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Afghanistan from a Y-chromosome perspective

Harlette Lacau^{1,2}, Tenzin Gayden^{1,3}, Maria Regueiro¹, Shilpa Chennakrishnaiah^{1,2}, Areej Bukhari^{1,3}, Peter A Underhill⁴, Ralph L Garcia-Bertrand⁵ and Rene J Herrera^{*,1}

Central Asia has served as a corridor for human migrations providing trading routes since ancient times. It has functioned as a conduit connecting Europe and the Middle East with South Asia and far Eastern civilizations. Therefore, the study of populations in this region is essential for a comprehensive understanding of early human dispersal on the Eurasian continent. Although Y- chromosome distributions in Central Asia have been widely surveyed, present-day Afghanistan remains poorly characterized genetically. The present study addresses this lacuna by analyzing 190 Pathan males from Afghanistan using high-resolution Y-chromosome binary markers. In addition, haplotype diversity for its most common lineages (haplogroups R1a1a*-M198 and L3-M357) was estimated using a set of 15 Y-specific STR loci. The observed haplogroup distribution suggests some degree of genetic isolation of the northern population, likely due to the Hindu Kush mountain range separating it from the southern Afghans who have had greater contact with neighboring Pathans from Pakistan and migrations from the Indian subcontinent. Our study demonstrates genetic similarities between Pathans from Afghanistan and Pakistan, both of which are characterized by the predominance of haplogroup R1a1a*-M198 (> 50%) and the sharing of the same modal haplotype. Furthermore, the high frequencies of R1a1a-M198 and the presence of G2c-M377 chromosomes in Pathans might represent phylogenetic signals from Khazars, a common link between Pathans and Ashkenazi groups, whereas the absence of E1b1b1a2-V13 lineage does not support their professed Greek ancestry.

European Journal of Human Genetics (2012) 20, 1063–1070; doi:10.1038/ejhg.2012.59; published online 18 April 2012

Keywords: Afghanistan; Pathans/Pashtuns; Y-SNP; phylogenetic analyses; haplogroup; haplotype

INTRODUCTION

The Islamic Republic of Afghanistan is a landlocked nation in southwest Central Asia. Its central highlands include the Hindu Kush Mountains that extend from the northeast of the country to the southwest, separating Afghanistan into northern and southern provinces (Figure 1). Afghanistan has served as crossroads for human migrations and pilgrimages, including an important stop along the Silk Road. Mesolithic artifacts, Neolithic pottery (about 7.2 kya), bones of domesticated animal and tools (eg, sickle blades used to collect wild grasses) have been uncovered during excavations in the Ghar-i-Mar (Snake Cave) site, in north Afghanistan, indicating early cultivation of wheat and barley (9-11 kya) and domestication of animals (7–9 kya).¹ More recent archeological discoveries include Buddhist relics transported northward from India along the Silk Road, as well as inscriptions engraved on rocks in ancient Hebrew dating from the eleventh to the thirteenth centuries.² Although it is still unknown when urban civilizations began in what is now Afghanistan, historians speculate that it possibly occurred between 4 and 4.5 kya.3 Historical records indicate that Aryans from Iran were the first to occupy Afghanistan (eighth century B.C.E.) followed by Persians (sixth century B.C.E), Greeks (fourth century B.C.E.), Maurvans from India (ushered Buddhism into the region), Greco-Bactrians, Arabs (introduced Islam) and Mongols among others.³

The official languages of Afghanistan, Dari and Pashto, are of Indo-European origin and are spoken by the Tajiks and Pathans or Pashtuns, respectively. Pathans live primarily south of the Hindu Kush Mountains and comprise the most prevalent ethnic group in Afghanistan (42%) while representing approximately 15% of the population of neighboring Pakistan. Although different origins for Pathans, including Greek and Jewish ancestry, have been suggested,^{3,4} these affiliations have not been demonstrated and only a few studies characterizing the genetic structure of this group in Pakistan,^{5–9} Afghanistan¹⁰ and India¹¹ have been published.

Given the limited amount of genetic data available, the present study was undertaken to ascertain, for the first time, the genetic diversity of Pathans in Afghanistan utilizing high-resolution Y-chromosome binary markers. The data generated were subsequently compared with previously published geographically and ethnically targeted worldwide populations to explore paternal signals of modern humans' dispersals across Central Asia. In addition, we assessed genetic affinities between Pathans from Afghanistan and Pathans from neighboring Pakistan, as well as their hypothesized phylogenetic relation to Greek and Jewish populations.

MATERIALS AND METHODS

Sample collection and DNA isolation

Blood samples were obtained with informed consent from 190 unrelated males representing the general population of Pathans from Afghanistan. The collection was subsequently divided along the Hindu Kush Mountain Range into north Afghanistan (n = 44) and south Afghanistan (n = 146) to assess the role of the massif as a barrier and/or corridor to gene flow between the two regions (Figure 1). Donors provided information about their paternal ancestry

¹Department of Molecular and Human Genetics, College of Medicine, Florida International University, Miami, FL, USA; ²Department of Chemistry and Biochemistry, Florida International University, Miami, FL, USA; ⁴Department of Psychiatry and Behavioral Sciences, Stanford University, Palo Alto, CA, USA; ⁵Department of Biological Sciences, Colorado College, Colorado Springs, CO, USA

^{*}Correspondence: Dr RJ Herrera, Department of Molecular and Human Genetics, College of Medicine, Florida International University, University Park, OE 304, Miami, FL 33199, USA. Tel: +1 305 348 1258; Fax: +1 305 348 1259; E-mail: herrerar@fiu.edu

Received 10 November 2011; revised 20 February 2012; accepted 29 February 2012; published online 18 April 2012



Figure 1 Geographic location of the populations analyzed for CA, ML and AMOVA. For population description see references in Table 1.



Figure 2 Hierarchical phylogenetic relationships of Y-chromosome haplogroups and their percentages in Afghanistan. The following 77 markers were typed but were not polymorphic in the populations: C3a-M93, C3b-P39, C3c-M77, C3d-M407, C3e-P53.1, C3f-P62, DE-YAP, F1-P92, F2-M427, F2-M428, F3-P96, F3-M282, F4-P254, G1-M285, G2a1a-P18, G2a2-M286, H1a1-M197, H1a2-M97, H1a3-M39, H2-Apt, I-M258, J1a-M62, J1b-M365, J1c-M390, J1d-P56, J1e-PAGE8, J2a1-M47, J2a2-M67, J2a2a-M92, J2a2b-M163, J2a4-M137, J2a6-M289, J2a7-M318, J2a8-M319, J2a9-M339, J2a10-M340, J2a11-M419, J2a12-P81, J2a13-P279, J2b2a-M99, J2b2b-M280, J2b2c-M321, J2b2d-P84, K1-M147, K2-P60, K3-P79, K4-P261, L2-M317, M-P256, NO-M214, O-M175, Q1a1-M120, Q1a2-M25, Q1a3a-M3, Q1a4-P48, Q1a5-P89, Q1a6-M323, Q1b-M378, R1a1a1-M56, R1a1a2-M157, R1a1a3-M204, R1a1a4-P98, R1a1a5-PK5, R1a1a6-M434, R1a1a7-M458, R1b1b1a-M412, R1b1b1a1a-M405/U106, R1b1b1a1b1-U152, R1b1b1a1b2a-M222, R1b1b1a1b2b-M37, R1b1b1a1b2c1-P66, R1b1b1a1b3-M65, R1b1b1a1b4-M153, R1b1b1a1b5-M167, R1b1b-M335, S-M230 and T-M184.

for at least two previous generations. The samples were collected in accordance with the ethical guidelines of the institutions involved in this study. Genomic DNA was extracted and stored as previously described.^{12,13}

Y-SNP and Y-STR genotyping

A total of 116 bi-allelic markers (Figure 2) were hierarchically genotyped as previously described.^{14,15} The nomenclature followed for the Y-SNP haplogroup assignment is in accordance with the Y Chromosome

Consortium¹⁵ and subsequent updates.^{16,17} The 17-loci Y-STR haplotype data from our previous report¹⁸ were utilized in the current investigation to perform pertinent Y-STR-based analyses.

Statistical and phylogenetic analyses

To assess the phylogenetic relationships between the two Afghan collections and 16 geographically targeted reference populations (Table 1),^{19–22} a correspondence analysis (CA)²³ plot, Maximum-Likelihood (ML)²⁴ dendrogram and Analysis of

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AFN

AFS

Geographic region	Population	Abbreviation	n	Language family	References
West Central Asia (WCA)	North Afghanistan	AFN	44	Indo-European	Present study
	South Afghanistan	AFS	146	Indo-European	Present study
South Central Asia (SCA)	India	IND	728	Indo-European	Sengupta <i>et al</i> ⁸
	Pakistan (Balochi)	BAL	25	Indo-European	Sengupta <i>et al</i> ⁸
	Pakistan (Brahui)	BRA	25	Akin to Dravidian	Sengupta <i>et al</i> ⁸
	Pakistan (Burusho)	BUR	20	Burushaski	Sengupta <i>et al</i> ⁸
	Pakistan (Hazara)	HAZ	25	Indo-European	Sengupta <i>et al</i> ⁸
	Pakistan (Kalash)	KAL	20	Indo-European	Sengupta <i>et al</i> ⁸
	Pakistan (Makrani)	MAK	20	Indo-European	Sengupta <i>et al</i> ⁸
	Pakistan (Pathan)	PAT	20	Indo-European	Sengupta <i>et al</i> ⁸
	Pakistan (Sindhi)	SIN	21	Indo-European	Sengupta <i>et al</i> ⁸
South East Europe (SEE)	Greece (Athens)	GRE	92	Indo-European	Battaglia <i>et al</i> ³¹
	Heraklion Prefecture (Crete)	HER	104	Indo-European	Martinez et al ²⁰
	Lasithi Plateau (Crete)	LS1	41	Indo-European	Martinez et al ²⁰
	Lasithi Prefecture (Crete)	LS2	23	Indo-European	Martinez et al ²⁰
Middle East (ME)	North Iran	NIR	33	Indo-European	Regueiro et al ²¹
	South Iran	SIR	117	Indo-European	Regueiro et al ²¹
Levant (LEV)	Jewish (Ashkenazi)	ASH	442	Afro-Asiatic and Semitic	Behar et al ²²

Table 1 Populations analyzed

Molecular Variance (AMOVA)²⁵ were generated. In order to make informative comparisons, the above-mentioned analyses were performed at the highest resolution available across reference populations (Table 1). Pair-wise genetic distances (Rst)^{26,27} for populations included in Supplementary Table 1 were calculated using the 7-loci R1a1a-M198 haplotypes with the 'Microsat' software (http://hpgl.stanford.edu/projects/microsat/). The Rst distances calculated were subsequently utilized to construct a multi-dimensional scaling (MDS) projection with the statistical software package SPSS ver. 14.0 (SPSS, Chicago, IL, USA, 2001). In addition, spatial frequency maps based on individuals in the L-M20, L3-M357, R1a1a-M198 and R1a1a*-M198 haplogroups were generated using the Surfer software version 9 (Golden software Inc., Cold Spring Harbor, NY, USA; http://www.goldensoftware.com/). Haplotypes of individuals belonging to the above haplogroups were employed to generate median-joining (MJ) networks using NETWORK 4.5.1.6 (http://www.fluxus-engineering.com).28 All reference populations included in the network analyses are provided in Supplementary Table 1 (L-M20, L3-M357, R1a1a-M198 and R1a1a*-M198) and for the contour maps in Supplementary Tables 2 (R1a1a-M198 and R1a1a*-M198) and 3 (L-M20 and L3-M357). Supplementary Table 4 presents haplogroup and haplotype data for north and south Afghanistan.

The Y-STR markers were weighted inversely to their repeat variance²⁰ and the Maximum Parsimony option was selected to obtain the least complex topology.

Expansion times based on Y-STR variation within L-M20, L3-M357, R1a1a-M198 and R1a1a*-M198 lineages present in the reference populations (Supplementary Table 5) were recalculated using the method described elsewhere.^{8,29,30} In addition, haplotype variance for the above-mentioned haplogroups were calculated using the Vp formula provided in Kayser *et al.*³¹

RESULTS

Distribution of Y-chromosome haplogroups

From a total of 116 binary markers typed, we identified 25 paternal lineages in 190 Afghan males, 11 of which were observed in north Afghanistan and 22 in the southern collection, (Figure 2) indicating a relatively higher degree of genetic diversity in the region south of the Hindu Kush range with respect to the north. However, this finding should be tempered considering the difference in sample size between the two collections. Overall, only haplogroups R1a1a*-M198 (62.1%), L3*-M357 (7.4%) and G2c-M377 (5.3%) display frequencies >5% and collectively comprise nearly three-fourths of the Afghanistan paternal gene pool.

The majority of the Afghan individuals examined in the current investigation belong to haplogroup R-M207 (67.4%), with sub-haplogroup R1a1a*-M198 predominating in both the north (50%) and south (65.8%) collections (Figure 2). This is also reflected in the contour maps of R1a1a-M198 (Supplementary Figure 1a) and R1a1a*-M198 (Supplementary Figure 1b). It is interesting to note that the paralagous branch, R2a-M124, is observed at a considerably higher proportion in north Afghanistan (11.4%) as compared with the south (2.1%) and that haplogroups R2*-M479 (0.7%) and R1b1a2a-L23 (0.7%) are detected, albeit at low frequencies, exclusively in the southern region of the country.

Haplogroup L-M20 (9.5%), the second most abundant in this study, exhibits substantial disparity in its distribution on either side of the Hindu Kush range, with 25% of the northern Afghans belonging to this lineage, compared with only 4.8% of males from the south. Specifically, paragroup L3*-M357 accounts for the majority of the L-M20 chromosomes in both the north (20.5%) and south (4.1%) collections as evident in the contour maps provided in Supplementary Figures 2a (L-M20) and 2b (L3-M357) in Afghanistan and surrounding regions.

Haplogroup G-M201, which occurs at an average frequency of 7.9% in the Afghan gene pool, is reported at high frequencies in the Caucasus (eg, 74% in Ossetians from Digora)³² and it is thought to be associated with the Neolithic expansion throughout the region.³³ Although G2c-M377 is the second most frequent haplogroup in south Afghanistan (6.2%), it is found at substantially lower levels in the northern population (2.3%). With the exception of sub-lineage G2a*-P15, observed in the north (2.3%) and south (1.4%), the remaining G derivatives (G2a1*-P16 (0.7%) and G*-M201 (0.7%)) are found exclusively in the southern provinces. Likewise, haplogroups C-M216 (3.4%) and H1a*-M82 (4.1%) are restricted to individuals from the south. Other informative haplogroups that are shared between the north and south include J-M304 and Q-M242. In the north, the aforementioned haplogroups are defined by the J2a*-M410 (2.3%), J2a3-M68 (2.3%) and Q1a3*-M346 (4.5%) mutations. Southern Afghans, in contrast, are characterized by five sub-clades within haplogroup J-M304 (ie, J1*-M267, J2a*-M410, J2a5-M158, J2b1-M205 and J2b2-M241) collectively at frequency of <3%, as well





Figure 3 (a) CA constructed utilizing the NTSYSpc 2.02i software (Applied Biostatistics Inc., Setauket, NY, USA) and based on Y-chromosome haplogroups. Population abbreviations are given in Table 1. (b) MDS plot based on *Rst* distances of R1a1a-M198 haplotype frequencies of Pathans, Ashkenazi Jews, Greeks, Indians and Pakistani samples. For the MDS analysis Stress = 0.06142. Population codes are: NAFG = Pathans from MDK Afghanistan, SAFG = Pathans from south Afghanistan, PTHP = Pathans from Pakistan, RUS = Russians (South Central Russia, South Western Russia, North Western Russia and Uralic Russia), GRE = Greeks, ASH = Ashkenazi Jews, IND = India and PAK = Pakistan general population. Populations used in the analysis are listed in Supplementary Table 1.

as by sub-haplogroups Q1a3*-M346 (1.4%), Q1*-P36.2 (0.7%) and Q1a*-MEH2 (0.7%).

Population relationships

In the CA plot (Figure 3a), the collections from north and south Afghanistan are part of a loose aggregate in the right half, near the center of the graph, with several neighboring populations from Pakistan. The Greeks, Ashkenazi Jews and north and south Iranians partition separately from the Pakistani and Afghani collections on the upper-left quadrant. Interestingly, Hazara from Pakistan is an outlier, possibly reflecting its unique ancestry. The genetic affinities of the 18 populations from Table 1 are illustrated in the ML tree (Supplementary Figure 3), where geographic structuring is apparent, with a few exceptions. For example, unlike the partitioning observed in the CA plot, the ML dendrogram illustrates a separation between north and south Iran, with the former branching together with Hazara from Pakistan, an outlier in the CA plot. AMOVA (Table 2) was performed to investigate potential geographic correlations with the genetic profiles of the north and south Afghanistan populations, as well as reference collections. As anticipated, the highest fraction of variability was found within populations. Variance components

Table 2 Analysis of molecular variance (AMOVA) using Y-SNPs

		Fixation indices		
Classification criterion	Total variation (%)	ΦST	ΦSC	ΦCT
Geography (five groups)				
Among groups	6.02			0.06^{*}
Among populations within groups	5.35		0.06*	
Within population	88.63	0.11^{*}		

Geographic Partitioning (five groups). 1 Middle East: Northern Iran and Southern Iran. 2 Levant: Jewish-Ashkenazi. 3 West Central Asia: North Afghanistan, South Afghanistan. 4 South Central Asia: India, Balochi, Hazara, Kalash, Makrani, Pathan, Sindhi, Barhui, and Burusho. 5 South East Europe: Greece-Athens, Heraklion, Prefecture-Crete, Lasithi Plateau-Crete and Lasithi Prefecture-Crete. Φ ST = variance within populations; Φ SC = variance among populations within groups; Φ CT = variance among groups. **P*-value < 0.001.

indicate a greater degree of geographical correlation among groups of populations as compared with among populations within groups.

The MDS plot based on R1a1a-M198 (Figure 3b) haplotypes was executed to examine the genetic affinities among Pathans from Afghanistan and Pakistan in addition to several other pertinent reference populations (Supplementary Table 1). In Figure 3b, Pathans from Afghanistan and Pakistan plot close to each other in the lower-right quadrant, whereas populations from India, Pakistan and Russia loosely cluster in the upper-right section of the graph. In contrast, the Ashkenazi Jewish collection lies in the upper-left quadrant partitioning distant from all other populations included in the analysis.

Network projections and time estimates

Figure 4a illustrates MJ network based on the eight-loci Y-STR profile of R1a1a*-M198 individuals from different regions throughout Eurasia. The most abundant haplotype (DYS19*16, DYS389I*13, DYS389II*17, DYS390*24, DYS391*11, DYS392*11, DYS393*13 and DYS439*10) is observed in 5 (11.3%) individuals from north Afghanistan and 35 (24%) Afghans from the southern collection. In contrast, the second most common haplotype (DYS19*16, DYS389I*13, DYS389II*17, DYS390*25, DYS391*11, DYS392*11, DYS393*13 and DYS439*10), is prevalent in north Afghanistan (6.8%) compared with the south collection (4.8%) and it is also found among Russia, Ukraine, Slovakia and Turkey. Interestingly, the third most frequent haplotype (DYS19*16, DYS389I*13, DYS389II*17, DYS390*25, DYS391*10, DYS392*11 and DYS393*13) in the R1a1a-M198 projection (Supplementary Figure 4a) is mainly shared among Ashkenazi Jews from western and eastern Europe (4.1%) and south Afghans (3.4%), but absent in individuals from north Afghanistan.

In Figure 4b, the most abundant haplotype (*DYS*19*14, *DYS*389I*12, *DYS*389II*16, *DYS*390*22, *DYS*391*10, *DYS*392*14, *DYS*393*11 and *DYS*439*13) among the L-M20 individuals examined comprises 7 (2.3%) samples from southern India and 12 (13.2%) from south Pakistan. Only one individual (2.3%) from North Afghanistan shares this haplotype. The other prevalent L-M20 haplotype (*DYS*19*14, *DYS*389I*12, *DYS*389II*16, *DYS*390*22, *DYS*391*10, *DYS*392*14, *DYS*393*11 and *DYS*439*12) is shared exclusively among 12 (4.0%) individuals from southern India and 3 (3.3%) from southern Pakistan. A network analysis based on L3-M357 was not informative because of the limited number of individuals and populations available for comparison (Supplementary Figure 4b).

Supplementary Table 5 presents coalescence time estimations based on Y-STR diversity as well as haplotype variances associated with haplogroups L-M20, L3-M357, R1a1a-M198 and R1a1a*-M198 in



Figure 4 (a) Network projection of populations analyzed for R1a1a*-M198 at the resolution of eight-loci: *DYS*389I, *DYS*389I, *DYS*3990, *DYS*391, *DYS*3992, *DYS*393, *DYS*19 and *DYS*439. (b) Network projection of populations analyzed for L-M20 at the resolution of eight-loci: *DYS*389I, *DYS*

Afghanistan and other reference populations. In addition, for the studied populations of north and south Afghanistan, we performed time estimate and haplotype variance calculations for L3*-M357. In the case of R1a1a-M198, the evolutionary time estimates based on seven Y-STR loci is 7.8 ± 2.2 kya for the Afghanistan populations. The oldest dates for this haplogroup are observed in Pakistan (19.1 ± 2.9 kya) and India (17.9 ± 3.7 kya). Time estimates for Haplogroup L-M20 in north (14.6 ± 7.3 kya) and south (17.8 ± 8.4 kya)

Afghanistan are more recent than in Pakistan (26.3 ± 5.3 kya). When L3-M357 and Ra1a1*-M198 are considered, the same pattern of relative ages are seen across the above-mentioned populations, with the older dates detected in Pakistan and in the Indian subcontinent.

DISCUSSION

Although both Afghan populations are characterized predominantly by R1a1a*-M198 lineages, the two differ in the distribution of their npg

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remaining haplogroups. South Afghanistan is represented by 22 polymorphic binary markers, 14 of which are absent from the northern region (Figure 2), suggesting reduced genetic diversity in the latter population. A plausible explanation for the reduced diversity might be the presence of the Hindu Kush Mountain range, which may have served as a directional barrier to gene flow.

South Central Asian influence in Afghanistan

Haplogroup R1a1a*-M198, unlike other haplogroups, which are more geographically confined, is present at informative frequencies in a wide range of populations extending from Northern and Eastern Europe to Eastern Siberia, Central and Southern Asia.¹⁶ Eastern Europe (Ukraine)³⁴ and South Central Asia^{16,35} are proposed as the most likely places of origin because in both regions R1a1a-M198 is observed at frequencies >50%. According to Underhill *et al*,¹⁶ the highest Y-STR haplotype diversity for the R1a1a*(xM458) lineage is observed in South Central Asia with a coalescent time of 14 kya, suggesting that this region is the likely source of the dispersal for the M198 mutation. Alternatively, Klyosov³⁶ claims that haplogroup R1a1a-M198 originated in South Siberia about 20 kya. Unfortunately, discrepancies between the methods used to compute TMRCA in these previous studies^{16,35,36} hinder fair comparisons with those generated in the present report. The high R1a1a haplotype diversity observed in the network projection (Supplementary Figure 4a), as well as the substantially older time estimates and high haplotype variance (Supplementary Table 5) obtained for South Central Asia (Pakistan and India) as compared with South Central Russia (Shors and Tuvinians) and Afghanistan, argues for the South Central Asian origin of this marker. Because TMRCA calculations depend on sample size, mutation rate and the number of loci examined, the data generated should be interpreted with caution and alternative approaches and/or interpretation should not be ignored.

The R1a1a-M198 network (Supplementary Figure 4a) indicates that the Afghans exhibit the highest haplotype sharing (total of 69 individuals) with populations from South Central Asia and particularly with Pathans from Pakistan. When the analysis was performed at R1a1a*-M198 (Figure 4a), we see a similar haplotype sharing with South Central Asia. Unfortunately, Pathans from Pakistan were not included in the analysis because they were not typed at the R1a1a*-M198 resolution. Genetic affinities among the above-mentioned populations that share high frequency of R1a1a*(xM458) are also observed in the CA plot (Figure 3a), which illustrates both Afghan populations clustering with most of the neighboring groups from Pakistan, whereas India plots more distantly. This finding is consistent with the AMOVA results, which indicates the impact of geography on population differentiation. However, the second most common seven-loci R1a1a-M198 haplotype (DYS389I*13, DYS389II*17, DYS390*25, DYS391*11, DYS392*11, DYS393*13 and DYS19*16) possessed by 66 individuals in the projection (Supplementary Figure 4a) differs from the modal haplotype by one repeat at locus DYS390 and corresponds to the most abundant haplotype of Russia/Ukraine and the Balkans reported by Klyosov.³⁶ This basal haplotype is also the second most common haplotype in the R1a1a*-M198 network (Figure 4a) with similar groups of populations. The sharing of similar seven-loci haplotypes (mostly differing by a single repeat) along with relatively close age estimates (Supplementary Table 5) among populations with significant R1a1a-M198 frequencies may be indicative of common ancestry.

Haplogroup L-M20, on the other hand, was hypothesized to have originated in India or the Middle East³⁷ approximately 30 kya.

This marker, which is found at 25% in north Afghanistan and 4.8% in the south, has also been previously reported at high frequencies (48%) in the Kallar community of South India³³ as well as in the Druze (35%) population from Israel.³⁸ Time estimates generated based on seven Y-STR loci within L-M20 lineages for north $(14.6 \pm 7.3 \text{ kya})$ and south $(17.8 \pm 8.4 \text{ kya})$ Afghanistan populations are intermediate to those of Pakistan $(26.3 \pm 5.3 \text{ kya})$ and India $(7.5 \pm 1.7 \text{ kya})$ (Supplementary Table 5). Furthermore, Pakistan displays higher haplotype variance (0.548) than India (0.118), suggesting that L-M20 most likely originated in what is today Pakistan rather than in India. In addition, evolutionary time estimates (Supplementary Table 5) generated for L3-M357 lineages when combining north and south Afghanistan populations $(11.4 \pm 5.0 \text{ kya})$ are comparable to those computed for Pakistan (10.8 ± 3.8 kya) and higher than India (5.2 ± 2.4 kya) at the seven-loci resolution. This finding supports the notion that the L3-M357 sub-clade is of Afghani or Pakistani origin and subsequently spread southernly to India.

Similarity between Pathans from Afghanistan and Pakistan

Our study demonstrates close genetic ties between Pathans from Afghanistan and Pakistan. The affinity between Pathans from these two regions is expected considering that the Durand Line, which established Afghanistan southeastern limits since 1893, was imposed without concern to tribal and ethnic affiliations. The CA (Figure 3a) and ML analysis (Supplementary Figure 3) support this genetic similarity since both north and south Afghanistan cluster proximal to Pakistan. The inclusion of additional haplotype data from 41 Pathans residing in the city of Peshawar in northwestern Pakistan⁵ (available in the YHRD database) indicates that their most abundant seven-loci haplotype (29.3%) is identical to our modal haplotype, confirming the genetic similarities between Pathans from Afghanistan and Pakistan. This phylogenetic relationship is similarly evident in the MDS graph (Figure 3b), where all three Pathan populations group close to each other. Also, in the network projection of R1a1a-M198 (Supplementary Figure 4a), these populations share various haplotypes. Moreover, the presence of haplogroups L3-M357 and Q1a3-M346 in both Pathans from Afghanistan and Pakistan^{8,9} further supports genetic relationships among Pathans from these two regions.

On the origins of Pathans

Although Greeks and Jews have been proposed as ancestors to Pathans,^{3,4} their genetic origin remains ambiguous. The Lasithi Plateau isolate, in the highlands of eastern Crete, partitions relatively close to the Afghanistan populations in the CA graph (Figure 3a), which could be attributed to the elevated proportion of R1a1a chromosomes²⁰ shared among them. However, the absence of the predominantly Greek E1b1b1a2 -V13 lineage³⁹ in Pathans does not argue for genetic contributions from Greece. Ashkenazi Levites and our north and south Afghanistan populations, on the other hand, exhibit significantly higher frequency of the R1a1a-M198 haplogroup (51.7%, 50.0% and 65.8%, respectively), whereas lower frequencies are detected in Ashkenazi Cohanims (1.3%) and Ashkenazi Israelites (4.0%).⁴⁰ Overall, Ashkenazi Jews exhibit a frequency of 15.3% for haplogroup R1a1a-M198.37 Nebel et al41 proposed, on the basis of moderate to high frequency of R1a1a in Central Asia and southern Russia/Ukraine, that these R1a1a chromosomes in Ashkenazi groups might represent vestiges of Khazars. It is thought that this Turkic tribe from Central Asia had converted to Judaism at the end of the Eighth century C.E. and, after the fall of their kingdom, they spread and were assimilated by the emerging Ashkenazi community in Eastern Europe.

We envision a plausible scenario in which the converted Khazars could have been absorbed by the early Pathans and that R1a1a-M198 drifted to high frequency in Afghanistan, with the Khazars being the common nexus between Ashkenazi Jews and Pathans. In addition, the Jewish traditions (particularly circumcision, a talith prayer shawl, shabbat, praying in the direction of Jerusalem during the Day of Atonement or Yom Kippur and the Magen David symbol in their houses, among several others)² observed among Pathans from Afghanistan and the presence of haplogroup G2c-M377, a lineage commonly observed among the Ashkenazi Jewish population (\sim 7%),⁴² are congruent with the above-stated hypothesis.

CONCLUSION

The diversity of Y-chromosome haplogroups in the Afghanistan populations examined in this study reflects the history of the area signaling multiple migrations over thousands of years. Our results indicate that Afghanistan exhibits patrilineal elements from the Neolithic revolution dispersal across Central Asia. The observed haplogroup distribution suggests some degree of genetic isolation of the northern population, likely because of the Hindu Kush Mountain range acting as a barrier and separating them from the southern Afghans who have greater contact with neighboring Pakistan and migrants from the Indian subcontinent and Central Asia. Our study demonstrates genetic similarity among Pathans from Afghanistan and Pathans from Pakistan, supported by a significant number of individuals belonging to haplogroup R1a1a-M198 and sharing the same modal haplotype. Genetic signatures of South Central Asian populations are evident in their patrilineages and may represent gene flow associated with the Silk Road migrations. In addition, the high frequencies of R1a1a-M198 chromosomes and the sharing of seven Y-STR loci haplotypes among Pathans, Ashkenazi Jews and west Russians may represent vestiges from the Khazars of Central Asia that converted to Judaism at the end of the Eighth century C.E., connecting Pathans with certain Ashkenazi Jews by the way of a common Khazar ancestry.

CONFLICT OF INTEREST

The authors declare no conflict of interest.

ACKNOWLEDGEMENTS

The authors express their appreciation to the original DNA donors who made this study possible. We gratefully acknowledge Viviana Arce, Priscilla Torres, David Perez and Silvia Calderon for their contribution to the manuscript and we thank Tanya Simms for her valuable comments.

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Supplementary Information accompanies the paper on European Journal of Human Genetics website (http://www.nature.com/ejhg)