Genetic Diversity and Population Structure Assessed by SSR in a Peruvian Germplasm Collection of Loche Squash (*Cucurbita moschata*, Cucurbitaceae) †

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Abstract: Loche is an ancient landrace of squash from Northern Peru, notable for its vegetative reproduction and lack of seeds in fruits. To date, very little is known about its genetics. Here, we used 21 simple sequence repeats to assess the genetic diversity and population structure of a collection of 100 samples of loche from three localities in Peru, and 10 samples of related species, *C. pepo* and *C. maxima* (110 accessions in total). A total 85 bands were manually scored, obtaining an average of 4.05 alleles per locus. UPGMA clustering method and principal coordinate analysis showed a clear identification between the three species of *Cucurbita*. Population structure analysis clustered the 110 accessions into five populations: (i) three of loche, (ii) one of *C. pepo*, and (iii) one of *C. maxima*. Genetic diversity estimation was conducted considering only the three groups (populations) of loche identified, which was 0.024 as an average. AMOVA revealed the greatest variation between populations (79.66%) and indicated that variability within populations is 20.33%. Vegetative propagation by means of stem cuttings and cultivation in a very restricted geographical area would explain the rather low diversity of loche. This in turn would suggest that the apparent variation observed in fruit shape may be explained by somatic mutation and/or environmental factors.

Keywords: germplasm; microsatellites; genetic resources

1. Introduction

The loche squash (*Cucurbita moschata*) is a vegetable that is grown exclusively and traditionally in the north coast of Peru, mainly in the department from Lambayeque and is practically unknown elsewhere [1]. However, it is necessary to promote this landrace since it constitutes one of the crops of the Moche culture and can become a new product for international markets, recognizing the gastronomy of Lambayeque in the world. Currently, agro-export crops threaten to displace this exceptional pumpkin from its production area [1]. There are no studies on the genetic diversity of the loche squash in Peru. This study constitutes a fundamental factor to understand the way loche is structured. In addition, the estimation of genetic diversity is crucial for genetic improvement, conservation of species and gain in selection. To date, genetic diversity of loche cultivated in northern Peru is unknown. Furthermore, we also do not know its phenotypic diversity, that is, we do not we know how many cultivars are planted by farmers in northern Peru.

We here successfully employed 21 SSRs for the genetic characterization of loche from three departments of Peru, Amazonas, Lambayeque and Pasco, to initiate a modern
genetic improvement program and also to promote better conservation strategies for this species.

2. Materials and Methods

2.1. Samples Examined and DNA Amplification

Young fresh leaves from 100 individuals of loche from Amazonas, Lambayeque and Pasco departments in Peru were collected, considering their natural distribution range, and 10 samples of related species, *C. pepo* and *C. maxima* (110 accessions in total) as control. Samples were handled according to the protocol followed by Saldaña et al. [2,3]. Loche DNA extraction was performed by the method of Doyle and Doyle [4] with minor modifications. We used lower concentration of ethanol in washes (both at 70%) and shorter centrifugation time. To determine the genetic diversity, we tested 21 SSRs. The amplification procedure was conducted in a final volume of 10 µL containing 5 ng of DNA. PCR reaction followed the same steps of Saldaña et al. [2]. Amplified products size was tested with 100 bp marker (New England Biolabs, MA, USA).

2.2. Data Analysis

SSR band patterns were inspected to score the presence (1) or absence (0) of these. To construct a dendrogram considering the UPGMA clustering algorithm, we employed R to calculate the Nei’s genetic distances by using the *poppr* package v2.9.2 [5]. Also, 1000 bootstrap replicates were conducted. A principal coordinate analysis (PCoA) and a bayesian approach to infer the genetic population structure was performed in R and STRUCTURE software [6], respectively. We used the R package *poppr* considering the number of clusters inferred by STRUCTURE to conduct an analysis of molecular variance (AMOVA).

3. Results and Discussion

3.1. Data Analysis

The 21 SSR primers utilized for the molecular analysis revealed 85 fragments in 110 samples of *Cucurbita* spp., with 18.6 fragments as average with an average of 4.05 alleles per locus.

3.2. Genetic Diversity Estimates and Population Structure Analysis

With the 85 scored fragments, we constructed a 110 × 95 presence-absence data set. The Provesti’s genetic distances based UPGMA tree did not clearly discriminate loche samples according to their geographic locality (Amazonas, Lambayeque, Pasco). A total of two clusters of loche are supported by our dendrogram, but they present a bootstrap support lower than 70% (Figure 1). The first two axis of the PCoA explained 60.81% of the variation in agreement with our dendrogram, and showed that some samples of loche are intermingled. As expected, a clear separation among Cucurbita species was observed (Figure 2).
Figure 1. Dendrogram of 110 samples of *Cucurbita* using 21 SSR markers based on UPGMA clustering method and Provesti’s genetic distance. Numbers above the branches represent bootstrap values, with only values higher than 70% shown.

Figure 2. Principal coordinates analysis (PCoA) of 110 *Cucurbita*. Symbols and colors refer to clusters assigned by STRUCTURE software.

The Evanno method [7] depicted that the best K value (number of populations) is five for our data set: (i) cluster 1 includes loche from Pasco, (ii) cluster 2 is composed of loche from Lambayeque and Amazonas, (iii) cluster 3 contains samples of *C. maxima*, (iv) cluster 4 corresponds to samples of *C. pepo*, and (v) cluster 5 is composed of loche from Lambayeque and Amazonas. STRUCTURE analysis demonstrated admixture for few samples. This analysis also confirmed that loche samples are not clustered based on the geographic origin (Figure 3).
Figure 3. Population structure analysis of 110 samples of a Peruvian collection of loche and related species estimated by the software STRUCTURE employing 21 SSR.

The genetic diversity indices were estimated considering the five cluster (K = 5) inferred by STRUCTURE program. The Shannon-Wiener index ranged from 1.33 to 2.91, and Simpson’s index varied from 0.72 to 0.92. The expected heterozygosity (Nei’s genetic diversity index) was 0.012 for cluster 2 and 0.144 for cluster 4 (Table 1). The AMOVA revealed the greatest variation between populations (79.66%) and indicated that variability within populations is 20.33% (Table 2).

Table 1. The genetic diversity parameters based on 21 SSR markers in five clusters.

<table>
<thead>
<tr>
<th>Cluster</th>
<th>N</th>
<th>H</th>
<th>Lambda</th>
<th>He</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>2</td>
<td>68</td>
<td>2.16</td>
<td>0.721</td>
<td>0.012</td>
</tr>
<tr>
<td>3</td>
<td>5</td>
<td>1.61</td>
<td>0.8</td>
<td>0.089</td>
</tr>
<tr>
<td>4</td>
<td>5</td>
<td>1.33</td>
<td>0.72</td>
<td>0.144</td>
</tr>
<tr>
<td>5</td>
<td>31</td>
<td>2.91</td>
<td>0.92</td>
<td>0.035</td>
</tr>
<tr>
<td>Total</td>
<td>110</td>
<td>3.26</td>
<td>0.886</td>
<td>0.0973</td>
</tr>
</tbody>
</table>

N: population size, H: Shannon-Wiener index of diversity, lambda: Simpson’s index, He: Nei’s 1978 expected heterozygosity.

Table 2. Analysis of molecular variance (AMOVA) for the five clusters identified in STRUCTURE analysis.

<table>
<thead>
<tr>
<th>Source</th>
<th>df</th>
<th>SS</th>
<th>MS</th>
<th>Est. Var.</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Between clusters</td>
<td>4</td>
<td>365.25</td>
<td>91.31</td>
<td>6.11</td>
<td>79.66%</td>
</tr>
<tr>
<td>Within clusters</td>
<td>105</td>
<td>163.75</td>
<td>1.56</td>
<td>1.56</td>
<td>20.34%</td>
</tr>
<tr>
<td>Total</td>
<td>109</td>
<td>528.99</td>
<td>4.85</td>
<td>7.67</td>
<td>100.00%</td>
</tr>
</tbody>
</table>

Loche is an orphan landrace as it has been under-researched during recent years, even though it possesses traits of high agronomic and economic potential. Genetic studies of this landrace are null. In Peru, most research were focused on its botany [8] Ferriol et al. [9] used AFLP and SRAP to study the genetic diversity of a collection of pumpkin germplasm (C. moschata) maintained in Spain, finding that the accessions studied are
grouped in according to their geographical origin. This suggests the existence of two independent centers of domestication in the American continent, and/or introgression of species related to *C. moschata*. Wu et al. [10] studied the genetic diversity of this species using AFLP markers and 74 accessions from China, and 15 accessions from other countries. Accessions from China were classified into two groups, differing from the accessions from Mexico, Guatemala, Honduras, and Ecuador. The authors concluded that the American accessions present a higher number of loci than those from China. Genetic diversity of accessions of *C. moschata* from Mesoamerica were studied by chloroplast DNA sequences, demonstrating a high level of genetic diversity especially in the Mexican germplasm [11]. Most of the genetic variation among squash germplasm collections was attributed to variations between individuals within of the respective localities. In the present research study, we observed that the genetic diversity of the loche in the Peru is quite low. These results appear to be consistent with populations of almost homogeneous plants observed in commercial fields of the provinces of Lambayeque. Since loche is vegetatively propagated, very few clones closely related are being cultivated in the communities of Pacora and Illimo, as well as in other localities of Peru. Therefore, there is no opportunity for genetic recombination. Most likely the morphological variation has arisen from somatic mutation, which is maintained and propagated by cuttings. On the other hand, the “loche de montaña” that is cultivated in Bagua, Amazonas does not have genotypic variation of the loche grown in Lambayeque. The main reason for this is that the cuttings used by the farmers in Bagua would be the same as those employed in Lambayeque because farmers who decided to live in Bagua took their vegetative seeds from loche from there in order to continue cultivating loche. Therefore, the little morphological difference that exists between these fruits of loche cultivated in these two localities can be explained by environmental factors. It was also possible to confirm the transferability of the microsatellite markers within the genus *Cucurbita*, as described for other genera including *Phaseolus* and *Vigna* [12], among others.

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**References**


