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The final authenticated version is available online at 10.1016/j.scienta.2021.110405

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| 1 | Genetic diversity of common bean (Phaseolus vulgaris L.) germplasm from Serbia, as revealed |
|----|---|
| 2 | by single sequence repeats (SSR) |
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| 18 | |
| 19 | Abstract |
| 20 | Genetic diversity and structure of common bean (Phaseolus vulgaris) germplasm from Serbia, |
| 21 | comprising 118 landraces and 18 cultivars, was assessed with the application of 27 Single Sequence |
| 22 | Repeats (SSR) markers. Thirteen accessions from Agricultural Institute of Slovenia were used as |
| 23 | references for gene pool determination. Main parameters of genetic diversity were calculated for each |
| 24 | SSR loci, i.e. number of different and rare alleles, number of effective alleles, Shannon's information |
| 25 | index, observed and expected heterozygosity and polymorphic information content. A total of 445 |
| 26 | allelic variants, with 16.5 alleles per locus on average, were detected. Mean gene diversity (He= 0.79) |
| 27 | indicated sufficient reservoir of genetic variation preserved in studied bean germplasm. Landraces |
| 28 | displayed higher variability compared to cultivars (405 in relation to 233 allelic variants). Genetic |

29 structure and relatedness of accessions was assessed by model-based method and hierarchical clustering method in combination with genetic distance calculation. The Bayesian clustering model implemented 30 in STRUCTURE software, on the primary level (K=2), revealed clear separation of accessions into two 31 groups, corresponding to gene pool affiliation. Mesoamerican gene pool (M) was represented with 32 33 23.5% of accessions, while Andean (A) was larger, composed of 68.4% of studied germplasm. Small group (8.1%) showed admixed genetic structure between two gene pools. Additional variation in respect 34 35 to two recognized gene pools was revealed (K=3), whose basis was acknowledged to be within Andean 36 gene pool. Further subdivision of accessions (K=8), mainly according to the seed forms, was observed. 37 Genetic distance analysis associated with Neighbour-joining clustering method revealed grouping 38 pattern of landraces and cultivars corresponding to the gene pool and their seed phenotypes. 39 Classification and structuring of the bean accessions according to and beyond the gene pool of origin 40 should facilitate conservation strategies and breeding of this material. Combining the information of 41 phenotypic variation obtained in previous research and molecular data reveled in this study will assist in selection of parental components for breeding, or in the choice of smaller sample in order to further 42 43 acknowledge their breeding value. In addition, obtained results of this work should serve as an 44 additional information on common bean germplasm variation in Western Balkans and beyond, in 45 Europe. 46

- 47 Keywords: *Phaseolus vulgaris*, SSR, genetic diversity, gene pool
- 48

49 1. Introduction

50

The common bean (*Phaseolus vulgaris* L.) is one of the most valuable vegetable crops for human consumption since it is rich in proteins, fibres, vitamins, minerals and many other antioxidant compounds (Broughton et al., 2003; Maras et al., 2015; Sitohy et al., 2020). Being a diverse crop in terms of cultivation methods, use, phenotypic diversity and range of environments to which it is adapted, the common bean is grown worldwide (De Ron et al., 2016). In many European countries, *Phaseolus vulgaris* is a significant component of a traditional diet and life. 57 Phaseolus vulgaris has undergone two independent domestication events in primary centres of origin: one in Middle America, and one in Andes. As a result, two highly differentiated gene pools were 58 59 formed: Mesoamerican and Andean, which are distinguished and recognized according to their phenotypic, biochemical and genotypic differences (Gepts et al., 1986; Gepts, 1999; Šuštar-Vozlič et 60 61 al., 2006; Raggi et al., 2013; Carović-Stanko et al., 2017; Gioia et al., 2019; Savić et al., 2020). These 62 two gene pools are also characterized by geographic and partial reproductive barriers (Gepts and Bliss, 1985; Gioia et al., 2013). Furthermore, within each gene pool, there is subdivision of this species to 63 64 many eco-geographic races and seed forms (market classes). Inter-gene pool and interracial crosses of genotypes can exhibit negative combining ability and lethality problems, which aggravates the breeding 65 of common bean (Singh et al., 1991; Kelly et al., 1998; Blair et al., 2007). 66

67 Gene pool affiliation has usually been determined based on variation of the main storage protein 68 of the common bean, phaseolin (Gepts and Bliss, 1988; Šuštar-Vozlič et al., 2006; Logozzo et al., 2007; Carović-Stanko et al., 2017; Savić et al., 2020). Mesoamerican origin of common bean genotypes is 69 70 associated with phaseolin types S (Sanilac), M (Middle America) and B (Boyaca), while genotypes with 71 phaseolin types T (Tendergreen), C (Contender) and H (Huevo de Huanchaco) belong to the Andean 72 gene pool. However, since Nani et al. (2011) identified three indel spanning markers SHP1-A, SHP1-73 B and SHP1-C, newer researches more rely on these marker systems for gene pool identification (Maras 74 et al., 2015, 2016; Pipan and Meglič, 2019).

75 From primary centres of origin and domestication in America, common bean spread worldwide 76 (Zeven, 1997; Maras et al., 2015). It is believed that the common bean arrived in Europe on two 77 occasions; Mesoamerican beans through Spanish and Portuguese exploration of the Americas around 78 1506; and Andean beans somewhere later, in 1528, during Pizzaro's expeditions in Peru (Gioia et al., 79 2013). In Europe, common bean landraces and cultivars evolved under diverse environments, cropping systems and farmers preferences (Zeven, 1997; Carović-Stanko et al., 2017; Pipan and Meglič, 2019). 80 In addition, outcrossing among Andean and Mesoamerican genotypes facilitated in the development of 81 high genotypic and phenotypic diversity of the European common bean (Rodino et al., 2006; Gioia et 82 83 al., 2013).

Angioi et al. (2010) and Gioia et al. (2013) focused their research on hybridization phenomena of Andean and Mesoamerican germplasm in Europe. The presence of hybrid genotypes in high proportion within the European germplasm was revealed with chloroplast microsatellites (cpSSRs) and two unlinked nuclear loci (phaseolin and PvSHP1). Hegay et al. (2013) and Sinkovič et al. (2019) observed signs of introgression on the phenotypic level. Zhang et al. (2008) believe that identification of putative hybrids is of special interest since these genotypes are a reflection of growing regions and are adapted to ecological conditions within that region.

91 In Serbia, people have grown common bean for centuries, establishing it as an important part 92 of their diet and traditional life. Even though commercial cultivars have been developed and are widely 93 represented in production, common bean landraces are still maintained and used by farmers and people 94 in rural and marginal areas (Scarano et al., 2014; Mallor et al., 2018; Savić et al., 2020). In Europe 95 (including Serbia) development of new cultivars was instituted with the aim to maintain phenotypes 96 within each seed form (market class), usually by selecting parental components among elite material. 97 This resulted in narrowing the genetic base of this elite germplasm, compromising long-term genetic 98 gain (McClean and Lee, 2007; Gioia et al., 2019). In order to address these problems, breeders sought 99 to incorporate new variability, most commonly by exploring the existing diversity found among 100 landraces and unadapted germplasm.

101 Genetic collections of the common bean in Serbia, consisting of seeds of traditional and modern 102 cultivars and landraces, are maintained within breeding institutes (Vasić et al., 2009). Knowledge of 103 genetic diversity preserved in these collections is crucial for proper conservation, further research, 104 selection of parental components and for defining breeding strategies. Landraces are described as 105 genetically diverse material with traits specific for growing regions. They are traditionally grown in low 106 input systems, adapted to local agro-climatic conditions and display a high level of phenotypic diversity. 107 All of this makes landraces interesting material for conservation, research and implementation in 108 breeding programs (Carović-Stanko et al., 2017; Gioia et al., 2019).

Selected set of landraces and commercial cultivars from Serbia have already been characterized
 for diversity on a phenotype level, by assessing morphological traits chosen according to international
 descriptors. In addition, gene pool of origin was identified based on variation of phaseolin types (for

| 112 | more information see Savić et al., 2020). However, in order to better understand genetic variation and |
|-----|--|
| 113 | relationships among landraces and cultivars on a molecular level, complementary study on previous |
| 114 | research was performed. Single sequence repeats (SSR) markers were chosen for the analysis, because |
| 115 | they are abundant and widely distributed in the genome, codominantly inherited, highly polymorphic |
| 116 | and repeatable (Yu et al., 1999; Maras et al., 2015; Pipan and Meglič, 2019). Therefore, the aim of this |
| 117 | study was to: (i) assess the allelic diversity of the common bean germplasm from Serbia and determine |
| 118 | relationships among the accessions, and (ii) investigate the genetic structure and organization of genetic |
| 119 | diversity of the studied germplasm within and beyond gene pools of origin. |
| | |

- 120
- 121 2. Material and Methods
- 122

124

A total of 136 accessions from the Serbian common bean genetic collection maintained at the 125 126 Institute of Field and Vegetable Crops, Novi Sad (IFVCNS) were analysed in this paper. This included 127 118 local landraces collected from 53 sites in Serbia (Supplementary Material 1) in timescale of 1970-2014 and 18 commercially available cultivars from Serbia: Rozalija (C1), Žutotrban (C2), Sremac (C3), 128 129 Balkan (C4), Slavonski žutozeleni (C5), Pasuljica P-1 (C6), Biser (C7), Medijana (C8), Oplenac (C9), 130 Panonski gradištanac (C10), Panonski tetovac (C11), Aster (C12), Poboljšani gradištanac (C13), Galeb 131 (C14), Zlatko (C15), Dvadesetica (C16), Belko (C17) and Maksa (C18), Table 1. Part of the studied material was collected during field expeditions, while the rest was acquired via seed exchange with 132 other institutions. Landraces and cultivars were classified according to seed traits in several forms, most 133 commonly grown in Serbia: Roseus (pink seed colour), Versicolor (seed coat pattern), Griseus 134 (greenish-yellow seed colour), Aureus (yellow and golden-yellow seed coat colour), Albus (white seed 135 colour) and other (red, cream, brown, black seed colour). 136

As references for gene pool determination, accessions of familiar phaseolin type (type T –
PHA131, PHA306, PHA309, PHA318, PHA336; type C – PHA181, PHA222, PHA29, PHA315,
PHA390; type S – PHA245, PHA368, PHA371) from Agricultural Institute of Slovenia (AIS) were

^{123 2.1.} Plant material

included in the study (Supplementary Material 1). Phaseolin types T and C indicate germplasm origin
from Andean, while the Mesoamerican gene pool is determined with type S. For detailed phenotypic
characterization, see Savić et al. (2020).

143

144 2.2. SSR analysis

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Plants were grown in greenhouse conditions until the phase of the first true leaves at
Agricultural Institute of Slovenia. DNA was extracted from a total of 60 to 100 mg of bulked fresh plant
tissue (4 plants per accession), using a BioSprint 15 DNA Plant Kit (Qaigen) on KingFisher (Thermo)
isolation robot according to optimized manufacturer's instructions.

A set of 27 genome-specific SSR (single sequence repeats; microsatellites) markers distributed across all linkage groups was used for genetic diversity and genetic structure analysis of selected material (Supplementary Material 1). For the identification of genotypes gene pool affiliation (Mesoamerican/Andean) three indel spanning markers (SHP1-A, SHP1-B and SHP1-C) developed by Nanni *et al.* (2011) were used.

The final volume of PCR reaction was 11.5 μ L, which included: 8.4 ng genomic DNA, 1 μ L 10x PCR buffer (Biotools), 0.2 μ L of each 10 mM dNTP (Sigma-Aldrich), 0.5 μ L of 50mM MgCl₂ (Biotools), 0.1 μ L of 10 μ M forward primer (Sigma-Aldrich), 0.25 μ L of 10 μ M reverse primer (Sigma-Aldrich), 0.183 μ L of 10 μ M 5'-fluorescently labelled primer (6-FAM, NED or HEX; Omega), and 0.5 μ L of 5 U Taq DNA polymerase (Biotools). The forward primer of each SSR had an added 18-bp tail sequence of 5'-TGTAAAACGACGGCCAGT-3' (M13(-21)).

PCR analyses were performed on a thermal cycler (Veriti, ThermoFisher Scientific) under touch-down conditions: 94°C for 4 min; 15 cycles at 94 °C for 1 min; decreased temperature from 60 (62) °C to 49.5 (51.5) °C at 0.7 °C per cycle for 30 s; 72 °C for 1 min; followed by 23 cycles at 94 °C for 30 s; 53 °C for 30 s; 72 °C for 1 min; and final extension for 5 min at 72 °C, as described by Pipan and Meglič (2019). PCR conditions were dependent on each primer pair. Fragment analysis was performed on a genetic analyser (3130XL; Applied Biosystems). Allele lengths were determined by

- 167 comparison with an internal size standard (GeneScan-350 ROX; Applied Biosystems) using the
- 168 GeneMapper 4.0 software (Applied Biosystems).
- 169

| Accession | Туре | Seed form | Accession | Туре | Seed form |
|-----------|------|------------|-----------|------|------------|
| L1 | III | other | L56 | Ι | Versicolor |
| L2 | III | Aureus | L57 | Ι | Griseus |
| C1 | Ι | Roseus | L58 | Ι | Albus |
| L3 | Ι | Roseus | L59 | ΙΙ | Albus |
| L4 | Ι | Roseus | L60 | Ι | Griseus |
| L5 | Ι | Roseus | C14 | Ι | Albus |
| L6 | Ι | Roseus | C15 | Ι | Aureus |
| L7 | Ι | Roseus | C16 | Ι | Albus |
| L8 | Ι | Roseus | C17 | Ι | Albus |
| L9 | II | Griseus | C18 | Ι | Albus |
| L10 | Ι | Griseus | L61 | Ι | Griseus |
| L11 | II | Albus | L62 | Ι | Griseus |
| L12 | Ι | Albus | L63 | Ι | Griseus |
| C2 | Ι | Versicolor | L64 | Ι | Griseus |
| L13 | Ι | Versicolor | L65 | Ι | Aureus |
| L14 | Ι | Versicolor | L66 | Ι | other |
| L15 | Ι | Versicolor | L67 | Ι | other |
| L16 | Ι | Versicolor | L68 | Ι | Griseus |
| L17 | Ι | Versicolor | L69 | Ι | Griseus |
| L18 | Ι | Versicolor | L70 | Ι | Versicolor |
| L19 | Ι | Versicolor | L71 | Ι | other |
| L20 | Ι | Griseus | L72 | Ι | other |
| L21 | Ι | Griseus | L73 | Ι | Griseus |
| L22 | Ι | Griseus | L74 | Ι | Griseus |

170 Table 1. List of accessions from Serbian common bean genetic collection, used for genetic analysis

| L23 | Ι | Griseus | L75 | Ι | Griseus |
|-----------|----|---------|------|-----|------------|
| L24 | Ι | Griseus | L76 | Ι | Versicolor |
| L25 | Ι | Griseus | L77 | Ι | Albus |
| L26 | I | Griseus | L78 | Ι | Albus |
| L27 | I | Griseus | L79 | III | Albus |
| L28 | Ι | Aureus | L80 | Ι | Griseus |
| L29 | Ι | other | L81 | Ι | Griseus |
| L30 | Ι | other | L82 | Ι | Versicolor |
| L31 | Ι | Griseus | L83 | Ι | Albus |
| L32 | Ι | other | L84 | Ι | Griseus |
| L33 | Ι | other | L85 | Ι | Versicolor |
| L34 | Ι | Griseus | L86 | Ι | Griseus |
| L35 | Ι | other | L87 | Ι | other |
| L36 | Ι | other | L88 | Ι | Albus |
| L37 | Ι | Griseus | L89 | III | Albus |
| L38 | Ι | Roseus | L90 | Ι | other |
| L39 | Ι | Roseus | L91 | Ι | Albus |
| L40 | Ι | other | L92 | Π | Aureus |
| L41 | Ι | other | L93 | III | Albus |
| L42 | Ι | Roseus | L94 | II | other |
| L43 | II | Aureus | L95 | Ι | Griseus |
| L44 | Ι | Aureus | L96 | Ι | Versicolor |
| L45 | Ι | Aureus | L97 | Ι | other |
| L46 | Ι | Aureus | L98 | III | Albus |
| C3 | Ι | Griseus | L99 | Ι | Versicolor |
| C4 | Ι | Albus | L100 | Ι | other |
| C5 | Ι | Griseus | L101 | Ι | Versicolor |
| C6 | II | Albus | L102 | Ι | Griseus |
| C7 | Ι | Albus | L103 | Ι | Griseus |
| C8 | II | Albus | L104 | III | Albus |

| L47 | Ι | Albus | L105 | Ι | Albus |
|-----|-----|------------|------|-----|------------|
| L48 | Ι | Albus | L106 | Ι | Albus |
| L49 | Ι | Albus | L107 | II | Albus |
| L50 | Ι | Griseus | L108 | Ι | Versicolor |
| L51 | Ι | Albus | L109 | Ι | Albus |
| С9 | Ι | Albus | L110 | Ι | Versicolor |
| L52 | Ι | Albus | L111 | Ι | Versicolor |
| L53 | Ι | Albus | L112 | Ι | Albus |
| C10 | Ι | Albus | L113 | Ι | Albus |
| C11 | Ι | Albus | L114 | Ι | Griseus |
| L54 | II | Aureus | L115 | Ι | Versicolor |
| C12 | Ι | Albus | L116 | III | Albus |
| C13 | III | Albus | L117 | Ι | Albus |
| L55 | Ι | Versicolor | L118 | Ι | Albus |



Type – plant growth habit (I – determinate bush, II – indeterminate bush,

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III – indeterminate prostrate or vining)

173 2.3. Data analysis

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For each SSR locus, main parameters of genetic diversity were calculated in GenAlEx 6.1
(Peakall and Smouse, 2006) and Microsatellite-Toolkit (Park, 2001) software. They included number
of alleles (Na), number of alleles with frequency ≥ 5%, number of rare alleles, number of effective
alleles (Ne), Shannon's information index (I), observed heterozygosity (Ho), expected heterozygosity
(He) and polymorphic information content (PIC).

STRUCTURE 2.3.3 software was employed to establish genetic structure of common bean collection. STRUCTURE uses Bayesian clustering approach, applying Markov Chain Monte Carlo (MCMC) algorithm, to study difference in accessions distribution among population by placing accessions into clusters that share similar variation patterns. Bayesian clustering approach is described by the posterior probability that each accession belong to each genetic cluster, while each cluster (K) is characterized by a subset of allelic frequencies identified in the data. Beside determining and assigning 186 accessions to genetic clusters, this method is used to identify admixed accessions by investigating hybridization zones of individuals from different clusters that give genetically recombined offspring 187 188 (Pritchard et al., 2009). In the study of common bean collection from Serbia, the most probable number of clusters (Q value) was determined with ten independent runs for each K (1 to 10) for the admixture 189 190 model, with burning period of 50.000 followed by 500.000 Markov Chain Monte Carlo repeats. MCMC 191 algorithm initiates by randomly assigning accessions to a pre-determined number of clusters. After that, 192 allelic frequencies are estimated for each cluster and accessions are re-assigned based on those 193 frequency assessments. According to Evanno delta K statistics (Evanno et al., 2005), implemented in 194 the software Structure Harvester (Earl and von Holdt, 2011), the real K value was determined based on 195 the increase in the likelihood rations between the runs. An accession is assigned to a specific cluster 196 when the percentage of membership was $Q \ge 80\%$, while the accessions with membership coefficient 197 Q < 80% are believed to be of admixed origin (putative hybrids).

For more detailed analysis of genetic structure and relationships among genotypes, DARwin
software (https://darwin.cirad.fr/) was applied to perform cluster analysis based on similarity matrix
and construct dendrogram using Neighbour-joining method (NJ).

201

202 **3. Results**

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204 The whole set of 27 SSR markers chosen for the diversity study of the Serbian common bean genetic collection proved to be polymorphic, producing more than one allelic variants. A total of 445 205 alleles were scored for the studied collection, which included cultivars, landraces and reference 206 accessions. The average allele number per SSR was 16.5, ranging from six alleles for loci BM155, 207 BM210 and BMd044 to 25 alleles for loci GATS91 and ATA002. The highest number of effective 208 alleles (9.35) and the most alleles with frequency over >5% were found in locus BMd001. Total number 209 of rare alleles was 303, which accounted for 68.1% of all alleles detected in the studied germplasm 210 211 (Table 2).

212

Table 2. Parameters of genetic variability of Serbian common bean collection and reference cultivars

| Locus | Na | Allele | Number | Ne | Ι | Но | He | PIC |
|----------|----|-----------|---------|------|------|------|------|------|
| | | frequency | of rare | | | | | |
| | | ≥ 5% | alleles | | | | | |
| ATA003 | 19 | 6 | 13 | 6.87 | 2.26 | 0.74 | 0.85 | 0.84 |
| ATA004 | 15 | 4 | 11 | 3.82 | 1.77 | 0.44 | 0.74 | 0.71 |
| ATA005 | 20 | 6 | 14 | 6.60 | 2.28 | 0.93 | 0.85 | 0.83 |
| ATA007 | 17 | 5 | 12 | 3.42 | 1.74 | 0.30 | 0.71 | 0.68 |
| ATA016 | 12 | 5 | 7 | 4.41 | 1.78 | 0.91 | 0.77 | 0.74 |
| GATS91 | 25 | 5 | 20 | 8.38 | 2.49 | 1.00 | 0.88 | 0.87 |
| ATA002 | 25 | 5 | 20 | 5.31 | 2.09 | 0.85 | 0.81 | 0.79 |
| BM172 | 22 | 6 | 16 | 8.43 | 2.45 | 0.92 | 0.88 | 0.87 |
| BMd001 | 24 | 9 | 15 | 9.35 | 2.46 | 0.99 | 0.89 | 0.88 |
| ATA020 | 21 | 5 | 16 | 6.52 | 2.27 | 0.65 | 0.85 | 0.83 |
| Pv-ag004 | 21 | 5 | 16 | 5.80 | 2.14 | 0.94 | 0.83 | 0.81 |
| ATA010 | 13 | 5 | 8 | 4.25 | 1.75 | 0.31 | 0.77 | 0.73 |
| BM155 | 6 | 4 | 2 | 3.53 | 1.36 | 0.87 | 0.72 | 0.67 |
| BM170 | 20 | 6 | 14 | 8.07 | 2.40 | 0.95 | 0.88 | 0.87 |
| BM183 | 8 | 5 | 3 | 3.26 | 1.41 | 0.93 | 0.69 | 0.64 |
| BM210 | 6 | 2 | 4 | 2.14 | 0.85 | 0.97 | 0.53 | 0.42 |
| BMd044 | 6 | 4 | 2 | 2.69 | 1.23 | 0.45 | 0.63 | 0.58 |
| ATA009 | 22 | 8 | 14 | 8.18 | 2.45 | 0.75 | 0.87 | 0.87 |
| ATA145 | 20 | 3 | 17 | 3.68 | 1.80 | 0.31 | 0.73 | 0.69 |
| GA16 | 13 | 6 | 7 | 6.12 | 2.04 | 0.68 | 0.84 | 0.82 |
| ATA006 | 22 | 7 | 15 | 8.98 | 2.52 | 0.79 | 0.89 | 0.88 |
| BM157 | 14 | 6 | 8 | 5.88 | 2.00 | 0.60 | 0.83 | 0.81 |
| BMd042 | 18 | 4 | 14 | 6.08 | 2.18 | 0.98 | 0.84 | 0.82 |
| ATA289 | 17 | 4 | 13 | 3.87 | 1.74 | 0.97 | 0.74 | 0.71 |
| PvSHP1-A | 13 | 6 | 7 | 5.52 | 1.93 | 0.95 | 0.82 | 0.79 |
| PvSHP1-B | 11 | 6 | 5 | 5.89 | 1.89 | 1.00 | 0.83 | 0.81 |
| PvSHP1-C | 15 | 5 | 10 | 5.45 | 2.03 | 0.92 | 0.82 | 0.79 |

| average | 16.5 | 5.2 | 11.2 | 5.65 | 1.97 | 0.78 | 0.79 | 0.77 |
|---------|------|-----|------|------|------|------|------|------|
| total | 445 | 142 | 303 | | | | | |

214 215 Na – number of alleles, Ne – number of effective alleles, I - Shannon's information index, Ho –observed heterozygosity, He - expected heterozygosity, PIC - polymorphic information content

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Loci BMd001 and ATA006 generated the highest values of expected heterozygosity (0.89) and PIC (0.88). The locus ATA006 scored the highest value of Shannon's information index (2.58). Observed heterozygosity ranged from 0.30 (locus ATA007) to 1.00 (loci GATS and PvSHP1-B). Average values of Shannon's information index (1.97), expected heterozygosity (0.79) and PIC (0.77) indicated that all SSR markers showed sufficient polymorphism and are suitable for common bean diversity study.

223 Main genetic diversity parameters were calculated for each common bean form determined according to seed traits (Table 3). Overall, landraces and cultivars from the Albus group showed the 224 225 greatest diversity for all parameters, except number of alleles with a frequency higher than 5%. On 226 contrary, the lowest diversity was found among accessions from the *Rosues* group, with slightly larger 227 expected heterozygosity (0.76). The highest average number of alleles with a frequency higher than 5% 228 was observed in the Aureus form. Rare alleles were not found among common bean genotypes from 229 Roseus and Aureus groups, while for the other groups ranged from 3 (Versicolor) to 4.96 (Albus). Gene 230 diversity (He) ranged from 0.67 (Griseus) to 0.78 (Albus). Observed heterozygosity was the highest among accessions from other (0.82) and Aureus (0.81) groups. Almost twice as many alleles were 231 scored among landraces (405) compared to cultivars (233). In addition, the percentage of rare alleles 232 233 was much larger in landraces (67.6%) in relation to cultivars (29.6%). Average values of all the other parameters of genetic diversity (number of effective alleles, Shannon's information index, observed and 234 235 expected heterozygosity) were similar between the two groups.

236

Table 3. Genetic diversity calculated for 27 SSR loci considering six groups defined according to the
 seed form; cultivars and landraces separately

| | Total | | • • | Allele frequency | Number of rare | Ne | I | Но | Не |
|------------|------------|-----|---------|---------------------|-------------------|------|------|------|-----|
| Forms | number of | Na | Na | | | | | | |
| | accessions | | uveruge | ≥5% | alleles | | | | |
| Roseus | 10 | 144 | 5.33 | 5.33 | 0 | 3.39 | 1.33 | 0.66 | 0.′ |
| Versicolor | 21 | 205 | 7.59 | 4.59 | 3 | 4.22 | 1.57 | 0.72 | 0.7 |
| Grisues | 35 | 231 | 8.55 | 4.52 | 4.03 | 3.88 | 1.49 | 0.79 | 0.0 |
| Aureus | 10 | 175 | 6.48 | 6.48 | 0 | 4.33 | 1.56 | 0.81 | 0.7 |
| Albus | 42 | 294 | 10.89 | 5.93 | 4.96 | 5.17 | 1.84 | 0.77 | 0.7 |
| other | 18 | 242 | 8.96 | 5.06 | 3.52 | 5.06 | 1.76 | 0.82 | 0.7 |
| cultivars | 18 | 233 | 8.63 | 6.07 | 2.56 | 5.28 | 1.79 | 0.79 | 0.2 |
| landraces | 118 | 405 | 15.00 | 4.85 | 10.15 | 5.32 | 1.91 | 0.78 | 0. |



heterozygosity, He - expected heterozygosity

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According to the Bayesian clustering model implemented in STRUCTURE software, the most 242 informative number of subgroups was two (K=2). The second largest peak of Delta K value was 243 observed for three subgroups (K=3), while studied accessions also displayed classification at eight 244 245 subgroups (K=8) (Fig. 1).

Affiliation to the gene pools was identified when maximum likelihood and Delta K values were 246 247 two (K=2), assigning accession to Mesoamerican (M; marked red in Fig. 2a) or Andean (A; marked 248 green) group. This was confirmed based on allocation of reference accessions for gene pool 249 classification. Total of 91.9% genotypes had membership coefficients higher than 0.80, implying that the majority of samples were strongly assigned to the groups. In addition, 11 genotypes (8.1%) were 250 251 with membership coefficients lower than 0.80, showing admixed genetic structure between two groups 252 (gene pools) (Supplementary Material 1). First group (M) included 32 genotypes (23.5% of studied 253 germplasm) and reference accessions with mainly Mesoamerican phaseolin type. It comprised 21 254 landraces and 11 cultivars. A majority of genotypes in group M were from the Albus seed form (84%) 255 with primarily medium (56%) and large (34%) seed weight. Second group (A) comprised 93 genotypes (68.4% of studied germplasm) and reference accessions with mainly Andean phaseolin types (Fig. 2).
It included 88 landraces and five cultivars. Genotypes with coloured and medium to large seeds
predominated in this group (*Griseus, Aureus, Roseus* and *Versicolor* forms). Only 13 genotypes within
second group had white seed coats with large seed weight in general. Genotypes identified as potential
hybrids between the gene pools (11 accessions) according to STRUCTURE analysis were primarily
landraces (5 *Albus, 3 Aureus, 1 Grisues, 1 Versicolor* and 1 *other*) with phaseolin type T.

For K = 3 grouping pattern, further division of the Andean group into two additional subgroups 262 263 was observed. In this scenario, 88.9% of genotypes had membership coefficient higher than 0.80, while the rest showed admixed origin between the three subgroups (11.1%). First subgroup, M (marked red 264 in Fig. 2b), remained the same, comprising 23.5% of studied germplasm. Second subgroup, A1 (marked 265 266 green), was composed of 45 landraces that belonged to mainly Griseus, Roseus and Versicolor forms, 267 two cultivars (Oplenac and Aster) of large white seeds and six reference accessions. Third subgroup, 268 A2 (marked blue), comprised 34 landraces of largely Griseus and Versicolor common bean forms, 4 cultivars (Rozalija, Žutotrban, Sremac and Slavonski žutozeleni) and three reference accessions. 269

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Figure 1. Estimation of most likely subgroups number according to Evanno's method (Evanno et al.,

2005)

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analysis (Figure 3). The Neighbour joining-based dendrogram divided 136 genotypes and reference

293 accessions into two main clusters (gene pools), Mesoamerican and Andean, with additional subclusters



294 identified within each main cluster.

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Figure 3. Neighbour joining tree of 136 landraces and cultivars and 13 reference accessions based on
 similarity matrix (simple matching coefficient)

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In total, 32 genotypes (23.5%) were classified in Mesoamerican (M) cluster. Three subclusters are recorded within cluster M. Subcluster M1 included 9 landraces and 3 cultivars (Pasuljica P1, Panonski gradištanac, Belko) with seed weights from small to large. Subcluster M2 consisted of 12 landraces and 6 cultivars (Balkan, Biser, Panonski tetovac, Poboljšani gradištanac, Galeb, Maksa) with medium to large seeds. Two landraces and one cultivar (Medijana) clustered in subcluster M3. All accessions in this group had medium seed weight. Great majority of accessions in cluster M belonged to the *Albus* form.

The Andean (A) cluster comprised 75% of studied landraces and cultivars in total. Subluster A1 consisted of 31 landraces and 2 cultivars (Sremac and Slavonski žutozeleni). Accessions largely belonged to *Grisues*, followed by *Versiclor* in form. In total, 79% of genotypes in subcluster A1 corresponded to subgroup A2 identified in STRUCTURE analysis when K=3, while the rest were of admixed origin. Therefore, it can be suggested that genotypes comprising these two subclustersrepresent novel variation, which was created in this region or was introduced from different sources.

Andean subcluster A2 in NJ dendrogram consisted of 40 landraces and 2 cultivars (Oplenac, Zlatko). Landraces and cultivars predominantly belonged to *Grisues* form (48%); however, other seed forms were observed in smaller number – *Roseus*, *Versicolor* and *Aureus*. Only two landraces had a white seed coat. Subcluster A3 comprised 22 landraces, three cultivars (Rozalija, Žutotrban, Aster) and seven reference accessions. The great majority of accessions belonged to *Versicolor* (32%) and *Albus* (28%) forms. Five landraces (L2, L54, L59, L109 and L110) and two reference accessions (PHA390 and PHA131) deviated from this grouping pattern and were separated on the dendrogram.

319

320 4. Discussion

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322 For proper conservation, assessment of breeding value and organization and structuring of breeding programs, it is essential to identify levels of phenotypic and genetic diversity preserved in the 323 324 germplasm of any crop species (Mhlaba et al., 2018). Common bean accessions examined in this paper 325 have already been characterized for their phenotypic variability, while origin and dissemination of local 326 bean germplasm from Serbia was assessed based on variation of phaseolin types (see Savić et al., 2020). 327 Therefore, in order to further reveal genetic diversity preserved in studied germplasm, relatedness of 328 landraces and cultivars on a molecular level, as well as their structure within and beyond gene pools of 329 origin, the present study was performed, which should serve as complementary to the previous one.

330 Results of this research indicated that substantial allelic diversity was preserved in the germplasm from Serbia and reference accessions. Total number of alleles (Na) and gene diversity (He) 331 were notably higher when compared to germplasm from Portugal, Italy and Croatia (Leitao et al., 2017; 332 Carović-Stanko et al., 2017; Gioia et al., 2019). On the other hand, average number of alleles per locus 333 was comparable to those reported in common bean core collection by Blair et al. (2009). Even though 334 our material was collected from a relatively narrow geographic region and was smaller in size compared 335 336 to the other research, obtained result could be due to high percentage of rare alleles (68.1%) identified 337 in this paper. In addition, this research material included samples, mainly landraces, from both gene

pools belonging to various seed forms that represented great morphological diversity, which could have
affected allelic variability recorded in present study. In addition, the geographic position of Serbia,
which is a familiar trade and migration crossroad from east to west, might have been significant in
shaping common bean diversity found in this region (Vasić et al., 2009).

342 Genetic variability was also measured as the amount of actual or potential heterozygosity. Observed heterozygosity (Ho) represents the level of heterozygous individuals in the population 343 344 compared with expected heterozygosity (He) which reflects the genetic diversity at the specific loci 345 along genotypes due to the degree of its out-crossing potential (Stainer, 2010; Pipan et al., 2013). In the 346 case when Ho is equal to He, it means that the population is in Hardy-Weinberg equilibrium, among 347 other random crosses. In addition, the small deviations between average Ho and He (0.01, Table 2) can 348 indicate uniform abundance of alleles along Serbian common bean germplasm that could reflect their 349 common genetic origin. Moreover, there are some loci where the deviation between Ho and He is higher 350 than 0.3, i.e. loci ATA004, ATA007, ATA010, BM210 and ATA145 (Table 2). Those loci could be 351 highly applicable to evaluate cross-pollination potential of common bean under filed conditions.

Gene diversity (He) recorded in this study, which is not dependable on the sample size, was 352 353 much higher compared to the results of Leitao et al. (2017) and Carović-Stanko et al. (2017). On the 354 other hand, it mostly corresponded to that found in bean germplasm from Western Balkan countries (former Yugoslav republics) by Maras et al. (2015). This could be due to frequent material exchange, 355 gene flow between the countries that constituted former Yugoslavia, and a similar set of markers used. 356 357 It was also revealed that gene diversity (He) of each bean group generated according to seed form in this work was larger compared to that observed for common bean market classes from USA by Gioia 358 et al. (2019). Bearing in mind that mentioned authors investigated elite advanced cultivars compared to 359 360 landraces analysed in this research, it could be suggested that breeding interventions have narrowed the 361 genetic basis of elite material in comparison to landraces. This was also proven with more allelic 362 variants observed among landraces (405) compared to cultivars (233) in our study. Another interesting fact is that a larger percentage of rare alleles were found in landraces (67.6%) in relation to cultivars 363 364 (29.6%), even though other parameters of genetic diversity were quite similar for these two groups, 365 which also corresponded to values found for the entire collection. This revelation is important from a

breeding perspective, allowing breeders to use this unexplored variability preserved among landraces and cultivars in their advantage. Conversely, allelic variability differed among the groups. Accessions from *Albus* and *Griseus* forms proved to be the most variable, which could also be related to a proportionally larger number of accessions in these seed forms. Therefore, differences in allelic variability found in various other research could be in line with nature, number and variability of accessions assessed, geographic origin of studied material, DNA isolation and detection methods.

372 The genetic structure of the studied germplasm, on primary level, corresponded to familiar 373 differentiation of common bean accessions according to gene pool affiliation, Mesoamerican or 374 Andean. These results are in accordance with various investigations of *Phaseolus vulgaris* worldwide 375 (Sicard et al., 2005; Kwak and Gepts, 2009; Blair et al., 2010; Raggi et al., 2013; Bitocchi et al., 2017). 376 Separation of Serbian accessions into two recognized gene pools have already been shown based on 377 phaseolin and phenotypic variation, with a considerably larger proportion of accessions belonging to 378 the Andean gene pool (Savić et al., 2020), which is in accordance with results presented in this paper. 379 The prevalence of Andean in contrast to Mesoamerican accessions, along with congruency in bean 380 accessions clustering according to gene pool of origin (estimated at 95%) using different methods in 381 these two research, was determined. It was also shown that a chosen set of SSR markers, including the 382 combination of the PvSHP1 markers, proved their usefulness and efficiency in discriminating bean 383 accessions according to the gene pool affiliation, as suggested by Nanni et al. (2011), Maras et al. (2015, 384 2016), Pipan and Meglič (2019).

385 Moreover, application of molecular versus phenotypic markers revealed further subdivision of the studied germplasm from Serbia. Additional variation in respect to two recognized gene pools was 386 identified in both STRUCTURE analysis when K=3 (subgroup A2) and based on genetic distance 387 analysis combined with NJ dendrogram (subcluster A1). Accessions with membership coefficients of 388 389 these two groups largely corresponded to each other. It was also acknowledged that this additional 390 variation is concentrated within the Andean gene pool, which is in line with findings of Maras et al. (2015) for bean germplasm from Western Balkan countries, Raggi et al. (2013) for Italian and Leitao 391 392 et al. (2017) for Portuguese beans. It is believed that potential sources of this distinctive variation could 393 be accessions with admixed genetic bases derived from inter-gene pool crosses. Apart from that, it is

probable that unique variation was generated in this geographic area as a result of genotypic and phenotypic adaptation to local growing conditions during long period of cultivation. For the Serbian common bean germplasm, it was observed that mainly landraces with greenish-yellow (*Griseus*) and mottled (*Versicolor*) seeds comprised the mentioned clusters, and were marked as new variation. Since these seed types, apart from white-seeded cultivars and landraces, are favourite among the bean producers, it is possible that new variation is a result of farmer's selection towards most tolerant and high yielding landraces in those seed forms.

401 The theoretical speculation mentioned above is supported by the revelation of putative hybrid 402 genotypes in STRUCTURE analysis. In the case where phaesolin type T predominated among these 403 accessions, it was determined that genetic bases of this material was also within the Andean gene pool. 404 Observed frequency of putative hybrids in this study was quite low (8.1%), but nonetheless in 405 accordance with results of Blair et al. (2010), Gioia et al. (2013) and Maras et al. (2015). On the 406 contrary, Angioi et al. (2010) observed larger percentage (44%) of accessions derived from inter-gene 407 pool hybridization indicating a high contribution of admixed genotypes in the European common bean. 408 Santalla et al. (2002), Logozzo et al. (2007) and Gioia et al. (2013) highlighted the significance of 409 identification of such accessions in certain bean germplasm collections and their breeding value. 410 Putative hybrids might possess new and interesting combination of traits created in inter-gene pool 411 crosses, which could be related to higher adaptability to environmental stress, tolerance to pests and 412 pathogens, better productivity and overcome negative correlation between seed weight and yield 413 potential.

414 Relatedness of landraces and cultivars was discussed based on genetic distance analysis associated with Neighbour-joining clustering method, while subsequent genetic structure was revealed 415 when K=8 was considered in STRUCTURE analysis. In both methods, it was obvious that landraces 416 417 and cultivars formed subgroups according to their phenotype (with several exceptions). Arguably, 418 however, a genetic distance-based method might better assist in differentiation among accessions of similar phenotypes and in selection of more genetically distant parenting components for breeding. In 419 420 the case of common bean, it would support breeding of cultivars in specific seed type (form) with 421 improved productivity. Different types of clustering methods in combinations with genetic distance

422 calculations were applied. Observing the fact that both landraces and cultivars were used within 423 calculations, the combination of NJ and simple matching coefficient was the best choice to evaluate and 424 visualise the genetic relations and distribution of Serbian genotypes according their genetic origin (gene 425 pool determination). Moreover, the distribution of genotypes among clusters (A1-A3; M1-M3) on 426 Figure 3 is also correlated with seed characteristics for both, Andean and Mesoamerican group, 427 respectively.

428 A larger number of groups distinguished among landraces indicate existent diversity of this 429 material, despite the fact that farmers have maintained landraces during longer period on the farms close 430 to each other, exchanged seeds with surrounding farms and among themselves. Our results are in 431 accordance with research of Masi et al. (2009) and Raggi et al. (2013) who made similar observations. 432 Even though landraces clustered largely according to the seed traits, some deviations of this pattern 433 were observed. The largest discrepancy was detected among accessions belonging to the Albus seed 434 form. Even though these accessions are mainly of Mesoamerican origin, large white-seeded types were also observed in Andean groups. There are two possible explanations of this finding. Firstly, breeding 435 436 intervention in Serbia were mostly done within white-seeded beans, which could have resulted in 437 hybridization between the gene pools. Secondly, common bean landraces with white large seeds, 438 belonging to Nueva Granada race were introduced and are grown in Serbia. In addition, accessions of 439 Roseus and Versicolor group often clustered together. Although these accessions are usually 440 distinguished by their phenotype, they were not well separated on molecular level in this research.

Distribution of Serbian cultivars in distinctive, genetically diverse subclusters in NJ 441 dendrogram is a result of various selection and breeding criteria over time. It is well known that there 442 is accepted tendency to satisfy market demands in term of making cultivars that phenotypically 443 correspond to specific market class, or seed form (Geravandi et al., 2020). In various timeframes of 444 common bean breeding in Serbia, different available material was used, which also included 445 introduction of foreign germplasm for breeding purposes. This, together with adaptation of newly 446 created material to environmental and growing conditions at the time, resulted in genetic divergence of 447 448 Serbian common bean assortment. On the other hand, there were cases when two or more cultivars were

more closely positioned on the dendrogram. The most probable cause of this phenomenon is thecommon genetic origin they share, belonging to the same ancestral line.

In a conclusion, examined landraces and cultivars displayed marked genotypic variation, which 451 allowed detailed description of diversity present in common bean germplasm from Serbia, 452 453 accompanying the information on phenotypic variability previously assessed. Classification and structuring of the accessions in accordance within and beyond gene pool of origin should facilitate 454 conservation strategies and breeding of this material. A combination of phenotypic and molecular data 455 should allow researchers to make a selection of smaller number of accession for the core collections, 456 which will further be assessed for their breeding value (productivity, nutritional value, tolerance to 457 biotic and abiotic stress). As a result of the information presented in this research, it is hoped that the 458 459 study of common bean genetic diversity in the Western Balkans and the rest of Europe has been 460 purposefully expanded.

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462 Acknowledgement

This research was supported by the Ministry of Education, Science and Technological Development of the Republic of Serbia, Grant Number: 451-03-9/2021-14/200032, which covered experimental design, statistical data processing, manuscript writing and partially material analysis. It was also funded partially by the Slovenian Ministry of Agriculture, Forestry and Food and the Slovenian Research Agency, grants number P4–0072 and L4–7520, respectively. Under the applicative research project L4– 7520 we have designed the study and performed analyses. The funding under Agrobiodiversity research pregame P4–0072 covered the writing procedure of the manuscript.

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