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Screening chloroplast regions in *Calophyllum brasiliense* (Calophyllaceae) for suitability for population genetic analyses

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Abstract The uniparental inheritance and conserved structure of the chloroplast genome among many angiosperms makes it suitable for constructing gene lineages and inferring population histories based on these relationships. However, the analysis of a single chloroplast region to solve population issues has not yielded the desired amount of variability due to the low mutation rate. Therefore, it is necessary to screen and carefully analyze chloroplast intergenic or intronic regions to select the most appropriate for studying the genetic variability in the chloroplast genome of a particular taxon. In this context, the aim of this

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work was to characterize 8 regions in *Calophyllum brasiliense* using 11 selected primer pairs and to evaluate their usefulness for phylogeographic analysis. Intergenic spacers *petA-psbJ*, *petG-trnP* and *rpl32-trnL* and the *trnL* intron demonstrated potentially informative sites. The combination of these 4 regions will allow the analysis of genetic population structure and the integration of historical aspects to inform strategies for conserving *C. brasiliense*, especially for highly impacted populations and those at risk of local extinction.

Keywords cpDNA · cpSSRs · Noncoding regions

Introduction

Calophyllum brasiliense (Calophyllaceae) is a native tree species from Central and South America. Its distribution range extends from southern Mexico to northeast Argentina (Reitz and Klein 1978; Rodríguez et al. 2009). It grows exclusively in hygrophyle forests and in lowlands that are almost permanently flooded (Marques et al. 2003). It offers many ecosystem services as a component of riparian forests, and several active compounds of medicinal interest have been recently isolated from different plant tissues. These compounds have inhibitory activity against some tropical pathogens that cause regional and worldwide diseases such as schistosomiasis, Chagas disease and tuberculosis (Gasparotto et al. 2005; Rea et al. 2013; Pires et al. 2014).

The chloroplast genome (cpDNA) is now the most widely used to understand phylogenetic issues among plant families and genera (Shaw et al. 2014). In the past, the use of cpDNA for population analysis was discouraged because of its low mutation rate within species. However, since

low-cost sequencing technologies have become routine, many phylogeographic studies in plant species have been based on a variable number of regions (Olmstead and Palmer 1994; Dumoulin-Lapègue et al. 1997; Huang et al. 2004). At first, the few cpDNA regions commonly used for higher taxonomic levels sometimes did not serve the desired purpose, but once the first complete genomes were sequenced, the conservative nature and cpDNA synteny (Palmer 1987; Dumoulin-Lapègue et al. 1997) allowed researchers to test many potential regions. Shaw et al. (2005, 2007, 2014) compared the variability rates of chloroplast regions among distinct clades and found that the cpDNA could be much more informative than earlier thought. Nevertheless, the most suitable chloroplast regions for a population genetic analysis usually will depend on the target species. In this context, the aim of this study was to characterize 8 cpDNA regions among C. brasiliense samples derived from populations from Argentina, Paraguay and Mexico by using 10 previously reported primer pairs and 1 specific primer pair to evaluate their utility in phylogeographic analysis, information that could also help in conservation decisions within the most reduced populations.

Materials and methods

Material and study site

From 2 to 10 individuals from each of the populations in three regions were sampled and stored in Ziploc bags: (1) Misiones (San Ignacio POP—SI) and Corrientes (Puerto Valle POP—PV; Rincón Ombú POP—RO) Argentinean Provinces; (2) Misiones Paraguayan Department (Ayolas POP—AY and Isla Yacyretá POP—IY); and (3) a Mexican sample from the Ecological Reserve Jaguaroundí Park in Veracruz-Llave (Mexico POP—MX). Table 1 summarizes the geographic coordinates and the number of individual collected per site.

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DNA extraction

Once leaves were completely dried, total genomic DNA was isolated from these tissues following the protocol of Stange et al. (1998) with modifications made by Percuoco et al. (2014), which involved the addition of antioxidant reagents (2% w/v polyvinylpyrrolidone [PVP], 5 mM ascorbic acid, 4 mM sodium diethyldithiocarbamatetrihydrate [DIECA] and 1.2% β -mercaptoethanol) to the digestion buffer and the extension of incubation steps (digestion step: 0.5 vs. 3 h; precipitation step: 0.5 vs. 1 h).

cpDNA amplification and sequencing

Seven chloroplast intergenic regions were amplified: *trnHpsbA*, *psbI-trnS* (*ccmp2*), *trnS-trnG-trnG*, *psbC-trnS*, *petApsbJ*, *petG-trnP* and *rpl32-trnL*. The eighth region was the *trnL* intron. Eight heterologous and 2 universal primer pairs obtained from the literature and an intraspecific pair that we designed were used for the PCR reactions (Table 2). The PCR mix composition and cycling profiles are detailed in Tables 3 and 4, respectively. PCR products were purified using GFX PCR DNA #28-9034-70 (GE) and sequenced by Macrogen Inc. using both forward and reverse primers.

Sequence analysis

The quality control of each electropherogram and all sequence editing were done using BioEdit v.7.0.9.0 (Hall, 1999). Consensus sequences were created for each individual, and their identity was verified using BLASTn 2.2.25 (Zheng et al. 2000).

C. brasiliense cpDNA descriptive data

The percentage sequence coverage for *C. brasiliense* cpDNA was estimated using the genome of *Jatropha curcas* (NC_012224) as reference. We chose this genome arbitrarily among four available within the order Mapighiales. We also mapped the approximate physical position of

 Table 1 Location of Calophyllum brasiliense populations sampled

POP ID	Locality	Geographic reference	Ν
San Ignacio (SI)	San Ignacio (Misiones—ARG)	27°16′34.4″S-55°34′11.9″W	11
Rincón Ombú (RO)	Ituzaingó (Corrientes—ARG)	27°24′54.42″S-56°29′43″W	11
Puerto Valle (PV)	Ituzaingó (Corrientes—ARG)	27°35′23.84″S-56°31′34.17″W	10
Yacyretá Island (IY)	Misiones (PY)	27°24v53.88"S-56°45'21.25"W	3
Ayolas (AY)	Misiones (PY)	27°24′6.48″S–56°51′17.75″W	2
Jaguaroundi Reserve (MX)	Coatzacoalcos (Veracruz Llave- MX)	18°7'20.27"N-94°21'45.15"W	6

POP ID identity of the population, N number of samples collected at each site

1		T			
Region	Primers sequences $(5' - > 3')$	Reference data		Data for C. I study)	brasiliense (present
		References	Expected size	Sequence length	GenBank Accession
trnH(GUG)-psbA	ACT GCC TTG ATC CAC TTG GC CGA AGC TCC ATC TAC AAA TGG	Hamilton (1999)	495 bp (Nicotiana tabacum)	409	°1
psbl-trnS-(GCU)	GAT CCC GGA CGT AAT CCT G	Weising and Gardner (1999)	189 bp (<i>Nicotiana tabacum</i> , dicotyledons)	194	ام
	ATC GTA CCG AGG GTT CGA AT	H omilton (1000)	QIA hrs (N references)	600 ^a	υ
	GAA CGA ATC ACA CTT TTA CCAC		of a mount	000	
trnS(GCU)-tmG-intron- tmG	AAC TCG TAC AAC GGA TTA GCA ATC	Shaw et al. (2007)	1434 bp (angiosperms, Eurosid I)	1897	KJ093667
	GAA TCG AAC CCG CAT CGT TAG				
psbC-trnS(UGA)	GGT CGT GAC CAA GAA ACC AC	Demesure et al. (1995)	1680 bp (Quercus robur, universal)	1514	JN368484, JN596278
	GGT CCG AAT CCC TCT CTC TC				
trnL intron	CGA AAT CGG TAG ACG CTA CG	Taberlet et al. (1991)	200-1000 bp (universal, angiosperms)	611	JQ219659
	GGG GAT AGA GGG ACT TGA AC				
	AAA ATG GGC AAT CCT GAG C	Percuoco et al. (2015)	450 bp (C. brasiliense)	450	KF854307– KF854309
	ATG GGA CTC TAT CTT TAT TCT CG				
petA-psbJ	GGA GAT GCA GAG ATA GTA C	Huang et al. (2004)	752 bp (Trochodendron araliodes)	1100 ^b	٥I
	CTC TTT GGT TGA TAG GTA CTG				
	ATA GGT ACT GTA ACT GGT ATT	Shaw et al. (2007)	734–1261 bp (angiosperms)	1137-1174	KJ093668, KJ093669
	GGT CTA ATT CCT ATA ACT TTG GC				
petG-trnP	GGG ATG TGG CGC AGC TTG G	Huang et al. (2004)	350 bp (T. araliodes)	449	KJ093665, KJ093666
	AAC AGT TTG AGA AGG TTC AAT T				
rpl32-trnL(UAG)	CTG CTT CCT AAG AGC AGC GT	Shaw et al. (2007)	543-1417 bp (angiosperm)	811-901	KJ093670, KJ093671
	CAG TTC CAA AAA AAC GTA CTT C				
Total length				7022–7149	
For each cpDNA region, ex	pected size and tested species, based on published	l sources, are provided. Also listed	1 are sequences lengths and GenBank access	ion number for	C. brasiliense regions
"I wo other bands (1500, 15	100 bp) were always observed. Frimer not select	ed			

^bThese primers amplified this region only from some individuals

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Table 3 PCR mix composition for each of the 8 noncoding cpDNA regions amplified in Calophyllum brasiliense

Components (initial concentrations)	Final concent	Final concentrations										
	psbC-trnS	trnH-psbA	trnS-trnG	Spacers ^a	trnL intron	psbI-trnS (ccmp2)						
Buffer Taq SO ₄ (NH ₄) ₂ (10 ×)	$1 \times$	1×	$1 \times$	$1 \times$	1×	1×						
MgCl ₂ (25 mM)	2.5 mM	2.5 mM	2 mM	2 mM	2 mM	2 mM						
dNTPs (2 mM)	0.1 mM	0.2 mM	0.2 mM	0.2 mM	0.2 mM	0.2 mM						
Primer F (10 µM)	0.2 µM	0.3 μΜ	0.4 µM	0.1 µM	0.2 µM	0.2 µM						
Primer R (10 µM)	0.2 µM	0.3 μΜ	0.4 µM	0.1 µM	0.2 µM	0.2 µM						
BSA (10 mg/mL)	5 μg/μL	2.5 μg/μL	2.5 μg/μL	-	-	-						
TaqADN pol (5 U/µL)	1 U/25 μL	1 U/25 μL	1 U/25 μL	1 U/25 μL	1 U/25 μL	1 U/25 μL						
ADN (50 µg/µL)	5 μg/μL	5 μg/μL	5 μg/μL	5 μg/μL	5 μg/μL	5 μg/μL						
Ultrapure water to	25 µL	25 µL	25 µL	25 µL	25 µL	25 μL						

^atrnS-trnG-trnG, petA-psbJ, petG-trnP, rpl32-trnL

the 8 reported regions. Three other chloroplast sequences of *C. brasiliense* available in GenBank were included in this cpDNA illustration generated with DNA Plotter, free on-line software available at http://www.sanger.ac.uk/sci ence/tools/dnaplotter (Carver et al. 2009).

Results

The 8 regions amplified showed single fragments of approximately the expected size in a variable number of individuals and populations. The editing and analysis of the obtained sequences allowed the characterizing of different noncoding cpDNA regions in *C. brasiliense*. Three intergenic spacers (see below) and the *trnL* intron had putative informative sites for *C. brasiliense*. The length of the sequences obtained and the GenBank accessions for each of the 8 regions analyzed are described in Table 2.

- (1) *trnH-psbA* From the 500-bp products separated by agarose gel electrophoresis, a 409-bp consensus sequence was obtained with several microsatellite motifs included in it. These mono- and dinucleotide tandem repeats made the sequencing reaction difficult in some samples, even using microsatellite capillary electrophoretic conditions. The only alignment obtained showed 100% identity between 2 individuals from different Argentinean populations (SI and RO).
- (2) *psbI-trnS* (*ccmp2*) Amplicons of \approx 200-bp were obtained from samples from 2 Argentinean populations (SI, RO), whereas a smaller band size was observed in the Mexican samples. A 194-bp sequence was edited based on the SI and RO populations. This intergenic fragment corresponded to a chloroplast microsatellite that had an interrupted

motif in *C. brasiliense*. The sequence shared 63% identity with *Nicotiana tabacum* (Z00044) and maximum identity (73%) with *Mallotus paniculatum* (AY159390). The sequence could not be submitted to GenBank due to the minimal length of 200-bp required by the database (see http://www.ncbi.nlm. nih.gov/genbank/submit_types).

- (3) trnS-trnG and trnS-trnG intron-trnG Two primer pairs were tested to amplify this region: (a) Hamilton's (1999) primers amplified the expected band size of \approx 600-bp and two extra bands; and (b) The primers of Shaw et al. (2007) that amplified a unique, expected band of \approx 2000-bp. From the latter, an 1897-bp consensus sequence was edited based on an individual from the Isla Yacyretá population, which shared 86% identity with Jatropha curcas. However, most accessions in the BLAST alignment showed coverage only up to 43% of the query sequence, except when aligned with Licania heteromorpha (KJ414481), another tree species from the family Chrysobalanaceae within the order Malpighiales, with which coverage was 66%. Four more samples (9%) were sequenced and partially aligned, but no polymorphic site was identified.
- (4) *psbC-trnS* A unique amplicon of \approx 1500-bp was amplified in all individuals from SI and RO (22/ 51%) but failed to be amplified among samples derived from the other populations (21/49%). Once the reliable electropherograms were edited, we obtained a 1514-bp consensus sequence, from which 1307 bp matched the *psbC* gene 3' end encoding photosystem II CP43 chlorophyll apoprotein, and only 207-bp corresponded to the noncoding intergenic portion in *C. brasiliense*. Simultaneously, we carried out an additional digestion assay in this

 Table 4 PCR profiles for 8 noncoding cpDNA regions amplified in Calophyllum brasiliense

Step	psbC—trnS		trnH-psbA		trnS-trnG		rpl32-trnL		Spacers ^a		trnL intron		psbI-trnS (ccmp2)	
	Т	t	Т	t	Т	t	Т	t	Т	t	Т	t	Т	t
Initial denaturation	94	4 min	96	5 min	96	5 min	96	5 min	80	5 min	94	5	94	5 min
Denaturation	94	45 s	94	30 s	94	45 s	94	45 s	95	1 min	93	1	94	1 min
Annealing	57	45 s	62	30 s	55	1 min	55	1 min	50	1 min	62	1	60	1 min
Elongation	72	2 min	72	30 s	72	1 min	72	1.5 min	65	4 min	72	2	72	2 min
No. of cycles	30		35		30		30		35		30		35	
Final elongation	72	10 min	-	-	72	10 min	72	10 min	65	10 min	72	4	72	4 min
Final hold	4	∞	4	∞	4	∞	4	∞	4	∞	4	∞	4	∞

T temperature (°C), t time

^atrnS-trnG-trnG, petA-psbJ, petG-trnP, rpl32-trnL

region with MboI restriction endonuclease and compared the profiles after standard agarose electrophoresis. In this assay, 30 individuals of the SI and RO POP were analyzed, showing a unique profile for both populations.

- (5) *trnL intron* An amplicon of \approx 700-bp was obtained in all populations. A first 611-bp consensus sequence was obtained for this region, but difficulties in sequencing of most of the amplicons required us to design a new specific primer pair for the species, which improved the electropherogram quality. The new consensus sequence was 450-bp long, and the alignment of the 43 sequences (100%) showed 4 informative sites, 2 SNPs, a 2-bp inversion and 1 indel 7-bp long. Three haplotypes were then identified. SSRs previously reported for this intron in other species (Hale et al. 2004) were interrupted and monomorphic in *C. brasiliense*.
- (6) petA-psbJ Two primer pairs were assayed to amplify this region. Both pairs amplified the expected band of \approx 1200-bp, but the primers designed by Huang et al. (2004) yielded no amplicon in several individuals (7/18%); thus, we chose the primers published by Shaw et al. (2007). The primer petA yielded overlapped products and low signal after 140-bp in all individuals analyzed, although the reverse primer *psbJ* allowed us to obtain 1200-bp readings. Once editing was completed, sequences from 1137 to 1174-bp long were aligned. Two indels, a substitution, and a $(T)_n$ microsatellite allowed discriminating 2 haplotypes exclusive to the southernmost populations and Mexican populations respectively.
- (7) *petG-trnP* An amplification product of \approx 500-bp was obtained from all samples; 19 (44%) of these were sequenced and resulted in a 449-bp consensus sequence. An 11-bp inversion was the only

polymorphism found, which discriminated 2 monohaplotypic clusters, 1 from the Mexican population samples and the others from the other 5 populations.

(8) rpl32-trnL Bands of \approx 1000-bp were observed in all populations. Once the sequences were edited, we obtained a fragment in the range of 811–890-bp. The variation in length was partially due to the inferior quality of the consensus sequence obtained and to a variable number of the (AAATT)_n motifs that differentiated the Argentinean and Paraguayan from the Mexican populations.

Three other partial *C. brasiliense* coding sequences for cpDNA are available in GenBank. These correspond to *ndhF* (HQ331853), *matK* (HQ331550) and 8 accessions of *rbcL* (JQ591092–JQ591094; KC493366–KC493368; KC570911; KF981208). The alignment including all 8 *rbcL* sequences showed complete identity among them.

The 8 sequences reported here totaled 7149-bp, which increased to 9529 bp when *ndhF*, *matK* and *rbcL* (JQ591094) available sequences were added. Considering that the *Jatropha curcas* genome is 163,856-bp, approximately 6% of *C. brasiliense* cpDNA is now accessible, of which $\approx 4.5\%$ corresponds to noncoding regions. The 11 regions (our 8 regions plus the 3 previous deposited in GenBank) now available for *C. brasiliense* were mapped over the reference genome (Fig. 1).

Discussion

All 8 regions analyzed in this work have been previously used in studies of several tree species (Dumoulin-Lapègue et al. 1997; Huang et al. 2004; Ramos et al. 2007), showing different informative rates. Eight heterologous, 2 universal and 1 specific primer pair developed in this work allowed amplicons to be obtained from many of the *C. brasiliense*

samples analyzed. Three intergenic regions (*petA-psbJ*, *petG-trnP*, *rpl32-trnL*) and the *trnL* intron were informative for the species. Particularly, two indels, a substitution and a mononucleotide cpSSR in *petA-psbJ*, an 11-bp inversion in *petG-trnP* and a 5-bp repeat motif within *trnL-rpl32* were the polymorphisms that allowed 2 haplotypes to be identified, 1 common to the southernmost *C. brasiliense* populations and the other restricted to the samples derived from Mexico. We can thus affirm that they are suitable chloroplast noncoding regions for population genetic approaches in *C. brasiliense*.

In the case of the *trnL* intron a 7-bp indel, one 2-bp inversion and 2 transversions were analyzed. These discriminated 3 haplotypes among *C. brasiliense* populations, 1 common haplotype distributed in Argentina and Paraguay, and 2 other haplotypes, exclusive to Paraguayan and Mexican populations respectively. Therefore, this region was used in a phylogeographic analysis (Percuoco et al. 2015). The *trnL* intron was used for several phylogenetic analyses (Hale et al. 2002; Hollingsworth et al. 2011) although some authors have discarded it for intraspecific studies due to its low resolution at low taxonomic levels (Shaw et al. 2005). This region was also considered as a putative barcode region in plants, because of its high conservation rate within species and the robustness of the amplification process (Taberlet et al. 2007).

The primers used to amplify the *psbC-trnS* intergenic spacer in C. brasiliense showed that this region contains a long *psbC* fragment, while the spacer is very short for the species. Although we did not sequence the fragment in many samples, the enzymatic digestion revealed no difference among the C. brasiliense samples, in accordance with the expected low level of variation because of the selection pressure exerted over coding regions. However, Ramos et al. (2007) used psbC-trnS in an intraspecific study of 17 populations of Hymenaea stigonocarpa from Brazil and found 23 haplotypes based on 13 substitutions and 4 indels in the alignment of a 524-bp fragment. Unfortunately, these sequences were not deposited in GenBank, and the electropherograms were not included as supplementary material for comparison with our results to enrich the present analysis.

The *trnH-psbA* intergenic spacer showed low-quality electropherograms due to several polyA/T repeats. There is, however, only one published phylogeographic analysis of *C. brasiliense* populations from Brazil where the use of this intergenic spacer seems to have been successful (Salgueiro et al. 2011). Although these authors used the same primers that we tested in this work, they reported a 263-bp sequence, a fragment much shorter than the one we obtained (409 bp). Surprisingly, they identified 28 informative sites within the 263 bp of the *trnH-psbA* intergenic spacer based on 192 individuals of *C. brasiliense*; however,



Fig. 1 Sequences of *C. brasiliense* cpDNA herein described mapped over the *Jatropha curcas* genome (NC_012224) used as reference

again, the electropherograms are not available to compare with our data.

The high number of repeats observed in the *trnH-psbA* region of *C. brasiliense* is in accordance with the common mono- and dinucleotide SSRs usually found in cpDNA, but these informative polymorphisms could easily arise as parallelism or homoplasies (Petit and Vendramin 2007), leading to wrong phylogeographic conclusions. Therefore, this type of mutation is often discarded in phylogeographic analysis (Huang et al. 2004; Lihová et al. 2010).

The *trnH-psbA* spacer has also been proposed as a potential DNA barcoding region (Kress et al. 2005; Chase et al. 2007), its high variability in length across larger angiosperm families and the ambiguity added by the obligatory manual editing because of multiple cpSSRs, as those observed in *C. brasiliense*, make it a problematic if it were the standard plant barcode alone (Chase et al. 2007).

Primer mismatches, a frequent problem when heterologous or universal primers are used, likely were responsible for the difficulties in the sequencing reaction for the *trnHpsbA*, *psbI-trnS*, *trnS-trnG-trnG* and *psbC-trnS* intergenic spacers amplified in *C. brasiliense*.

The chloroplast genome remains the most appropriate target in plant phylogeography studies due to its uniparental inheritance and its haploid nature. In this sense, cpDNA "records" historical fluctuations in populations for each species with high fidelity, providing a historical glance to improve conservation decisions. Although many cpDNAs have been completely sequenced (Malé et al. 2014), various nonmodal tropical tree species remain to be studied. Moreover, universal primers are frequently tested in several species but only in a few individuals within each species; therefore, the intraspecific performance and features of the primers are unknown. Particularly for our native tree, all cpDNA regions were amplified with diverse levels of difficulty in the subsequent analysis of the sequences obtained. We improved our results when we designed a pair of primers for C. brasiliense as we showed for the trnL intron. Thus, we strongly recommend designing specific primer pairs to ensure the quality of the data even for those regions difficult to analyze. Finally, using more than one cpDNA region increases the number of informative sites and takes advantage of genome fitness. Thus, the 8 regions described in this work contribute to the knowledge of C. brasiliense cpDNA, but based on our results, we suggest petA-psbJ, petG-trnP, rpl32-trnL and the *trnL* intron analyzed in this work as suitable noncoding regions that could be useful for phylogeographic studies of this species.

Conclusions

All 8 chloroplastic regions were amplified, and a sequence of 7149-bp, which represents approximately 6% of *C. brasiliense* cpDNA is now available in GenBank. Three intergenic regions (*petA-psbJ, petG-trnP, rpl32-trnL*) and the *trnL* intron were informative and will be useful for population analyses of *C. brasiliense*.

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