Autosomal Short Tandem Repeat (STR) Variation Based on 15 Loci in a Population from the Central Region (Riyadh Province) of Saudi Arabia

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Abstract

Introduction: The small size of Short Tandem Repeats (STRs), their ubiquitous genome-wide distribution and polymorphic nature enhances their value in human forensic/population genetics applications.

Objectives: This study aims to investigate the short tandem repeat variation based on 15 loci in a population from the central region of Saudi Arabia.

Methods: Allele frequency variation for 15 Short Tandem Repeat (STR) loci was examined in 190 unrelated Saudi volunteers.

Results: This study summarizes the allele distribution in the Saudi population and compares them to other populations located in Asia, Africa, the Middle East and Europe. The standard forensic parameters of Observed Heterozygosity (Ho), Expected Heterozygosity (He) and Gene Diversity Index (GD) were determined for the following 15 STR loci: D8S1179, D21S1, D7S820, CSF1PO, D3S1358, TH0, D13S317, D16S539, D2S1338, D19S433, vWA, TPOX, D18S51, D5S818 and FGA. The most frequent alleles in the Saudi population were: 8 repeats (0.558) at TPOX, 12 (0.411) at D13S317, 12 (0.385) at CSF1PO, 11 (0.382) at D16S539 and 10 (0.356) at D7S820. The 15 markers utilized in this study are highly informative as evidenced by their high power of discrimination (PD) values with D2S1338, D19S433 and FGA having the highest PD values. The relationship between the Saudi population and other geographically distributed populations, assessed by a Multidimensional Scaling (MDS) plot, showed that the Saudi population clustered with groups from Yemen, Iraq, Qatar, Oman and Bahrain.

Conclusion: TPOX, D13S317, CSF1PO, D16D539 and D7S820 markers were found suitable for forensic analysis, paternity testing and can also be used for chimerism study after allogenic bone marrow transplantation for Saudi population. On the other hand, the population admixture with other ethnic origins might explain the variable degree of genetic distances of this population and other Arab-related groups.

Keywords: Short tandem repeat; Saudi Arabia; Allele frequencies

Abbreviations: STR(s): Short Tandem Repeat(s); MDS: Multidimensional Scaling; HWE: Hardy–Weinberg Equilibrium

Background

Short Tandem Repeats (STRs) are nucleotide sequences with repeat motifs of variable lengths (2 to 8bp) that are polymorphic (i.e. number of repeats varies between individuals) [1,2]. STRs represent about 3% of human DNA and occur approximately one time in every 10,000 nucleotides. STRs are an invaluable tool and their unique sequences used in genetic finger printing during forensic investigations involve significant biological evidence. Their high degree of polymorphisms makes them informative [3], especially when considering multiple loci simultaneously. The new STR kits now contain over 20 STR loci that are amplified in a single multiplex reaction. Additionally, STR markers have been widely used in medical applications such as for the assessment of allogenic bone marrow transplantation engraftment in calculating the ratio of donor/patient DNA presence and in the study of population genetics. The use of STRs depends on the allele frequencies distributions that vary between various populations [1-4].

Various DNA-based techniques have been used to identify the genetic differences in human populations. STR loci are useful and preferred because of their small size, relatively low incidence of mutation and wide spread distribution [5]. Genetic studies and in particular those based on STR applications have been very important towards developing an appreciation of the extent of genetic polymorphisms that exists between different populations.

Throughout history, society had been stratified on the basis of caste, class, clan, race, region, religion, ethnicity, gender, age and socioeconomic status. It is ethnicity and racial discrimination that distinguishes one nation from the other. Ethnicity is, as defined by Macionis, “a shared cultural heritage and people define themselves or others as members of an ethnic category based on common ancestry, language or religion that gives them a distinctive social identity” [6].

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same is the case within the Arab world, which has maintained a unique ethnographic identity, historical background, ancestry, cultural traits, social norms, moral values, religious beliefs and genealogy. At present, the total Arab population is estimated at about 325 million, increasing at a rate of approximately 2.3% annually. The Arab population of the Middle East and North Africa is distributed throughout 17 different countries, namely: Algeria, Bahrain, Egypt, Iraq, Jordan, Kuwait, Lebanon, Libya, Morocco, Oman, Palestine, Qatar, Saudi Arabia, Syria, Tunisia, the United Arab Emirates, and Yemen.

In Saudi Arabia, approximately 87% of the population is of Arab descent, a group of people who have inhabited this region for thousands of years. The remaining 13% of inhabitants of this region migrated into this country during the last, 400 years [7]. The rate of consanguineous marriage is relatively high and has been estimated to at around 57.7% [8]. The Saudi population can be characterized by a specific ethnic subset which presumably reflects a unique set of allele frequencies and genotypes and that could differentiate them from other populations in near geographical areas and worldwide i.e. the alleles frequencies will be different from other populations and the genetic distances between these populations will be variable depending on the time of divergence from the common human ancestor.

Therefore, this study was conducted to investigate the genetic variation of populations from Saudi Arabia by using 15 autosomal STR markers - currently used in forensic and paternity testing. DNA was collected from a sample of 190 unrelated volunteers residing in the central region (Riyadh Province) of Saudi Arabia. The allele frequencies of the 15 STR markers were calculated using gene counting methods and compared with frequencies from other populations previously studied.

Methods

Population

A total of 190 healthy unrelated individuals from the central region (Riyadh Province) of Saudi Arabia were randomly recruited at King Fahad Medical City (KFMC) (Riyadh, Saudi Arabia) from the donors of patients under preparation for Bone Marrow Transplantation. According to KFMC hospital record system, the selection of the study subjects precluded having any two individuals or their father/mother descending from the same parent. Thus, first and second cousins are not included in the study. The age of the study population ranged between 10 and 45 years (median of 25) at the time of blood collection. There were 131 (68.9 %) males and all were of Saudi origin. The sample size was selected to provide sufficient analytical power in terms of degrees of freedom to determine allele frequencies that will allow testing of the Hardy-Weinberg Equilibrium (HWE) assumption based on a Fisher’s Exact Test. The study received ethical approval from the Institutional Review Board at KFMC. DNA extraction, PCR and Fragment analysis for STR markers

Genomic DNA was extracted from whole blood samples in EDTA anticoagulant using a MagNa pure compact instrument (Roche Diagnostics GmbH, Mannheim, Germany). PCR amplification was performed according to manufacturer instruction on GeneAmp® PCR Diagnostics GmbH, Mannheim, Germany). PCR amplification was performed according to manufacturer instruction on GeneAmp® PCR System 9700 Thermal Cycler (Applied Biosystems, Foster City CA, USA) using AmpFISTR™ Identifier PCR amplification kit (Applied Biosystems, Foster City CA, USA) that include D8S1179, D21S11, D7S820, CSF1PO, D3S1358, THO, D13S317, D16S539, D21S338, D19S433, vWA, TPOX, D18S55, D5S818, FGA loci. The PCR product was separated on the 3130xL Genetic Analyzer (Applied Biosystems, Foster City, CA, USA) and analyzed using GeneMapper® Software v3.2 (Applied Biosystems, Foster City CA, USA). For quality assurance purposes, multiple DNA samples with known combined STR genotypes were run in parallel with our study samples and appropriate results were obtained for these control samples on every occasion.

Statistical analysis

The allele frequencies were calculated based on the number of the detected alleles for each specific locus. To assess departures from Hardy-Weinberg Equilibrium (HWE) (observed heterozygosity (Ho), expected heterozygosity (He), p-value; using Fisher’s Exact test) and Gene Diversity indices (GD), the Arlequin software version 3.5.1.3 was used [9]. Forensic and population genetic parameters including power of discrimination (PD), Polymorphic Information Content (PIC), Matching probability (MP) and probability of exclusion (PE) were derived utilizing Powerstats software version 1.2 [10]. An overall significance level for testing HWE hypothesis was set at 0.05 and adjusted for multiple comparisons using Bonferonni Criteria. Genetic relationships between groups were assessed through multidimensional scaling (MDS) plot of the genetic distance matrix using NTSYS 2.1. Software to determine the genetic differences and similarities between this study sample and published data from other populations [11].

Results and Discussion

The 15 autosomal STR loci from all samples tested were amplified successfully. A total of 150 alleles were identified for the 15 STR loci. Their corresponding frequencies were calculated (Table 1). The highest allele frequencies observed were: allele 10 of TPOX (55.8%) and allele 12 of D13S317 (41.1%) with the most polymorphic loci observed in this study being D18S55, FGA, D21S11--defined by 17, 15 and 15 alleles respectively. However, the Polymorphic Information Content (PIC) of the 15 loci were greater than 0.57 through all loci that suggesting that the markers are highly polymorphic and would be useful as informative markers for differentiating individuals of Saudi descent. The degree of polymorphism at each locus can also be expressed in terms of heterozygosity along with the PIC value [12]. The highest values were observed for the D19S433 locus (He=0.86977; PIC=0.85) while the lowest values were identified in TPOX (He=0.6212; PIC=0.57).

After applying the Bonferonni corrections to the Fisher’s exact test a deviation from Hardy–Weinberg equilibrium (HWE) was detected for the THO, D5S818 and FGA loci. A deviation from HWE was also detected in a previous Saudi study of HLA allele frequencies [13]. This deviation was attributed to the high rates of consanguineous marriage among Saudis or to the Wahlund effect resulting in the reduction of heterozygosity. Findings in a previous Saudi study observed that 2 out of 8 STR loci did not conform to HWE which is consistent with our data [14]. As shown in Table 2, the p values for HWE of the available STR loci data that were obtained from previous Arab related populations reports were compared with the data from this study. It is worth noting that some alleles such as 34.2 (D21S11), 16.2 (D18S51), 12.2 (D16S539), 29 and 22.2 (FGA) and 7.3 and 8.3 (THO1) that were considered to be variant alleles in previous reports were also identified in frequencies ranging between 0.3 to 1.6% within the Saudi population studied in this project.

This study was based on a random sample of unrelated healthy volunteers of Arab descent who reside in the Riyadh district of Saudi Arabia. Allelic frequencies of 15 STR markers and their population genetic/forensic parameters were analyzed and results showed that the polymorphic nature of loci examined were sufficient to allow the
Table 1: Allele Frequency Distribution for the 15 STR loci in a Sample of Saudi population (n=190)
characterization of individuals from Saudi Arabia (markers D21S1, THO, D2S1338, D19S433, D18S58 and FGA). This study showed that D2S1338, D19S433 and FGA were the most polymorphic makers as evidenced by their high PD values. It is worth noting that the 4 markers with high PD values (vWA, THO1, CSF1PO and TPOX) were also identified by a previous study that used 8 STR loci to study unrelated subjects in Saudi Arabia (14).

In other studies, the most frequent allele for forensic STR loci can vary depending on the population, for example 12 (0.4093) at D5S818 in Slovenian population [15], 15 (0.4143) at D3S1358 in Bolivians [16], 8 (0.4989) at TPOX among Wallachians in South Romania [17] and 8 (0.424) at TPOX in Aidama community from Egypt [18]. In this study the most frequent alleles were 8 (0.558) at TPOX, 12 (0.411) at D13S317, 12 (0.385) at CSF1PO, 11 (0.382) at D16D539 and 10 (0.358) at D7S820. The most polymorphic marker in this study was D18S51; spanning 17 tandem repeat alleles. Allele 13 of this locus was the most predominant one with a frequency of 0.268. Interestingly, a study of Tunisians found markers D19S433 and D21S11 to be the most polymorphic; each spanning 18 alleles [4]. In contrast, a lesser degree of polymorphism was reported for Iraqi individuals as compared to Saudis and Tunisians [19] due to less admixture.

For the sample used in this study, the observed values of heterozygosity ranged from 0.621 at TPOX to 0.869 at D19S433. This lower degree of homozygosity suggests presence of a random mating in the study population and less consanguinity. Despite prolific rates of consanguineous marriage in the Saudi population [8], this study was able to clearly identify seven markers that can be used for characterizing the genetic makeup among individuals of Arab descent in Saudi Arabia.

Furthermore, it can also be argued that Independent Inheritance property could apply to two cases for markers collocated on the same chromosome, namely CSF1PO and D5S818 on chromosome 5, and TPOX and D2S1338 on chromosome 2. This was reflected by the disparate statistical values regarding HWE assumption that rendered in each case one marker as statistically significant while the other one was not.

It is also interesting to note that variant alleles of 34.2 (D21S11), 16.2 (D18S51), 12.2 (D16S539), 29 and 22.2 (FGA) as well as 7.3 and 8.3 (THO1) have also been identified by other studies. The 15 markers studied here were highly informative as illustrated by their high PD values (Table 1). Consistently higher PD values were also reported studies that examined populations from South Romania at locus FGA (PD=0.970), Tunisian at locus D19S433 (PD=0.976), sub-population of Botswana at locus D21S11 (PD=0.974) and Turkish at locus D18S51 (PD=0.979) [3,4,17,20].

A Multidimensional Scaling (MDS) plot was constructed to illustrate the genetic distances between 14 populations (Figure 1). The analysis of these populations showed clustering into four groups: (1) the Asian subpopulation, (2) the Caucasians, (3) African and (4) Middle East subpopulations. The Saudi group in this study clustered with populations from Yemen, Iraq, Qatar, Oman and Bahrain. In contrast, African populations like the Hutu and Kenyans as well as the Pakistani, Bangladesh, Punjabi and those of Caucasian origin like the Belgium, Croaitans and Georgians were located furthest from the samples of this study.

Conclusions

TPOX, D13S317, CSF1PO, D16D539 and D7S820 markers were found suitable for forensic analysis, paternity testing and can also be used for chimerism study after allogenic bone marrow transplantation for Saudi population. On the other hand, the variable degree of genetic distances of this population and other Arab related groups might be explained by the populations admixture with other ethnic origins. Further analyses of Arab population are needed to understand the interrelationship between the different ethnic groups of the region.

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