Polymorphism in Random Amplified and Nuclear rDNA Sequences Assessed in Certain Apple (*Malus × domestica* Borkh.) Cultivars

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**Abstract**

Eighth apple (*Malus × domestica* Borkh.) cultivars: 'Delikates', 'Cortland', 'James Grieve', 'Lired', 'Jonathan', 'Golden Delicious', 'Jonagold' and 'Idared' were characterized by two different molecular tools. These included analysis of the distribution of RAPD markers and length variability of the SSU, 5.8S, LSU and ITS region of the nuclear rRNA genes assessed in PCR reactions with different combinations of ‘universal’ primers. RAPD analysis was performed with 17 out of 24 RAPD primers tested. Those amplified a total of 183 loci (872 amplicons) out of which 34 (18.5%) were monomorphic, 128 (69.5%) were polymorphic and 22 (12%) cultivar-specific. Cultivar-specific RAPD products were detected for each apple cultivar.

Amplification of the rDNA sequences showed variability. Fifty-four amplicons were generated in the experiment including 14 monomorphic, 26 polymorphic, and 14 cultivar-specific products. Altogether 232 amplicons were generated, whose length ranged from 220 to 940 bp. The analysis of dendrograms constructed on the basis of the analysis of RAPD genetic profiles and profiles amplified on rDNA matrices showed their significant correlation (Mantel test: \(r_{AB} = 0.430;\) p-value (Two-tailed) = 0.024), which proves that the used methods correctly presented variability within the examined cultivars, and the molecular markers identified in the study can be considered appropriate.

**Keywords**: apple cultivars, *Malus*, Mantel test rDNA, RAPD, variability

**Introduction**

Apple (*Malus × domestica* Borkh.), together with many economically significant fruit plants growing in the temperate zone, belongs to the family *Rosaceae* (King *et al.*, 1991).

The genus *Malus* Mill. comprises from 25 to as many as 47 species, and the number has been changing and depends on the possibility of approval of new species or hybrids (van Treuren *et al.*, 2010; Yan *et al.*, 2008).

Apple is a species that is highly diversified in terms of morphological, phenological and genetic traits (among other things genotype polyploidy) (Zhou and Li, 1999). Many studies on the genus *Malus* have been conducted. They included studies on the use of molecular tools to identify cultivars and rootstocks, to construct genetic maps, and in marker assisted selection (MAS), using various types of markers: RAPD (Royo and Itoiz, 2004; Sesstras *et al.*, 2009; Zhou and Li, 2000), SSR (van Treuren *et al.*, 2010) or ISSR (Goulão and Oliveira, 2001; Smolik and Krzysztozsek, 2010).

rDNA sequences have become a subject of numerous experiments. Nuclear rDNA are grouped into arrays consisting of hundreds to thousands of tandem repeats. This region includes two spacers (ITS1 and ITS2) that separate the SSU (18S), 5.8S and LSU (26S) of nuclear ribosomes genes (Baldwin, 1992). ITS sequence data have been extensively documented to provide insights into phylogenetic history, polyploidy ancestry, genetic relationships and fingerprinting (Álvarez and Wendel, 2003).

There are many studies in the scientific literature where authors describe genetic variability of various types of sequences (including rDNA) and base the interpretation of results on the significance of correlation coefficient between the genetic similarity matrices calculated for the variability described within the examined sequences. In the case of a significant correlation, the described results gain in value. During analysis of rDNA sequences, variability per se within them should be taken into account together with variability resulting from the occurrence of rDNA fragments polymorphic in length (e.g. additional ITS amplicons), especially in polyploid genotypes (Smolik *et al.*, 2010, 2011). Using different pairs of ‘universal’ primers in the PCR analysis, it is possible to amplify products of different length in a reaction with only one pair of ‘universal’ primers. It may be difficult to choose an amplicon representative for rDNA sequences for sequencing and further analyses. Therefore, a question arises whether selecting a definite amplicon for sequencing and disregarding others introduces a too considerable error into the results of the analyses. The scientific literature on the taxonomy of subjects based on the analysis of rDNA sequences does not provide full explanation if one reaction product, characteristic of the genus (e.g. ITS), or a mixture of e.g. ITSs of different length, was taken into consideration in the sequencing of regions/products.
Therefore, the aims of the present study were; to identify the genotypic variability between eight apple cultivars (‘Delikates’, ‘Cortland’, ‘James Grieve’, ‘Lired’ and ‘Jonathan’, ‘Golden Delicious’, ‘Jonagold’ and ‘Idared’) using RAPD technique, to describe the variability between the cultivars on the basis of DNA profiles obtained as a result of amplification of additional rDNA products, generated in reactions with various combinations of pairs of ‘universal’ primers, and to conduct the analysis of correlation of the genetic similarity matrices obtained in the analysis of variability of random DNA and rDNA sequences.

Materials and methods


DNA preparation

Total DNA was extracted from fresh leaves. Plant material (±100 mg) was ground in liquid nitrogen with a mortar and pestle and incubated at 37°C with Protease K. The DNA extraction was described in Genomic Mini AX Plant protocol (A&A Biotechnology Gdynia-Poland). RNaseA (Sigma-Aldrich) was added (2 µl) to the DNA samples to eliminate RNA contaminations by incubation at 37°C for 10 min. Genomic DNA was quantified (GeneQuant DNA/RNA Calculator-Pharmacia LKB) to obtain 50 ng DNA pro 1 µl of DNA template.

RAPD amplification

PCR amplifications were performed with a set of 24 random primers-part of Operon kits (Operon Technologies USA). Reaction mixture (25 µl) contained 10× PCR buffer with (NH₄)₂SO₄, 2 mM dNTPs and 25 mM MgCl₂, 2.5 µM of primer, 1 u of Taq DNA polymerase enzyme (Fermentas MBI), 2.5 µM of each primer and 100 ng of template DNA. The reactions were performed in Mastercycler 5333 (Eppendorf). The program of thermal cycling was as follows: initial activation step at 95°C for 1.5 min followed by 40 cycles of 1 min at 94°C, 1 min at 36°C and 2 min at 72°C, with a final extension step at 72°C for 10 min. The amplification was concluded by a final extension at 72°C for 5 min.

rDNA. Oligonucleotides for PCR amplifications

The analysis encompassed nuclear rDNA, including the following sequences: SSU, ITS1; 5.8S; ITS2 and LSU. The general structure and the arrangement of the analyzed sequences encoding the formation of rRNA genes are presented in Fig. 1. The figure also includes names, approximate locations and directions of the hybridizations of primers used for PCR reactions. The origin of primers, sequences and their authors are presented in Tab. 1.

PCR amplification of rDNA sequences

PCR mixtures (25 µl) contained 10× PCR buffer with (NH₄)₂SO₄, 2 mM dNTPs and 25 mM MgCl₂, 1.0 units of Taq DNA polymerase (Fermentas MBI), 2.5 µM of each primer and 100 ng of template DNA. DNA was amplified using a Mastercycler 5333 (Eppendorf) thermocycler and using the following program: initial denaturation at 94°C for 7 min, 40 cycles of 30s at 94°C, annealing temperature, 2 min at 72°C and 7 min at 72°C for a final extension. The annealing temperature was usually adjusted according to the Tm of the primers being used in the reaction (Tab. 1). Amplifications were performed according to the touch-down PCR protocol by raising the temperature of the first singles cycles by 6°C above the annealing temperature, adopted from the preliminary experiments for both primers.

![Fig. 1. rDNA unit. Arrows indicate approximate positions of primers used to generate amplicons in different rDNA regions. Primer’s names, sequences and they authors were listed in Tab. 1](image)

<table>
<thead>
<tr>
<th>rDNA fragment amplified</th>
<th>Primer’s name</th>
<th>Author(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SSU</td>
<td>NS1, NS2, NS4, NS3</td>
<td>White et al., 1990</td>
</tr>
<tr>
<td>ITS1</td>
<td>NS5, ITS2, ITS5, ITS1</td>
<td>White et al., 1990</td>
</tr>
<tr>
<td></td>
<td>5.8S, SR6R</td>
<td>Vilgalys and Hester, 1990</td>
</tr>
<tr>
<td></td>
<td>ITS1-F</td>
<td>Gardes and Bruns, 1993</td>
</tr>
<tr>
<td>ITS2</td>
<td>ITS3, ITS4</td>
<td>White et al., 1990</td>
</tr>
<tr>
<td></td>
<td>ITS4-B, 5.8SR</td>
<td>Vilgalys and Hester, 1990</td>
</tr>
<tr>
<td>LSU</td>
<td>LR7, LR3R</td>
<td>Vilgalys and Hester, 1990</td>
</tr>
</tbody>
</table>

*Vilgalys unpublished [http://www.botany.duke.edu/fungi/mycolab]
Electrophoresis and data analysis

PCR products were separated by electrophoresis on a 1.5% agarose (Basica LE Prona) in 1× TBE buffer, then stained with EtBr and visualized (MiniBIS Pro-DNR Bio-Imaging System-Israel). O’RangeRuler 200 bp DNA Ladder (Fermentas MBI) was used as a size marker (3000-200 bp). RAPD and rDNA products were scored as 1 (present) or 0 (absent) for the cultivar and similarity index matrices were generated based on number of shared fragments. The similarity coefficient between accessions compared was determined by Jaccard’s coefficient (Nei and Li, 1979). Similarity matrices were generated using the Phytool software (Buntjer, 2001) and were compared.

Cultivar-specific RAPD products (Tab. 2) were amplified for the eight apple cultivars. Their length ranged from ~2780 (OPM_05) to ~220 bp (OPM_03).

Results

Randomly amplified DNA (RAPD) and rDNA length polymorphism showed that the eight apple cultivars exhibited a relatively wide range of variability. The analysis of genetic profiles revealed that the range of variability, determined with the use of each of the methods separately, was similar. In the both methods, polymorphic and cultivar-specific products were amplified. Variability (length polymorphism) was found within rDNA sequences, presented as amplicons polymorphic in terms of length, generated in reactions with different pair combinations of ‘universal’ primers.

RAPD–Random Amplified Polymorphic DNA

Twenty-four primers were used in reactions, generating 17–465 amplicons. The range of variability was assessed on the basis of 232 products, of which 112 were polymorphic, 114 were cultivar-specific, and 6 were monomorphic. On average from 43 (‘Idared’) to 71 (‘Jonagold’) products were generated for the examined apple cultivars. Their length ranged from ~2780 (OPM_05) to ~220 bp (OPM_03).

Polymorphic loci

The greatest number of products (22) was amplified in a reaction with primer OPA_03, and the least–5, respectively: with primers OPJ_13, OPW_07, and 7 in reactions with each of the primers: OPM_02 and OPM_01_1.

Cultivar–specific loci

One hundred fourteen polymorphic and cultivar-specific RAPD products (Tab. 2) were amplified for the eight apple cultivars (Fig. 2).
The greatest number (32) was amplified in amplifications with 13 primers for 'Jonathan', and the least (4) — with 4 primers for the cultivar 'Idared' (Tab. 2).

**rDNA sequences**

In the initial experiments, 28 universal primer pairs were used, of which in amplifications with 17 primer pairs combinations, distinct products were generated that differed in length and hybridization sites within the rDNA sequences (Tab. 1, Fig. 1). A total of 44 PCR products were generated of which 20 (46%) were polymorphic, 13 (29%) were monomorphic and 11 (25%) were cultivar-specific (data not shown). Amplicons (193) of the length ranging from ~1000 (NS5+ITS2) to ~220 bp (SR6R+5.8S) were generated for the examined cultivars. The greatest number of polymorphic products (4) was found after using primer pair SR6R+5.8S. The amplified products were from ~640 to ~220 bp long. Cultivar-specific products (11) were generated in reactions with the different primer combinations and were listed in Tab. 3 and Fig. 3. More than 12 monomorphic products were amplified, present in all the examined apple cultivars. They were obtained by the amplification of each of characterized rDNA sequences (Fig. 1, 3).

**SSU**

Two primer pairs (NS1+NS2, NS3+NS4) and partially NS5+5.8S were used for amplification of SSU region. In PCR with primer combinations: NS1+NS2 and NS3+NS4 one (530 bp) and two monomorphic products were amplified, respectively (Tab. 3, Fig. 3).

Tab 3. Apple cultivar-specific products generated in PCR reaction on rDNA templates

<table>
<thead>
<tr>
<th>Cultivar</th>
<th>Primer’s pair and lengths of amplified amplicons (bp)</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>'Cortland'</td>
<td>NS5 + 5.8S, SR6R + 5.8S, ITS5 + ITS4</td>
<td></td>
</tr>
<tr>
<td>'James Grieve'</td>
<td>ITS1 + ITS2</td>
<td></td>
</tr>
<tr>
<td>'Lired'</td>
<td>ITS1F + ITS2[340], ITS5 + ITS4</td>
<td></td>
</tr>
<tr>
<td>'Jonathan'</td>
<td>ITS1_F + ITS2[440], ITS5 + ITS2</td>
<td></td>
</tr>
<tr>
<td>'Golden Delicious'</td>
<td>ITS1F + ITS2[360, 270]</td>
<td></td>
</tr>
<tr>
<td>'Jonagold'</td>
<td>NS5 + ITS2</td>
<td></td>
</tr>
</tbody>
</table>

ITS1

Combinations of eight primer pairs were used for amplification of the ITS1 sequence (Tab. 3). As a result, monomorphic (5), polymorphic (12) and cultivar-specific (10) products were amplified (Tab. 3, Fig. 3), presented in detail in Tab. 4.

**ITS (ITS1+5.8S+ITS2)**

The ITS sequences were amplified using combination of three primer pairs: ITS5+ITS4, ITS1_F+ITS4 and SR6R+ITS4 (Tab. 3). As a result, three monomorphic, polymorphic and one 'Lired' cultivar-specific product were amplified, respectively (Tab. 3, 4).

**ITS2**

The sequences of ITS2 spacer were amplified using three primer pairs combination (Tab. 3). As a result four polymorphic and two monomorphic products were generated (Tab. 3, Fig. 3).

**LSU**

Monomorphic fragment (810 bp) of the LSU sequence was amplified in a reaction with primer pair LR3R and LR7 (Tab. 4).

**Genetic relationships. RAPD**

The analysis of topology of the constructed dendrogram enabled to distinguish two similarity groups corresponding to the pedigrees of two examined cultivars: 'Jonagold' and 'Delikates', respectively grouped as clusters 'a' and 'b' (Fig. 3). Group 'a' included the cultivars: 'Jonathan', 'Golden Delicious', 'Jonagold' and 'Idared', and group 'b': 'Cortland', 'Delikates', 'James Grieve' and 'Lired' (Fig. 3). The phylogenetic similarity between them ranged from 0.48 between the cultivar 'Idared' and 'Golden Delicious' to 0.19 between the cultivars 'Jonathan' and 'Cortland' (data not shown). The phylogenetic similarity of 0.32 was found between the cultivars 'Jonagold' and 'Jonathan', and the lowest similarity (0.25) was found in the pedigree 'a' ('Jonagold') between the cultivars 'Idared' and 'Jonathan' (Fig. 3). The analysis of the topology of groups for pedigree

MW-Molecular Weight (O’RangeRuler 200bp DNA Ladder), 1-'Cortland', 2-'Delikates', 3-'James Grieve', 4-'Lired', 5-'Jonathan', 6-'Golden Delicious', 7-'Jonagold', 8-'Idared'

Fig. 3. PCR products profiles generated in amplifications primers pairs designed for investigation of rDNA variability. Arrows mark selected monomorphic or polymorphic products
rDNA

Similar genetic relationships were found between the cultivars in the analysis of genetic profiles obtained after amplification of the rDNA sequences (Fig. 3). Genotype similarity ranged respectively from 63% between the cultivars 'Cortland' and 'James Grieve' and 'Cortland' to 86% between the cultivars 'Jonagold' and 'Idared'. In addition, significant correlation coefficient was found, assessed with the Mantel test (r(AB) = 0.430, p-value (two-tailed = 0.024) for the matrices of phylogenetic similarity calculated for the examined cultivars with the use of the RAPD technique and on the basis of the analysis of rDNA length polymorphism.

Discussion

The results of the present study did not differ from those obtained from such type of previous studies on various apple accessions, including cultivars, in terms of both the number of the used primers, and the range of length and structure of the amplified products: poly-, monomorphic or cultivar-specific (Sestras et al., 2009; Zhou and Li, 2000).

Hence a high percentage of polymorphic products, including cultivar-specific (~70%) was observed by Zhou and Li (2000) in their study of apple, Royo and Itoiz (2004) noted 82%, and Goulão et al. (2001)-60%. Forte et al. (2002) found 52% of polymorphic RAPD products (loci) for 72 apple accessions. It resulted in relatively low genetic similarity coefficients, presented by Zhou and Li (2000) for 21 apple accessions.

Royo and Itoiz (2004) presented similarity coefficients ranging from 0.3 to 1.0 and Adebayo et al. (2009) on average amounted to 0.65. In the present study, genetic similarity coefficients for RAPD markers ranged from 0.27-0.40 for 'Delikates' and 0.25-0.48 for 'Jonagold' pedigrees, respectively. Additionally, it was proved that the dendrogram of phylogenetic similarity properly presented genetic relationships between the examined apple cultivars. The grouping of cultivars enabled to divide them depending on the origin to a described pedigree ('Delikates' and 'Jonagold'). The cultivar-specific RAPD products amplified in the described experiment can be used together with the products described for ISSR (Smolik and Krzysztozszek, 2010) as markers supporting determination of genetic identity of cultivars, and also in tree nursery production in order to control varietal purity, which, as it is proved by a study by Mogdil et al. (2005), is not without significance.

Multicopy structure of rDNA makes amplification highly efficient, especially on templates of diploid genomes, and becomes interesting on templates of polyploid genomes (including Malus). Amplified fragments were not too long, so, for example, the ITS sequence (ITS1+5.8S+ITS2) on average was about 700 bp long, apart from numerous exceptions, in whose genetic profiles there were additional amplification products probably resulting from the presence in the genome of, for example, shorter copies of rDNA e.g. as in the tetraploid forms of Fagopyrum or accessions of Syringa (Smolik et al., 2010; 2011; Yasuo and Ohnishi, 1998; ).

The present study showed the possibility of amplification of additional products of reactions with pairs of 'universal' primers, intentionally combined in different ways, on the matrices of rDNA sequences. In reactions with ITS-flanking primers (ITS5+ITS4, ITS1_F+ITS4 and SR6R+ITS4), products of the length ~700 bp and additional amplicons were amplified, whose characterization is presented in Tab. 4. Among them one specific product was amplified-for the cultivar 'Lired' (ITS5+ITS4)728);

According to many authors, the ITS sequences, due to their variability level, provide enough information that can be used in identification of molecular markers or in phylogenetic analyses. It is not without significance that the sequences are inherited from both parents, they are
variable in polyploids (Forte et al., 2002; Nalini et al., 2007; Saini et al., 2008) and with their help hybridity and introgression can be assessed (Baldwin, 1992).

Shen et al. (1998) stated that assessment of genetic variability in the ITS1 region can be used in the identification of species of a given organism and in determination of genetic origin of a plant. Studying on beet, they used 18 primers in PCR reactions that generated products of the length from 740 to 610 bp. The longest products were generated in reactions with the primers ITS2, ITS4 and ITS5 (740 bp each). Similar results were obtained from the present study. Eighteen universal primer pairs, out of 24 used, generated products easily identified on electrophoregrams. Three primers (ITS2, ITS4 and ITS5), used by Shen et al. (1998) in their study on beet, were also used in a study of apple, only in different combinations. Primer ITS2 was used in combinations with 4 other primers (NS5, ITS1, ITS_1F and ITS5). Primer ITS4-in combinations with 5 other primers (ITS5, ITS_1F, SR6R, 5.8SR and ITS3), while primer ITS5 with the following primers: ITS2 and ITS4. The combinations comprising the above-mentioned primer pairs generated products of the length from 220 to 720 bp in PCR reactions. In a PCR reaction, where primer pair ITS5 + ITS4 was used, the longest generated product was the band of the length of 720 bp, which corresponded to the length of the products generated by these primers in a study of beet (740 bp).

The analysis of genetic similarity dendrogram constructed on the basis of genetic profiles obtained in PCR reactions (RAPD and rDNA) showed that, similarly as in the ISSR technique (Smolik and Krzysztoforschek, 2010), the examined cultivars were divided into two groups. One pedigree included the cultivar ‘Delikates’ and its parental forms ‘Cortland’, ‘James Grieve’, and ‘Jonathan’, and the second one ‘Jonagold’ and its parental forms ‘Golden Delicious’, ‘Idared’ (a hybrid of the cultivars ‘Jonathan’ and ‘Wagner’), and the cultivar ‘Lired’ (a mutant deriving from the cultivar ‘James Grieve’).

It was proved that there is a deviation in the grouping of the cultivars presented on the dendrogram (rDNA). Two cultivars ‘Jonathan’ and ‘Lired’ were exchanged in the similarity group ‘a’ and ‘b’. Thus the cultivar ‘Jonathan’ was grouped together with the cultivars forming the pedigree of ‘Delikates’, and the cultivar ‘Lired’ was grouped with the cultivars of the ‘Jonagold’ pedigree. The other cultivars were grouped according to their origin. Different classification of the cultivars ‘Jonathan’ and ‘Lired’, as opposed to their already known origin, presumably is a result of the limited analyses. Their only results are of the analysis of genetic profiles obtained in reactions with 18 combinations of pairs of various universal primers. The values of genetic similarity are comparable to those presented in a study by Forte et al. (2002) for 72 apple accessions. These authors analysed polymorphism of rDNA sequences collected in the GeneBank database, and demonstrated a relatively wide range of variability. However, the range of variability in relation to grouping of genotypes on the dendrogram was consistent with the range of variability simultaneously determined using the RAPD technique.

Significant correlation coefficient was found, [r(AB) = 0.430, p-value; two-tailed = 0.024)] assessed with the Mantel test, for genetic similarity matrices calculated on the basis of the analysis of RAPD and generated rDNA profiles. It should be mentioned that, according to Sneath and Sokal (1973), the calculated value of the correlation coefficient (significant) is too low. The authors state that in order to describe genetic similarity matrices as significantly correlated, the value of ‘r’ coefficient should be higher than 0.8. On the other hand, Lapointe and Legendre (1992) consider the matrices strongly correlated when r ≥ 0.5.

Conclusions

The results of the present study confirm the suggestion of many authors on the existence of great variability potential of the genus Malus. However, it should be noted that statistically significant correlation between the genetic similarity matrices enables to state that the variability within the cultivars of two known pedigrees (’Delikates’ and ’Jonagold’) was correctly determined. Hence the already known (even though discussed) usefulness of the RAPD technique for determination of varietal distinctness was confirmed. What is more, the amplification of additional products on the rDNA templates demonstrated their length variability that is probably the presence of rDNA sequence copies of different length, and also the genetic potential of the genus Malus. The dendrogram constructed on the basis of the analysis of rDNA genetic profiles, despite it is significantly correlated with the RAPD dendrogram, may be a source of erroneous conclusions. The study confirms only the genetic potential of Malus and indicates the need for including ITS sequences also differing in copy length in the most frequent analyses.

References


