



Citation: Costa MF, Morales-Marroquín JA, de Araújo Batista CE, Alves-Pereira A, de Almeida Vieira F, Zucchi MI (2022) Population genomics of the neotropical palm *Copernicia prunifera* (Miller) H. E. Moore: Implications for conservation. PLoS ONE 17(11): e0276408. https://doi.org/10.1371/journal.pone.0276408

Editor: Branislav T. Šiler, Institute for Biological Research, University of Belgrade, SERBIA

Received: June 10, 2022

Accepted: October 5, 2022

Published: November 3, 2022

Copyright: © 2022 Costa et al. This is an open access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Data Availability Statement: All relevant data are within the paper and its <u>Supporting Information</u> files.

Funding: This work was supported by CNPq (Conselho Nacional de Desenvolvimento Científico e Tecnológico) - process 431226/2018-0 and 311811/2019-1 and FAPESP (São Paulo Research Foundation) for the Project Grant (2021/10319-0). The funders had no role in study design, data

RESEARCH ARTICLE

Population genomics of the neotropical palm *Copernicia prunifera* (Miller) H. E. Moore: Implications for conservation

Marcones Ferreira Costa 61,2*, Jonathan Andre Morales-Marroquín 2°, Carlos Eduardo de Araújo Batista 4°, Alessandro Alves-Pereira 2², Fábio de Almeida Vieira 4†, Maria Imaculada Zucchi^{2,5}*

- 1 Campus Amílcar Ferreira Sobral, Federal University of Piauí, Floriano, Piauí, Brazil, 2 Graduate Program in Genetics and Molecular Biology, Institute of Biology, State University of Campinas, São Paulo, Brazil,
- 3 Department of Genetics, "Luiz de Queiroz" College of Agriculture, University of São Paulo, Piracicaba, São Paulo, Brazil, 4 Academic Unit Specialized in Agricultural Sciences, Federal University of Rio Grande do Norte, Macaíba, Brazil, 5 Paulista Agency of Agrobusiness Technology, Piracicaba, São Paulo, Brazil
- These authors contributed equally to this work.
- ‡ These authors also contributed equally to this work.
- * marconescosta@ufpi.edu.br (MFC); mizucchi@sp.gov.br (MIZ)

Abstract

Copernicia prunifera (Miller) H. E. Moore is a palm tree native to Brazil. The products obtained from its leaf extracts are a source of income for local families and the agroindustry. Owing to the reduction of natural habitats and the absence of a sustainable management plan, the maintenance of the natural populations of this palm tree has been compromised. Therefore, this study aimed to evaluate the diversity and genetic structure of 14 C. prunifera populations using single nucleotide polymorphisms (SNPs) identified through genotypingby-sequencing (GBS) to provide information that contributes to the conservation of this species. A total of 1,013 SNP markers were identified, of which 84 loci showed outlier behavior and may reflect responses to natural selection. Overall, the level of genomic diversity was compatible with the biological aspects of this species. The inbreeding coefficient (f) was negative for all populations, indicating excess heterozygotes. Most genetic variations occurred within populations (77.26%), and a positive correlation existed between genetic and geographic distances. The population structure evaluated through discriminant analysis of principal components (DAPC) revealed low genetic differentiation between populations. The results highlight the need for efforts to conserve C. prunifera as well as its distribution range to preserve its global genetic diversity and evolutionary potential.

Introduction

Habitat reduction and deforestation resulting from human activities have had adverse effects on forest populations, contributing to high rates of species extinction, particularly in the Neotropical region [1]. With the exception of some natural areas, most tropical species occur in anthropogenic landscapes, where the previously continuous forest has now been reduced to

collection and analysis, decision to publish, or preparation of the manuscript.

Competing interests: The authors declare no conflict of interest.

smaller and isolated patches [2]. This modification of landscape composition and structure leads to habitat fragmentation, contributing to the loss of alleles, reduction of heterozygosity, and increase in inbreeding [3].

The palm tree *Copernicia prunifera* (Mill.) H. E. Moore (Arecaceae; subfamily: Coryphoideae), known as carnaúba, generally forms monodominant populations known as "carnaubais" [4]. The species has multiple inflorescences, which are made up of yellowish and hermaphroditic flowers [5]. Flowering is more intense between November and February, and the fruiting period is between January and March [6]. Fruits are likely dispersed by sanhaçudo-coqueiro (*Tangara palmarum*) [5].

C. prunifera is endemic to the Caatinga biome [7], which is one of the largest seasonally dry tropical forest areas in South America [8]. The Caatinga is an exclusively Brazilian biome, covering an area of approximately 900,000 km² in northeast Brazil. The climate in this region is characterized by a long dry season with irregular rainfall, representing a xeric, semi-deciduous shrubland and forest vegetation [9]. This palm tree also grows in the Restinga region, which contains vegetation of the coastal plain under marine influence and established on sandy soil composed of physiognomic variations from the beach towards the interior of the coastal plain [10,11]. Caatinga and Restinga are the major vegetation units in northeast Brazil [12].

Local populations use this species as a source of employment and income. Leaf extraction is responsible for sustaining several families during the period of drought that extends from July to December in northeast Brazil [13]. The fruits serve as food for animals, the stems can be used in the construction of houses, and the fasciculated roots have medicinal properties [14]. Due to its versatility and usefulness, this palm tree is known as "the tree of life" [15]. The main product of economic value obtained from this tree is carnaúba wax, which is extracted from young leaves and is of interest in the pharmaceutical and automotive industries [16]. Carnaúba populations suffer from intense exploitation because the method used to extract carnauba wax, which consists of practically removing all the leaves of the plant to obtain ceriferous powder [17].

However, unsustainable harvesting practices and the absence of sustainable management programs pose major threats to the long-term survival of this palm tree species. *C. prunifera* populations show signs of intense exploitation with visible signs of anthropogenicity, such as fire, extraction and cutting of leaves, soil impacted by livestock, and low or absence of regeneration (Fig 1). In addition, anthropogenic disturbances in the last century, mainly due to deforestation, agricultural expansion, and modernization of agriculture, have led to a rapid decline in these populations [17].

The maintenance of genetic diversity is a powerful conservation strategy for preserving the adaptive potential of species in neotropical regions [18]. In addition to configuring the ability of species to adapt to various changing environments, genetic diversity is the driving force behind evolution and speciation. [19]. Consequently, maintenance of genetic diversity within populations ensures that the species can remain biologically active and adaptable to structural changes caused by anthropogenic actions [20,21].

The genetic diversity and structure of forest populations evaluated based on molecular markers is a widely used strategy in conservation genetics [22,23]. With the advent of next-generation sequencing, it is possible now to identify thousands of molecular markers of single nucleotide polymorphism (SNP) throughout the genome. This provides a genomic approach to evaluating genetic diversity [24]. A larger SNP sample size facilitates the identification of regions that show signs of selection and can serve as a starting point for the identification of adaptive differences between populations, which is fundamental for optimizing biological conservation efforts [25,26]. These markers enable the identification of outlier and neutral loci. Specifically, outlier loci show differentiated behavior regarding genetic variation and offer an



Fig 1. Signs of anthropization in the evaluated populations of *C.prunifera.* a) wood cutting; b) leaf extraction and cutting; c) cattle raising; and d) burns.

opportunity to evaluate local adaptation patterns; neutral loci are similarly affected by the demographic and evolutionary history of populations [27].

Genetic diversity studies based on molecular markers of natural populations of *C. prunifera* in tropical areas such as Caatinga and Restinga are still scarce [17,28–30]. In addition, no studies on *C. prunifera* have applied next-generation sequencing technology for data acquisition in population genomics. Due to the importance of this neotropical palm tree for local communities and considering the rapid and recent increases in the exploitation of its populations, the present study employed next-generation sequencing to evaluate the genetic diversity and structure of 14 natural populations of *C. prunifera* in two environments (Caatinga and Restinga) in Brazil using SNP markers to provide information that can help in the design of efficient strategies for the conservation and sustainable use of this species.

Material and methods

Plant material and DNA extraction

In the present study, 160 individual plants from 14 populations of *C. prunifera* were evaluated. Out of the samplings collected, 10 populations came from the Caatinga (RUS, LGP, SER,

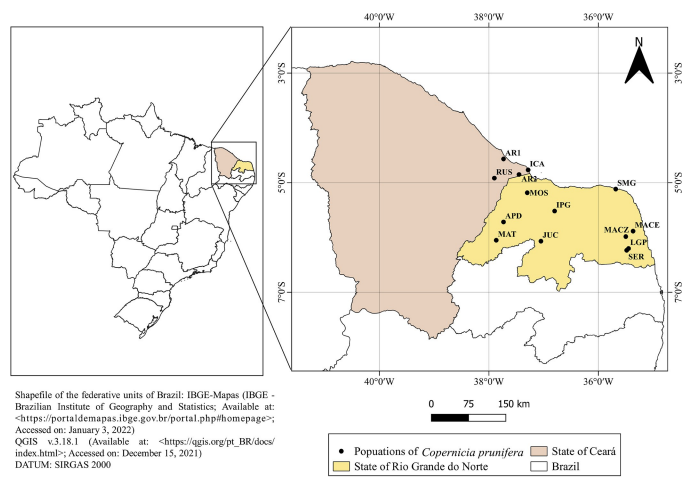


Fig 2. Map of the collection sites of *C. prunifera* populations in the states of Ceará and Rio Grande do Norte, Brazil. Distribution map of the evaluated populations was drawn using the software QGIS v3.18.1. (Open Access Geographic Information System, https://qgis.org/pt_BR/site). This figure is licensed under CC BY 4.0.

MACZ, MACE, JUC, APD, IPG, MOS, and MAT) and four from the Restinga (ICA, SMG, AR1, and AR2) regions in the states of Ceará and Rio Grande do Norte, Brazil (Fig 2 and S1 Table). The distance between the plants evaluated within the 14 populations was 15–20 m, with a minimum height of 6–10 m; regenerating and young plants were not collected. The IPG population is composed of a different type of carnaúba, known as "white carnaúba," which is phenotypically distinct from the "common carnaúba" due to the presence of a light stipe, smaller fruits, and the absence of thorns in the petiole in addition to limited occurrence in the region [14].

Small pieces of leaves were cut using a tree trimmer, placed in plastic tubes containing 2 mL of hexadecyltrimethylammonium bromide (CTAB 2X), labeled, and stored in a freezer at -20°C until DNA extraction. This study was conducted according to the recommendations of the Brazilian Ministry of the Environment and registered in the National System of Management of Genetic Heritage and Associated Traditional Knowledge (SISGEN; Sistema Nacional de Gestão do Patrimônio Genético e do Conhecimento Tradicional Associado) with the number A411583.

Genomic DNA was extracted from the processed leaves according to the protocol described by Doyle and Doyle [31]. DNA quality was evaluated in a 1% agarose gel and stained with

SYBR Safe™ (Life Technologies Corporation) for visualization under ultraviolet light, using lambda phage DNA of known concentrations as a reference. Quantification of the samples was performed using a Qubit 3.0 fluorometer with the dsDNA BRKitt (Life Technologies), and the DNA was standardized to a concentration of 30 ng.µl⁻¹

GBS library preparation and high-performance sequencing

To obtain SNPs, genomic libraries were developed using the genotyping-by-sequencing technique (GBS) with two restriction enzymes, according to the protocol described by Poland et al. [32] with modifications. First, 7 μl of genomic DNA from each sample was digested at 37°C for 12 h using the restriction enzymes *NsiI* and *MspI*. Subsequently, 0.02 μM barcode-specific adapters for Illumina technology were ligated to the ends of the digested fragments. Binding reaction was performed at 22°C for 2 h, 65°C for 20 min, 10°C indefinitely. After the adapters were ligated, the samples were purified using a QIAquick PCR Kit (Qiagen). The library was enriched by PCR (Polymerase Chain Reaction) using the following amplification program: 95°C for 30 s, followed by 16 cycles of 95°C for 10 s, 62°C for 20 s, and 72°C for 30 s, and ending at 72°C for 5 min. Finally, the library was purified using a QIAgen[®] QIAquick PCR Purification Kit. The Agilent DNA 12000 kit and Agilent[®] 2100 Bioanalyzer System were used to verify the average size of the DNA fragments. Sequencing was performed using the Illumina HiSeq 2500 Mid Output Kit v4 (50 cycles) (Illumina Inc., San Diego, CA, USA) in a single-end configuration.

Identification of SNPs

The identification of SNPs was performed using Stacks software v.1.42 [33,34]. The first step comprised filtering and demultiplexing with the $process_radtag$ module. In the absence of a reference genome for C. prunifera, the DeNovo Stacks pipeline was used, starting with the ustacks module to identify putatively homologous read stacks (putative loci). This step was performed for each sample separately using the following parameters: minimum stack depth (-m=3) and maximum distance between stacks (-M=2). The loci of each sample were grouped into a catalog using the cstacks module, allowing a maximum distance of two nucleotides (-n 2) between the loci of each sample. Loci with lower probability values $(-\ln l_{-} lim - 10)$ were eliminated using the rxstacks correction module. Finally, the population module was used to filter the SNP markers using the following parameters: only one marker per sequenced tag, frequency of least frequent allele $(MAF \ge 0.01)$, minimum stack depth 3X, and minimum occurrence in 75% of saplings in each population.

Loci determination under selection

Two complementary tests were performed, pcadapt and fsthet, were performed to detect outlier loci (hypothetically under selection). The pcadapt method [35] was used to identify loci associated with the genetic structure revealed by a principal component analysis (PCA), that is, without any underlying genetic model. The analysis was performed using the pcadapt package [35] on the R platform [36] by retaining the first eight principal components of the PCA and considering the loci with q-values ≤ 0.1 as outlier SNPs. The fsthet method [37] was used to identify loci with F_{ST} values that were excessively high or low compared with what was expected under neutrality. The analysis was performed using the fsthet package [37] on the R platform [36] by considering the loci below or above the 95% confidence intervals constructed with 1000 bootstraps for the expected relationship between H_E and F_{ST} as outlier SNPs. This test was performed by considering the estimates of F_{ST} in two different scenarios: i) comparing the Restinga and Caatinga populations and ii) comparing the samples of the white morphotype

with those of conventional morphotype. The final set of SNP markers hypothetically under selection consisted of loci identified as outliers in at least two of the three tests performed. Thus, the outlier SNP loci may reflect the action of selection on different types of vegetation.

Sequences containing outlier SNPs were searched with the BLASTX tool against the genomic data set of the National Center for Biotechnology Information (NCBI) using blast2go [38]. This analysis was performed to identify the similarities between the protein-coding data deposited in the NCBI database and the loci with outlier SNPs identified. For sequences with significant BLASTX hits, the functional annotation associated with characterized and/or described coding sequences was performed using the gene ontology system (GO terms). GO terms summarize information on cellular components, molecular functions, and biological processes in which the gene products are involved.

Population genomic analyses

Genetic diversity was estimated based on the number of alleles (A), number of private alleles (Ap), observed heterozygosity (H_O), and expected heterozygosity (H_E). Inbreeding coefficients (f) were also estimated, and their confidence intervals were obtained using 1000 bootstraps. Estimates of diversity and inbreeding were obtained using the diveRsity [39] and PoPPr [40] packages of the R software [36]. The distribution of genetic variation within and between populations of C. prunifera was evaluated using analysis of molecular variance (AMOVA), and its significance was tested with 10,000 permutations using the PoPPr program [40].

Genetic differentiation was estimated using pairwise F_{ST} values with confidence intervals of 1000 bootstraps, using the diveRsity package [39] of R software [36]. The population structure was evaluated using discriminant analysis of principal components (DAPC) with Adegenet [41,42] for R software [36]. This analysis was performed for neutral loci, and *priori* groups were defined from the 14 sampling sites. DAPC does not presuppose the underlying population genetic processes (e.g., binding equilibrium and Hardy–Weinberg equilibrium) common to other methods used to detect population structure, and as it is based on principal component analysis, this method can analyze genomic datasets relatively efficiently [43].

Genetic relationships and divergence between individuals were investigated by constructing a dendrogram generated based on the distance of Nei using the neighbor-joining method [44]. The final dendrograms were formatted using MEGA version 7 [45].

Results

Identification of SNPs and determination of loci under selection

Sequencing of the genomic libraries resulted in 566,922,165 reads, and after quality control, the total number of reads retained was 397,047,980. In total, 1,013 SNPs (average depth of 21X) were identified. A total of 391 outlier SNPs were identified using the pcadapt method, 70 using the fsthet method to compare morphotypes, and 47 using the fsthet method to compare vegetation types (Fig 3). Of these, 84 SNP markers were identified in at least two of the three tests and were considered hypothetically under selection, whereas the other 929 markers were considered as neutral loci. Among the outlier loci, 55 were putatively under positive selection and 29 were putatively under balancing selection. Only six outlier loci were found in the sequences similar to the annotated proteins (S2 Table). Considering the results of the GO terms, the most frequent annotations for these proteins were the molecular functions of "binding" and "catalytic activity," and the biological process of cell metabolism (S1 Fig).

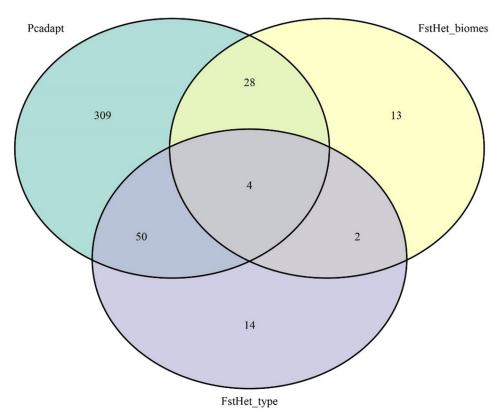


Fig 3. Venn diagram with the number of outlier loci detected for the fsthet and Pcadapt tests with the overlap between them.

Population genomic analyses

The genomic diversity estimates were based on 929 neutral SNP markers. The number of alleles (A) ranged from 1,059 to 1,497. The IPG population (white carnaúba) had the lowest number of alleles, probably because of the small sample size of the population. Expected heterozygosity (H_E) ranged from 0.201 to 0.265 (Table 1). The APD population had higher genetic diversity (H_E = 0.265) and the highest number of private alleles (Ap = 40) compared with that of the other populations. The inbreeding coefficients (f) were similar and negative for all populations, indicating an excess of heterozygotes.

 F_{ST} values 0–0.05 and 0.05–0.15 indicate low and moderate genetic differentiation, respectively, whereas values > 0.15 indicate high differentiation (Hartl, Clark 1997). In the present study, F_{ST} estimates suggested low to high genetic differentiation between populations of C. prunifera (Table 2). In general, there was a greater differentiation between the population from MACE and those from the other sites (F_{ST} ranged from 0.118 to 0.20). In addition, the SMG and IPG populations showed moderate levels of differentiation. A low genetic structure was observed for the populations from LGP and SER (0.18) and AR1 and AR2 (0.19), suggesting a genetic flow between these localities.

The low genetic divergence suggested by the pairwise F_{ST} was also observed in DAPC, which retained 28.7% of the total variation in the first two principal components (Fig 4). This analysis also showed greater genetic differentiation of the population from MACE in comparison with that of others in addition to pointing out an overlap between individuals from almost all populations, especially AR1, AR2, and RUS.

Table 1. Estimates of genomic diversity and inbreeding based on 929 neutral SNP markers for populations of *C. prunifera*.

Population	N	A	AP	H _O	H_E	f	f (95% CI)
ICA	7	1439	2	0.375	0.257	-0.367	(-0.4730.308)
AR1	17	1497	35	0.387	0.260	-0.391	(-0.4390.357)
AR2	11	1484	3	0.388	0.258	-0.454	(-0.5260.394)
RUS	21	1495	27	0.383	0.260	-0.287	(-0.5560.326)
SMG	10	1353	16	0.370	0.235	-0.272	(-0.340.211)
LGP	6	1059	5	0.378	0.229	-0.553	(-0,6420.473)
SER	10	1235	6	0.381	0.246	-0.438	(-0,5560.326)
MACZ	10	1462	9	0.387	0.259	-0.405	(-0,4610.365)
MACE	14	1227	7	0.369	0.201	-0.102	(-0,1260.081)
JUC	12	1466	3	0.376	0.247	-0.447	(-0,5640.38)
APD	12	1317	40	0.372	0.265	-0.238	(-0,3830.0127)
IPG	5	1249	1	0.375	0.224	-0.123	(-0,2090.049)
MOS	18	1404	28	0.380	0.255	-0.112	(-0.1740.065)
MAT	7	1361	8	0.376	0.237	-0.353	(-0.4370.306)

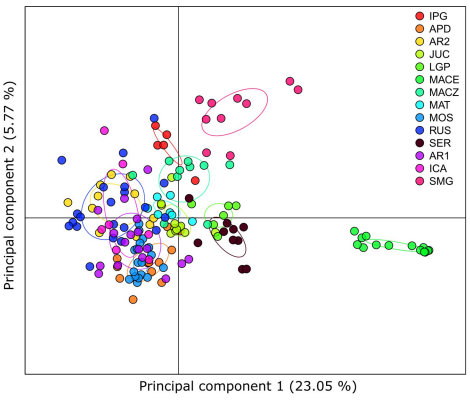
Number of individuals (N); total number of alleles (A); number of private alleles (Ap); observed heterozygosity (H_O); expected heterozygosity (H_E); inbreeding coefficient (f); f (CI of 95%) = lower and upper confidence interval of 95% of the inbreeding coefficients.

https://doi.org/10.1371/journal.pone.0276408.t001

Table 2. Estimates of pairwise F_{ST} between populations of *C. prunifera* (lower diagonals). Upper diagonals contain the lower and upper limits of the confidence interval.

	IPG	APD	AR2	JUC	LGP	MACE	MACZ	MAT	MOS	RUS	SER	AR1	ICA	SMG
IPG		(0.05- 0.14)	(0.02- 0.11)	(0.05- 0.20)	(0.05– 0.17)	(0.17- 0.27)	(0.03- 0.12)	(0.06- 0.19)	(0.04- 0.14)	(0.03- 0.13)	(0.05- 0.16)	(0.04- 0.13)	(0.02- 0.13)	(0.05- 0.16)
APD	0.092		(0.01- 0.07)	(0.01- 0.12)	(0.03- 0.12)	(0.12- 0.19)	(0.02- 0.08)	(0.06- 0.13)	(0.02- 0.08)	(0.01- 0.06)	(0.02- 0.09)	(0.02- 0.07)	(0.00- 0.09)	(0.07- 0.14)
AR2	0.053	0.036		(0.02- 0.11)	(0.03- 0.10)	(0.13- 0.20)	(0.01- 0.05)	(0.03- 0.10)	(0.01- 0.06)	(0.00- 0.04)	(0.02- 0.08)	(0.00- 0.04)	(-0.03- 0.03)	(0.04- 0.09)
JUC	0.109	0.062	0.057		(0.03- 0.15)	(0.11- 0.18)	(0.01- 0.09)	(0.01- 0.15)	(0.05- 0.14)	(0.03- 0.11)	(0.02- 0.10)	(0.03- 0.11)	(0.02- 0.12)	(0.06- 0.14)
LGP	0.096	0.064	0.054	0.072		(0.10- 0.16)	(0.02- 0.09)	(0.06- 0.15)	(0.06- 0.13)	(0.04- 0.09)	(-0.01- 0.07)	(0.02- 0.08)	(0.01- 0.11)	(0.05- 0.13)
MACE	0.210	0.150	0.162	0.144	0.123		(0.09- 0.15)	(0.17- 0.23)	(0.15- 0.21)	(0.13- 0.17)	(0.09- 0.15)	(0.12- 0.16)	(0.14- 0.21)	(0.12- 0.17)
MACZ	0.063	0.046	0.027	0.042	0.051	0.118		(0.04- 0.10)	(0.03- 0.09)	(0.01- 0.05)	(0.02- 0.08)	(0.02- 0.06)	(0.00- 0.07)	(0.03- 0.08)
MAT	0.113	0.087	0.058	0.073	0.094	0.195	0.063		(0.06- 0.12)	(0.04- 0.09)	(0.06- 0.12)	(0.04- 0.10)	(0.03- 0.10)	(0.08- 0.15)
MOS	0.077	0.047	0.032	0.084	0.086	0.172	0.050	0.084		(0.02- 0.07)	(0.05- 0.10)	(0.03- 0.07)	(0.00- 0.07)	(0.06- 0.11)
RUS	0.068	0.033	0.016	0.060	0.057	0.148	0.030	0.054	0.041		(0.04- 0.09)	(0.01- 0.04)	(0.00- 0.05)	(0.05- 0.11)
SER	0.094	0.053	0.045	0.054	0.018	0.118	0.046	0.080	0.069	0.063		(0.03- 0.07)	(0.02- 0.08)	(0.05- 0.10)
AR1	0.074	0.041	0.019	0.062	0.043	0.138	0.034	0.064	0.045	0.019	0.048		(0.01- 0.07)	(0.05- 0.10)
ICA	0.060	0.036	-0.003	0.058	0.050	0.171	0.028	0.055	0.032	0.022	0.044	0.029		(0.05- 0.11)
SMG	0.090	0.101	0.064	0.098	0.081	0.142	0.048	0.102	0.079	0.077	0.070	0.073	0.072	

https://doi.org/10.1371/journal.pone.0276408.t002



AR2 JUC LGP MACE MACZ MAT MOS RUS SER AR1 ICA SMG

Fig 4. a) Discriminant analysis of principal components (DAPC) representing the genetic structure of *C. prunifera* populations based on 929 SNPs. b) Bar graph representing the coefficients of DAPC, where each bar delimits one individual.

https://doi.org/10.1371/journal.pone.0276408.g004

IPG APD

1.0 0.8 0.6 0.4

Analysis of molecular variance (AMOVA) indicated that most of the variation was found within populations (77.26%), and the genetic differentiation between populations was high and significant (ϕ = 0.227) (Table 3). The Mantel test revealed a positive and significant correlation between the geographical and genetic distances based on the F_{ST} values (r = 0.0612; p = 0.002).

According to the dendrogram (Fig 5), the MACE population was the most genetically distant, corroborating the results observed in the DAPC population. Individual saplings from the LGP and SER populations exhibited similar levels of genetic similarity. In addition, there was a

Table 3. Analysis of molecular variance (AMOVA) based on 929 neutral SNP markers for fourteen natural populations of C. prunifera.

	Degrees of freedom	Sum of the squares	Mean square	Variance	Percentage	F _{ST} global (φ)	p-value
Between the populations	13	3307.963	254.45867	17.32471	22.7366	0.227366	0.00005
Within the population	146	8595.417	58.87272	58.87272	77.2634		
Total	159	11903.38	74.86402	76.19743			

https://doi.org/10.1371/journal.pone.0276408.t003

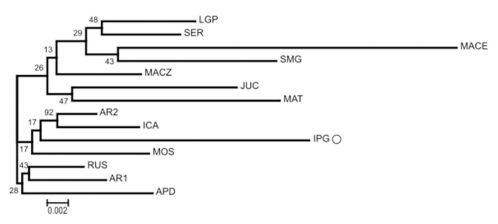


Fig 5. Dendrogram obtained by the neighbor-joining method based on SNP markers for the 14 populations of *C. prunifera*.

clear distinction among the three groups: the first group was formed by the populations from LGP, SER, MACE, SMG, MACZ, JUC, and MAT; the second group consisted of AR2, ICA, IPG (white carnaúba), and MOS; and the third group consisted of RUS, AR1, and APD. Populations from AR2 and ICA had the highest bootstrap value, which indicates a statistically well-supported grouping.

When the genetic diversity and structure of the *C. prunifera* populations were evaluated based on the type of vegetation (Caatinga and Restinga), similar levels of genetic diversity were observed (Table 4).

The populations from the Caatinga had the largest number of private alleles (Ap = 253) compared to the Restinga populations and this result is probably associated with the sample size. The inbreeding coefficients (f) are both similar and negative. In addition, the F_{ST} estimates suggested low genetic differentiation between the Caatinga and Restinga populations (F_{ST} = 0.008). When considering only two morphotypes of C. prunifera (white carnaúba and common carnaúba), similarities were observed in the estimates of diversity in addition to low genetic differentiation (F_{ST} = 0.008) (Table 4). Analysis of molecular variance (AMOVA) among the vegetation types (Caatinga and Restinga) produced small genetic differentiation (ϕ = 0.024). It revealed 97.522% of the genetic variation within the vegetation types whereas, 2.478% of the total genetic variation was observed between types of vegetation (Table 5).

Discussion

Loci putatively under selection

The large number of SNP markers obtained in this study allowed for the identification of loci with deviations from the expected neutral behavior, which are putatively under selection

Table 4. Estimates of genomic diversity and inbreeding based on 929 neutral SNP markers for populations of *C. prunifera*, considering the different types of vegetation (Caatinga and Restinga) and morphotypes (common carnaúba and white carnaúba).

Population	N	A	AP	H_{O}	H_E	f	f(95% CI)
Caatinga	126	1802	253	0.379	0.270	-0.400	(-0.4170.382)
Restinga	34	1603	54	0.380	0.266	-0.432	(-0.4680.397)
Common carnaúba	155	1855	607	0.348	0.271	-0.400	(-0.4170.384)
White carnaúba	5	1249	1	0.345	0.224	-0.123	(-0.2090.048)

Number of individuals (N); total number of alleles (A); number of private alleles (Ap); observed heterozygosity (H_O); expected heterozygosity (H_E); inbreeding coefficient (f); f (CI of 95%) = lower and upper confidence interval of 95% of the inbreeding coefficients.

https://doi.org/10.1371/journal.pone.0276408.t004

Table 5	5. Molecular analysis of v	ariance (AMOVA) consid	lering the Caatinga and	Restinga for the po	pulations of <i>C.1</i>	orunifera.

	Degrees of freedom	Sum of the squares	Mean square	Variance	Percentage	F _{ST} global (φ)	<i>p</i> -value
Between types of vegetation	1	175.234	175.23395	1.886185	2.478	0.02478074	0.00005
Within types of vegetation	158	11728.146	74.22877	74.22877	97.522		
Total	159	11903.38	74.86402	76.114954			

(outlier loci). The identification of outlier loci is an important step in understanding local adaptation and evaluating the evolutionary potential of a species [46]. The palm tree *C. prunifera* has no annotated reference genome, and probably for this reason, most sequences with outlier loci are similar to uncharacterized proteins. Regarding the results obtained from the annotation, most loci are associated with genes involved in metabolic processes, which have been regularly found under selection in a variety of organisms because the gene functionality correlates with environmental stressors [47].

Interestingly, some annotated loci were associated with genes of transposable elements (S2 Table). According to Gogvadze and Buzdin [48], transposable elements promote changes in the genome, which is an important evolutionary mechanism for the adaptation of organisms to changes in environmental conditions. This is expected in *C. prunifera* because the palm trees grow in different environments such as seasonally flooded areas in the semi-arid region [49]. In addition, outlier loci may be associated with environmental differences in the collection sites, especially as sampling areas are scattered over the Restinga and Caatinga.

It is important to highlight that the analyses performed in this study are unable to indicate associations between genomic and functional variation; therefore, it is not possible to associate generic molecular functions or biological processes with any adaptive traits involved in the diversification of the evaluated populations. Therefore, studies with larger sample sizes with better representation of the different geographical habitats are needed to generate information on the evolution and diversification of *C. prunifera*. Small sample sizes belonging to populations with relatively small geographical distances, which enable gene flow to quickly spread new adaptations to surrounding areas, reduce the capacity to detect recent evolutionary changes [50]. However, the identified outlier loci can be used as candidates in association mapping studies. Thus, integrative approaches of association genetics, genome-wide scans, and measurements of phenotype selection are necessary to understand the adaptive nature of a given allele [51].

Genetic diversity, inbreeding, and structure

Genetic diversity is one of the three classes of biodiversity recognized as a global conservation priority and plays a decisive role in conservation efforts. Genetic diversity has a substantial effect on both individual fitness and the adaptive capacity of the population, playing a vital role in maintaining the capacity of species to withstand various biotic and abiotic stressors and evolve under altered environmental conditions [52]. The present study provides the first estimates based on SNPs for genetic diversity in *C. prunifera*. The GBS approach used in this study produced a large number of SNP loci for the genomic evaluation of this palm tree without the need for a reference genome. This has resulted in robust estimates of diversity and patterns of genetic structure.

The results of genetic diversity and population structure were similar based on the results of the analysis according to population (among the 14 localities), type of vegetation (Caatinga and Restinga), and morphotype (common carnaúba and white carnaúba). In all situations, the populations showed a negative *f* value, suggesting limited inbreeding with reduced self-

pollination capacity under environmental conditions. Therefore, individual plants are less related than expected under conditions of random mating. Genetic diversity and population structure is influenced by biological characteristics of the species, including the mating system [53]. Therefore, the reproductive biology of this species may explain the observed patterns of genetic variation. The mating system of *C. prunifera* is mixed and preferably allogamous [5], which favors the crossing between unrelated individuals. Thus, inbreeding coefficients are reduced, and the maintenance of genetic diversity within populations is ensured.

Although the evaluated populations were susceptible to anthropogenic threats, it is possible that they had high genetic diversity (H_E). High levels of genetic diversity led to an increase in long-term survival of a species; therefore, a strong positive correlation exists between heterozygosity and population fitness, which is important for populations to adapt to new environmental conditions [54]. This high level of diversity is expected in forest species that are largely not domesticated as a result of local adaptation and neutral evolutionary processes in heterogeneous environments [22].

The identification of private alleles is useful for genetic conservation [55]. In the present study, the populations from APD (Ap = 40), AR1 (Ap = 35), and MOS (Ap = 28) had the highest number of private alleles and diversity was not found in the other localities; therefore, these populations deserve special management because the levels of private alleles are indicative of individual fitness and explain the evolutionary potential of populations and their ability to adapt to the adverse environmental conditions [21]. Therefore, this information can be used to increase the genetic representation in germplasm banks. and to convey the need to explore seed collection *in situ* to ensure future replacement.

Genetic variation in plant species is strongly affected by several historical and demographic factors, including geographic distribution, life form, and population size [56]. The results of AMOVA showed that most of the genetic diversity was found within *C. prunifera* populations (Table 3). Similarly, Santos et al. [17] analyzed the genetic differentiation of this palm tree in the northeast region of Brazil and found that 62.86% of molecular variance was accounted for by differences within populations. These results agree with those of different studies conducted on forest species that reproduce by allogamy, seeing as these species have maintained most of their genetic variability within populations [57].

Genetic structure analyses indicated that the 14 collection sites did not belong to a single homogeneous population, and the geographically closest populations showed low values of pairwise F_{ST} and overlap in the DAPC. Greater genetic similarity was found between the populations from LGP and SER and between AR1 and AR2. In addition, low genetic differentiation was observed when the populations were evaluated according to vegetation type and morphotype.

The low level of global genetic differentiation found between the populations studied here (supported by F_{ST} , cluster analysis, and DAPC) and the higher proportion of genetic diversity within populations with only fewer partitions between them could result from the combined effect of different factors, such as cross rate, reproductive system, and high genetic flow rate in this species. The Mantel test corroborates this result. Since geographically close populations tend to be genetically similar, this indicates a pattern of isolation by distance. However, the MACE population had the lowest level of diversity ($H_E = 0.201$) and the highest degree of structuring, being the most genetically divergent population compared with the others. This differentiation was supported by the F_{ST} value, which is an indirect estimator of the population connectivity between subpopulations (Table 2). The population from MACE corresponds to a small population with the lowest number of plants in an area of approximately 0.9 hectares, and spatially isolated from other populations. Furthermore, anthropogenic factors are observed in higher intensity in this population, this fact is probably related to the intense

exploitation of carnauba wax in this area [14]. These factors can lead to a reduction in genetic diversity within population, likely as a result of genetic drift [58]. Santos et al. [17], using ISSR markers, also indicated that MACE population has a high differentiation, and genetic discontinuities were observed between this population and the others, with indications of a recent genetic bottleneck. Therefore, the impact of human activities may have contributed to the levels of genetic differentiation observed in the MACE population.

Implications for conservation

Conservation genomics is an extension of conservation genetics that seeks to apply genomic techniques to the practical management of natural populations [59]. In this context, evaluations of genetic variations in the entire genome are powerful approaches to gain an understanding of the processes that lead to molecular diversification and inform effective management and conservation strategies [60]. However, application in real-time has been slow and a persistent gap exists between theory and practice.

In Brazil, the legislation that guides forest management does not clearly describe the importance of genetic evaluation within natural populations; therefore, information that seeks to associate genetic data with the formulation of sustainable management plans is unfortunately not mentioned [61]. Although *C. prunifera* is not listed as an endangered species, the expansion of agricultural activities over time has contributed to a reduction in its natural population [17]. Therefore, conservation measures are necessary to minimize the additional loss of alleles and to ensure the maintenance of genetic resources.

Conserving genetic diversity within a population should be the cornerstone of any conservation strategy aimed at ensuring the long-term persistence of species and habitats [62]. *In situ* and *ex situ* conservation strategies are considered promising alternatives for the conservation of forest genetic resources (FGR) and aim to maintain the genetic diversity of species over time, preserving the evolutionary processes and adaptive potential of populations [63]. Although *ex situ* approaches have the potential to conserve much of the biological diversity, they do have a limitation of being more suited and efficient for conservation in plants that have orthodox seeds. Therefore, *in situ* conservation is recommended for *C. prunifera* because this species contains recalcitrant seeds [64]. However, active management, including the establishment of *in vivo* seed banks and the promotion of natural regeneration, can prevent the decrease of population size, loss of genetic variability, and ensure long-term conservation [17].

The high genetic diversity observed in the evaluated populations of C. prunifera indicates the need for large areas of land dedicated to in situ conservation for capturing the existing genetic diversity of these populations. F_{ST} values estimated in the present study could help in recommending the optimal number of populations for sampling, including populations that had the highest estimates of diversity and the largest number of private alleles.

C. prunifera exploitation is an important source of employment and income for local communities in the semi-arid region of Brazil. In this context, the rational management of palm tree products should be a principal strategy in the efforts to conserve the natural habitats of the species. Another strategy aimed at conservation and sustainable use would be the development of a community and family forest management (CFFM) plan [65], which consists of the planning and management of actions and appropriate techniques for the sustainable use of forest resources aimed at traditional communities and family farmers [66].

In addition, practical measures aimed at successful plant regeneration, such as the pause of extractive activity during reproductive periods and the introduction of rotation cycles for leaf harvesting in the explored areas, need to be implemented for the sustainable management of carnaúba. However, the current social and economic conditions of workers employed in the

activity of extraction and production of carnaúba wax must be considered. Workers in poorer areas need to be provided additional support, including investments, to maintain the balance between socioeconomic demand and conservation, which would pave the way to a more sustainable supply of resources while reducing the pressure of uncontrolled harvesting.

A third approach would be to preserve populations and divergent genetic groups identified in this study throughout their geographic distribution range through effective long-term genetic and ecological monitoring, stimulating the development of ecological corridors between fragments and natural forests, and avoiding the reduction of genetic variability. In addition, interdisciplinary programs that study different aspects of *C. prunifera* populations (e.g., habitat quality, impact of extractive activity on individuals, and genetic diversity) throughout their distribution would be fundamental for the successful implementation of species conservation management.

Supporting information

S1 Fig. Genetic ontology assignment graph (GO). GO Annotations are summarized into three main categories: cellular location, biological process and molecular function for carnaúba (*Copernicia prunifera*). (DOCX)

S1 Table. Collection sites of the evaluated populations of Copernicia prunifera. (DOCX)

S2 Table. Similarity with proteins and Gene Ontology classifications obtained in blast2go for outlier SNPs putatively under selection in carnaúba (Copernicia prunifera). H_E = expected heterozygosity of the locus; F_{ST} = genetic divergence among groups of accessions estimated based on the locus; e-value = number of hits expected by chance (E = x 10); Sim (%) = BLASTX percentage of similarity between SNP tags and annotated proteins. (DOCX)

Acknowledgments

The first autor would like to thank Universidade Federal do Piauí for granting leave from doctorate degree at Universidade Estadual de Campinas. AA-P thanks São Paulo Research Foundation for a post-doctoral scholarship (FAPESP 2018/00036-9).

Author Contributions

Conceptualization: Marcones Ferreira Costa, Maria Imaculada Zucchi.

Data curation: Jonathan Andre Morales-Marroquín, Carlos Eduardo de Araújo Batista, Alessandro Alves-Pereira.

Formal analysis: Marcones Ferreira Costa, Jonathan Andre Morales-Marroquín, Carlos Eduardo de Araújo Batista, Alessandro Alves-Pereira.

Funding acquisition: Maria Imaculada Zucchi.

Investigation: Marcones Ferreira Costa, Fábio de Almeida Vieira.

Methodology: Marcones Ferreira Costa, Carlos Eduardo de Araújo Batista, Alessandro Alves-Pereira.

Project administration: Maria Imaculada Zucchi.

Resources: Marcones Ferreira Costa, Fábio de Almeida Vieira, Maria Imaculada Zucchi.

Supervision: Maria Imaculada Zucchi.

Writing - original draft: Marcones Ferreira Costa, Fábio de Almeida Vieira.

Writing – review & editing: Marcones Ferreira Costa, Fábio de Almeida Vieira, Maria Imaculada Zucchi.

References

- Chacón-Vargas K, García-Merchán VH, Sanín MJ. From keystone species to conservation: conservation genetics of wax palm Ceroxylon quindiuense in the largest wild populations of Colombia and selected neighboring ex situ plant collections. Biodivers Conserv. 2020; 29: 283–302. https://doi.org/10.1007/s10531-019-01882-w
- Soares LASS, Cazetta E, Santos LR, França DDS, Gaiotto FA. Anthropogenic disturbances eroding the genetic diversity of a threatened Palm tree: A multiscale approach. Frontiers in Genetics. 2019; 10:1090. https://doi.org/10.3389/fgene.2019.01090 PMID: 31788000
- Santos AS, Cazetta E, Dodonov P, Faria D, Gaiotto FA. Landscape-scale deforestation decreases gene fow distance of a keystone tropical palm, *Euterpe edulis* Mart (Arecaceae). Ecol Evol. 2016; 6:6586–6598. https://doi.org/10.1002/ece3.2341 PMID: 27777732
- Arruda GMT, Calbo MER. Effects of flooding on carnaúba growth, gas exchange and root porosity (Copernicia prunifera (Mill.) H.E. Moore). Acta Bot Bras. 2004; 18(3):219–224 https://doi.org/10.1590/ S0102-33062004000200002
- Silva RAR, Fajardo CG, Vieira FA. Mating system and intrapopulational genetic diversity of *Copernicia prunifera* (Arecaceae): a native palm from Brazilian semiarid. Genet Mol Res. 2017; 16 (3) 1:12. https://doi.org/10.4238/gmr16039764 PMID: 28973744
- Rocha TGF, Silva RAR, Dantas EX, Vieira FA. Phenology of Copernicia prunifera (Arecaceae) in a caatinga area of Rio Grande do Norte. Cerne. 2015; 21:673–681. https://doi.org/10.1590/01047760201521041758
- Tomchinsky B, Ming LC. As plantas comestíveis no Brasil dos séculos XVI e XVII segundo relatos de época. Rodriguésia. 2019; 70:1–16. https://doi.org/10.1590/2175-7860201970040
- Siegmund-Schultze M. A multi-method approach to explore environmental governance: A case study of a large, densely populated dry forest region of the neotropics. Environ Dev Sustain. 2021; 23(2):1539– 1562. https://doi.org/10.1007/s10668-020-00635-y
- Silva JMC, Leal IR, Tabarelli M. Caatinga: the largest tropical dry forest region in South America. Springer, New York, p 482 2017.
- Correia BEF, Alencar MM, Almeida- Júnior E B. Lista florística e formas de vida da vegetação de uma restinga em Alcântara, litoral ocidental do Maranhão, Nordeste do Brasil. Revista Brasileira de Geografia Física. 2020; 13(5):2198–2211.
- Correia BEF, de Almeida EB, Zanin M. Key Points about North and Northern Brazilian Restinga: a Review of Geomorphological Characterization, Phytophysiognomies Classification, and Studies' Tendencies. Bot Rev. 2020; 86(3), 329–337. https://doi.org/10.1007/s12229-020-09230-2
- Nascimento ELDL, Maia LC, Caceres MEDS, Lücking. Phylogenetic structure of lichen metacommunities in Amazonian and Northeast Brazil. Ecol Res. 2021; 36(3), 440–463. https://doi.org/10.1111/1440-1703.12206
- Almeilda JAS, Feitosa NA, Sousa LDC, Silva RNO, Morais RF, Monteiro JM, et al. Use, perception, and local management of *Copernicia prunifera* (Miller) HE Moore in rural communities in the Brazilian Savanna. J Ethnobiol and Ethnomed. 2021; 17: 1–13. https://doi.org/10.1186/s13002-021-00440-5 PMID: 33752732
- Sousa RFD, Silva RAR, Rocha TGF, Santana JADS, Vieira FDA. Etnoecologia e etnobotânica da palmeira carnaúba no semiárido brasileiro. Cerne. 2015; 21:587–594.
- Costa MF, Francisconi AF, Vancine MH, Zucchi MI (2022) Climate change impacts on the Copernicia alba and Copernicia prunifera (Arecaceae) distribution in South America. Braz J Bot. 2022; 45:807– 818. https://doi.org/10.1007/s40415-022-00801-8
- Ferreira CDS, Nunes JAR, Gomes RLF. Manejo de corte das folhas de Copernicia prunifera (Miller) HE Moore no Piauí. Revista Caatinga. 2013; 26: 25–30.
- Santos JRM, Vieira FA, Fajardo CG, Brandão MM, Silva RAR, Jump AS. Overexploitation and anthropogenic disturbances threaten the genetic diversity of an economically important neotropical palm. Biodivers Conserv. 2021; 30: 2395–2413. https://doi.org/10.1007/s10531-021-02200-z

- Mijangos JL, Pacioni C, Spencer PB, Craig MD. Contribution of genetics to ecological restoration. Mol Ecol. 2015; 24: 22–37. https://doi.org/10.1111/mec.12995 PMID: 25377524
- Luo Q, Li F, Yu L, Wang L, Xu G, Zhou Z. Genetic diversity of natural populations of *Taxus mairei*. Conserv Genet. 2021; 1–12. https://doi.org/10.1007/s10592-021-01403-9
- 20. Duarte JF, de Carvalho D, de Vieira FA. Genetic conservation of Ficus bonijesulapensis R.M. Castro in a dry forest on limestone outcrops. Biochem Syst Ecol. 2015; 59:54–62. https://doi.org/10.1016/j.bse. 2015.01.008
- Siqueira MVBM, Bajay MM, Grando C, Campos JB, Toledo JAM, Domingues GT, et al. Genetic diversity of reintroduced tree populations of Casearia sylvestris in Atlantic forest restoration sites. For Ecol Manag. 2021; 502:119703. https://doi.org/10.1016/j.foreco.2021.119703
- Nunes VV, Silva-Mann R, Souza JL, Calazans CC. Geno-phenotypic diversity in a natural population of Hancornia speciosa Gomes: implications for conservation and improvement. Genet Resour Crop Evol. 2021; 68: 2869–2882. https://doi.org/10.1007/s10722-021-01160-1
- 23. Díaz BG, Zucchi MI, Alves-Pereira A, de Almeida CP, Moraes ACL, Vianna SA, et al. Genome-wide SNP analysis to assess the genetic population structure and diversity of Acrocomia species. PLoS One. 2021; 16: 1–24. https://doi.org/10.1371/journal.pone.0241025 PMID: 34283830
- 24. Xia W, Luo T, Zhang W, Mason AS, Huang D, Huang X, et al. Development of high-density snp markers and their application in evaluating genetic diversity and population structure in *Elaeis guineensis*. Front Plant Sci. 2019; 1–11. https://doi.org/10.3389/fpls.2019.00130 PMID: 30809240
- Supple MA, Shapiro B. Conservation of biodiversity in the genomics era. Genome biology. 2019; 19: 1– 12. https://doi.org/10.1186/s13059-018-1520-3 PMID: 30205843
- Dalapicolla J, Alves R, Jaffé R, Vasconcelos S, Pires ES, Nunes GL, et al. Conservation implications of genetic structure in the narrowest endemic quillwort from the Eastern Amazon. Ecol Evol. 2021; 11: 10119–10132. https://doi.org/10.1002/ece3.7812 PMID: 34367563
- 27. Rajora OP. Population Genomics. Concepts, Approaches and Applications. Cham: Springer. 2019.
- Vieira FA, Sousa RF, Fajardo CG, Brandão MM. Increased relatedness among the neighboring plants from seedling to adult stages in carnaúba wax palm. Genet. Mol. Res. 2016; 15: 1–10. https://doi.org/10.4238/gmr15049100 PMID: 28002596
- Pinheiro LG, Chagas KPT, Freire ASM, Ferreira MC, Fajardo CG, Vieira FA. Anthropization as a determinant factor in the genetic structure of *Copernicia prunifera* (Arecaceae). Genet Mol Res. 2017; 16(3). https://doi.org/10.4238/gmr16039768 PMID: 28973747
- 30. Fajardo CG, Silva RAR, Chagas KPT, Vieira FA. Genetic and phenotypic association of the carnauba palm tree evaluated by inter-simple sequence repeat and biometric traits. Genet Mol Res. 2018; 17(3). https://doi.org/10.4238/gmr18018
- 31. Doyle JJ, Doyle JL. Isolation ofplant DNA from fresh tissue. Focus. 1990; 12(13):39-40.
- **32.** Poland JA, Brown PJ, Sorrells ME, Jannink J-L. Development of high-density genetic maps for barley and wheat using a novel two-enzyme genotyping-by-sequencing approach. PloS one. 2012; 7(2): e32253. https://doi.org/10.1371/journal.pone.0032253 PMID: 22389690.
- 33. Catchen JM, Amores A, Hohenlohe P, Cresko W, Postlethwait JH. Stacks: building and genotyping loci de novo from short-read sequences. G3: Genes genom genet. 2011; 1(3):171–82. https://doi.org/10.1534/q3.111.000240 PMID: 22384329.
- Catchen JM, Hohenlohe P, Bassham S, Amores A Cresko WA. Stacks: an analysis tool set for population genomics. Mol Ecol. 2013; 22(11):3124–3140. https://doi.org/10.1111/mec.12354 PMID: 23701397
- Luu K, Bazin E, Blum MG. pcadapt: an R package to perform genome scans for selection based onprincipal component analysis. Mol Ecol Resour. 2017; 17(1):67–77. https://doi.org/10.1111/1755-0998. 12592 PMID: 27601374.
- R Development Core Team. R: A language and environment for statistical computing. Vienna, Austria; 2018.
- Flanagan SP, Jones AG. Constraints on the F_{ST}-heterozygosity outlier approach. J Hered. 2017; 108 (5):561–73. https://doi.org/10.1093/jhered/esx048 PMID: 28486592.
- Gotz S, Garcia-Gomez JM, Terol J, Williams TD, Nagaraj SH, Nueda MJ, et al. High- through-put functional annotation and data mining with the Blsat2Go suite. Nucleic Acids Res. 2008; 36(10):3420–3435. https://doi.org/10.1093/nar/gkn176 PMID: 18445632
- **39.** Keenan K, McGinnity P, Cross TF, Crozier WW, Prodo PA. diveRsity: An R package for the estimation and exploration of population genetics parameters and their associated errors. Methods in ecology and evolution. 2013; 4(8):782–8.

- 40. Kamvar ZN, Brooks JC, Grünwald NJ. Novel R tools for analysis of genome-wide population genetic data with emphasis on clonality. Frontiers in genetics. 2015; 6:208. https://doi.org/10.3389/fgene.2015. 00208 PMID: 26113860.
- Jombart T, Ahmed I. adegenet 1.3–1: new tools for the analysis of genome-wide SNP data. Bioinformatics. 2011; 27(21):3070–1. https://doi.org/10.1093/bioinformatics/btr521 PMID: 21926124.
- **42.** Jombart T, Devillard S, Balloux F. Discriminant analysis of principal components: a new method for the analysis of genetically structured populations. BMC Genet. 2010; 11:94. https://doi.org/10.1186/1471-2156-11-94 PMID: 20950446
- **43.** Miller JM, Cullingham CI, Peery RM. The infuence of a priori grouping on inference of genetic clusters: simulation study and literature review of the DAPC method. Heredity.2020; 124(5): 269–280. https://doi.org/10.1038/s41437-020-0348-2 PMID: 32753664
- Nei M. Estimation of average heterozygosity and genetic distance from a small number of individuals. Genetics. 1978; 89(3):583–90. https://doi.org/10.1093/genetics/89.3.583 PMID: 17248844.
- 45. Kumar S, Stecher G, Tamura K. MEGA7: molecular evolutionary genetics analysis version 7.0 for bigger datasets. Mol Biol Evol. 2016; 33(7):1870–1874. https://doi.org/10.1093/molbev/msw054 PMID: 27004904
- 46. Brancalion PH, Oliveira GC, Zucchi MI, Novello M, van Melis J, Zocchi SS et al. Phenotypic plasticity and local adaptation favor range expansion of a Neotropical palm. Ecol Evol. 2018; 8:7462–7475. https://doi.org/10.1002/ece3.4248 PMID: 30151163
- 47. Rhode C, Bester-van der Merwe AE, Roodt-Wilding R. An assessment of spatio-temporal genetic variation in the South African abalone (*Haliotis midae*), using SNPs: implications for conservation management. Conserv Genet. 2017; 18: 17–31. https://doi.org/10.1007/510592-016-0879-5
- Gogvadze E, Buzdin A. Retroelements and their impact on genome evolution and functioning. Cell Mol Life Sci CMLS. 2009; 66(23):3727–3742. https://doi.org/10.1007/s00018-009-0107-2 PMID: 19649766
- **49.** Holanda SJ, Araújo FSD, Gallão MI, Medeiros Filho S. Impacto da salinidade no desenvolvimento e crescimento de mudas de carnaúba (*Copernicia prunifera* (Miller) HE Moore). Revista Brasileira de Engenharia Agrícola e Ambiental. 2011; 15: 47–52.
- Cordeiro EMG, Macrini CM, Sujii PS, Schwarcz KD, Pinheiro JB, Rodrigues RR, et al. Diversity, genetic structure, and population genomics of the tropical tree Centrolobium tomentosum in remnant and restored Atlantic forests. Conserv Genet. 2019; 20: 1073–1085. https://doi.org/10.1007/s10592-019-01195-z
- Barrett RDH, Hoekstra HE (2011) Molecular spandrels: tests of adaptation at the genetic level. Nat Rev Genet 12:767–780. https://doi.org/10.1038/nrg3015 PMID: 22005986
- Oliveira IS, Machado T, Banci KRDS, Almeida-Santos SM, Silva MJDJ. Genetic variability, management, and conservation implications of the critically endangered Brazilian pitviper Bothrops insularis. Ecol Evol. 2020; 10:12870–12882. https://doi.org/10.1002/ece3.6838 PMID: 33304500
- Kireta D, Christmas MJ, Lowe AJ, Breed MF. Disentangling the evolutionary history of three related shrub species using genome-wide molecular markers. Conserv Genet. 2019; 20:1101–1112. https://doi.org/10.1007/s10592-019-01197-x
- 54. Stojni S, Avramidou EV, Fussi B, Westergren M, Orlovi S, Matovi B, et al. Assessment of genetic diversity and population genetic structure of norway spruce (*Picea abies* (L.) Karsten) at its southern lineage in Europe: Implications for conservation of forest genetic resources. Forests. 2019; 10:258. https://doi.org/10.3390/f10030258
- 55. Laviola BG, Santos A, Rodrigues EV, Teodoro LPR, Teodoro PE, Rosado TB, et al. Structure and genetic diversity of macauba [Acrocomia aculeata (Jacq.) Lodd. ex Mart.] approached by SNP markers to assist breeding strategies. Genet Resour Crop Evol. 2021; 1–13. https://doi.org/10.1007/s10722-021-01295-1
- 56. Crispim BDA, Fernandes JDS, Bajay MM, Zucchi MI, Batista CEDA, Vieira MDC, et al. Genetic Diversity of *Campomanesia adamantium* and Its Correlation with Land Use and Land Cover. Diversity. 2021; 13: 160. https://doi.org/10.3390/d13040160.
- 57. Tong Y, Durka W, Zhou W, Zhou L, Yu D, Dai L. Ex situ conservation of *Pinus koraiensis* can preserve genetic diversity but homogenizes population structure. For Ecol Manag. 2020; 465:117820. https://doi.org/10.3390/f11010039
- Cobo-Simón I, Méndez-Cea B, Jump AS, Seco J, Gallego FJ, Linares JC. Understanding genetic diversity of relict forests. Linking long-term isolation legacies and current habitat fragmentation in Abies pinsapo Boiss. For Ecol Manag. 2020; 461:117947. https://doi.org/10.1016/j.foreco.2020.117947.
- Andrews KR, Good JM, Miller MR, Luikart G, Hohenlohe PA (2016) Harnessing the power of RADseq for ecological and evolutionary genomics. Nat Rev Genet 17:81. https://doi.org/10.1038/nrg.2015.28 PMID: 26729255

- **60.** Ferrante JA, Smith CH, Thompson LM, Hunter ME. Genome-wide SNP analysis of three moose subspecies at the southern range limit in the contiguous United States. Conser Genet. 2021; 1–13. https://doi.org/10.1007/s10592-021-01402-w
- Pádua JAR, Rocha LF, Brandão MM, Vieira FA, Carvalho D. Priority areas for genetic conservation of *Eremanthus erythropappus* (DC.) MacLeish in Brazil. Genet Resour Crop Evol. 2021; 68(6): 2483– 2494. https://doi.org/10.1007/s10722-021-01144-1
- 62. Fussi B, Westergren M, Aravanopoulos FA, Baier R, Kavaliauskas D, Finžgar D, Alizoti P, Božič G, Avramidou E, Konnert MW, Kraigher Het al. Forest genetic monitoring: an overview of concepts and definitions. Environm Monit Assessm. 2016; 188:493–504. https://doi.org/10.1007/s10661-016-5489-7 PMID: 27473107
- **63.** Fady B, Aravanopoulos FA, Alizoti P, Mátyás C, von Wühlisch G, Westergren M, Belletti P, Cvjetkovic B, et al. Evolution-based approach needed for the conservation and silviculture of peripheral forest tree populations. For Ecol Manage. 2016; 375:66–75. https://doi.org/10.1016/j.foreco.2016.05.015
- 64. Araújo LHB, Silva RAR, Dantas EX, Sousa RF, Vieira FA. Germinação de sementes da Copernicia prunifera: biometria, pré-embebição e estabelecimento de mudas. Enciclopédia Biosfera. 2013; 9 (17):1517–1528.
- 65. Brasil. Portaria N° 313, de 30 de dezembro de 2019. Brasília, 2019.
- 66. Miranda K, Neto MA, Sousa R, Coelho R. Manejo Florestal Sustentável em Unidades de Conservação de uso comunitário na Amazônia. Sociedade & Natureza. 2020; 32: 778–792. https://doi.org/10.14393/SN-v32-2020-51621