

## Croatian national reference Y-STR haplotype database

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**Abstract** A reference Y-chromosome short tandem repeat (STR) haplotype database is needed for Y-STR match interpretation as well as for national and regional characterization of populations. The aim of this study was to create a comprehensive Y-STR haplotype database of the Croatian contemporary population and to analyze substructure between the five Croatian regions. We carried out a statistical analysis of the data from previously performed genetic analyses collected during routine forensic

work by the Forensic Science Centre “Ivan Vučetić”. A total of 1,100 unrelated men from eastern, western, northern, southern and central Croatia were selected for the purpose of this study. Y-STRs were typed using the AmpFISTR Yfiler PCR amplification kit. Analysis of molecular variance calculated with the Y chromosome haplotype reference database online analysis tool included 16 population samples with 20,247 haplotypes. A total of 947 haplotypes were recorded, 848 of which were unique (89.5%). Haplotype diversity was 0.998, with the most frequent haplotype found in 9 of 1,100 men (0.82%). Locus diversity varied from 0.266 for DYS392 to 0.868 for DYS385. Discrimination capacity was 86.1%. Our results

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suggested high level of similarity among regional sub-populations within Croatia, except for mildly different southern Croatia. Relative resemblance was found with Bosnia and Herzegovina and Serbia. Whit Athey's Haplogroup Predictor was used to estimate the frequencies of Y-chromosome haplogroups. I2a, R1a, E1b1b and R1b haplogroups were most frequent in all Croatian regions. These results are important in forensics and contribute to the population genetics and genetic background of the contemporary Croatian population.

**Keywords** Y-chromosome · Short tandem repeat · Population genetics · Haplotypes · Haplogroups · Croatian population

## Introduction

Croatia is located in the south-eastern Europe, sharing boundaries with Slovenia on the north-west, Hungary on the north, Serbia on the east, Montenegro on the south-east, Bosnia and Herzegovina on the south and east, and the Adriatic Sea on the west. Genetic structure of contemporary populations is interesting for a number of reasons, including genetic epidemiology, forensics, and enlightenment of the historical events. The use of Y-STRs has a central role in a number of such attempts, since it provides a good proxy of paternal lineage [1]. In forensics, reference Y-STR database within local population is needed for Y-STR match interpretation [2]. For this reason, Ministry of Interior of the Republic of Croatia created the reference Croatian Y-STR haplotype database. Variant alleles which contain incomplete repeat units are increasing the power of discrimination in DNA comparisons and can reveal phylogenetic substructure within the Y-chromosome phylogeny [3].

Creation of a national Y-STR haplotype database is one of the principal steps in a detailed genetic characterization. This is interesting not only for population genetics and forensics, but also for historical analyses, since genetic material can be used to infer information on demographic events.

Several previous studies were published in the Y-STRs in Croatia, mostly focusing on regional samples or rather small number of total samples involved in the study [4–9]. Recently, we announced population data of Y-STR haplotyping in eastern, northern and central Croatia [10–12] to emphasize the importance of Y-STR variations on regional level in Croatia.

The aim of this study was to create a large and comprehensive Y-STR haplotype database of the Croatian contemporary population and to analyze intrapopulation variability between the five Croatian regions. This intrapopulation variability may be mirrored in Y-chromosomal haplogroups distribution.

## Subjects and methods

### Study sample

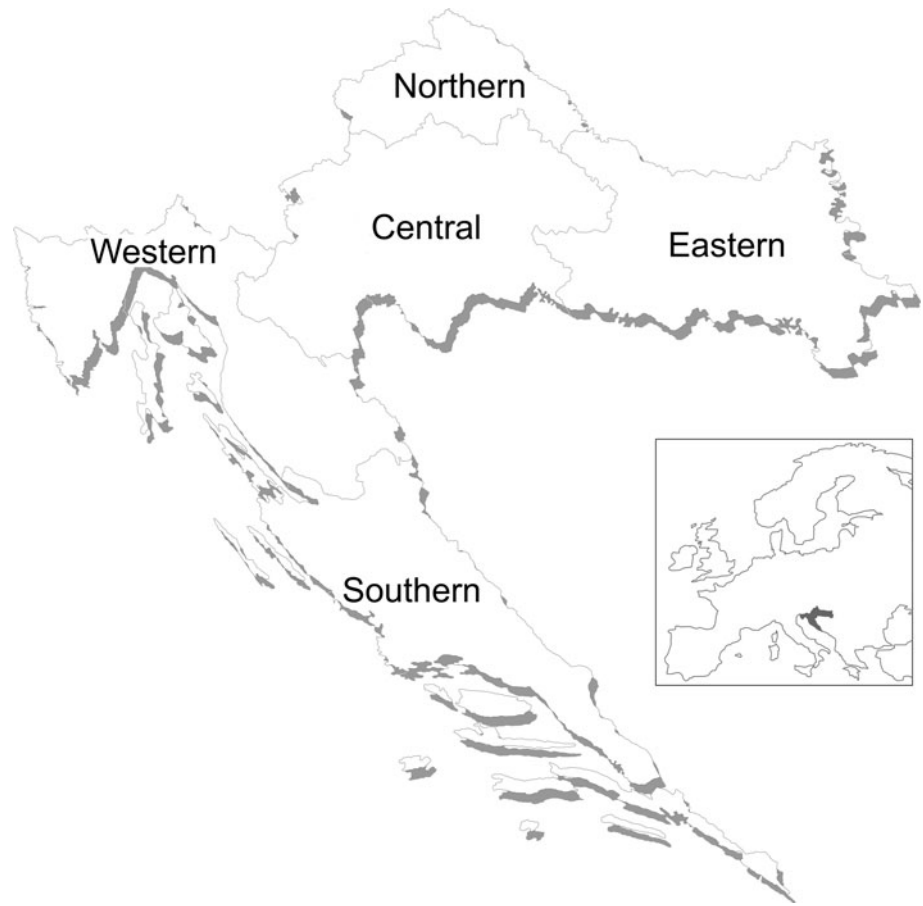
Statistical analysis of the data from previously performed genetic analyses collected during routine forensic work by the Forensic Science Centre “Ivan Vučetić” was performed in this study. Based on geographical and historical information we divided Croatia into five regions: eastern, western, southern, northern and central (Fig. 1). A total of 1,100 samples from eastern Croatia [ $n = 220$ ] (Virovitičko-podravka, Požeško-slavonska, Brodsko-posavska, Osječko-baranjska and Vukovarsko-srijemska counties), western Croatia [ $n = 220$ ] (Primorsko-goranska, Ličko-senjska and Istarska counties), northern Croatia [ $n = 220$ ] (Krapinsko-zagorska, Varaždinska, Koprivničko-križevačka and Međimurska counties), southern Croatia [ $n = 220$ ] (Zadarska, Šibensko-kninska, Splitsko-dalmatinska and Dubrovačko-neretvanska counties) and central Croatia [ $n = 220$ ] (Zagrebačka, Sisačko-moslavačka, Karlovačka, Bjelovarsko-bilogorska counties and city of Zagreb) were collected. The donors had no known familial relationships among one another and were of sufficient quality to be included in the statistical analysis. The study was approved by the Ethics Committee of the Institute for Medical Research and Occupational Health, Zagreb, Croatia.

### DNA analysis

Genomic DNA from all samples of the materials expertise was extracted from FTA cards (Whatman, Maidstone, Kent, UK) using Chelex [13]. The AmpFISTR Yfiler PCR amplification kit was used for the amplification of 17 Y-STRs: DYS19, DYS389I, DYS389II, DYS390, DYS391, DYS392, DYS393, DYS385, DYS437, DYS438, DYS439, DYS448, DYS456, DYS458, DYS635 and GATAH4 according to manufacturer's instruction (Applied Biosystems Foster City, California, USA). Y-STR amplification products were analyzed on 3130xl Genetic Analyzer (Applied Biosystems). Analysis of the data was performed using Genemapper<sup>®</sup> software (version 3.2, Applied Biosystems). Amplicon sizing was performed using an internal size standard (GeneScan-500 LIZ, Applied Biosystems), and the amplicons were compared with the AmpFISTR Yfiler allelic ladder for unambiguous allele designation.

### Quality control

Blind testing and evaluation was performed according to the quality assessment scheme provided by the Y Chromosome Haplotype Reference Database (YHRD, [www.yhrd.org](http://www.yhrd.org)) [14] and a certificate for successful testing was issued at November 23, 2009 for the Forensic Science Centre “Ivan

**Fig. 1** Map of Croatian regions

Vučetić". The contributing Centre (YC000198) received the following accession numbers after evaluation of the data: central (YA003593), eastern (YA003594), northern (YA003595), southern (YA003596) and western (YA003601) Croatia.

#### Statistical analysis

Allelic and haplotype frequencies were estimated by direct counting. Locus and haplotype diversities were calculated using Arlequin 3.1 software [15]. Furthermore, for analysis of molecular variance (AMOVA) Professor Lutz Roewer and Sascha Willuweit, MSc used the AMOVA tool provided by the YHRD [14]. Due to the limited number of markers reported in other publicly available samples, the comparative analysis was performed on a minimal European Y-STR haplotype comprising nine loci: DYS19, DYS389I, DYS389II, DYS390, DYS391, DYS392, DYS393 and DYS385ab [16]. A total of 16 population samples with 20,247 haplotypes were included in this analysis. The samples came from the following areas (the YHRD designation is given in parentheses): 150 haplotypes from Croatia including Zagreb, Croatia [Croatian,  $n = 150$ ], 1,100 haplotypes from Croatia including central Croatia [Croatian,

$n = 220$ ], east Croatia [Croatian,  $n = 220$ ], north Croatia [Croatian,  $n = 220$ ], south Croatia [Croatian,  $n = 220$ ], west Croatia [Croatian,  $n = 220$ ]; 141 haplotypes from Albania including Albania [Albanian,  $n = 111$ ], Tirana, Albania [Albanian,  $n = 30$ ]; 485 haplotypes from Austria including Graz [Austrian,  $n = 65$ ], Oberösterreich [Austrian,  $n = 178$ ], Vienna [Austrian,  $n = 66$ ], Salzburg [Austrian,  $n = 176$ ]; 100 haplotypes from Bosnia and Herzegovina including Doboј-Banja Luka-Bjeljina [Bosnian,  $n = 31$ ], Mostar, [Bosnian,  $n = 34$ ], Sarajevo [Bosnian,  $n = 35$ ]; 122 haplotypes from Bulgaria [Bulgarian]; 2,226 haplotypes from Czech Republic including Prague [Czech,  $n = 645$ ], Southern Bohemia [Czech,  $n = 111$ ], Karlovy Vary [Czech,  $n = 31$ ], Plzen [Czech,  $n = 62$ ], Liberec [Czech,  $n = 45$ ], Usti nad Labem [Czech,  $n = 87$ ], Central Bohemia [Czech,  $n = 455$ ], Hradec Kralove [Czech,  $n = 49$ ], Pardubice [Czech,  $n = 84$ ], Southern Moravia [Czech,  $n = 216$ ], Moravia-Silesia [Czech,  $n = 115$ ], Zlin [Czech,  $n = 64$ ], Olomouc [Czech,  $n = 53$ ], Vysocina [Czech,  $n = 40$ ], Czech Republic [Czech,  $n = 169$ ]; 6,119 haplotypes from Germany including Cologne [German,  $n = 738$ ], Berlin [German,  $n = 657$ ], Bonn [German,  $n = 90$ ], Chemnitz [German,  $n = 833$ ], Greifswald [German,  $n = 208$ ], Hamburg [German,  $n = 275$ ], Freiburg [German,  $n = 433$ ],

Halle [German,  $n = 234$ ], Dresden [German,  $n = 88$ ], Leipzig [German,  $n = 811$ ], Mainz [German,  $n = 104$ ], Magdeburg [German,  $n = 283$ ], Muenster [German,  $n = 196$ ], Munich [German,  $n = 281$ ], Rostock [German,  $n = 243$ ], Stuttgart [German,  $n = 613$ ], Palota, Bihor, Romania [German,  $n = 32$ ]; 515 haplotypes from Hungary including Budapest [Hungarian,  $n = 200$ ], Hungary [Hungarian,  $n = 215$ ], Szeged [Hungarian,  $n = 100$ ]; 302 haplotypes from Macedonia including Macedonia [Macedonian,  $n = 250$ ], Skopje [Macedonian,  $n = 52$ ]; 2,023 haplotypes from North Italy including Bologna [Italian,  $n = 51$ ], Brescia [Italian,  $n = 106$ ], Liguria [Italian,  $n = 81$ ], Modena [Italian,  $n = 130$ ], Lombardy [Italian,  $n = 182$ ], Marche [Italian,  $n = 162$ ], Umbria [Italian,  $n = 51$ ], Verona [Italian,  $n = 153$ ], Tuscany [Italian,  $n = 218$ ], Veneto [Italian,  $n = 120$ ], Ravenna [Italian,  $n = 384$ ], Val Marecchia [Italian,  $n = 65$ ], Latium [Italian,  $n = 222$ ], Rimini [Italian,  $n = 98$ ]; 4,131 haplotypes from Poland including Bydgoszcz [Polish,  $n = 411$ ], Bialystok [Polish,  $n = 185$ ], Krakow [Polish,  $n = 207$ ], Gdansk [Polish,  $n = 942$ ], Limanowa [Polish,  $n = 53$ ], Nowy Sacz [Polish,  $n = 114$ ], Lublin [Polish,  $n = 246$ ], South Eastern Poland [Polish,  $n = 161$ ], Szczecin [Polish,  $n = 105$ ], Suwalki [Polish,  $n = 82$ ], Wroclaw [Polish,  $n = 715$ ], Warsaw [Polish,  $n = 393$ ], Nowy Targ [Polish,  $n = 52$ ], Zakopane [Polish,  $n = 7$ ], Southern Poland [Polish, 380], Northern Poland [Polish,  $n = 78$ ]; 399 haplotypes from Romania including Constanta [Romanian,  $n = 36$ ], Romania [Romanian,  $n = 104$ ], Moldavia [Romanian,  $n = 40$ ], Transylvania, Romania [Romanian,  $n = 14$ ], Ploiesti [Romanian,  $n = 36$ ], Wallachia [Romanian,  $n = 96$ ], Oradea, Bihor [Romanian,  $n = 73$ ]; 215 haplotypes from Serbia including Novi Sad [Serbian,  $n = 215$ ]; 1,108 haplotypes from Slovakia including Bratislava [Slovakian,  $n = 164$ ], Eastern Slovakia [Slovakian,  $n = 629$ ], Slovakia [Slovakian,  $n = 80$ ], Sinteu, Bihor, Romania [Slovakian,  $n = 35$ ], Novi Sad, Serbia [Slovakian,  $n = 200$ ]; 931 haplotypes from Italy Alcamo [Italian,  $n = 23$ ]; Caccamo [Italian,  $n = 19$ ], Puglia [Italian,  $n = 70$ ], Pantelleria, Italy [Italian,  $n = 21$ ], Northern Sardinia [Italian,  $n = 100$ ], Ragusa [Italian,  $n = 29$ ], Mazara del Vallo [Italian,  $n = 25$ ], Santa Ninfa [Italian,  $n = 34$ ], Troina [Italian,  $n = 21$ ], Trapani [Italian,  $n = 32$ ], Sciacca [Italian,  $n = 20$ ], Sicily [Italian,  $n = 314$ ], Piazza Armerina [Italian,  $n = 30$ ], Reggio di Calabria, Italy [Italian,  $n = 97$ ], Cosenza, Italy [Italian,  $n = 37$ ], Catanzaro [Italian,  $n = 59$ ], and 180 haplotypes from Slovenia including Ljubljana [Slovenian,  $n = 180$ ] [17–57]. In this study we calculated population pair-wise genetic distances ( $R_{ST}$ ). This is an extension of the commonly used  $F_{ST}$  measure [58], defined as  $R_{ST} = (S_b - S_w) / S_b$ , where  $S_w$  is the sum over all loci of twice the weighted mean of the within-population variances  $V(A)$  and  $V(B)$ , and  $S_b$  is the sum over all loci of twice the variance  $V(A + B)$  of

the combined population [58].  $R_{ST}$  values were calculated by AMOVA using on-line tools of the YHRD,  $P$  values were obtained with 10,000 permutations (significance set at  $P < 0.05$ ).

We had estimated the Y-chromosomal haplogroups from a set of acquired Y-STR data. Regarding the fact that Y-chromosomal single-nucleotide polymorphism (Y-SNP) data had not been available for this study all predictions of haplogroups were performed by use of the web-accessible program, Whit Athey's Haplogroup Predictor ([www.hprg.com/hapest5/](http://www.hprg.com/hapest5/)), which is based on a Bayesian allele frequency approach. We considered all limitations and biases [59–61]. Therefore, we included in our study only the samples with haplogroup probability over 90%. This is very often the only approach in forensic practice to found out the supplementary information for particular unknown haplotype. The haplogroup estimation was performed for every haplotype and the haplogroup frequencies were estimated by direct counting.

Median-Joining Network (MJN) was constructed with Network 4.60 software ([www.fluxus-engineering.com](http://www.fluxus-engineering.com)) [62, 63]. We utilized all 17 loci (DYS19, DYS389I, DYS389II, DYS390, DYS391, DYS392, DYS393, DYS385, DYS437, DYS438, DYS439, DYS448, DYS456, DYS458, DYS635 and GATAH4) for the MJN analysis. For the complete set of 1,100 Croatian samples, firstly, we reduced the amount of haplotypes in the computation by star contraction pre-processing algorithm ( $\Delta = 5$ ). Subsequently, the reduced median algorithm ( $r = 2$ , frequency >1 criterion active) and median-joining algorithm ( $\epsilon = 0$ ) were applied. Before drawing the network, maximum parsimony post-processing procedure was performed. MJN analysis for the single haplogroups included only the reduced median ( $r = 2$ ) algorithm.

Software Splitstree 4.10 ([www.splitstree.org](http://www.splitstree.org)) was used for constructing the Neighbor-Joining (NJ) phylogenetic tree [64]. As input data we used the matrix of  $R_{ST}$  pair-wise distances of the Croatian population focused on in this study (central, western, northern, southern and eastern) and neighboring populations as described previously, including population from Zagreb, Croatia.

## Results

A total of 1,100 samples from five regions of Croatia were included in the analysis. Locus diversity varied from 0.266 for DYS392 to 0.868 for DYS385. Variant alleles have been found at DYS458 locus. Allele frequencies and locus diversity values are shown in Table 1. A total of 947 haplotypes were recorded, 848 of which were unique (89.5%). Total haplotype diversity was 0.998, with the most common recorded haplotype found in 9 of 1,100 men (0.82%). Discrimination capacity was 86.1%.

**Table 1** Allele frequencies and locus diversity values at 17 Y-STR loci in Croatian population

Allele	STR locus on Y chromosome																Genotype	DYS385* (%)	
	DYS456	DYS389I	DYS390	DYS389II	DYS458	DYS19	DYS393	DYS391	DYS439	DYS635	DYS392	YGATAH4	DYS437	DYS438	DYS448				
8							0.09	0.36									10-10	0.18	
9							1.55	0.45										10-13	0.09
10							49.73	13.00								9.00		10-14	2.00
11		0.27					47.27	26.27								59.91		10-15	0.09
12	0.73	14.09			0.09	0.45	1.36	32.09					0.09			7.27		10-17	0.09
13	1.73	70.09			0.73	79.73		24.73					0.18			0.18		10-17.1	0.09
14	10.45	15.27			1.82	19.45	7.27	2.73					42.82					10-20	0.09
15	53.55	0.27			23.18	24.00	0.27	0.27					46.91					11-11	0.91
16	20.55				24.91	37.45		0.09					9.27					11-12	0.27
17	11.00				30.27	9.45							0.73			0.09		11-13	3.36
18	2.00				14.09	0.27									1.36			11-14	17.09
19					3.00										41.09			11-15	3.63
20					0.45						0.09				48.82			11-16	0.73
21			0.45								4.18				7.36			11-18	0.09
22			7.64								15.27				1.09			12-12	0.18
23			13.73								22.00				0.18			12-14	1.73
24			55.82								48.36							12-15	0.82
25			20.73								8.73							12-16	0.27
26			1.45								1.36							12-17	0.09
27			0.18															12-18	0.45
28						0.82												13-13	0.27
29						9.09												13-14	4.18
30						17.45												13-15	2.27
31						31.00												13-16	1.00
32						34.00												13-17	0.64
33						7.18												13-17.2	0.27
<b>12.2</b>						0.45												13-18	0.18
<b>15.1</b>																		13-19	0.36
<b>16.2</b>																		13-20	0.27
<b>17.2</b>																		14-14	3.36
<b>18.2</b>																		14-15	30.00
<b>19.2</b>																		14-16	4.55
<b>15,16</b>																		14-17	1.55
																		14-18	0.82
																		14-19	0.27
																		15-15	3.09
																		15-16	0.36
																		15-17	2.00
																		15-18	0.82
																		15-19	0.91
																		15-20	0.36



**Table 2** Analysis of molecular variance pair-wise distances based on  $R_{ST}$  values between the Croatian subpopulations in this study and selected comparison populations

Population samples <sup>a</sup> Population samples <sup>b</sup>	Central Croatia, Croatia (Croatian)	East Croatia, Croatia (Croatian)	North Croatia Croatia (Croatian)	South Croatia Croatia (Croatian)	West Croatia, Croatia (Croatian)	Zagreb, Croatia (Croatian)	Albanian	Austrian	Bosnian	Bulgarian	Czech
Central Croatia, Croatia [Croatian]	–	0.8105	0.6582	0.0322	0.6352	0.9354	0.0001	0.0000	0.0446	0.0001	0.0000
East Croatia, Croatia [Croatian]	–0.0027	–	0.2100	0.1037	0.9564	0.7050	0.0001	0.0000	0.1719	0.0010	0.0000
North Croatia, Croatia [Croatian]	–0.0021	0.0018	–	0.0044	0.1296	0.5236	0.0000	0.0000	0.0098	0.0002	0.0000
South Croatia, Croatia [Croatian]	0.0084	0.0042	0.0171	–	0.0699	0.0522	0.0000	0.0000	0.0195	0.0000	0.0000
West Croatia, Croatia [Croatian]	–0.0019	–0.0036	0.0035	0.0055	–	0.6297	0.0000	0.0000	0.1218	0.0005	0.0000
Zagreb, Croatia [Croatian]	–0.0043	–0.0028	–0.0015	0.0085	–0.0023	–	0.0007	0.0000	0.0621	0.0008	0.0000
Albanian	0.0539	0.0473	0.0598	0.0904	0.0510	0.0490	–	0.0000	0.0148	0.1331	0.0000
Austrian	0.0761	0.0902	0.0682	0.1354	0.0942	0.0746	0.1088	–	0.0000	0.0000	0.0000
Bosnian	0.0125	0.0040	0.0226	0.0162	0.0061	0.0118	0.0295	0.1362	–	0.0092	0.0000
Bulgarian	0.0394	0.0312	0.0475	0.0717	0.0325	0.0383	0.0063	0.0977	0.0265	–	0.0000
Czech	0.0860	0.1066	0.0697	0.1335	0.1113	0.0877	0.1818	0.0372	0.1591	0.1678	–
German	0.1070	0.1267	0.0926	0.1623	0.1296	0.1054	0.1679	0.0105	0.1773	0.1544	0.0185
Hungarian	0.0189	0.0301	0.0120	0.0572	0.0322	0.0203	0.0796	0.0235	0.0640	0.0644	0.0289
Macedonian	0.0357	0.0257	0.0454	0.0508	0.0262	0.0341	0.0084	0.1217	0.0101	–0.0011	0.1851
North Italian	0.0938	0.1050	0.0844	0.1500	0.1090	0.0924	0.0908	0.0135	0.1364	0.0875	0.0651
Polish	0.1409	0.1652	0.1215	0.1780	0.1709	0.1454	0.2825	0.1221	0.2244	0.2655	0.0270
Romanian	0.0150	0.0113	0.0195	0.0357	0.0106	0.0154	0.0229	0.0794	0.0138	0.0019	0.1286
Serbian	0.0125	0.0061	0.0186	0.0281	0.0072	0.0125	0.0177	0.1005	0.0011	0.0046	0.1425
Slovakian	0.0461	0.0626	0.0344	0.0791	0.0666	0.0472	0.1510	0.0527	0.1074	0.1363	0.0113
Slovenian	0.0690	0.0879	0.0550	0.1193	0.0966	0.0695	0.1618	0.0533	0.1516	0.1711	0.0067
South Italian	0.0584	0.0594	0.0569	0.1002	0.0587	0.0575	0.0295	0.0460	0.0701	0.0169	0.1218
Population samples <sup>a</sup> Population samples <sup>b</sup>	German	Hungarian	Macedonian	North Italian	Polish	Romanian	Serbian	Slovakian	Slovenian	South Italian	
Central Croatia, Croatia [Croatian]	0.0000	0.0003	0.0000	0.0000	0.0000	0.0026	0.0154	0.0000	0.0000	0.0000	0.0000
East Croatia, Croatia [Croatian]	0.0000	0.0000	0.0001	0.0000	0.0000	0.0068	0.0616	0.0000	0.0000	0.0000	0.0000
North Croatia, Croatia [Croatian]	0.0000	0.0043	0.0000	0.0000	0.0000	0.0013	0.0037	0.0000	0.0000	0.0000	0.0000
South Croatia, Croatia [Croatian]	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0001	0.0000	0.0000	0.0000	0.0000
West Croatia, Croatia [Croatian]	0.0000	0.0000	0.0004	0.0000	0.0000	0.0072	0.0502	0.0000	0.0000	0.0000	0.0000
Zagreb, Croatia [Croatian]	0.0000	0.0008	0.0000	0.0000	0.0000	0.0060	0.0235	0.0000	0.0000	0.0000	0.0000
Albanian	0.0000	0.0000	0.0591	0.0000	0.0000	0.0022	0.0131	0.0000	0.0000	0.0004	0.0000
Austrian	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000
Bosnian	0.0000	0.0000	0.0602	0.0000	0.0000	0.0223	0.2920	0.0000	0.0000	0.0000	0.0000
Bulgarian	0.0000	0.0000	0.4661	0.0000	0.0000	0.2156	0.1396	0.0000	0.0000	0.0000	0.0043



Table 2 continued

Population samples <sup>a</sup> Population samples <sup>b</sup>	German	Hungarian	Macedonian	North Italian	Polish	Romanian	Serbian	Slovakian	Slovenian	South Italian
Czech	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0171	0.0000
German	-	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000
Hungarian	0.0384	-	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0001	0.0000
Macedonian	0.1836	0.0765	-	0.0000	0.0000	0.0201	0.1624	0.0000	0.0000	0.0000
North Italian	0.0275	0.0433	0.1172	-	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000
Polish	0.0828	0.0863	0.2741	0.1619	-	0.0000	0.0000	0.0000	0.0000	0.0000
Romanian	0.1291	0.0378	0.0074	0.0824	0.2119	-	0.3228	0.0000	0.0000	0.0000
Serbian	0.1494	0.0476	0.0024	0.1009	0.2209	0.0003	-	0.0000	0.0000	0.0000
Slovakian	0.0457	0.0174	0.1437	0.0859	0.0278	0.0949	0.1019	-	0.2328	0.0000
Slovenian	0.0368	0.0265	0.1649	0.0873	0.0213	0.1180	0.1296	0.0010	-	0.0000
South Italian	0.0956	0.0469	0.0391	0.0360	0.2264	0.0242	0.0379	0.1134	0.1216	-

<sup>a</sup> *P* values are shown above the diagonal and *R*<sub>ST</sub> values below

<sup>b</sup> Names in parentheses refer to the YHRD designations ([www.yhrd.org](http://www.yhrd.org))

graphs) to found out intra haplogroup diversity. Some very rare European haplogroups (Q,T) were detected in the sample that are very useful for forensic application.

To examine the substructure of Croatian population and further investigate its connections with European populations, we constructed the phylogenetic tree based on *R*<sub>ST</sub> pair-wise genetic distances with NJM. The topology of NJ tree was in concordance with previous genetic studies and known historic layout of European populations.

Length of the branches correlated with genetic distances between particular populations (Fig. 4).

The mutual relationships between populations were displayed without the dependence on the genetic distances in the cladogram (Fig. 5).

The topology of NJ tree also confirmed the substructure in Y-chromosomal variation in Croatian population. The major dichotomy in the tree divided the Croatian subpopulations into southern and central European clusters. The second cluster was further divided into Germanic populations (German, Austrian, and North Italian), Slavic populations (Czech, Polish, Slovenian and Slovak) and Hungarian. Interestingly, subpopulation from northern Croatia was, regarding to the *R*<sub>ST</sub> genetic distance, closer to this cluster than to the rest of Croatia.

Southern European cluster consisted mostly of the populations with putative Illyrian (Albanians, Serbians, Bosnians and disputably Macedonians) and Roman (Latin) roots (Southern Italians and Romanians, although here is the influence of Dacians probably more prevalent) [65, 66]. Subpopulations from eastern, western and southern Croatia were close to this cluster, but at the same time exhibited a degree of isolation from it.

Remarkable is the position of central Croatian subpopulation as a part of the deepest dichotomy of the NJ tree, and although it lay closer to the northern populations cluster, it was isolated both from northern as well as from eastern, southern and western Croatian subpopulations. It is positioned exactly on the boundary between the northern Croatian subpopulation with its affinity to Slavic/Germanic cluster and the rest of the Croatian subpopulations with their affinity to Balkan/southern European cluster.

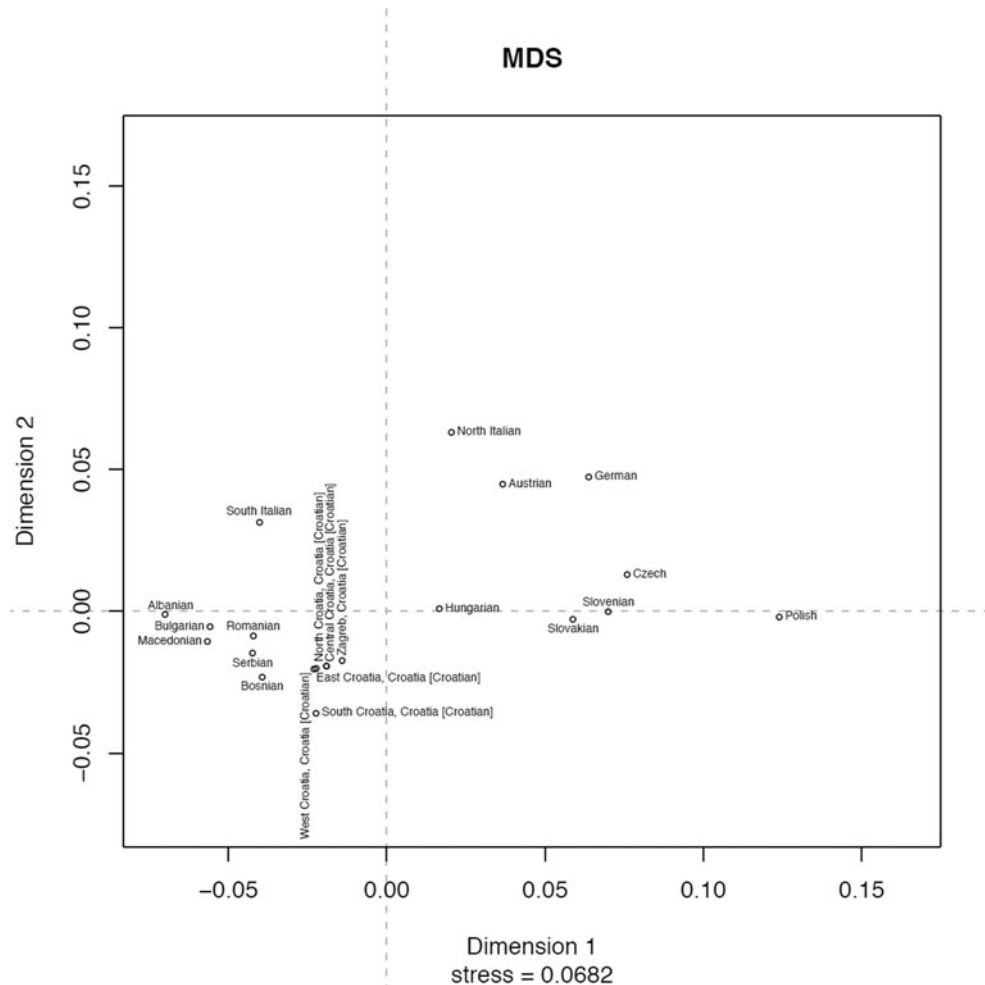
## Discussion

The results of this study are based on the first attempt to create a national reference Y-STR haplotype database on a sufficient number of samples for both national and regional characterization.

We do acknowledge the previous studies in this light and should also state that the results of this study show a strong similarity with the existing data, suggesting that the results are likely valid. This paper follows the



**Fig. 2** Multidimensional scaling (MDS) based on pair-wise  $R_{ST}$  genetic distances between Croatia and neighboring countries



recommendations of the ISFG on the use of Y-STRs in forensic analysis [67]. Y-STR population data have been submitted to the YHRD as stated in “[Subjects and methods](#)” section.

In population genetics the sample size does not confirm the quality of population sampling [14, 17, 25, 68]. Therefore, the important aspect of population sampling is the good representation of different population specific haplotypes. On the basis of these facts we created our database. The high value of haplotype diversity (0.998) confirms the sampling quality. Moreover, large data sets typically identify a larger number of rare alleles as more individuals in a population are included in the analysis, as was shown in our case for DYS458 locus. At this locus we reported for the first time variant alleles 12.2, 16.2, 17.2 and 19.2 in eastern [10], 17.2, 18.2 and 19.2 in central [12] and 15.1 in southern Croatia.

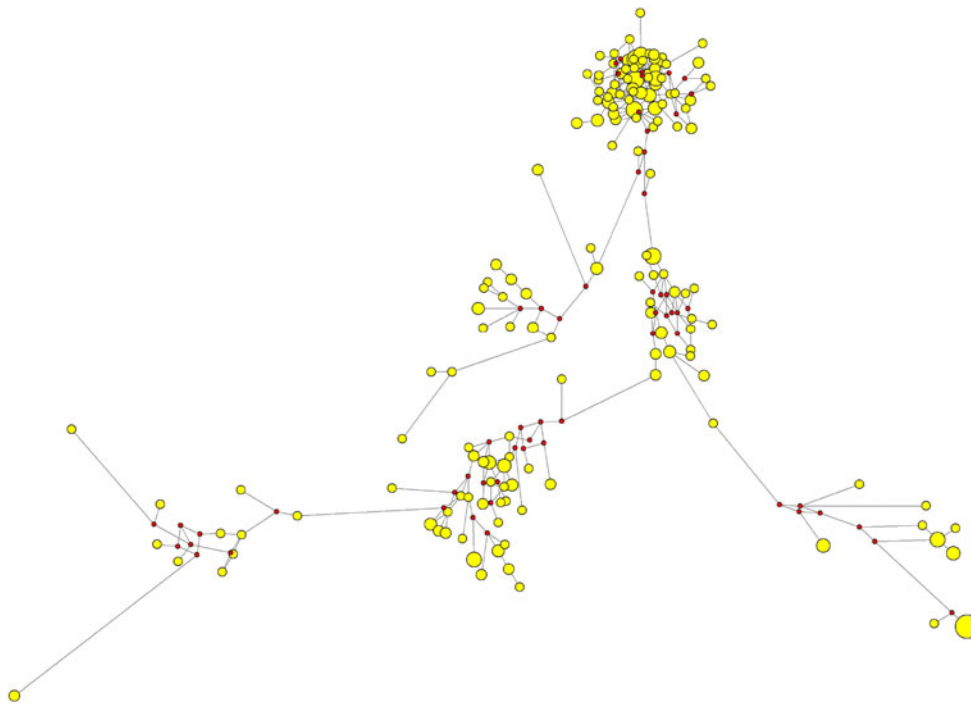
Barać et al. [6] published population genetic study, where eight Y-STRs were analyzed in the sample of 457 participants; 109 from mainland Croatia and the rest of them from four Adriatic islands; Krk, Brač, Hvar and

Korčula. Our results are in concordance to those published by Barać et al. concerning the allelic frequency distribution in Croatia, when taking into account the identical ones, except DYS391. Moreover, they also concluded that majority of their loci showed the similar allelic frequency distribution to those published for other European populations [16]. The exception was found at DYS19 and DYS389II loci. Allele 16 at DYS19 locus was the most frequent in Barać et al. as well as in our study, opposite to allele 14 in Europe [16]. In addition, both Croatian studies showed alleles 30 and 31 as the most common at DYS389II locus, opposite to allele 29 at the same locus in Europe.

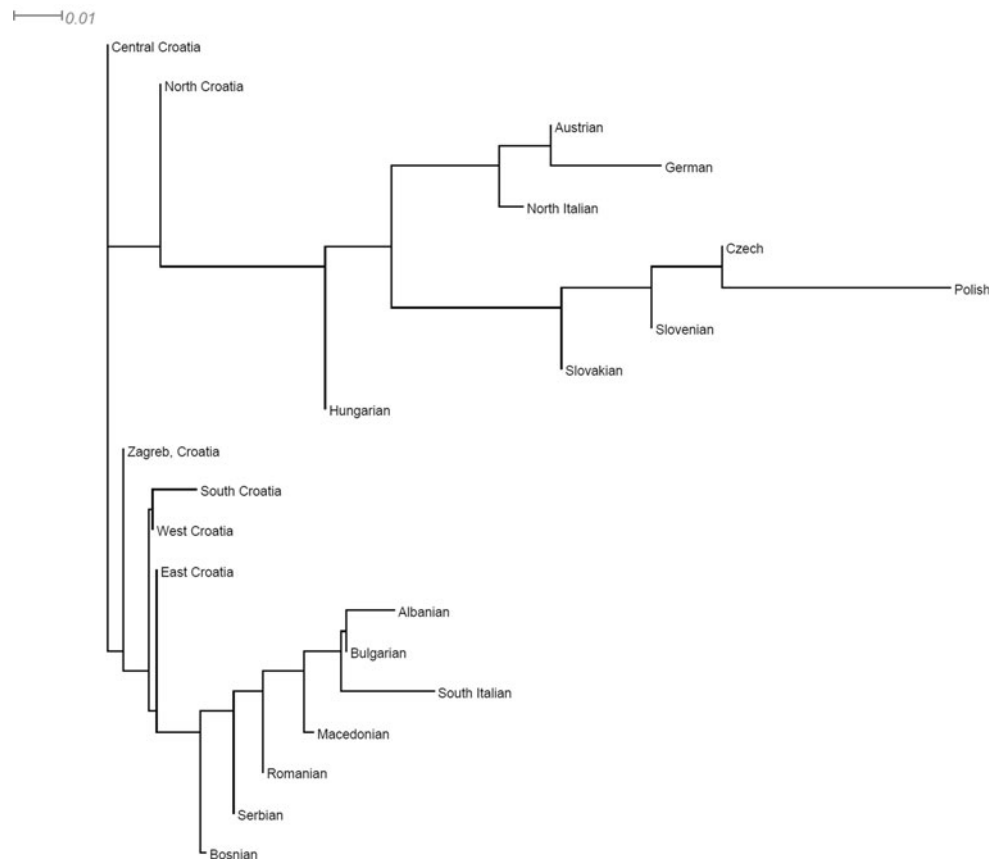
The greatest publicly available YHRD is very important in forensic community, because it provides information on Y-chromosome haplotypes from world-wide distributed populations [14]. Enlargement of Y-STR databases sample size allows detection of variant alleles that are informative for Y-chromosome diversity [69]. European minimal haplotype set consisting of nine loci with high levels of variability in worldwide populations was used for analysis of molecular variance pair-wise distances based on  $R_{ST}$  values

**Table 3** Haplogroup composition in single Croatian regional dataset

Haplogroup	Central Croatia Number of haplotypes	North Croatia Number of haplotypes	East Croatia Number of haplotypes	South Croatia Number of haplotypes	West Croatia Number of haplotypes	Pooled frequency (%)
I2a(×I2a1)	70	56	88	120	81	37.7
R1a	52	64	41	42	44	22.1
E1b1b	26	24	25	14	28	10.6
R1b	23	23	18	10	13	7.9
I1	11	9	13	12	19	5.8
J2b	11	11	6	6	7	3.7
G2a	8	7	4	4	7	2.7
H	3	11	2	1	4	1.8
J2a1h	1	1	5	2	4	1.2
J1	5		6		1	1.1
J2a1b	2	1	3	1	4	1
E1b1a		1				<1
G2c					1	<1
I2a1	1					<1
I2b1	2	5	1	1	1	<1
I2b(×I2b1)	2	2				<1
J2a1-bh			1	2	1	<1
L	1	1		1		<1
N	2	1	2		2	<1
Q			4	3	1	<1
T		3	1	1	2	<1

**Fig. 3** Median-Joining network of the pooled Y-chromosomal haplogroups in Croatian population ( $n = 1,100$ )

**Fig. 4** Phylogram of Neighbor-Joining tree of the examined populations

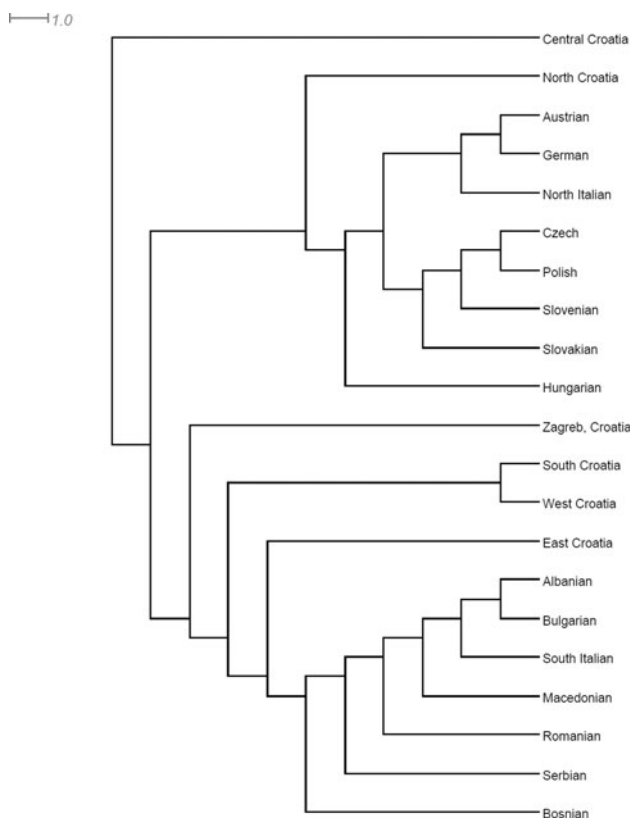


between the Croatian subpopulations and neighboring countries. We compared a total of 16 population samples with 20,247 haplotypes. Only small genetic distances were documented between the Croatian regions, except for southern Croatia, indicating that Croatian population is quite homogeneous in terms of paternal genetic contribution. The relative genetic similarity was evident between the Croatian subpopulations when compared to Bosnian and Serbian populations. This similarity was less pronounced with Hungarian population. In this comprehensive study, we also showed that Croatian subpopulations are genetically distant from Czech, Polish and Slovakian populations. Strong genetic difference was observed from Slovenia concerning the paternal lineage. This result is very interesting in several historical dimensions. Croatian history is vivid and marked with strong dynamics. Initial historical records indicate the presence of old Illyrian tribes in the region, who were slowly or abruptly being assimilated into numerous newly arrived groups which include at least some Greek colonies in the coastal parts, Romans across entire territory, Slavic tribes in VIth century and later on, followed by, intrusions of Franks, and other Western-European groups, the long-lasting occupations by Ottoman Empire, agglomeration within Austrian-Hungarian empire and lastly substantial population changes

imposed by the World War I and II, and most recently changes during the Croatian Homeland War in 1991–1995 [70].

Considering the fact that the extent of Y-STR variability may be influenced by the haplogroup composition, we firstly analyzed raw haplogroup structure in the large Croatian sample set diversified regionally. As it was mentioned earlier, by use the web-accessible program, Whit Athey's Haplogroup Predictor we estimated the frequencies of Y-chromosome haplogroups from a set of acquired Y-STR data. The results of our study identified four main haplogroups in all five Croatian regions: I2a (37.7%), R1a (22.1%), E1b1b (10.6%) and R1b (7.9%). Each of the detected haplogroup had different level of haplotype diversification except R1a and E1b1b: I2a (0.31), R1a (0.382), E1b1b(0.383) and R1b (0.402). The lowest haplotypic diversity was measured in haplogroup I2a, which seemed to be the most homogenous haplogroup in all Croatian regions. On the other hand, differences in haplogroup frequencies between northern and southern samples were statistically significant what could be important in terms of forensic applications.

MJN graphs for the eight most frequent haplogroups were performed. This form of visual displays enabled to study remote relationships between the single haplotypes



**Fig. 5** Cladogram of Neighbor-Joining tree of the examined populations

inside particular haplogroup and to explore mutation events that might cause the present state of this affinity. In analyzed population sample 10 rare subhaplogroups were found (E1b1a, G2c, I2a1, I2b1, I2b(xI2b1), J2a1-bh, L, N, Q and T). The haplogroup rarity in the population sample had some benefits for the forensic applications, because in rare haplogroups the probability of finding haplotypes identical by descent is higher than finding haplotypes identical by state. This fact can be applicable in methods of familial searching where paternal lineages of special interest could be determined.

The high frequency of haplogroup I2a was observed in Bosnia and Herzegovina (70%), and decreasing frequencies were spreading from the southern Dinaric Alps to northern Croatia [71]. The same authors reported lower I2a1 haplogroup frequency in Croatia (32%).

We would like to emphasize the previously published studies based on haplogroup analysis using Y-SNPs in Croatian population [8, 9]. Barać et al. showed that the most prevalent haplogroups in Croatia are I (49%), R1a (27%) and R1b (7%), while haplogroups E, G, J, F, K and P are represented with 20% [9]. In addition, Peričić et al. analyzed 451 individuals, 348 from four Adriatic islands (Krk, Hvar, Brač, Korčula) and 109 from Croatia mainland (Osijek, Zabok, Zagreb, Delnice, Pazin and Dubrovnik)

[9]. They showed that the most frequent Y-chromosomal haplogroups in the overall Croatian population are I1b-P37 (41.7%) and R1a-SRY1532 (25%) indicating the Slavic gene pool component. Moreover, Peričić et al. stress out that R1b lineage is present at relatively low frequency (8.5%) in southeastern Europe and Croatia (7.4%). Moreover, Semino et al. [72] data indicated that Croatian men can be subdivided into three major haplogroups, I (45%), R1a1 (30%) and R1b (10%). However, haplogroups E3b, J2a, G, LT are included all together with 15%. Although our haplogroup analysis was based on prediction; the frequencies of I2a (37.7%), R1a (22.1%) and R1b (7.9%) haplogroups were in concordance to those previously published for Croatian population [8, 9, 71, 72].

The main goals of our project were to create a national reference Y-STR haplotype database, as well as to investigate the Croatian intrapopulation Y-STR variability as well as genetic contribution from neighboring countries. These results may serve several important tasks, including that of forensics [73], but also contribute to our knowledge and understanding of the genetic background of contemporary Croatian populations, in terms of paternal lineages.

Future lofty goal will include analysis of informative biallelic markers in non-recombining region of Y-chromosome on this representative sample to further explore Croatian genetic heritage and to write additional chapter in the book of past European migration processes [71, 72, 74–77].

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