Genetic Variation and Structure in Remnant Population of Critically Endangered Melicope zahlbruckneri

Raji JA1,2* and Atkinson CT3

1Hawai'i Cooperative Studies Unit, University of Hawai'i at Hilo, HI, USA
2U.S. Geological Survey, Pacific Island Ecosystems Research Center, HI, USA
3Agronomy Department, Iowa State University, Ames, IA, USA

Corresponding author: Raji, JA, Hawai'i Cooperative Studies Unit, University of Hawai'i at Hilo, 200 W. Kawili St. Hilo, HI, USA, Tel: +8089330759; E-mail: j.raji@outlook.com; jbr80@iastate.edu

Received date: November 16, 2016; Accepted date: December 13, 2016; Published date: December 19, 2016

Abstract

The distribution and amount of genetic variation within and between populations of plant species are important for their adaptability to future habitat changes and also critical for their restoration and overall management. This study was initiated to assess the genetic status of the remnant population of Melicope zahlbruckneri-a critically endangered species in Hawaii, and to determine the extent of its genetic variation and diversity in order to propose rational conservation approaches. Molecular marker allele frequencies identified genetic groups with low overall differentiation but identified the most genetically diverse individuals within the population. Analysis of Amplified Fragment Length Polymorphic (AFLP) marker loci in the population based on Bayesian model and multivariate statistics classified the population into four subgroups. We inferred a mixed species population structure based on Bayesian clustering and frequency of unique alleles. The percentage of Polymorphic Fragment (PPF) ranged from 18.8 to 64.6% for all marker loci with an average of 54.9% within the population. Inclusion of all surviving M. zahlbruckneri trees in future restorative planting at new sites are suggested, and approaches for longer term maintenance of genetic variability are discussed. To our knowledge, this study represents the first report of molecular genetic analysis of the remaining population of M. zahlbruckneri and also illustrates the importance of genetic variability for conservation of a small endangered population.

Keywords: Endangered species; Genetic conservation; Melicope zahlbruckneri; Molecular markers

Introduction

Rare and endangered species are generally vulnerable to environmental changes and genetic risks. Due to their often fragmented and small population sizes [1], they are more likely to have limited fitness along with a fixation of recessive and deleterious mutations. This increases the risk of genetic variability loss through random genetic drift when compared with larger and more diverse populations [2]. Preservation of natural genetic diversity levels and structure is thus critical for the long term management of these endangered and threatened species to prevent further loss of diversity that could arise from unexpected fluctuations in their habitat and the environment [3].

Melicope zahlbruckneri is endemic to the Island of Hawai'i with a small and declining population. It belongs to the Rutaceae family and is currently considered as a single population at the Hawaii Volcanoes National Park with less than 30 individuals left in the wild. While there is little published information on the reproductive ecology of M. zahlbruckneri, it is believed to be outcrossing, with pollination and seed dispersal by birds. Species in this family generally have flowers that divide into four or five parts. In terms of size, individual plants range from herbs to shrubs and small trees. M. zahlbruckneri grows up to about 10-12 m tall with mature leaves that are about 6-24 cm long and 4-12.5 cm wide, with well-defined lateral veins [4]. There is currently no information available on the genetics of the species, also, very little is known on their historic distribution. Thus, it is important to determine the current level of genetic variability and structure in the remnant population to aid development of sampling strategies for population restoration and propose informed recommendations to assist their ongoing management.

Molecular markers have proved to be efficient means of investigating population structure, genetic diversity, and are important components in genetic conservation studies of different species [5,6]. Molecular markers are able to provide a precise understanding of genetic diversity through the identification of genomic segments that differentiate individuals or population without the need for genetic information about the genome [7,8]. Neutral markers in combination with geographic information and appropriate statistical tools have been used frequently for the study of genetic diversity in a wide range of plant species [9-12] and have been especially useful for studies involving intra and inter-species genotypic variation in both large and small populations [13].

Several molecular markers are available with the choice often determined by availability of information for the species, the population type and size, and available resources. Microsatellites and Single Nucleotide Polymorphic (SNP) markers are recognized as the most efficient at revealing genetic diversity within and between species and can distinguish homzygous and heterozygous individuals when pedigree information is desired. However, the use of these co-dominant markers requires the availability of genomic sequence information about the species as well as high cost consideration for the
initial marker development particularly in studies involving natural populations where several species families are involved. Amplified Fragment Length Polymorphism (AFLP) on the other hand has the potential to screen a large number of genetic loci and does not require prior information about the genome of the species under investigation. Here we report an AFLP-based molecular assessment of *M. zahlbruckneri* to evaluate the genetic structure of the remnant population actively managed at the Hawai‘i Volcanoes National Park.

**Materials and Methods**

**Sample collection and DNA isolation**

The *M. zahlbruckneri* population was sampled by collecting leaf tissue from all known and accessible individual trees of the species. For each individual, a small sized healthy leaf was chosen and stored on ice in a sealed plastic bag. Leaf samples were taken to the laboratory for immediate processing or stored at -70°C until DNA extraction. A total of 28 samples were collected from the Kipukapuaulu (Table 1), the only remaining site where the species is known to be extant [except for a previously reported individual tree back in 1997 at Laupahoehoe [14].

DNA was extracted using a modified plant DNA isolation protocol by Dellaporta et al. with minor modifications [15]. The concentration and purity of DNA was measured using the nanodrop ND-1000 spectrophotometer and quality was assessed on agarose gel.

**SSR marker transferability test**

Simple Sequence Repeats (SSRs) have had limited use in studies involving native plant populations in Hawaii because they have to be developed specifically for each plant species. However, because several studies have demonstrated the successful transferability of gene based microsatellites across related genera in crop plants [11,16,17], we decided to test available genomic or gene based microsatellites that have been developed for other taxa close to *M. zahlbruckneri* to explore the prospects of transferability of such markers to genotype *M. zahlbruckneri* population.

Primer sequences of seven compound microsatellite loci developed for *Melicope quadrilocularis* [18] were synthesized (Integrated DNA Technologies, Inc. Corvallis, IA) for testing. We optimized PCR reaction components and cycling parameters for each primer pair and tested them for successful amplification and polymorphism detection in *Melicope zahlbruckneri* population. PCR reactions for SSR marker assay were performed in a Bio Rad DNA Engine (Peltier Thermal Cycler); PCR products were first separated on 1.5% TAE agarose gels and visualized under UV after ethidium bromide staining and then analyzed by capillary electrophoresis using an automated Genetic Analyzer ABI 3730.

**AFLP restriction digestion and amplification**

The AFLP procedure followed the original method described by Vos et al. with a few modifications, and unless otherwise indicated, Life Technologies (Invitrogen and Applied Biosystems) reagents were used in all reaction procedures [19]. Total genomic DNA (500 ng) was digested with 4.0 units of EcoRI and MseI restriction enzymes at 37°C for 4 h in a 25 μl reaction volume that included 2.5 μl of 5X reaction buffer and 0.2 μl of 10 mg/ml bovine serum albumin (BSA). After digestion, a 5 μl aliquot of digested DNA from each sample was run on TAE 1% agarose gel to check for complete and uniform digestion. Ligation reaction was performed in a 40 μl solution and contained 20 μl of the digested DNA, 5 pmol of the EcoRI adapter, 50 pmol of the MseI adapter, 1 units T4 DNA ligase, 0.10 μl of 10 mg/ml BSA (1 μg), and 4 μl DNA Ligase buffer. The reaction was left to incubate for 3 h at 20°C.

Pre amplification reaction was performed in a 30 μl volume using 1 μl of the ligation reaction as the template, and combined with 21.5 ng each of the EcoRI+A and MseI+A primers, 3.0 μl Amplitaq 360 DNA buffer (500 mMKCl), 15 mM MgCl$_2$, 10 mM dNTPs (each at 2.5 mM), and 1 unit of Taq DNA polymerase. The reaction was processed in a thermal cycler using the following cycling parameters: an initial denaturation step at 94°C for 60 s, 20 cycles of 94°C/30 s, 56°C/60 s, 72°C/60 s and a final extension at 72°C for 10 min. Selective amplification was performed using 36 EcoRI/Msel primer combinations to genotype the population. The primer pairs were chosen based on the number, clarity, and reproducibility of fragments produced in a preliminary screening of 60 primer combinations. Selective amplification was performed in a 10 μl reaction volume containing 2.5 μl of diluted pre amplification reaction (diluted 1:20 in 1x Tris-ethylenediaminetetraacetic acid buffer), 5 ng of EcoRI+3 and 15 ng of Msel+3 selective primers, 1 μl Amplitaq 360 buffer, 0.24 μl dNTPs (10 mM), and 0.2 units of Amplitaq 360 DNA polymerase. All EcoRI+3 selective primers were fluorescently labeled with FAM, NED and JOE using the DS-30 Matrix standard (Applied Biosystems, Foster City, CA). Amplification was performed with a touch down cycling procedure as follows: An initial denaturation step at 94°C for 2 min, 1 cycle of 30 s at 94°C, 30 s at 65°C, and 60 s at 72°C, followed by 11 cycles in which the annealing temperature decreases 0.7°C per cycle, and 22 cycles of 30 s at 94°C, 30 s at 56°C and 60 s at 72°C, and a final extension step at 72°C for 5 min.

**Sample genotyping and pre-analysis of data**

AFLP fragment analysis of all individuals was performed by capillary electrophoresis using automated Genetic Analyzer ABI 3730. ROX labeled size standard that range in size from 35-500 bp was used for fragment sizing. AFLP fragment data were normalized based on local southern size-call algorithm, peak saturation, baseline saturation, pull-up correction, and spike removal corrections. Following normalization, allele sizing and call procedures were performed using the GENEMAPPER (ver. 4.1, Applied Biosystems) software program. Individual peaks were called on the basis of the total signal intensity and a peak was scored only if its intensity exceeded a fixed threshold. Data from a standardized template was used for each primer combination to ensure that peak sizing and position was precise and reproducible in all samples.

**Data analysis**

Marker fragments were scored as either present or absent to create a binary matrix for each individual. The resulting binary matrix was used as input data to generate Jaccard’s similarity coefficient and then transformed into a dissimilarity measure as (1–Jaccard’s similarity) coefficient using the SIMQUAL module of NTSYSpc ver. 2.20d program [20–21]. Cluster analysis was performed using the unweight pair group method with arithmetic averages (UPGMA). The resulting tree branching pattern was evaluated by bootstrapping based on 1000 replicates (PHYLIP software package v.3.66) [22]. Genetic variation was further assessed by using the TREECON program to generate Neighbor joining cladograms. Additionally, genetic diversity and allele frequency
measures for the population were calculated using the AFLP-SURV program [23]. Hardy-Weinberg equilibrium (FIS=0) was assumed for the estimation of allele frequencies as an essential requirement for dominant markers where the absent fragment frequency is q2 (q being the null allele frequency). Furthermore, Nei’s unbiased expected heterozygosis was used to calculate genetic diversity within the population as the percentage of within-population polymorphic loci relative to overall polymorphic loci. This is based on the assumption that the presence of a fragment indicates either a heterozygote or homozygote for the dominant allele with frequencies estimated based on the null allele frequency. To further examine the genetic structure of *M. zahlbruckneri* population, a model based Bayesian analysis was performed using the program STRUCTURE ver. 2.4 [24-26]. A non-uniform prior distribution of allelic frequencies was assumed with its parameters derived from the observed distribution of fragment frequencies among AFLP loci [27]. Log likelihood and estimates for allele frequency divergence as a result of random drift (FST) were computed using the admixture model and correlated allele frequencies options with a burn-in period of 80,000 and 100,000 Markov Chain Monte Carlo (MCMC) iterations after burn in. Likely genetic clusters (K) groups were set over a range from K=1-5 with 10 replicate runs at each K. The posterior probability P (X|K) was estimated to give an indication of the true likely number of groups without prior information about the population. Results from STRUCTURE were further analyzed and collated with the structure harvester program [28]. The CLUMPP program was used for permutations of the most likely observations among replicate runs for each K, and results were visualized in the DIUSTRICT program [29,30].

Principal Coordinate Analysis (Pcoa) was used to explore genetic relatedness of individuals based on the modified Roger’s distance matrices generated from the binary data populations. PcoA computations were performed with the MVSP (Ver.3.21) program. A plot of the population was produced to show distances among samples in the plot as likely illustration of genetic distances within the population. The Mantel statistics was used to assess correlation between observed spatial and genetic distances among individuals.

**Results**

**Microsatellites marker analysis**

Twenty-eight plant samples from the *M. zahlbruckneri* population were genotyped using the seven compound microsatellite primers developed for *Melicope quadrilocularis* [18]. Five of the seven SSR's amplified expected allele sizes for each primer pair as described for *M. quadrilocularis*. Although transferability of the SSRs was confirmed in *M. zahlbruckneri*, the markers were unable to detect polymorphism in *M. zahlbruckneri* samples. Consequently, we decided to focus the rest of the study on the AFLP marker analysis results.

<table>
<thead>
<tr>
<th>Selective Sequence E/M Pairs</th>
<th>Total No of Loci</th>
<th>Polymorphic Loci</th>
<th>Polymorphic Fragments (%)</th>
<th>Size Range (Base pair)</th>
</tr>
</thead>
<tbody>
<tr>
<td>E-AAC/M-CTC</td>
<td>130</td>
<td>84</td>
<td>64.6</td>
<td>50-480</td>
</tr>
<tr>
<td>E-ACT/M-CTT</td>
<td>189</td>
<td>114</td>
<td>60.3</td>
<td>50-500</td>
</tr>
<tr>
<td>E-AAG/M-CAC</td>
<td>145</td>
<td>85</td>
<td>58.6</td>
<td>45-450</td>
</tr>
<tr>
<td>E-ACA/M-CAC</td>
<td>172</td>
<td>96</td>
<td>55.8</td>
<td>50-500</td>
</tr>
<tr>
<td>E-ACC/M-CAC</td>
<td>136</td>
<td>75</td>
<td>55.1</td>
<td>65-480</td>
</tr>
<tr>
<td>E-ACC/M-CAT</td>
<td>130</td>
<td>80</td>
<td>61.5</td>
<td>55-500</td>
</tr>
<tr>
<td>E-AGC/M-CTA</td>
<td>142</td>
<td>81</td>
<td>57.0</td>
<td>50-480</td>
</tr>
<tr>
<td>E-AGC/M-CAC</td>
<td>106</td>
<td>66</td>
<td>62.3</td>
<td>50-500</td>
</tr>
<tr>
<td>E-AGG/M-CTA</td>
<td>132</td>
<td>80</td>
<td>60.6</td>
<td>60-500</td>
</tr>
<tr>
<td>E-AGG/M-CAG</td>
<td>135</td>
<td>73</td>
<td>54.0</td>
<td>50-500</td>
</tr>
<tr>
<td>E-ACT/M-CAG</td>
<td>134</td>
<td>72</td>
<td>53.7</td>
<td>50-490</td>
</tr>
<tr>
<td>E-AGM/M-CTC</td>
<td>167</td>
<td>95</td>
<td>56.8</td>
<td>50-475</td>
</tr>
<tr>
<td>E-ACA/M-CTG</td>
<td>155</td>
<td>87</td>
<td>56.1</td>
<td>55-480</td>
</tr>
<tr>
<td>E-AAC/M-CTT</td>
<td>102</td>
<td>59</td>
<td>57.8</td>
<td>50-500</td>
</tr>
<tr>
<td>E-AGM/M-CTT</td>
<td>103</td>
<td>39</td>
<td>37.9</td>
<td>40-500</td>
</tr>
<tr>
<td>E-ACT/M-CTC</td>
<td>150</td>
<td>85</td>
<td>56.6</td>
<td>50-500</td>
</tr>
<tr>
<td>E-AAG/M-CTC</td>
<td>137</td>
<td>78</td>
<td>56.9</td>
<td>45-500</td>
</tr>
<tr>
<td>E-AAC/M-CAG</td>
<td>140</td>
<td>67</td>
<td>47.9</td>
<td>40-500</td>
</tr>
<tr>
<td>E-ACT/M-CTG</td>
<td>185</td>
<td>110</td>
<td>59.4</td>
<td>50-500</td>
</tr>
</tbody>
</table>
Table 1: AFLP marker pairs, characteristics and average variation in selective amplification of *M. zahlbruckneri*.

### Genetic distance and population structure

Thirty-six AFLP EcoRI/MseI combination primer pairs generated 2780 fragments (Table 2).

The percentage of polymorphic fragments per AFLP locus ranged from 18.8% to 64.6% with an average of 54.9% (Table 2).

Of the total fragments scored, 620 unique fragments (22.3%) were detected in a few samples within the population. Coefficients of similarity for all possible pair-wise comparisons ranged from 0.621-1.171.

Hierarchical trees with bootstrap values revealed low to moderate sub structuring in the *M. zahlbruckneri* population. Main clusters that are subdivided into smaller groups are shown as well as 2 individual entries located on separate branches away from the rest of the groups (Figure 1). Neighbour joining tree derived from Jaccard's coefficient appeared to be less differentiated than expected considering the percentage of unique alleles detected in a small population. Many of the samples that accounted for the private alleles found in the *M. zahlbruckneri* population showed genetic profiles that varied substantially from the rest of the population. MZ23 and MZ27 in particular were noticeably the most diverged from the rest of the population (Figure 2). Cluster analysis of the population was reconstructed without MZ23 and MZ27 to examine possible outlier effect but no significant effect was detected in the overall topology of the trees.

<table>
<thead>
<tr>
<th>Cluster</th>
<th>Average distance*</th>
<th>Mean Fst value</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.3762</td>
<td>0.0117 (p &gt; 0.05)</td>
</tr>
<tr>
<td>2</td>
<td>0.3661</td>
<td>0.0150 (p &gt; 0.05)</td>
</tr>
<tr>
<td>3</td>
<td>0.2271</td>
<td>0.3471 (p &lt; 0.01)</td>
</tr>
<tr>
<td>4</td>
<td>0.2317</td>
<td>0.9157 (p &lt; 0.01)</td>
</tr>
</tbody>
</table>

### Parameter values

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value (clusters 1-4)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Estimated Ln Prob of Data (Ln P(D))</td>
<td>-33461.6</td>
</tr>
<tr>
<td>Mean ln likelihood</td>
<td>-31309.1</td>
</tr>
<tr>
<td>Variance of ln likelihood</td>
<td>4305</td>
</tr>
<tr>
<td>Mean value of alpha</td>
<td>0.1453</td>
</tr>
</tbody>
</table>

*Average distance (between individuals within a cluster)*

Ln Likelihood: log likelihood of the data given values of P (estimated allele frequencies) and Q (the estimated membership coefficient for each individual in a cluster).

Est. Ln P (D): current estimate of ln (P(X|K)), averaging over all iterations after burn in period.
Figure 1: Hierarchical cluster analysis of *M. zahlbruckneri* using UPGMA (unweighted pair group method with arithmetic mean). Bootstrap values (based on 1000 permutations) are indicated at each tree node.

Bayesian analysis using the STRUCTURE program partitioned the observed population genetic variation into four clusters (K) that are identical to those obtained from PCoA and unrooted NJ trees (Figures 2 and 3).

A correlated allele frequency with admixture model was indicated for the analysis. The optimal convergence of the MCMC algorithm was achieved by using a burn-in period of 80,000 steps, followed by 100,000 steps of data collection and 10 replicated runs to ensure efficiency in clustering computation and accurate estimates of the posterior distribution. The optimal sample resolution in each cluster was obtained at K=4. The graphical output showing the distribution and proportion of shared AFLP genomic segments is presented in Figure 4. The plot showed the assignment of individuals into different cluster groups denoted by vertical bars with different colours.

Two individuals (MZ23 and MZ27) were significantly diverged from the rest of the population. Similarly, eight individuals (MZ3, MZ13, MZ14, MZ16, MZ24, MZ25, MZ26 and MZ28) had significant numbers of rare alleles but also shared a significant number of alleles with the rest of the population. These 8 individuals are suspected to have mixed ancestry based on the combination of unique allelic profiles and characteristics that seemed unusual in shape and outside of the expected size range for *M. zahlbruckneri* (McDaniels S., Pers. Comm). They also showed substantial degree of genetic similarities to both MZ23 and MZ27.
0.012, P>0.05. The average distance between individuals within each cluster varied among the 4 clusters. The highest distance score was found within individuals of cluster 1 (0.3762) and the lowest with the members of cluster 3 (0.2271). The mean value of log likelihood for each cluster (K) is -13109, the mean value of alpha is 0.1453 and the estimated value of Ln probability of data is -33461.

Genetic relatedness and population variation

Principal coordinates analysis scores from the first three coordinates in the *M. zahlbruckneri* population showed that eigenvectors accounted for 65.82% cumulative variation of the total variance observed, with the first coordinate accounting for the most variation at 29.9%. The plot of genetic distances estimated from the coordinates showed genetic grouping and relationships between individual samples that is reflective of a high degree of common alleles in more than 60% of the individuals. It also highlights the effect of unique alleles on grouping along the PCO2 axis (Figure 4).

Two individuals, MZ 23 and MZ 27 varied significantly and formed a cluster away from the rest of the population. This result is consistent with the pattern of variation observed from Bayesian analysis, NJ cladogram and UPGMA trees constructed for the population. In addition, Mantel statistics with 1000 random permutations did not indicate a significant evidence of correlation between genetic differentiation (FST) and geographical distances among individuals within the location (R2=0.0333, P=0.708) (Figure 5).

![Figure 5: Mantel correlation of *M. zahlbruckneri* population computed from matrices of within population pairwise genetic differentiation (Nei and Li (1979)) and the natural logarithm of pairwise geographic distances.](image)

Analyses of AFLP data revealed moderate levels of genetic variation among individuals in our study population. Identical patterns of genetic clustering of *M. zahlbruckneri* were obtained from both the UPGMA and NJ cladograms. Despite low bootstrap values at some of the UPGMA tree nodes, there was consistency in the pattern of clustering as shown by other methods used in this study with several individuals assigned to the same clusters with only minor overlap of samples between clusters. Model based Bayesian analysis provided additional information about the genetic structure of the population. Bayesian assignment and patterns of genetic variation suggested the presence of a moderately varied gene pool and the possibility of hybridization with closely related *Melicope* species based on the proportion of shared versus unique alleles across individuals in different cluster groups. In addition, the significant average FST values shown by two out of four clusters in the population further increased the likelihood of some external gene flow and the presence of individuals from other unidentified closely related *Melicope* species. The observed pattern was quite unexpected given the small population size and endangered status of the population. A plausible explanation for this observation is the high percentage of unique alleles detected in the eight individuals that showed the possibility of having mixed ancestry and a few others (particularly, MZ 23 and MZ 27) that may possibly belong to another species. It is noteworthy to mention that samples that showed the presence of unique alleles also had distinctly different morphological characteristics compared to the rest of the population. Additional genetic analysis is recommended to clarify the identity of these samples to determine their pedigree and species assignment.

The extent of overall genetic diversity detected in the population could also be attributed to historic gene flow that may have occurred with other closely related taxa when populations were more abundant and continuous, or it might be a result of more recent gene flow events with other *Melicope* species. It remains unknown how wide spread the observed unique alleles are with respect to other existing *Melicope* species. Geographic information showed that *M. radiata* is the closest population to *M. zahlbruckneri* in terms of physical distance. Although the present work did not examine *M. radiata* by molecular methods, there are strong indications based on morphological information that some level of gene flow may be occurring between these two species. It would be informative in future studies to obtain the nucleotide sequence of these unique alleles and assess their fixation in other populations.

The results presented here demonstrate that AFLP analyses in combination with appropriate statistics can detect genetic variability and provide genetic information that are valuable for conservation purposes in natural populations. Similar observations of relatively high to moderate genetic diversity in small populations have been reported in other tropical species e.g. *Swietenia humilis* [34] and *Eucalyptus albans* [35] and endangered populations [36,37]. A number of the authors attributed the observed level of genetic diversity to the outcrossing mating system of the species, which enabled pollination across isolated and small populations to influence gene flow. This is mostly characteristic of tree species with long distance seed and pollen dispersal [38]. Also, long term adaptation of small remnant populations to local conditions is a factor that can influence genetic diversity. There are clearly some limitations with using dominant molecular markers for genetic analysis compared with co-dominant markers; for example, their inability to provide locus information and distinguish between hybrids is apparent drawbacks [39]. Nonetheless, their ease of application across taxa in natural populations, ability to
detect small genetic differences and genome-wide coverage has made them suitable and highly recommended as the markers of choice in circumstances where resources are limited to develop co dominant markers and where the study objective is to obtain initial information about overall genetic variation of a population. Moreover, their compatibility with the model based Bayesian approach for inferring genetic structure without prior information about the population has demonstrated that unbiased estimates of heterozygosity and genetic differentiation can be obtained.

Given the amount of variability detected in the M. zahlbruckneri population, it may be safe to reason that the genetic risk for this species is relatively moderate at present, but considering the small population size of the single remaining population, additional studies may be needed to examine the sustainability of the observed genetic diversity regarding longer term ecological impacts on the population distribution and habitat preservation. Furthermore, it is essential to understand that using only a few samples as founding individuals can considerably impact allelic diversity and result in higher genetic drift in the long run, therefore, representation of all identified M. zahlbruckneri individuals, including their putative hybrids in restorative planting at new sites would be beneficial to restoration success. Also, MZZ3 and MZZ7 may be excluded pending confirmation of their species identity.

Future studies to determine the level of available genetic variability across the species range would be desirable. This could be done by including other related taxa in the family Rutaceae for a comparative genetic analysis. Additionally, developing co-dominant marker systems such as Single Nucleotide Polymorphic (SNP) markers via the application of genotyping by sequencing technique as well as microsatellite markers may be useful for finer scale genetic studies of the endangered population to obtain phylogenetic information and comparative analysis of genomes between Melicope sp. and other closely related taxa. In addition, it would be beneficial to know genome sizes and ploidy levels of these species for a better understanding of the structure, organization and evolution of the species’ genome. This information will better assist resource managers in planning conservation of this critically endangered species.

Conclusion

The results presented in this study illustrates the current genetic variation status of federally endangered M. zahlbruckneri and provides useful baseline information based on genetic and geographic population evaluation that can assist ongoing conservation and management priorities for the only known remaining population.

Acknowledgement

This work was supported in part by the U.S. Geological Survey Natural Resource Preservation (NRPP) and Wildlife and Terrestrial Resources Programs. The authors are grateful to Linda Pratt, Josh Van Demark, and Sierra McDaniel for assistance with field sample collections and for providing location information; and USGS Pacific Island Ecosystems Research Center for additional funds to support completion of the project. Any use of trade, product, or firm names in this publication is for descriptive purposes only and does not imply endorsement by the U.S. Government.

References


