CHARACTERIZATION OF TUNISIAN CASTOR GENOTYPES 
(RICINUS COMMUNIS L.) USING RAPD MARKERS

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The aim of this work was to detect genetic variability among the set of 20 Tunisian castor genotypes using 5 RAPD markers. Amplification of genomic DNA of 20 genotypes, using RAPD analysis, yielded 42 fragments, with an average of 7.00 polymorphic fragments per primer. Number of amplified fragments ranged from 5 to 9, with the size of amplicons ranging from 200 to 1400 bp. The polymorphic information content (PIC) value ranged from 0.662 to 0.855 with an average of 0.780 and diversity index (DI) value ranged from 0.669 to 0.857 with an average of 0.785. The dendrogram based on hierarchical cluster analysis using UPGMA algorithm was prepared. Knowledge on the genetic diversity of castor can be used for future breeding programs for increased oil production to meet the ever increasing demand of castor oil for industrial uses as well as for biodiesel production.

Keywords: Castor, Genetic diversity, Molecular markers, RAPD technique

Introduction

Castor (Ricinus communis L., 2n = 2x = 20, Euphorbiaceae), is industrially important non-edible oilseed crop widely cultivated in the arid and semi-arid regions of the world (Govaerts et al., 2000). The seed of castor contain more than 45% oil and this oil is rich (80–90%) in an unusual hydroxyl fatty acid, ricinoleic acid (Jeong and Park, 2009). Castor oil is the only vegetable oil soluble in alcohol, presenting high viscosity, and requiring less heating than others oils during the production of biodiesel (Jeong and Park, 2009). Due to its unique chemical and physical properties, the oil from castor seed is used as raw material for numerous and varied industrial applications, such as: manufacture of polymers, coatings, lubricants for aircrafts, cosmetics, etc., and for the production of biodiesel (Jeong and Park, 2009). With more than 95% of the world’s castor production concentrated in limited parts of India, China, and Brazil (Sailaja et al., 2008), and because of the ever increasing world-wide demand of castor for industrial use, there is a pressing need to increase the hectarage and productivity of castor. Castor is a cross pollinated crop and is usually cultivated as a hybrid in India, as hybrids give significantly greater yields than pure lines or varieties (Birchler et al., 2003; Reif et al., 2007).

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Higher magnitude of heterosis and genetically superior hybrids can be obtained by combining diverse parents in hybrid development. Conventional diversity analysis methods, in the field, are time consuming, laborious, resource intensive and drastically affected by environmental factors, therefore, a technique that is rapid and not affected by environment is needed for assessment of genetic diversity and selection of parental lines for use in hybrid development programmes (Santalla et al., 1998). Genetic diversity assessment prior to developing hybrids can aid in better exploitation of heterosis. Assessment of genetic variation using molecular markers appears to be an attractive alternative to the conventional diversity analyses and can also aid in management and conservation of biodiversity. A large number of polymorphic markers are required to measure genetic relationships and genetic diversity in a reliable manner (Santalla et al., 1998).

The aim of this study was to detect genetic variability among the set of 20 Tunisian castor genotypes using 5 RAPD markers.

Material and methodology

Plant material and DNA extraction

Ricin lines (20) were obtained from the University of Carthage, National Institute of Research in Rural Engineering, Waters and Forests (INRGREF), Regional Station of Gabès, Tunisia. DNA of 20 genotypes of castor was extracted from 10 day old leaves using the Gene JET Plant Genomic DNA Purification Mini Kit.

RAPD amplification

Amplification of RAPD fragments was performed according to Gajeraa et al. (2010) (Table 1) using decamer arbitrary primers (Operon technologies Inc, USA; SIGMA-D, USA). Amplifications were performed in a 25 μl reaction volume containing 5 μl DNA (100 ng), 12.5 μl Master Mix (Genei, Bangalore, India), and 1 μl of 10 pmol of primer. Amplification was performed in a programmed thermocycler (Biometra, Germany) with initial denaturation at 94 °C for 5 min, 42 cycles of denaturation at 94 °C for 1 min, primer annealing at 38 °C for 1 min, extension at 72 °C for 1 min, and final extension at 72 °C for 5 min. Amplified products were separated in 1.5% agarose in 1× TBE buffer. The gels were stained with ethidium bromide and documented using gel documentation system Grab-It 1D pre Windows.

| RAPD primers | Primer sequence (5’-3’) | Molecular weight range (bp) |
|--------------|--------------------------|----------------------------|
| OPA-03       | AGTCAGCCAC               | 330–870                    |
| OPA-13       | CAGCACCAC                 | 370–1800                   |
| OPB-08       | GTCCACACGG               | 530–1550                   |
| OPD-02       | GGACCCCAACC              | 280–1850                   |
| SIGMA-D-14   | TCTCGCTCCA                | 350–900                    |
Data analysis

The RAPD bands were scored as present (1) or absent (0), each of which was treated as an independent character regardless of its intensity. A dendrogram based on hierarchical cluster analysis using the unweighted pair group method with arithmetic average (UPGMA) with the SPSS professional statistics version 17 software package was constructed. For the assessment of the polymorphism between genotypes ricin and usability RAPD markers in their differentiation we used diversity index (DI), the probability of identity (PI) and polymorphic information content (PIC).

Results and discussion

PCR amplifications using 5 RAPD primers produced 42 DNA fragments that could be scored in all genotypes. The selected primers amplified DNA fragments across the 20 genotypes studied, with the number of amplified fragments varying from 5 (OPD-02) to 9 (OPB-08), and the amplicon size varying from 200 to 1400 bp. Of the 42 amplified bands, all 42 were polymorphic, with an average of 7.00 polymorphic bands per primer. The polymorphic information content (PIC) value varied from 0.662 (OPD-02) to 0.855 (OPB-08), with an average of 0.780 and index diversity (DI) value varied from 0.669 (OPD-02) to 0.857 (OPB-08) with an average of 0.785 (Table 2). A dendrogram based on UPGMA analysis separated 20 Tunisian castor genotypes into three groups. First cluster contained 5 genotypes from the region Souassi (S-1 – S-5). Second cluster of 13 ricin genotypes (2) was divided into two main clusters (2a and 2b). Main cluster 2a was divided into subcluster 2aa and 2bb. Subcluster 2aa contained four genotypes from the region Bouthay (BT-1, BT-2, BT-3, BT-5) and one genotypes from the region Matmata (MT-1). Subcluster 2ab contained 3 Tunisian ricin genotypes from the region Ghomrassen (GH-2, GH-4, GH-5). Main cluster 2b contained 2 ricin genotypes from the region Ghomrassen (GH-1 and GH-3) and 3 genotypes from the region Sidi bou ali (BA-2, BA-3, BA-4). Third cluster contained 2 ricin genotypes from the region Sidi bou ali (BA-1 and BA-5). We could not distinguish two genotypes, S-4 and S-5, which can be caused due the same genetic background (Figure 1).

Table 2 The statistical characteristics of the RAPD markers used in castor

| Primers    | Number of alleles | DI    | PIC    | PI    |
|------------|-------------------|-------|--------|-------|
| OPA-03     | 8                 | 0.778 | 0.776  | 0.016 |
| OPA-13     | 7                 | 0.805 | 0.794  | 0.010 |
| OPB-08     | 9                 | 0.857 | 0.855  | 0.009 |
| OPD-02     | 5                 | 0.669 | 0.662  | 0.071 |
| SIGMA-D-04 | 6                 | 0.817 | 0.811  | 0.019 |
| Average    | 7,00              | 0.785 | 0.780  | 0.025 |

Note: DI – diversity index; PIC – polymorphic information content; PI – probability of identity.

Gajeraa et al. (2010) used 30 RAPD polymorphic primers for the analysis of 22 castor bean genotypes. RAPD analysis yielded 256 fragments, of which 205 were polymorphic, with an average of 6.83 polymorphic fragments per primer. Genetic diversity of 37 ricin genotypes
grown in China using RAPD markers was studied by Li et al. (2012). Using RAPD markers, together they detected 122 alleles, of which 71 were polymorphic, representing the percentage of polymorphism alleles 58.20%. In the study Machado et al. (2013) used 58 RAPD primers for the analysis of 15 castor bean cultivars. The genetic dissimilarity between cultivars was calculated by Jaccard’s index, using the unweighted pair-group method with arithmetic mean (UPGMA). Pecina-Quintero et al. (2013) study the diversity and genetic relationships among accessions of *R. communis* from the state of Chiapas, México using AFLP (amplified fragment length polymorphism) and SSR (simple sequence repeat) markers. Tomar Rukam et al. (2014) investigated the fingerprinting and phenotyping of 25 castor genotypes available in Gujarat and other States of India using RAPD and ISSR markers. One hundred thirty decamer RAPD primers from Operon series (OPA to OPZ – five from each series) were screened with the DNA of the 2 castor genotypes. Some researchers have considered RAPD markers to represent segments of DNA with noncoding regions and to be selectively neutral (Landergott et al., 2001), and some studies have shown that RAPD markers are distributed throughout the genome and may be associated with functionally important loci (Penner, 1996).

**Figure 1:** Dendrogram of 20 Tunisian castor genotypes prepared based on 5 RAPD markers. Regions of origin of analyzed genotypes of Tunisian ricin: S – Souassi; BT – Bouthay; GH – Ghomrassen; BA – Sidi bou ali; MT – Matmata

**Conclusions**

The analysis showed that the RAPD markers are very effective molecular markers for the assessment of the genetic diversity in castor bean. A dendrogram based on UPGMA analysis separated 20 Tunisian castor genotypes into three groups. Using 5 RAPD markers only two castor bean genotypes have not been distinguished. Our analysis proved utilization of RAPD markers for differentiation of used set of castor genotypes. For better discrimination of the analyzed ricin genotypes, it is necessary to use a higher number of RAPD markers.
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