

Molecular characterization and genetic relationships among most common identified morphotypes of critically endangered rare Moroccan species *Argania spinosa* (Sapotaceae) using RAPD and SSR markers

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Abstract

• The objective of this work is the molecular characterization of most common identified morphotypes of critically endangered rare Moroccan species *Argania spinosa*.
• Eighteen RAPD markers and twenty SSR markers have been assayed in 38 argan tree accessions from the three most commonly identified morphotypes: oval, spherical and spindle fruit types.
• A total of 140 polymorphic RAPD bands were detected out of 146 bands. The number of presumed alleles revealed by the SSR analysis ranged from 1 to 5 alleles per locus with a total number of 32 alleles identified. Results demonstrated an extensive genetic variability within the tested argan accessions. RAPDs presented a high level of polymorphism and greater information content than SSRs.
• Our results could indicate that *Vitellaria paradoxa* is genetically closer to argan than *Manilkara huberi*. In addition, the correlation between the clustering based on RAPD and SSR markers were in general low. The observed clustering could be better explained according to geographic proximity than morphotype. For this reason the traditional morphological characterization of the argan accessions in morphotypes (according to fruit phenotype) seem not to be in correlation to the real genetic background (genotype) of this specie. The implications of these results in the creation of effective germplasm core collection in argan have been also discussed.

Résumé – Caractérisation moléculaire et relation génétique entre les morphotypes les plus connus d'une espèce marocaine rare et menacée, *Argania spinosa* (Sapotaceae) à l'aide de marqueurs RAPD et SSR.

• L'objectif de ce travail consiste en la caractérisation moléculaire et l'étude des relations génétiques entre des morphotypes identifiés chez une espèce marocaine rare et menacée, l'arganier (*Argania spinosa*).
• Pour cela, dix-neuf marqueurs RAPD et vingt marqueurs SSR ont été testés dans 38 accessions d'arganier des trois morphotypes du fruit identifiés : ovale, sphérique et broche.
• En utilisant la technique d'amplification au hasard de l'ADN (RAPD), 140 fragments polymorphes ont été détectés parmi les 146 amplifiés. Le nombre d'allèle révélé par l'analyse des marqueurs microsatellites (SSRs) varie de un à cinq allèles par locus avec un total de 32 allèles identifiés. Les résultats démontrent une grande variabilité génétique au sein des accessions d'arganier testées. Les marqueurs RAPD présentent un taux de polymorphisme et une hétérozygotie attendue supérieurs aux SSR.
• Les résultats indiquent également que *Vitellaria paradoxa* est plus proche génétiquement de l'arganier que *Manilkara huberi*. La corrélation entre les groupements observés dépend plus de la proximité géographique que de la classification selon la forme du fruit (morphotypes). La classification traditionnelle par morphotypes n'est donc pas en accord avec l'analyse génotypique chez cette espèce. L'apport de ces résultats pour la création d'une collection de référence chez cette espèce est discuté.

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1. INTRODUCTION

Argan [*Argania spinosa* (L.) Skeels], an endemic tree species from Southwest Morocco, belongs to the *Sideroxyleae* tribe of the tropical family Sapotaceae. It is the only representative of this family in Morocco. *Argania* is a monospecific genus with *A. spinosa* as the unique representative (Swenson and Anderberg, 2005). *A. spinosa* develops under the hot arid Mediterranean climate and has very different distribution and ecological preferences compared to the other species of the *Sideroxylon* tribe (Emberger, 1939). It occupies a wide distribution range (800 000 ha) and has a great socio-economical and ecological role in these arid and semi-arid zones. The argan tree is valued for its highly unsaturated edible oil, wood, and forage. In addition, thanks to its powerful root system, it is valuable in land reclamation and fighting against desertification. However, a great zoo-anthropoc action led to the absence of a natural regeneration (Benchakroun and Buttoud, 1989). The three most commonly identified morphotypes of argan were oval, spherical and spindle fruit types (Bani-Aameur, 2004) (Fig. 1).

This species is genetically and molecularly very poorly known (Majourhat et al., 2007). Traditionally, characterization and identification of variability has been based on morphological (Bani-Aameur, 2004; Chernane et al., 2000) and biochemical (Chernane et al., 2000; El Mousadik and Petit, 1996a) markers. It has been suggested to use fruit form as criterion for varietal selection in this species (Bani-Aameur, 2004). However, such markers are not always available for analysis, and are affected by environmental conditions. DNA marker technology offers several advantages over the sole use of conventional markers. In the case of argan, only one work studying the genetic variability of argan populations using DNA amplification and restriction was published until now (El Mousadik and Petit, 1996b).

Random amplified polymorphic DNA (RAPD) markers are based on the PCR amplification of random locations in the genome (Messaoud et al., 2007; Welsh and McClelland, 1990). These markers are especially used in the molecular characterization of rare species including argan which are poorly known from the molecular point of view. The most crucial factor for this technique is the DNA quality and concentration offering the advantage of rapid screen for polymorphisms (Dax et al., 1993; Gérard et al., 2006). On the other hand, Simple sequence repeat (SSR) markers (microsatellites), also based on the PCR technique, are currently becoming the markers of choice for genetic fingerprinting studies because of their high polymorphism, co-dominance, multiallelism, abundance and uniform dispersion in plant genomes (de-Lucas et al., 2008; Gupta et al., 1996). However, to date, none of these markers have been described in argan although the ability to use the same microsatellite primers in different plant species from the same family, called transportability or transferability, has been well described (Decroocq et al., 2003).

The objective of this work is to present an appropriate set of RAPD and SSR markers suitable for the characterization of most common identified morphotypes of critically endangered rare Moroccan species *Argania spinosa*.



Oval

Spherical

Spindle

Figure 1. Most common identified morphotypes of the critically endangered rare species argan (*Argania spinosa* Skeel) assayed in this study: oval, spherical and spindle fruit types.

2. MATERIAL AND METHODS

2.1. Plant material and DNA extraction

Leaf material from 38 argan tree accessions growing in natural conditions was collected on May 2006 at Essaouira (Southwest of Morocco). The three most commonly identified morphotypes of argan tree were sampled, oval, spherical and spindle fruit types (Fig. 1), together with some accessions of intermediate morphotypes.

Well expanded young leaves were used for DNA isolation using the procedure described by Doyle and Doyle (1987) using CTAB extraction buffer. DNA was quantified using a Biophotometer (Eppendorf, Barcelona, Spain).

2.2. Development of RAPD markers

Nineteen RAPD universal primers purchased from Operon Technologies (Huntsville, USA) were assayed (see supplemental data available Online only, Tab. A). Amplifications were carried out in 20 μ L total volume containing 1 \times Buffer (New England, Biolabs, Ipswich, USA), 1 mM MgCl₂, 0.16 mM of dNTP, 0.4 μ mol of primer, 1.0 unit of *Taq* DNA polymerase (Biolabs), and 4 ng templates DNA. The amplification program consisted of a step of DNA melting of 4 min at 94 $^{\circ}$ C, followed by 35 cycles of 94 $^{\circ}$ C for 1 min, 35 $^{\circ}$ C for 1 min, and 72 $^{\circ}$ C for 1 min, and a final elongation step of 72 $^{\circ}$ C for 10 min. The amplified products were resolved in 2% agarose gels stained with ethidium bromide and visualized with UV transmitted light. A 1 Kb DNA Ladder (Invitrogen Life Technologies, Barcelona, Spain) was used as molecular size standard. RAPD amplifications were repeated at least twice in order to check the reproducibility of bands.

2.3. Development of SSR markers

Twenty SSRs developed in two different species inside the *Sapotaceae* family, 12 primer-pairs in *Manilkara huberi* (Ducke)

Standl (Azevedo et al., 2005) and 8 additional ones in *Vitellaria paradoxa* C.F. Gaertn (Cardi et al., 2005) were tested for their transferability to amplify microsatellite loci in the 38 argan samples studied (Tab. I). PCRs were performed in 20 μ L mix containing 0.18 μ M each primer, 1 \times *Taq* buffer (Biolabs), 2.5 mM MgCl₂, 0.1 mM of each dNTP, 1 unit of *Taq* DNA polymerase (Biolabs), and approximately 5 ng of DNA. The PCR program consisted of an initial melting step (94 °C, 4 min) followed by 30 cycles (94 °C, 30 s, annealing temperature, 30 s and 72 °C, 1 min) and a final extension step (72 °C, 7 min). PCR products were separated using 3% Metaphor Agarose gel electrophoresis (Biowittaker, Maine, USA). To confirm the results, SSR products smaller than 5 bp were also separated using 10% polyacrylamide gel electrophoresis (PAGE). A 1 kb DNA Ladder was also used as molecular size standard.

2.4. Data analysis

Polymorphic alleles were scored as present or absent (1/0). DNA band scoring was analyzed using GeneTools gel analysis software of SYNGENE (Beacon House, Nuffield Road, Cambridge, UK). The genetic diversity characterization was based on the number of alleles per locus in RAPDs and SSRs. The average polymorphic information content (PIC) was calculated for RAPD markers across assay units by applying the formula given by Powell et al. (1996).

The “genetic information content” of the codominant SSR markers was estimated by the observed heterozygosity (H_o), the expected heterozygosity (H_e) (Nei, 1978) using the POPGENE program (<http://www.ualberta.ca/~fyeh/>) and by power of discrimination (PD) (Kloosterman et al., 1993). On the other hand, mean character difference distances were calculated for all pairwise comparisons with the MEGA4 test (<http://www.megasoftware.net/>), which was used to construct UPGMA dendrograms (cluster analysis) (Nei and Li, 1979) depicting the phenetic relationship among the different accessions. Relative support for the branches in each dendrogram was assessed with 2000 replicates of UPGMA bootstrap.

3. RESULTS AND DISCUSSION

The analysis of the 19 RAPD primers showed that all primers generated polymorphic and reproducible patterns. A total of 140 polymorphic RAPD bands were detected out of 146 bands with a size range between 322 and 2009 bp. The number of RAPD bands detected by each primer depends on primer, sequence and the extent of variation in specific argan genotype. The number of bands varied from 2 (for OPA-16) to 14 (for OPW-13), with an average of 7.8 bands per primer. The mean PIC score over all loci was 0.84, ranging from 0.35 in OPA-16 to 0.96 in OPW-13 (see supplemental data, Tab. A and Fig. A).

In addition, amplification was successful with 4 of the 12 (33%) SSR markers developed in *Manilkara huberi* and 7 of the 8 (87%) SSR markers developed in *Vitellaria paradoxa*. A total of 32 polymorphic bands were scored for argan accessions. The number of presumed alleles revealed by the SSR analysis ranged from one to five with a size range of amplified bands between 129 and 278 bp. The observed heterozygosities ranged from 0.00 in Mh04, Mh07, Mh12, mVpCIRB03, and mVpCIRE04 to 0.60 in mVpCIRF05 SSR markers with

an average of 0.18. Mean expected heterozygosity was 0.59, ranging from 0.00 in Mh12 to 0.80 in mVpCIRH04 (Tab. I and supplemental data Fig. A). In general the low heterozygosity values (observed and expected) showed in this study can be a consequence of the low number of argan accessions assayed which can also be genetically close according with the geographic proximity of the samples collected. For all the loci studied the expected heterozygosity was greater than the observed heterozygosity, implying the presence of null alleles or a deficit of heterozygotes due to non-random mating. The mean PD value for all loci was of 0.45. The most informative loci were mVpCIRF11 and mVpCIRH04 with a PD of 0.83 and the least informative were Mh12 and mVpCIRE04 with a PD of 0.0 (Tab. I). These PD values are lower than those obtained in other assays with SSR (Shiran et al., 2007) as a consequence of the low polymorphism observed in the application of SSR in argan from relatively phylogenetically distant *Sapotaceae* species such as *Manilkara huberi*, as has been previously described in *Prunus* species by Martínez-Gómez et al. (2003). In the case of *Vitellaria paradoxa* PD are in general higher with higher level of SSR polymorphism indicating that this species is genetically closer to argan than *Manilkara huberi*. In addition, these PD values are in general lower than PIC values obtained in the other assays with RAPD as a consequence of the lower polymorphism observed in the application of SSR in comparison with RAPD, as has been previously reported in different species (de-Lucas et al., 2008; Lachenaud and Zhang, 2008; Messaoud et al., 2007; Shiran et al., 2007).

The level of polymorphism was slightly smaller in argan to that reported by Azevedo et al. (2005) and Cardi et al. (2005) in the original species in which the SSR markers were developed, i.e. *Manilkara huberi* and *Vitellaria paradoxa* respectively. However, the range of the amplified band sizes in argan was similar to those reported in the original species in which the SSR markers were developed. Overall, the results of the obtained polymorphism suggest a high degree of transportability of SSR in *Sapotaceae* family. These results agree with reports on the successful utilization of these markers in different species from the same genus (Martínez-Gómez et al., 2003) and the successful utilization of these markers in different species from the same family (Decroocq et al., 2003; de-Lucas et al., 2008).

Differences in amplification success for SSR markers observed among species are due to the genetic variability between the different species in *Sapotaceae* family where the SSRs were developed. A decrease in the amount of polymorphic SSR markers would be expected as genetic distance increases from the designated anchor species, as has been previously reported in *Prunus* by Martínez-Gómez et al., (2003). It has been shown that closely related species are more likely to share microsatellite priming sites than more distantly related ones, but it is possible to transfer functional microsatellite primers from more distantly related species in some cases (Lorieux et al., 2000). In this sense, these results also indicate that *Vitellaria paradoxa* is genetically closer to *Argania spinosa* than *Manilkara huberi*.

Using these two types of markers, the genetic distance among the argan accessions studied was studied. Phenetic

Table 1. Nuclear SSR markers obtained in *Manilkara huberi* and *Vitellaria paradoxa* assayed in the molecular characterization of the argan accessions assayed.

Marker (locus)	Repeat motif	Primers sequence	T ^a annealing (PCR)	No. of alleles		Size range (bp)		Ho	He	PD
				Original species	Argan	Original species	Argan			
Mh03 ^z	(CT) ₁₇	F: CACTTCTGTCCTCTCTCGT R: GGGTGATTTGCAGAGACGTA	56	7	na ^x	176–204	–	–	–	–
Mh04 ^z	(CT) ₁₂	F: GCACCTCCATGGTTCCAGT R: AAAGAGTCAATGGCGTGAGC	52	5	4	189–209	161–197	0.00	0.72	0.71
Mh06 ^z	(GA) ₁₄	F: ACACGCACAAAACAACCAA R: TTCTTGAAGGAGGGTTGCTC	56	7	na ^x	162–188	–	–	–	–
Mh07 ^z	(CT) ₂₃	F: ATTGCAGCATATCCACACCA R: GCAAAGGGTGTATGGGTAGA	56	4	3	153–187	138–143	0.00	0.67	0.66
Mh08 ^z	(CT) ₁₁	F: GTAATGGAGCCGTTTGAGA R: CTGGTAGCAITTTGTTGCAI	56	7	na ^x	172–202	–	–	–	–
Mh12 ^z	(CT) ₉ (AC) ₆	F: TGGGGAACGTGGAAAAGAGT R: ATCCACAGCAATGACTGACG	56	7	1	187–211	200	0.00	0.00	0.00
Mh17 ^z	(CT) ₁₃	F: CACGATGACCTCTCAGTGA R: CCTGTGTATGCGTTCCGATG	56	6	na ^x	240–274	–	–	–	–
Mh19 ^z	(CT) ₂₁	F: AATTACAACAAAAGCTCCACIT R: TGAGAGTCTTTTCGCACTTCA	56	7	na ^x	146–164	–	–	–	–
Mh20 ^z	(GA) ₁₃	F: GAAAGTTTTCACCAATTTGGGAAT R: GACATAACACTAACCCCTTCACGA	56	8	na ^x	134–166	–	–	–	–
Mh22 ^z	(CT) ₁₅	F: CCCATTATAGCCCTCCACCT R: AGAGAGCACATGCAAGCTCA	56	7	3	180–206	190–206	0.40	0.51	0.50
Mh24 ^z	(CT) ₁₇	F: CCACCTCTGCTCTCTCTCTCTCGT R: GACAITGTGGGTGATTCGAG	60	7	na ^x	181–209	–	–	–	–
Mh26 ^z	(CT) ₁₄	F: TGCTCAGACTGCTTCTTTTTG R: TGCAATAAGTGTGATTTGGAGAA	58	6	na ^x	224–250	–	–	–	–
mVpCIRB03 ^y	(GT) ₇ (GA) ₂ ... (GA) ₄	F: GCACCTCCATGGTTCCAGT R: AAAGAGTCAATGGCGTGAGC	52	3	4	150–153	143–151	0.00	0.66	0.65
mVpCIRB05 ^y	(CT) ₅ (TT)(CT) ₄	F: TATCCGTAGTGGGTGT R: TTTGGGTCTAITTCATCT	48	3	2	157–161	183–197	0.18	0.66	0.36
mVpCIRE04 ^y	(TG) ₈ (CG) ₅	F: CGGCAITTTGGTCTTACTT R: TTTTCCCTTTGGTCA	48	9	1	116–126	129	0.00	0.00	0.00
mVpCIRE05 ^y	(TG) ₇	F: GTTTGTATGGTTTCCGGTT R: CTTTCGTTTTCAGTAGGCTC	52	4	3	238–244	244–257	0.27	0.63	0.45
mVpCIRE06 ^y	(GT) ₄ (GA) ₆ (GT) ₂ (GA)	F: GCACCTCCATGGTTCCAGT R: AAAGAGTCAATGGCGTGAGC	52	5	na ^x	118–126	–	–	–	–
mVpCIRE11 ^y	(GT) ₁₀	F: TTGCTAITTCCTCTGT R: TTCATCACCTTCCCTC	48	3	4	224–230	225–235	0.27	0.71	0.83
mVpCIRF05 ^y	(TG) ₄ GG(GT) ₃	F: AGGACGGCGAATGGTG R: GCAGGGCAACTACAAAAGACA	52	4	3	190–209	223–230	0.60	0.70	0.50
mVpCIRH04 ^y	(TG) ₁₀ (AG) ₃ (TG)(AG) ₇	F: CTCTGTGCTTTATGTCGT R: ACTCACCAITTTGCTT	48	6	5	206–220	234–278	0.21	0.80	0.83

^z SSRs developed in *Manilkara huberi* by Azevedo et al. 2005.^y SSRs developed in *Vitellaria paradoxa* by Cardí et al., 2005.^x na: No amplification.

(taxonomy) relationships among argan morphotypes were analyzed with several UPGMA dendrograms (see supplemental data, Fig. B). In general, both markers showed some degree of similarity in dendrogram topologies. Relationships moderately supported were similar in both cases, although with some differences in the positioning of some morphotypes in the main groups. However, bootstrap values of UPGMA dendrogram obtained with the utilization of RAPDs were slightly higher than those obtained using SSRs, in line with the larger number of observed bands with a higher polymorphism (Tab. I and supplemental data, Tab. A and Fig. A). Finally, the general dendrogram, constructed using the combined data sets (RAPDs+SSRs), was very similar to those obtained separately with each marker although with some differences with similar bootstrap values than the RAPD dendrogram.

Dendrograms consist of some main clusters including some oval and spindle morphotypes which seem to be the two more important morphotypes from the molecular point of view. The remaining oval, spindle, spherical and intermediate morphotypes clustered out of these main clusters and in separately clusters. In addition, some well-supported clusters (according to bootstrap analysis) were established among some replications of some morphotypes using both markers (ie. spherical-1 and spherical-2; spindle-9 and spindle-10; spindle-4 and spindle-5; oval-11 and oval-12; oval-13 and oval-14; and oval-spindle-4 and oval-spindle-5).

In general no main groups were distinguished using these markers in relation to the morphotype groups. In some cases argan accessions from different morphotypes are closer than accessions from the same morphotype. This clustering could be better explained by geographic proximity than by morphotype as has been previously describe in a molecular study using chloroplast and mitochondrial DNA amplification (El Mousadik and Petit, 1996b). Traditional morphological characterization of the different argan accessions into morphotypes (according to fruit phenotype) does not correlate well with the real genetic background (genotype) of this species although main different morphotypes from the genetic point of view are oval and spindle fruit types. Morphotypes based on fruit morphology do not make up recognisable genetic entities based on molecular markers. On the other hand, the results obtained also confirm the great genetic diversity observed in argan using isoenzyme markers (El Mousadik and Petit, 1996a).

One of the goals of conservation programmes in these rare fruit species such as argan is to characterize and maintain existing level of genetic variation and genetic resources (Martínez-Gómez et al., 2007). Genetic resources not only provide the required raw material for suitable genetic crop improvement, but offer a unique gene combination to ensure adaptability and productivity (Lachenaud and Zhang, 2008; Wang et al., 2006). Designing of core collection involves an appropriate use of diversity, offering breeders an opportunity to work with a manageable number of accessions.

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Online Material

Supplemental data

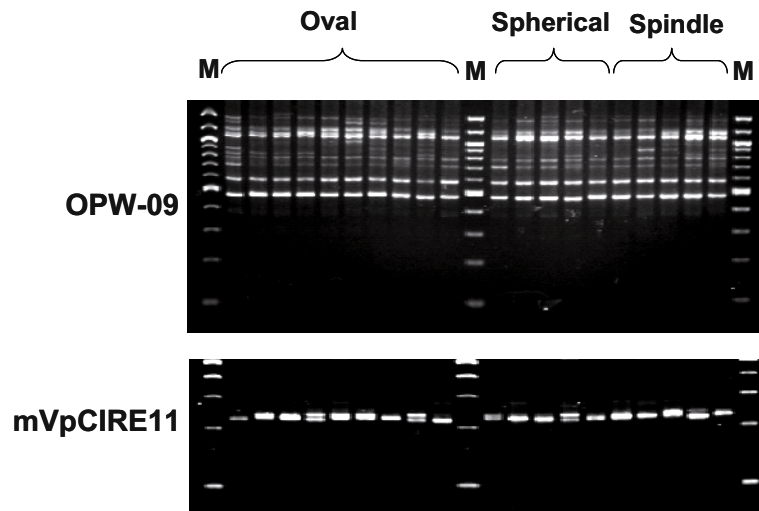


Figure A. Agarose gel (1.5%) showing the allelic segregation of the OPW-09 RAPD marker and Metaphor[®] agarose gel (3%) showing the allelic segregation of the mVpCIRE11 nuclear SSR marker. M, 1Kb DNA Ladder (Invitrogen, Madrid, Spain).

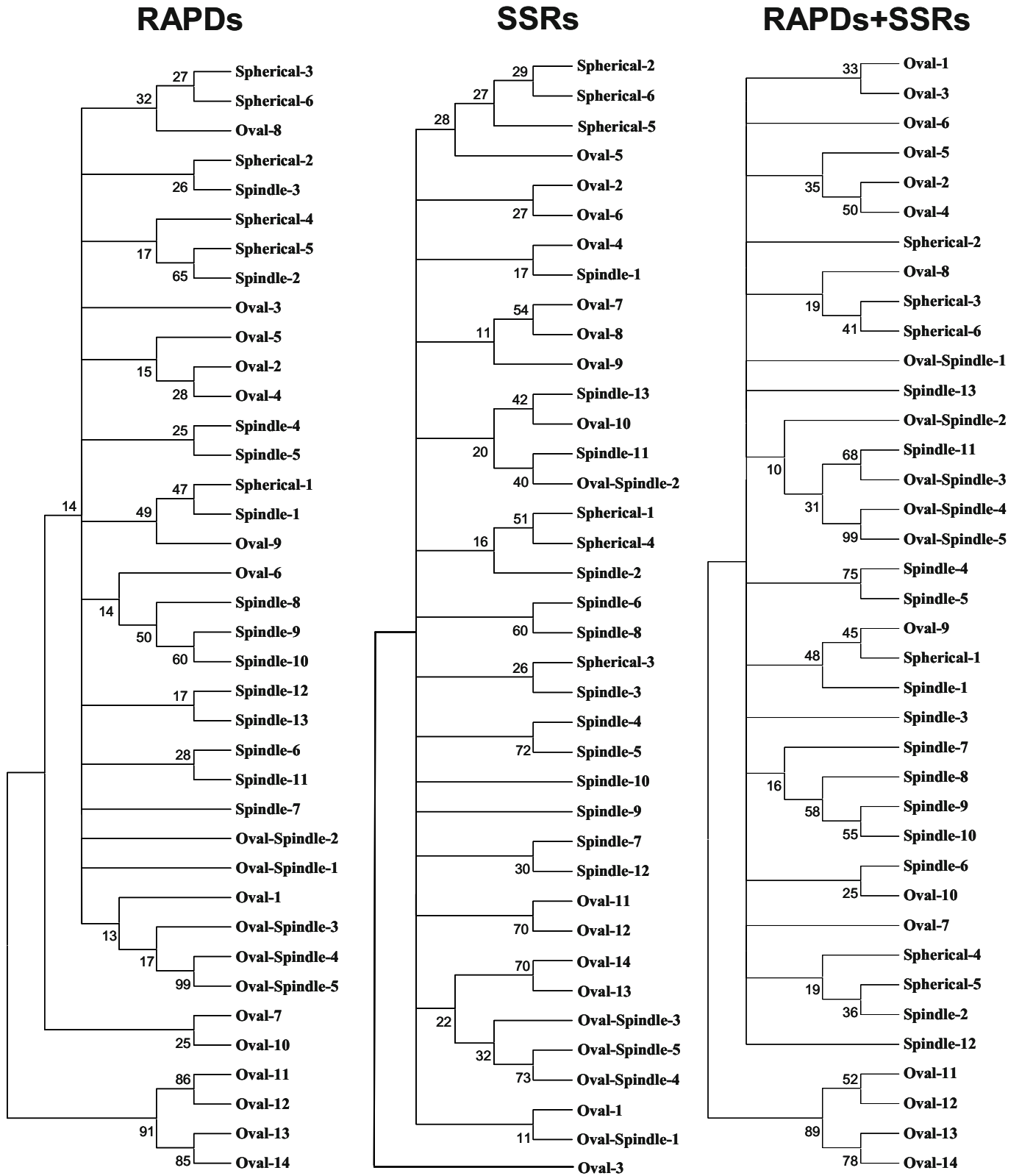


Figure B. Dendrograms obtained by UPGMA cluster analysis based on mean character differences among the argan morphotypes (oval, spherical and spindle types) assayed in this study using RAPD and SSR markers separately and the whole data set of RAPD and SSR markers. Numbers in the branches represent bootstrap values.

Table A. Universal primers (decamers) purchased from Operon Technologies Inc. (California, USA) used as RAPD markers molecular characterization of the argan accessions assayed.

Marker	Primers sequence	T ^a annealing (PCR)	Number of bands	Number of polymorphic bands	Size range (bp)	PIC
OPA-08	GTGACGTAGG	36	7	7	447-1413	0.93
OPA-10	GTGATCGCAG	36	7	7	620-1839	0.92
OPA-11	CAATCGCCGT	36	6	6	636-2514	0.90
OPA-16	AGCCAGCGAA	36	4	2	1000-2009	0.35
OPB-07	GGTGACGCAG	36	4	4	392-669	0.83
OPB-11	GTAGACCCGT	36	7	7	589-2090	0.88
OPG-13	CTCTCCGCCA	36	9	9	373-2553	0.95
OPN-14	TCGTGCGGGT	36	7	7	548-1363	0.91
OPN-15	CAGCGACTGT	36	9	8	349-2009	0.93
OPR-15	GGACAACGAG	36	7	7	615-1381	0.93
OPR-16	CTCTGCGCGT	36	9	9	317-1844	0.95
OPW-09	GTGACCGAGT	36	8	7	478-1075	0.79
OPW-12	TGGGCAGAAG	36	5	5	521-1244	0.83
OPW-13	CACAGCGACA	36	14	14	493-2001	0.96
OPX-03	TGGCGCAGTG	36	4	4	586-1875	0.54
OPX-06	ACGCCAGAGG	36	11	11	444-2098	0.92
OPY-06	AAGGCTCACC	36	8	8	447-1798	0.92
OPY-13	GGGTCTCGGT	36	12	12	322-1683	0.95
OPZ-09	CACCCAGTC	36	8	7	531-1485	0.89