#### Journal of Forensic and Legal Medicine 37 (2016) 45-49



Contents lists available at ScienceDirect

#### Journal of Forensic and Legal Medicine

journal homepage: www.elsevier.com/locate/jflm

#### Short report

## Baluchi and Pakhtun population data of 9 X-chromosomal short tandem repeat loci



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#### ARTICLE INFO

Article history: Received 15 April 2015 Received in revised form 8 September 2015 Accepted 24 October 2015 Available online 2 November 2015

Keywords: Forensic science DNA typing X-chromosome short tandem repeats Baluchi population Pakhtun population Population genetics

1. Introduction

#### ABSTRACT

Baluchistan is the largest province of Pakistan in terms of area, constituting approximately 44% of the country's total land mass, and the smallest in terms of population, being home to less than 5% of the country's population. Khyber Pakhtunkhwa (KPK) formerly called North-West Frontier Province is located in the north-west of Pakistan having an estimated 13.4% of total population of Pakistan in which Pakhtuns are the major ethnic group. A total of 250 samples from Baluchi population and 250 samples from Pakhtun population were typed for 9 X-chromosomal STR markers: DXS101, DXS6789, DXS7132, DXS7423, DXS7424, DXS8378, GATA31E08, GATA172D05 and HPRTB along with sex typing locus, Amelogenin. A total of 59 alleles were found in Baluchi population while 61 alleles were found in Pakhtun population. This is the first study of the two populations based on these markers and the population data can be used as reference database for Baluchi and Pakhtun populations.

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# Forensic scientists have developed, standardized and discarded many techniques for human identification, from fingerprinting to DNA profiling, until the data basing of core short tandem repeats (STR) loci. Since then, millions of profiles are generated and it is very likely that STRs will be the workhorses for the near future.<sup>1,2</sup>

To alleviate the problems associated with analyzing DNA from degraded samples a new set of STR primers known as Miniplexes were designed by moving the primers closer to the repeat region leaving the extra sequences out.<sup>3,4</sup> Using shorter amplicons in polymerase chain reaction (PCR), improvement has been reported in obtaining results from forensic evidence or a mass disaster site having degraded specimens.<sup>5</sup>

The DNA typing has played a pivotal role to establish the paternity of child which is utmost priority for support, inheritance right and other social benefits of a child. STRs located on X

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chromosome are powerful marker for complex kinship testing such as deficiency paternity testing when the disputed child is a female.<sup>6,7</sup> X-STRs are routinely used in parentage analysis and relationship investigations such as avuncular and first cousin relationships. X-STRs have also advantage over autosomal STRs for paternity cases involving close blood relatives as alternative putative fathers and in deficiency paternity cases, i.e. when the DNA sample from putative father is not available and DNA from paternal relative has to be analyzed instead.<sup>8</sup> Further, X-linked STRs can be used to solve sibling ship status, without using father's DNA, of two females having the same biological father.<sup>9,10</sup> X-STRs can determine the relationship of grandmother/granddaughter as granddaughter theoretically has to carry at least one allele in common with the grandmother.<sup>11</sup> In forensic analysis of mixed stains, X-STRs are helpful to identify the female DNA.<sup>12,13</sup> X-STR markers are low size markers and can efficiently be used for degraded DNA analysis.

This study reports 9 mini X-STRs (DXS101, DXS6789, DXS7132, DXS7423, DXS7424, DXS8378, GATA31E08, GATA172D05 and HPRTB) along with sex typing locus Amelogenin, data of Baluchi and Pakhtun populations of Pakistan using an ABI PRISM 3130xl Genetic Analyzer (Applied Biosystems).

http://dx.doi.org/10.1016/j.jflm.2015.10.006

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#### 1.1. General information about the populations

Baluchistan is a province of south-western Pakistan. It is the largest province of Pakistan in terms of area, constituting approximately 44% of the country's total land mass, and the smallest in terms of population, being home to less than 5% of the country's population. It has an eventful history dating back to the Stone Age. Recent research and archaeological excavations have revealed 9000 years old civilization (http://www.balochistan.gov.pk/index.php). Khyber Pakhtunkhwa (KPK) formerly called North-West Frontier Province is located in the north-west of Pakistan. The province has an estimated population of about 21 million, or 13.4% of total population of Pakistan (www.census.gov.pk). Pakhtuns or Pathans make up the largest ethnic group in province, historically have been living in the areas for centuries (http://www.khyberpakhtunkhwa. gov.pk/aboutus/History.php).

#### 2. Materials and methods

#### 2.1. Populations samples collection

After the study was approved by the Institutional Review Board of CEMB (Centre of Excellence in Molecular Biology), Blood samples were collection from 250 individuals (100 males, 150 females) from Baluchistan and 250 individuals (150 males and 100 females) from KPK after their informed consent were obtained. All the individuals in this study were unrelated with each other and were residents of the area since as far as 3 generations back. This was by selfdeclaration as no genealogy records are maintained in these parts of the world.

#### 2.2. DNA extraction and quantification

Phenol chloroform DNA extraction was performed on each of the blood sample as reported in the literature.<sup>14</sup> DNA concentration was estimated through NanoDrop ND-1000 (NanoDrop Technologies, Wilmington, DE).

Table 1

Allele frequency distribution of 9 X-STRs in Baluchi population.

#### 2.3. PCR amplification and genotyping

The STRs were amplified in multiplex set which included 9 STRs (DXS7132, DXS8378, DXS101, DXS6789, DXS7424, GATA31E08, DXS7423, GATA172D05 and HPRTB) plus Amelogenin.<sup>15–19</sup> The Polymerase chain reaction (PCR) was carried out on ABI GeneAmp<sup>®</sup> PCR System 2700 using temperature conditions as given in Supplementary Fig. 1. The reaction was carried out in a 10 µL reaction consisting of 1 unit of AmpliTaq Gold polymerase (ABI), 200 µM of each dNTP, 2 mM MgCl<sub>2</sub>, 1X (NH<sub>4</sub>SO<sub>2</sub>) PCR Buffer (ABI), 1 ng of genomic DNA, distilled water and appropriate amount of labeled primers (Supplementary Table 1).

The capillary electrophoresis was performed using 1  $\mu$ L of PCR product, 13.6  $\mu$ L Hi-Di<sup>TM</sup> Formamide (ABI), and 0.4  $\mu$ L GeneScan<sup>®</sup> 500-LIZ<sup>TM</sup> size standard (ABI). Using a 3130xl Genetic Analyzer (ABI), at 3 kV the samples were injected for 10 s. In a Performance Optimized Polymer (POP-4), electrophoresis was carried out for 25 min at 15 kV and run temperature of 60 °C. ABI 3130xl Data Collection Software application v3.0 was used for collection of data. Results were analyzed using GeneMapper software v3.7 (ABI).

#### 2.4. Data analysis

Allele frequencies for each locus were calculated for both males and females collectively using a spreadsheet program. For Hardy–Weinberg Exact Test and Linkage Disequilibrium, Arlequin v3.5 was used.<sup>20</sup> PowerMarker v3.1 was used for Gene Diversity,<sup>21</sup> Polymorphic Information Content (PIC), and Heterozygosity. The ChrX-STR website was used for calculation of Paternity Index, Power of Exclusion, Power of Discrimination in males (PDm) and females (PDf), and Mean Exclusion Chance (MEC) in trios involving a daughter and in father/daughter duos.<sup>7</sup>

Alleles were assigned as recommended by ISFG (International Society of Forensic Genetics) through comparison with standard DNA 9947A. Allele nomenclature and allele ranges were according to already reported literature.<sup>7</sup> This manuscript strictly follows the guidelines for publication of population data requested by the journal.

	5		11						
Alleles	DXS101	DXS6789	DXS7132	DXS7423	DXS7424	DXS8378	GATA172D05	GATA31E08	HPRTB
6							0.1850		
7								0.2475	
8							0.1400	0.0150	
9						0.0450	0.1025	0.1325	
10						0.3325	0.2425	0.2775	
11						0.5025	0.2350	0.2625	0.0075
12			0.1025	0.0300	0.0925	0.0750	0.0950	0.0650	0.1225
13			0.2850	0.4150	0.0725	0.0450			0.3475
14		0.1025	0.3850	0.3875	0.1750				0.3175
15		0.0575	0.1825	0.0950	0.2575				0.1375
16		0.0225	0.0375	0.0725	0.2650				0.0675
17			0.0075		0.0825				
18		0.0300			0.0550				
19		0.2625							
20	0.4075	0.1900							
21	0.1275	0.2400							
22	0.0425	0.0725							
23	0.1925	0.0150							
24	0.2850	0.0075							
25	0.1000								
20	0.1275								
∠/ ว⊽	0.0925								
20	0.0325								

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