the research would like to thank the Ethiopian Institute of Agricultural Research (EIAR) for the financial support to conduct this study. Ethiopian Biodiversity Institute (EBI) for tef accession provision and Dr. Zerihun Tadele from the University of Bern, Switzerland, for his technical support.

References

[1] L. Watson and M. J. Dallwitz, The Grass Genus of the World, CAB International, Wallingford, Oxon, UK, 1992.
[2] S. H. Costanza, Literature and Numerical Taxonomy of Tef (Eragrostis Tef), [MSc Thesis], Cornell University, Urbana, Illinois, 1974.
[3] G. Cannarozzi, S. Plaza-Wüthrich, K. Esfeld et al., “Genome and transcriptome sequencing identifies breeding targets in the orphan crop tef (Eragrostis tef),” BMC Genomics, vol. 15, no. 1, article 061102, p. 581, 2014.
[4] R. Van Buren, C. Wai, J. Keilwagen, and J. Pardo, “A chromosome-scale assembly of the model desiccation tolerant grass Oropetium thomaeum,” Plant Direct, vol. 2, no. 11, article e00096, 2018.
[5] K. Baye, “Tef: Nutrient Composition and Health Benefits,” ESPP Working Paper 67, International Food Policy Research Institute (IFPRI) and Ethiopian Development Research Institute (EDRI), Washington, D.C. and Addis Ababa, Ethiopia, 2014.
[6] L. Spaenij-Dekking, Y. Kooy-Winkelaar, and F. Koning, “The Ethiopian cereal tef in celiac disease,” The New England Journal of Medicine, vol. 355, no. 16, pp. 1748-1749, 2005.
[7] I. Alaunty, V. Stojceska, A. Plunkett, P. Ainsworth, and E. Derbyshire, “Improving the quality of nutrient-rich Tef (Eragrostis tef) breads by combination of enzymes in straight dough and sourdough baking,” Journal of Cereal Science, vol. 55, no. 1, pp. 22–30, 2012.
[8] S. Ketema, Tef: Eragrostis Tef (Zucc.) Trotter, International Plant Genetic Resources Institute (IPGRI), Rome, 1997.
[9] K. Assefa, J. K. Yu, M. Zeid, G. Belay, H. Tefera, and M. E. Sorrells, “Breeding tef (Eragrostis tef (Zucc.) trotter): conventional and molecular approaches,” Plant Breeding, vol. 130, no. 1, pp. 1–9, 2011.
[10] CSA, “Agricultural Sample Survey 2015/2016,” Report on Area and Production of Major Crops (Private Peasant Holdings, Meher Season), Statistical Bulletin 584, Vol. I, Addis Ababa, 2016.
[11] K. Assefa, S. Chanyalew, and Z. Tadele, “Tef Eragrostis tef (Zucc.) Trotter Millets and Sorghum,” in Biology and Genetic Improvement, J. V. Patill, Ed., 2017 John Wiley & Sons Ltd. Published by John Wiley & Sons Ltd, First edition, 2017.
[12] L. Mondini, A. Noorani, and M. A. Pagnotta, “Assessing plant genetic diversity by molecular tools,” Diversity, vol. 1, no. 1, pp. 19–35, 2009.
[13] K. Assefa, S. Ketema, H. Tefera et al., “Diversity among germplasm lines of the Ethiopian cereal tef [Eragrostis tef (Zucc.) trotter],” Euphytica, vol. 106, article 190531, 1999.
[14] K. Assefa, S. Ketema, H. Tefera, T. Kefyalew, and F. Hundera, “Trait diversity, heritability and genetic advance in selected germplasm lines of tef [Emmegistis tef(Zucc.) trotter],” Hereditas, vol. 133, no. 1, pp. 29–37, 2000.
[15] K. Assefa, H. Tefera, A. Merker, T. Kefyalew, and F. Hundera, “Variability, heritability and genetic advance in pheno-morphic and agronomic traits of tef [Eragrostis tef (Zucc.) trotter] germplasm from eight regions of Ethiopia,” Hereditas, vol. 134, no. 2, pp. 103–113, 2001.
[16] K. Assefa, A. Merker, and H. Tefera, “Qualitative trait variation in tef [Eragrostis tef (Zucc.) trotter] germplasm from western and southern Ethiopia,” Euphytica, vol. 127, no. 3, pp. 399–410, 2002.
[17] K. Assefa, A. Merker, and H. Tefera, “Multivariate analysis of diversity of tef (Eragrostis tef (Zucc.) trotter) germplasm from western and southern Ethiopia,” Hereditas, vol. 138, no. 3, pp. 228–236, 2003.
[18] T. Adnew, S. Ketema, T. Hailu, and H. Sridhara, “Genetic diversity in tef [Eragrostis tef (Zucc.) trotter] germplasm,” Genetic resource crop evolution, vol. 52, no. 7, pp. 891–902, 2005.
[19] S. Admas and G. Belay, “Drought-resistance traits variability in Eragrostis tef X Eragrostis pilosa recombinant inbred lines,” African Journal of Agricultural Research, vol. 6, pp. 3755–3761, 2011.
[20] H. Ayalew, T. Genet, T. Dessalegn, and L. Wondale, “Multivariate diversity, heritability and genetic advance in landraces in Ethiopia,” African Crop Science Journal, vol. 19, article 201212, 2011.
[21] W. Shiferaw, A. Balcha, and H. Mohammed, “Genetic variation for grain yield and yield related traits in tef [Eragrostis tef (Zucc.) trotter] under moisture stress and non-stress environments,” American Journal of Plant Sciences, vol. 3, no. 8, pp. 1041–1046, 2012.
[22] S. Plaza-Wuthrich, G. Cannarozzi, and Z. Tadele, “Genetic and phenotypic diversity in selected genotypes of tef Eragrostis tef (Zucc.)trotter,” African Journal of Agri-cultural Research, vol. 12, pp. 1041–1049, 2013.
[23] M. Ayale, Genetic diversity in tef (Eragrostis tef (Zucc.) trotter) for osmotic adjustment, root traits, and Amplified Fragment Length Polymorphism, [PhD Thesis], Texas Tech University, USA, 1999.
[24] G. Bai, M. Ayale, H. Tefera, and H. T. Nguyen, “Genetic diversity in tef [Eragrostis tef (Zucc.) trotter] and its relatives as revealed by Random Amplified Polymorphic DNAs,” Euphytica, vol. 112, no. 1, pp. 15–22, 2000.
[25] K. Assefa, A. Merker, and H. Tefera, “Inter simple sequence repeat (ISSR) analysis of genetic diversity in tef [Eragrostis tef (Zucc.) trotter],” Hereditas, vol. 139, pp. 174–183, 2003.
[26] S. Chanyalew, H. Tefera, S. Harjit, and M. E. Sorrells, “Comparison of AFLP, EST-SSR, ISSR and SSR markers for poly-morphism among recombinant inbred lines of tef (Eragrostis tef),” Journal of Genetic Breeding, vol. 61, pp. 27–34, 2007.
[27] M. Zeid, K. Assefa, A. Haddis, S. Chanyalew, and M. E. Sorrells, “Genetic diversity in tef (Eragrostis tef) germplasm
using SSR markers,” *Field Crops Research*, vol. 127, pp. 64–70, 2012.

[28] E. A. Desta, Pre-breeding of tef (Eragrostis tef (Zucc.) trotter) for tolerance to aluminum toxicity, 2015.

[29] M. Abraha, S. Hussein, M. Laing, and K. Assefa, "Performance of tef (Eragrostis tef (Zucc.) trotter) genotypes for yield and yield components under drought stressed and non-stressed conditions," *Crop Science*, vol. 56, pp. 1–8, 2016.

[30] T. Fikre, K. Tesfaye, and K. Assefa, "Genetic diversity of Ethiopian tef (Eragrostis tef (Zucc.) trotter) released and selected farmers' varieties along with two wild relatives as revealed by microsatellite markers," *Journal of Crop Science and Biotechnology*, vol. 21, no. 4, pp. 367–374, 2018.

[31] H. Jifar, K. Dagne, K. Tesfaye, K. Assefa, and Z. Tadele, "Genetic diversity and population structure of tef (Eragrostis tef (Zucc.) trotter) as revealed by SSR markers," *Advances in Crop Science and Technology*, vol. 8, no. 1, 2020.

[32] M. S. Mohan, A. B. Nair, A. Bhagwat et al., "Genome mapping, molecular markers and marker-assisted selection in crop plants," *Molecular Breeding*, vol. 3, no. 2, pp. 87–103, 1997.

[33] J. J. Doyle and J. L. Doyle, "Isolation of plant DNA from fresh tissue," *Focus*, vol. 12, pp. 13–15, 1990.

[34] M. Zeid, G. Belay, S. Mulkey, J. Poland, and M. E. Sorrells, QTL mapping for yield and lodging resistance in an enhanced SSR-based map for tef, Department of Plant Breeding and Genetics, Cornell University, 240 Emerson Hall, Ithaca, NY 14853, USA, 2011.

[35] J. G. K. Williams, A. R. Kubelik, K. J. Livak, J. A. Rafalski, and S. V. Tingey, "DNA polymorphisms amplified by arbitrary primers are useful as genetic markers," *Nucleic Acids Research*, vol. 18, no. 22, pp. 6531–6535, 1990.

[36] B. P. Ana and I. V. Cristian, "PyElph a software tool for gel images analysis and phylogenetics," *BMC Bioinformatics*, vol. 13, p. 9, 2012.

[37] P. E. Smouse, M. R. Whitehead, and R. Peakall, "An informational diversity framework, illustrated with sexually deceptive orchids in early stages of speciation," *Molecular Ecology Resource*, vol. 15, no. 6, pp. 1375–1384, 2015.

[38] K. J. Liu and S. V. Muse, "Power marker: an integrated analysis environment for genetic marker analysis," *Bioinformatics*, vol. 21, pp. 21–28, 2005.

[39] X. Perrier and J. P. Jacquemoud-Collet, "DARwin software," 2006, http://darwin.cirad.fr/.

[40] J. K. Pritchard, M. Stephens, and P. Donnelly, "Inference of population structure using multilocus genotype data," *Genetics*, vol. 155, no. 2, pp. 945–959, 2000.

[41] G. Evanno, S. Regnaut, and J. Goudet, "Detecting the number of clusters of individuals using the software STRUCTURE: a simulation study," *Molecular Ecology*, vol. 14, pp. 11–20, 2005.

[42] A. Dent and M. Bridgett, "STRUCTURE HARVESTER: a website and program for visualizing STRUCTURE output and implementing the Evanno method," *Conservation Genetics Resource*, vol. 4, pp. 359–361, 2012.

[43] N. M. Kopelman, J. Mayzel, M. Jakobsson, N. A. Rosenberg, and I. Mayrose, "CLUMPAK: a program for identifying clustering modes and packaging population structure inferences across K," *Molecular Ecology Research*, vol. 15, no. 5, pp. 1179–1191, 2015.

[44] M. A. Alam, A. S. Juraimi, M. Y. Rafii, A. A. Hamid, I. W. Arolu, and M. A. Latif, "Application of EST-SSR marker in detection of genetic variation among purslane (Portulaca oleracea L.) accessions," *Revista Brasileira de Botânica*, vol. 38, no. 1, pp. 119–129, 2015.

[45] P. Gepts and R. Papa, "Possible effects of (trans)gene flow from crops on the genetic diversity from landraces and wild relatives," *Environmental Biosafety Research*, vol. 2, no. 2, pp. 89–103, 2003.

[46] S. Wright, "The genetical structure of populations," *Annals of Eugenics*, vol. 15, no. 4, pp. 323–354, 1951.

[47] S. Chanyalew, H. Tefera, and S. Harjit, "Genetic variability, heritability and trait relation- ships in recombinant inbred lines of tef (Eragrostis tef (Zucc.) trotter) research," *Journal of Agriculture and Biological Science*, vol. 4, pp. 474–479, 2009.

[48] M. Slatkin and N. Takahata, “The average frequency of private alleles in a partially isolated population,” *Theoretical Population Biology*, vol. 28, no. 3, pp. 314–331, 1985.

[49] G. Keneni, E. Bekele, M. Imtiaz, and K. Dagne, "Genetic vulnerability of modern crop cultivars: causes, mechanism and remedies," *International Journal of Plant Research*, vol. 2, no. 3, pp. 69–79, 2012.