# DEHYDROGENASE ISOENZYME POLYMORPHISM IN GENUS Prunus, SUBGENUS Cerasus 

Slavica ČOLIĆ ${ }^{1}$, Vera RAKONJAC ${ }^{2}$, Milica FOTIRIĆ AKŠIĆ ${ }^{2}$, Dragan NIKOLIĆ ${ }^{2}$, Vladislav OGNJANOV ${ }^{3}$ and Dragan RAHOVIĆ ${ }^{1}$<br>${ }^{1}$ Institute for Science Application in Agriculture, Belgrade, Serbia<br>${ }^{2}$ University of Belgrade, Faculty of Agriculture, Belgrade, Serbia<br>${ }^{3}$ University of Novi Sad, Faculty of Agriculture, Novi Sad, Serbia

Čolić S., V. Rakonjac, M. Fortić. Akšić, D.Nikolić, V. Ognjanov, and D. Rahović (2012): Dehydrogenase isoenzyme polymorphism in genus prunus, subgenus cerasus. - Genetika, Vol 44, No. 3,619-632.

Dehydrogenase polymorphism was studied in 36 sour cherry (Prunus cerasus L.), sweet cherry (Prunus avuim L.), mahaleb (Prunus mahaleb L.), ground cherry (Prunus fruticosa Pall.), duke cherry (Prunus gondounii Redh.), Japanese flowering cherry (Prunus serrulata Lindl.) and four iterspecific hybrids (standard cherry rootstocks 'Gisela 5', 'Gisela 6', 'Max Ma' and 'Colt'). Inner bark of one-year-old shoots, in dormant stage, was used for enzyme extraction. Vertical PAGE was used for isoenzyme analysis: alcohol dehydrogenase (ADH), formate dehydrogenase (FDH),

Corresponding author: Slavica Čolić, Institute for Science Application in Agriculture, Bul. Despota Stefana 68b, 11000 Belgrade, Serbia, phone: ++ 38111 2751622, fax: ++ 381112752 959, e-mail: slavicacol@yahoo.com
glutamate dehydrogenase (GDH), isocitrate dehydrogenaze (IDH), malate dehydrogenase (MDH), phosphogluconate dehydrogenase (PGD), and shikimate dehydrogenase (SDH). All studied systems were polymorphic at 10 loci: Adh -1 (3 genotypes) and Adh-2 (5 genotypes), Fdh-1 (2 genotypes), Gdh-1 (3 genotypes), Idh-1 (4 genotypes) i Idh -2 (5 genotypes), Mdh-1 (3 genotypes), Pgd-1 (4 genotypes), Sdh-1 (1 genotype) i $S d h-2$ (3 genotypes). Cluster analysis was used to construct dendrogram on which four groups of similar genotypes were separated. Obtained results indicate that studied enzyme systems can be used for determination of genus Prunus, subgenus Cerasus. Among studied enzyme systems ADH, IDH and SDH were the most polymorphic and most useful to identify genetic variability. Polymorphism of FDH and GDH in genus Prunus, subgenus Cerasus was described first time in this work. First results for dehydrogenase variability of Oblačinska indicate that polymorphism of loci $I d h-2$ and $S d h-2$ can be useful for discrimination of different clones.

Key words: cluster analysis. dehydrogenase, electrophoresis, polymorphism, Prunus spp

## INTRODUCTION

Knowledge of the genetic diversity and relationships among the cultivated and wild species of Cerasus subgenus is important for recognizing gene pools, identifying pitfalls in germplasm collections and developing effective conservation and management strategies (KHADIVI-KHUB et al., 2012). Characterisation and identification of species and cultivars in genus Prunus within Cerasus subgenus has been mostly based on morphological traits (PEREZ-SANCHEZ et al., 2008, PEREZ et al., 2010, SHAHI-GHARAHLAR et al., 2010).

Isoenzyme were among the first mmolecular markers applied in horticultural science, because they allow the identification of plants in early stages of development and are not affected by the environmental conditions. Isoenzyme analysis is important technique in Prunus genetic and breeding for identification of cultivars (AGARWAL et al., 2001, MILATOVIĆ et al., 2009, NIKOLIĆ et al., 2010), detection of phylogenetic relationships among species (DAEIL, 2004), also for analysis of genetic variability in native populations (GAŠIĆ et al., 2001, ČOLIĆ et al., 2010) and construction of gene linkage maps (CLARKE et al., 2009). Although molecular DNK based markers were dominantly used in the last decade, studies of isoenzyme polymorphism conducted in the last twenty years in genus Prunus, subgenus Cerasus verify that this technique was efficient to detect polymorphism. According to DAEIL (2004) MDH and GPI were the most polymorphic and most valuable to identify genetic relationships among the taxa in subgenus Cerasus. Some genotypes with identical morphological characters and previously treated as one cultivar can be separated on the basis of isoenzyme genotype. Recently, CORTS et al. (2008) used extracts from young leaves of nine sweet and eight sour cherry varieties for analysis of five isoenzyme systems in order to characterize these varieties and
detect problems of synonymies and homonymies that frequently present and found that PGM and PGI had highest discrimination power.

Molecular DNA based markers become an essential tool in genus Prunus genetic studies (MARTÍNEZ-GÓMEZ et al., 2003). Different types of molecular markers RFLPs (BOUHADIDA et al., 2007), RAPDs (ZAMANI et al., 2012), AFLPs (TAVAUD et al., 2004) and SSRs (ERCISLI et al., 2011) have been used for the genetic characterization of germplasm and the establishment of genetic relationships between cultivars and species.

The objective of our study was to evaluate dehydrogenase isoenzyme polymorphism of six species in genus Prunus, subgenus Cerasus including cherry germplasm from the rich native flora of Serbia and four interspecific hybrids 'Gisela 5', 'Gisela 6', 'Max Ma' and 'Colt'. The goal was to establish usability of studied isoenzymes in identification of genetic diversity, relationships among genotypes so as intraspecies and intracultivar variability.

## MATERIALS AND METHODS

The plant material represents 36 sour (Prunus cerasus L.), sweet (Prunus avuim L.), mahaleb (Prunus mahaleb L.), ground (Prunus fruticosa Pall.), duke (Prunus gondounii Redh.) and Japanese flowering cherry (Prunus serrulata Lindl.) genotypes. Four widely-used standard cherry rootstocks 'Gisela 5', 'Gisela 6', 'Max Ma' and 'Colt' were included. Eight 'Oblačinska' (autochthonous and heterogeneous cultivar), four wild sweet cherry, six ground and one mahaleb genotypes was selected from native flora in Serbia. Selection of genotypes was done according to observed diversity of phenology and morphological traits of tree and fruits.

Seven isoenzyme systems - alcohol dehydrogenase (ADH), formate dehydrogenase (FDH), glutamate dehydrogenase (GDH), isocitrate dehidrogenaze (IDH), malate dehydrogenase (MDH), phosphogluconate dehydrogenase (PGD) and shikimate dehydrogenase (SDH) were analyzed. Inner bark of one-year-old shoots was used for enzyme extraction. Preparation of samples was done in accordance with the protocol given by BOŠKOvić et al. (1994) for stone fruit species. Vertical PAGE was used for isoenzyme analysis. Polyacrylamide gel containing $8 \%$ acrylamide was used for separation. Staining procedures were essentially based on the protocol for isoenzymes given by BOšKOviĆ et al. (1994). Gels were visually observed and the bands represent isoenzyme patterns, called zymograms, were analyzed. Genetic interpretations for regions attributed to polymorphic loci were proposed. Cluster analysis was done with all polymorphic loci using the UPGMA method. For cluster analysis the data of polymorphic loci were transformed into $0 / 1$ code. Statistical analysis was conducted with the program 'Statistica' (StatSoft, Inc., Tulsa, Oklahoma, USA).

## RESULTS AND DISCUSSION

All seven analyzed enzyme systems were polymorphic at 10 loci: $A d h-1$ and $A d h-2$, Fdh-1, Gdh-1, Idh-1 and Idh -2, Mdh-1, Pgd-1, Sdh-1 and i Sdh-2. The patterns of polymorphic systems for each of the studied genotype was presented in the Table 1.

Tab. 1. Zymogram patterns of seven dehydrogenase systems in genus Prunus, subgenus

| No | Cultivar/ genotype | Species <br> Interspecific hybrid | Zymogram patterns |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | ADH | FDH | GDH | IDH | MDH | PGD | SDH |
| 1 | Drogans yellow | P. avium | 2 | 1 | 1 | 3 | 1 | 1 | 1 |
| 2 | Celeste | P. avium | 2 | 1 | 1 | 3 | 1 | 1 | 1 |
| 3 | Victoria | P. avium | 2 | 1 | 1 | 5 | 1 | 2 | 1 |
| 4 | Early Star | P. avium | 2 | 1 | 1 | 3 | 1 | 1 | 1 |
| 5 | Vera | P. avium | 2 | 1 | 1 | 3 | 1 | 1 | 2 |
| 6 | Sara | P. avium | 2 | 1 | 1 | 6 | 1 | 2 | 1 |
| 7 | NS KK 6/10 | P. avium | 2 | 1 | 3 | 2 | 1 | 2 | 2 |
| 8 | DT X9 | P. avium | 5 | 1 | 1 | 5 | , | 1 | 7 |
| 9 | DT X3 | P. avium | 5 | 1 | 1 | 5 | 1 | 1 | 7 |
| 10 | DT X7 | P. avium | 5 | 1 | 2 | 5 | 1 | 1 | 7 |
| 11 | DT K9 | P. avium | 5 | 1 | 4 | 11 | 1 | 1 | 7 |
| 12 | MD | P. serrulata | 4 | 1 | 1 | 4 | 1 | 3 | 1 |
| 13 | BNS | P. serrulata | 4 | 2 | 1 | 6 | 2 | 1 | 3 |
| 14 | Amanogawa | P. serrulata | 4 | 1 | 1 | 1 | 1 | 1 | 3 |
| 15 | Lara | P. cerasus | 1 | 1 | 1 | 10 | 3 | 2 | 1 |
| 16 | Montmorency | P. cerasus | 1 | 1 | 1 | 12 | 3 | 5 | 1 |
| 17 | Rexelle | P. cerasus | 1 | 1 | 1 | 10 | 3 | 2 | 1 |
| 18 | Keleris 16 | P. cerasus | 1 | 1 | 1 | 12 | 3 | 5 | 1 |
| 19 | Maynard | P. cerasus | 3 | 1 | 3 | 9 | 3 | 2 | 2 |
| 20 | Oblačinska <br> UD 1 | P. cerasus | 1 | 1 | 1 | 8 | 2 | 2 | 2 |
| 21 | $\begin{aligned} & \text { Oblačinska } \\ & \text { UD } 8 \\ & \hline \end{aligned}$ | P. cerasus | 1 | 1 | 1 | 10 | 2 | 2 | 1 |
| 22 | Oblačinska UD 6 | P. cerasus | 1 | 1 | 1 | 7 | 4 | 2 | 2 |
| 23 | Oblačinska <br> D1 R | P. cerasus | 1 | 1 | 1 | 10 | 2 | 2 | 2 |
| 24 | $\begin{aligned} & \hline \text { Oblačinska D4 } \\ & \mathrm{R} \\ & \hline \end{aligned}$ | P. cerasus | 1 | 1 | 1 | 10 | 2 | 2 | 2 |
| 25 | Oblačinska $\mathrm{II} / 10 \mathrm{R}$ | P. cerasus | 1 | 1 | 1 | 10 | 2 | 2 | 2 |
| 26 | $\begin{aligned} & \text { Oblačinska } \\ & \text { XI/3 R } \\ & \hline \end{aligned}$ | P. cerasus | 1 | 1 | 1 | 10 | 2 | 2 | 2 |
| 27 | $\begin{aligned} & \text { Oblačinska D4 } \\ & \text { RS̆ } \\ & \hline \end{aligned}$ | P. cerasus | 1 | 1 | 1 | 10 | 2 | 2 | 2 |
| 28 | SV 1 | P. fruticosa | 1 | 1 | 3 | 6 | 2 | 5 | 2 |
| 29 | SV 2 | P. fruticosa | 1 | 1 | 3 | 6 | 2 | 5 | 2 |
| 30 | SV 3 | P. fruticosa | 2 | 1 | 1 | 10 | 2 | 2 | 2 |


| 31 | SV 4 | P. fruticosa | 2 | 1 | 1 | 10 | 4 | 5 | 2 |
| :--- | :--- | :--- | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 32 | SV 5 | P. fruticosa | 2 | 1 | 1 | 10 | 4 | 5 | 2 |
| 33 | SV 7 | P. fruticosa | 2 | 1 | 1 | 4 | 2 | 4 | 2 |
| 34 | Radmilovac | P. gondouinii | 8 | 1 | 1 | 5 | 2 | 3 | 5 |
| 35 | Uroš | P. gondouinii | 9 | 1 | 1 | 3 | 2 | 3 | 5 |
| 36 | TT | P. mahaleb | 6 | 1 | 1 | 10 | 3 | 3 | 3 |
| 37 | Colt | P. avium $x P$ P. <br> pseudocerasus | 2 | 1 | 2 | 2 | 1 | 2 | 4 |
| 38 | Gisela 5 | P.cerasus $x P$ (anescens <br> cal | 1 | 1 | 3 | 10 | 4 | 2 | 1 |
| 39 | Gisela 6 | P.cerasus $x$ P. <br> canescens | 1 | 1 | 1 | 7 | 4 | 2 | 1 |
| 40 | Max Ma | P. mahaleb $x P$. <br> avium | 1 | 1 | 1 | 4 | 1 | 1 | 6 |

Alcohol dehydrogenase. ADH analysis resulted in zymograms (Fig. 1) with two regions of activity $A d h-1$ and $A d h-2$. Contrary to SEKER (2008), who observed that $P$. avium showed polymorphism only for loci Adh-l we obtained variability in two regions. Wild sweet cherry was monomorphic and homozygous for both loci, while $P$. avium cultivars were monomorphic but heterozygous for loci Adh-1 and homozygous for Adh-2. Japanese flowering cherry was monomorphic for both loci. Locus Adh-1 was homozygous, while Adh-2 was heterozygous and showed three bands. For P. cerasus polymorphism was obtained for Adh-2 loci that had $b b$ and $a b$ genotypes. Genotypes of $P$. fruticosa showed activity in both regions, but polymorphism only for $A d h-2$ that showed homozygous genotype $a a$ and heterozygous genotype $a b$. No activity for $A d h-1$ and variability for presence or absence of activity in Adh-2 recorded for $P$. gonduinii. Among all samples $P$. mahaleb had unique zymogram with genotype $b b$ in loci $A d h-1$ and genotype $b c$ in Adh-2 loci. Zymograms of Max Ma, Gisela 5, Gisela 6 and Colt had the same heterozygous pattern $a b$ for loci Adh-1. Only difference that we found for loci $A d h-2$ showed that Colt (homozygous $a a$ ) can be distinguish from other rootstocks (heterozygous $a b$ ). Three bands for heterozygous genotype that indicate dimeric structure of ADH found for sour, wild, sweet, flowering and ground cherry and four iterspecific hybrids but not for duke cherry and mahaleb.

Formate dehydrogenase. Literature indicates that, so far, this system was not studied in subgenus Cerasus. This system was characterized with low activity and variability in one locus marked as $F d h-1$. Two types of zymogram were present (Fig. 2). Only one sample of $P$. serulatta had heterozygous genotype $a b$ with three bands characterized dimeric structure, described also in almond (ČOLIĆ et al., 2009). Other 39 genotypes had homozygous genotype $b b$.


Fig. 1. Zymogrames obtained for ADH


Fig. 2. Zymogrames obtained for FDH

Glutamate dehydrogenase. On GDH zymograms activity was observed in the region closer to the cathode (Fig. 3). Two alleles and three genotypes were proposed for the loci marked as $G d h-1$. The most genotypes were $a a$, two $P$. fruticosa, one $P$. avium and one $P$. cerasus had $a b$ genotype while one wild cherry and Colt had $b b$ allelic constitution. One genotype of wild cherry showed no activity for GDH. Presence of seven bands in heterozygote phenotypes indicates hexameric structure reported also for almond by ČOLIĆ et al. (2009). The greatest polymorphism and three genotypes ( $a a, a b$, and $b b$ ) for GDH showed $P$. avium This is the first report of

GDH polymorphism in subgenus Cerasus. GDH polymorphism was observed in the other stone fruit species such as peach (GAŠIĆ et al., 2001) and apricot (MILATOVIĆ et al., 2009).


Fig. 3. Zymogrames obtained for GDH
Isocitrate dehydrogenase. IDH showed twelve different zymograms (Fig. 4) being the most polymorphic system. Both $P$. avium and $P$. cerasus showed five patterns. In accordance with previously reported results of PASHKOULOV et al. (2000) activity was visible in two regions. All genotypes showed activity for loci $I d h-1$, with three alleles ( $a, b$ and $c$ ) and four genotypes ( $a b, b b, b c$ and $c c$ ). For loci $I d h-2$ we observed polymorphism and variabilty in presence or absence in activity. Three alelles and five genotypes ( $a a, a b, a c, b b$ and $b c$ ) were identified. From all observed loci in this study we found that $I d h-2$ was most polymorphic. The greatest polymorphism for IDH exhibited $P$. cerasus, with discriminated five genotypes. Unique patterns observed for $P$. serrulata cv Amanogava, one sweet cherry and one "Oblačinska" genotype. Our findings about one region of activity for wild cherry and mahaleb are in accordance to results of SEKER (2008).
Malate dehydrogenase. MDH analysis resulted with observable variability in one region of activity marked as $M d h-1$ (Fig. 5), and presence of alleles $a$ and $b$. Homozygous allelic constitution $a a$ observed for $P$. avium, P. serrulata, Colt and Max Ma. Heterozygous allelic constitution $a b$ had sour cherry, duke, ground cherry, Gisela 5 and 6. Variability was expressed in Oblačinska and $\quad P$. fruticosa. We discriminated genotypes with two and four bands. Homozygous allelic constitution $b b$ observed for $P$. cerasus and $P$. mahaleb. Contrary to our findings SEKER (2008)
reported about $M d h-1$ and $M d h-2$ loci. These differences can be attributed to different tissue for extraction (leaf), and different type of gel (starch).


Fig. 4. Zymogrames obtained for IDH


Fig. 5. Zymogrames obtained for MDH


Fig. 6. Zymogrames obtained for PGD


Fig. 7. Zymogrames obtained for SDH

Phosphogluconate dehydrogenase. One polymorphic region of activity was observed on PGD zymograms (Fig. 6) in the region closer to the cathode. Three alleles and four genotypes ( $\mathrm{aa}, \mathrm{ab}$, ac and bb ) were distinguished. The PGD system displayed five patterns. Homozygous allelic constitution $a a$ obtained for wild cherry, while $b b$ observed for $P$. gonduinii and $P$. mahaleb. Two allelic constitutions $a a$ and $a b$ obtained for sweet, $a a$ and $b b$ for flowering and $a b$ and $a c$ for ground and sour cherry. Zymograms of Gisela 5, Gisela 6 and Colt had the same heterozygous genotype $a b$ while Max Ma was homozygous (aa). That difference can distinguish Max Ma from other rootstocks. More variability in sweet cherry than CORTS et al. (2008) reported can be explained by wider genetic basis of genotypes in our research. Our findings indicate dimeric structure of loci for sour and ground cherry.

Shikimate dehydrogenase. On SDH zymograms (Fig. 7) two regions with bands was observed, but activity for loci $S d h-1$ was showed only for $P$. gonduinii. That difference can distinguish $P$. gonduinii from other taxa in this research. For wild cherry we observed no activity for this system, while sweet cherry showed greatest polymorphism with two alleles ( $a$ and $b$ ) and two types of zymograms. For $S d h-2$ two allelic constitution obtained for sour and flowering cherry ( $a a$ and $a b, a a$ and $b b$, respectively). Intraspecific variability was detected for Oblačinska sour cherry, where we distinguish $a a$ and $a b$ genotypes. Diversity of Oblačinska sour cherry clones on the morphological, chemical and pomological level has been approved in research of NIKOLIC et al. (2005) and RAKONJAC et al. (2010). No intraspecific variability for SDH and only $a b$ allelic constitution was detected within the $P$. fruticosa genotypes. Unique four band pattern for SDH represented Colt. Two unique bands were probably incorporated from $P$. pseudocerasus genome. Dimeric structure of loci $S d h-2$ was recorded for Max Ma.

Tab. 2. Polymorphic loci obtained for seven isoenzymatic systems

| Genotype | Species/Hybrid | No of polymorphic loci | Polyimorphic loci |
| :---: | :---: | :---: | :---: |
| Wild cherry | P. avium | 4 | Gdh-1, Idh-2, Pgd-1, Sdh-2 |
| Sweet cherry | P. avium | 2 | Gdh-1, Idh-1 |
| Flowering cherry | P. serrulata | 6 | Fdh-1, Gdh-1, Idh-1, Idh-2, Pgd-1, Sdh-2 |
| Sour cherry | P. cerasus | 6 | $\begin{gathered} \text { Adh-2, Gdh-1, Idh-1, Idh-2, } \\ \text { Pgd-1, Sdh-1 } \end{gathered}$ |
| Oblačinska | P. cerasus | 1 | Idh-2, Sdh-2 |
| Ground cherry | P. fruticosa | 5 | Adh-2, Gdh-1, Idh-1, Idh-2, Pgd-1 |
| Duke cherry | P. gonduinii | 2 | Adh-2, Idh-2 |

Greatest dehydrogenase variability showed $P$. serrulata and $P$. cerasus that had six polymorphic loci (Tab. 2). Of 40 studied cherry genotypes, nineteen showed unique zymogrames and can be distinguished from other genotypes. Also, on the basis of unique polymorphism in one locus species can be discriminate mutually. Among ten polymorphic loci $F d h-1$ was unique for $P$. serrulata and locus $S d h-1$ for P. cerasus.


Fig. 8. Dendrogram of 40 analyzed genotypes generated from the isoenzyme data by UPGMA cluster analysis
Group 1-SV1, SV2; Group 2- Oblačinska višnja: D1 R, D4 R, II/10 R, XI/3 R, D4
RŠ; Group 3- Lara, Rexelle; Group 4- Montmorency, Keleris 16; Group 5- Gisela 5, Gisela 6; Group 6- Drogans yellow, Celeste, Early Star; Group 7- DT X9, DT X3; Group 8-SV4, SV5

Application of cluster analysis on all polymorphic loci resulted in a dendrogram shown on Figure 8. Analyzed genotypes are connected on different hierarchical levels. Four clusters were identified and among them eight groups of genotypes with the same isoenzymatic profile. Dendrogram showed clear separation of Cluster A, represent $P$. mahaleb and two genotypes of $P$. gonduinii. Six genotypes that were split into subgroup of wild cherry and subgroup of flowering cherry represent cluster B includes. Ten genotypes grouped in cluster C: seven sweet cherry, one flowering cherry and rootstocks Colt and Max Ma (interspecies hybrids where $P$. avium is aone of the parents). Our results for different linkage between Victoria, Sara, Vera and Maynard than reported by LJUBOJEVIĆ et al. (2012) showed that morphological variability did not always correspond with molecular characterisation. Cluster D consisted of 15 P. fruticosa and P. cerasus genotypes.

Although we expected that all genotypes of Oblačinska will fall in the same group, results showed discrimination into three subgroups. Also three genotypes of Oblačinska had unique isoenzyme profile. This diversity corresponds to variability obtained for loci $I d h-2$ and $S d h-2$. Significant discrimination observed also for $P$. fruticosa, separated in two groups and two unique genotypes.

Classification into four clusters corresponds with origin of genotypes, while further discrimination into subgroups of related genotypes can not be clearly defined.

CONCLUSION
Polymorphism was observed in all seven studied enzyme systems. One polymorphic locus was identified for FDH, GDH, MDH and PGD while two polymorphic loci were observed for ADH, IDH and SDH. Polymorphism of ADH, IDH and SDH was the most useful to identify genetic variability. Nineteen examined genotypes showed unique izoenzyme profiles, while 21 separated into eight groups. Unique polymorphic locus was observed for $P$. serrulata (Fdh-1) and for $P$. cerasus (Sdh-1).

Dendrogram showed four clusters that correspond with the origin of the studied genotypes. Obtained results showed that presented enzyme systems can be useful for identification of genetic variability in genus Prunus, subgenus Cerasus, but it must be combined with the agronomical and morphological characterization in order to assess or reject identities among varieties especially among clones. Polymorphism of FDH and GDH in genus Prunus, subgenus Cerasus was described first time in this work. First results for Oblačinska indicate that polymorphism of loci $I d h-2$ and $S d h-2$ can be useful for discrimination of different clones.

## ACKNOWLEDGEMENT

This paper is a result of a study performed within the Project TR-31308 that was supported by the Ministry of Education, Science and Technological Development of the Republic of Serbia.

Received July30 ${ }^{\text {th }}, 2012$
Accepted November $30^{\text {th }}, 2012$

## REFERENCES

AGARWAL, S., A.K., NATH and D.R., SHARMA (2001): Characterisation of peach (Prunus persica L.) cultivars using isoenzymes as molecular markers. Sci Hort, 90, 227-242.
bOSKOVIC, R., K.R. TOBUTT, P. ARUS and R. MESSEGUER (1994): Isoenzymes. In: R. Messeguer (Editor). Methods of molecular marker analysis in Prunus. IRTA, Barcelona, p. 4-25.
BOUHADIDA M., J.P. MARTÍN, G. EREMIN, J. PINOCHET, M.Á. MORENO and Y. GOGORCENA (2007): Chloroplast DNA diversity in Prunus and its implication on genetic relationships. J Am Soc Hortic Sci, 132, 670-679.

CLARKE, J.B., D.J. SARGENT, R.I. BošKOVIC, A. BELAJ and K.R. TOButt (2009): A cherry map from the inter-specific cross Prunus avium 'Napoleon' $\times P$. nipponica based on microsatellite, genespecific and isoenzyme markers. Tree Genet Genomes, 1, 41-51.
CORTS, R.M., L.C. RODRIGUES, J.M.O. MARCIDE, R.P. SANCHES (2008): Characterization of sour (Prunus cerasus L.) and sweet cherry (Prunus avium L.) varieties with five isozyme systems. Rev Bras Frutic, 30: 154-158.
COLIC S., D. MILATOVIC, D. NIKOLIC, G. ZEC (2009): Dehydrgenase isoenzyme polymrphism in selected almond genotypes (Prunus amygdalus Batsch.). Bulg J Agric Sci, 15: 552-556.
ČOLIĆ S., D. MILATOVIĆ, D. NIKOLIĆ and G. ZEC (2010): Isoenzyme polymorphism of almond genotypes selected in the region of northern Serbia. Hortic Sci, 37, 56-61.
DAEIL, K. (2004): Genetic characterisation using isoezyme analysis in the genus Prunus. Korean J Hortic Sci, 9: 321-327.
ERCISLI, S, G. AGAR, N. YILDIRIM, B. DURALIJA, A. vOKURKA and H. KARLIDAG (2011): Genetic diversity in wild sweet cherries (Prunus avium) in Turkey revealed by SSR markers. Gen Mol Res, 10: 1211-1219.
GAŠIĆ, K., V. OGNJANOV, R. BOŠKOVIĆ, K. TOBUTT and C. JAMES (2001): Characterisation of vineyard peach biodiversity. Acta Hortic, 546, 119-125.
KHADIVI-KHUB, A., Z. ZAMANI and M.R. FATAHI (2012): Multivariate analysis of Prunus subgen. Cerasus germplasm in Iran using morphological variables. Genet Resour Crop Evol, 59: 909-926.
LJUBOJEVIĆ, M, V. OGNJANOV, D. BOŠNJAKOVIĆ, G. BARAĆ, M. OGNJANOV, E. MLADENOVIĆ and J. ČUKANOVIĆ (2012): Sweet and sour cherry decorative forms. Genetika, 44, 367-375.
MARTÍNEZ-GÓMEZ, P., G.O. SOZZI, R. SÁNCHEZ-PÉREZ, M. RUBIO and T.M. GRADZIEL (2003): New approaches to Prunus tree crop breeding. Food Agr Environ, 1, 52-63.
MILATOVIĆ, D., D. NIKOLIĆ, D. ĐUROVIĆ and J. MILIVOJEVIĆ (2009): Isoenzyme polymorphism in apricot cultivars. J Amer Pomolog Soc, 63, 14-23.
NIKOLIĆ, D., V. RAKONJAC, M. MILUTINOVIĆ and M. FOTIRIĆ (2005): Genetic divergence of Oblačinska sour cherry (Prunus cerasus L.) clones. Genetika, 37, 191-198.
NIKOLIĆ, D., D. MILATOVIĆ, V. RAKONJAC, D. ĐUROVIĆ and B. ĐORĐEVIĆ (2010): Izoenzimski polimorfizam sorti šljive. Voćarstvo, 44(169-170), 7-12.
PASHKOULOV, D.T., K.R. TOBUTT and R. BOŠKOVIĆ (2000): Comparison of isoenzymes in Prunus avium separated by two different electrophoretic techniques. Plant Breeding, 119, 153-156.
PEREZ, R, F. NAVARRO, M.A. SANCHEZ, J.M. ORTIZ, and R. MORALES (2010): Analysis of agromorphological descriptors to differentiate between duke cherry (Prunus x gondouinii (Poit. \& Turpin) Rehd.) and its progenitors: sweet cherry (Prunus avium L.) and sour cherry (Prunus cerasus L.). Chilean J Agri Res, 70: 34-49.
PEREZ-SANCHEZ, R, M.A. GOMEZ-SANCHEZ and R. MORALES-CORTS (2008): Agromorphological characterization of traditional Spanish sweet cherry (Prunus avium L.), sour cherry (Prunus cerasus L.) and duke cherry (Prunus x gondouinii Rehd.) cultivars. Spanish J Agri Res, 6: 4255.

RAKONJAC, V., M. FOTIRIĆ, D. NIKOLIĆ, D. MILATOVIĆ and S. ČOLIĆ (2010): Morphological characterization of 'Oblacinska' sour cherry by multivariate analysis. Sci Hortic, 25, 679-684.
SHAHI-GHARAHLAR, A., Z. ZAMANI, M.R. FATAHI and N. BOUZARI (2010): Assessment of morphological variation between some Iranian wild Cerasus sub-genus genotypes. Hort Environ Biotechnol, 51, 308-318.

SEKER, M. (2008): Investigation of isozyme polymorphism in open-pollinated sweet cherry and 'Mahaleb' seedlings. Acta Hortic, 795, 423-428.
TAVAUD, M., A. ZANETTO, J.L. DAVID, F. LAIGRET and E. DIRLEWANGER (2004): Genetic relationships between diploid and allotetraploid cherry species (Prunus avium, Prunus $x$ gondouinii and Prunus cerasus). Heredity, 93, 631-638.
ZAMANI Z., A. SHAHI-GHARAHLAR, R. FATAHI and N. BOUZARI (2012): Genetic relatedness among some wild cherry (Prunus subgenus Cerasus) genotypes native to Iran assayed by morphological traits and random amplified polymorphic DNA analysis. Plant Syst Evol, 298, 499-509.

## POLIMORFIZAM DEHIDROGENAZA RODA Prunus, PODROD Cerasus

Slavica ČOLIĆ ${ }^{1}$, Vera RAKONJAC ${ }^{2}$, Milica FOTIRIĆ AKŠIĆ ${ }^{2}$, Dragan NIKOLIĆ ${ }^{2}$, Vladislav OGNJANOV ${ }^{3}$ i Dragan RAHOVIĆ ${ }^{1}$<br>${ }^{1}$ Institut za primenu nauke u poljoprivredi, Beograd, Srbija<br>${ }^{2}$ Univerzitet u Beogradu, Poljoprivredni fakultet, Beograd, Srbija<br>${ }^{2}$ Univerzitet u Novom Sadu, Poljoprivredni fakultet, Novi Sad, Srbija

Polimorfizam dehidrogenaza proučavan je kod 36 genotipova višnje (Prunus cerasus L.), trešnje ( $P$. avuim L.), magrive ( $P$. mahaleb L.), stepske višnje ( $P$. fruticosa Pall.), marele ( $P$. gondounii Redh.), japanske ukrasne trešnje ( $P$. serrulata Lindl.) kao i četiri interspecies hibrida koji se koriste kao standardne podloge za trešnju: Gisela 5, Gisela 6, Max Ma i Colt.. Za pripremanje enzimskog ekstrakta korišćena je unutrašnja kora jednogodišnjih grančica, prikipljenih u fazi mirovanja. Metoda vertikalne poliakrilamidne gel elektroforeze (PAGE) korišćena je za razdvajanje proteina za analizu enzimskih sistema: ADH (alkohol dehidrogenaze), FDH (format dehidrogenaze), GDH (glutamat dehidrogenaze), IDH (izocitrat dehidrogenaze), MDH (malat dehidrogenaze), PGD (fosfoglukonat dehidrogenaze) i SDH (šikimat dehidrogenaze). Polimorfizam je utvrđen za sve proučavane sisteme. Ukupno je utvrđeno 10 polimorfnih lokusa i to Adh -1 (3 genotipa) i Adh-2 (5 genotipova), Fdh-1 (2 genotipa), Gdh-1 (3 genotipa), Idh-1 (4 genotipa) i Idh -2 (5 genotipova), Mdh-1 (3 genotipa), Pgd-1 (4 genotipa), Sdh-1 (1 genotip) i Sdh-2 (3 genotipa). Primenom klaster analize dobijen je dendrogram na kome se mogu izdvojiti četiri grupe srodnih genotipova. Dobijeni rezultati ukazuju da se proučavani sistemi mogu uspešno koristiti u determinaciji roda Prunus, podroda Cerasus. Za utvrđivanje genetičke varijabilnosti najveći značaj imaju enzimski sistemi ADH, IDH i SDH. Polimrfizam FDH i GDH roda Prunus, podroda Cerasus je po prvi put opisan u ovom radu. Prvi rezultati o varijabilnosti dehidrogenaza kod Oblačinske višnje ukazuju da polimorfizam lokusa $I d h-2$ i $S d h-2$ može biti koristan za identifikaciji klonova.

