

Deep common ancestry of Indian and western-Eurasian mitochondrial DNA lineages

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About a fifth of the human gene pool belongs largely either to Indo-European or Dravidic speaking people inhabiting the Indian peninsula. The 'Caucasoid share' in their gene pool is thought to be related predominantly to the Indo-European speakers.

A commonly held hypothesis, albeit not the only one, suggests a massive Indo-Aryan invasion to India some 4,000 years ago [1]. Recent limited analysis of maternally inherited mitochondrial DNA (mtDNA) of Indian populations has been interpreted as supporting this concept [2,3]. Here, this interpretation is questioned. We found an extensive deep late Pleistocene genetic link between contemporary Europeans and Indians, provided by the mtDNA haplogroup U, which encompasses roughly a fifth of mtDNA lineages of both populations. Our estimate for this split is close to the suggested time for the peopling of Asia and the first expansion of anatomically modern humans in Eurasia [4–8] and likely pre-dates their spread to Europe. Only a small fraction of the 'Caucasoid-specific' mtDNA lineages found in Indian populations can be ascribed to a relatively recent admixture.

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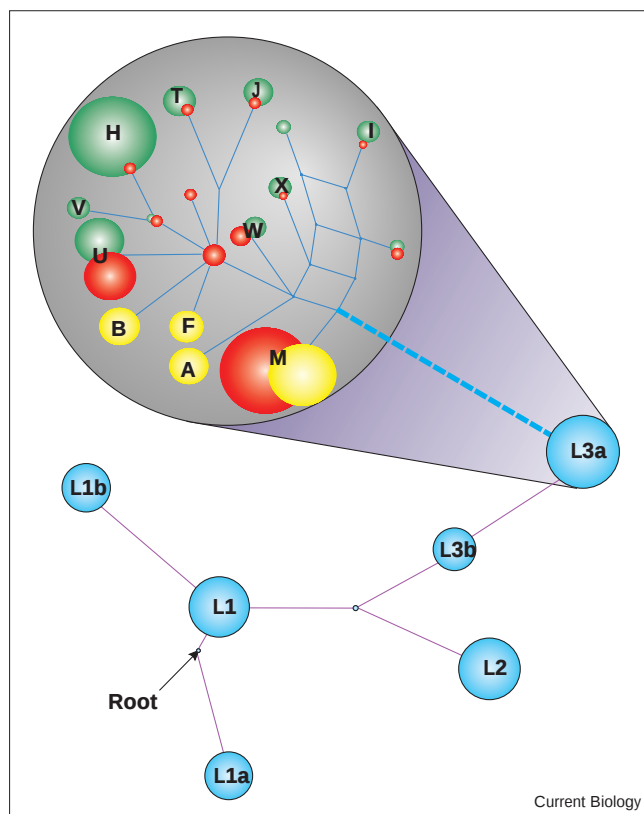
Results and discussion

The recent African origin of modern humans is now supported by palaeoanthropological, as well as sex-specific and autosomal genetic, evidence (for recent reviews, see [8,9]). The concordance between the interpretation of data obtained by mtDNA, Y-chromosomal and most of the

autosomal markers is encouraging and suggests that, irrespective of the differences in the mode of inheritance, these three genetic approaches produce consistent overall findings in this central issue.

We sequenced the mitochondrial hypervariable region I (HVR I) and performed extensive restriction fragment length polymorphism (RFLP) analysis of 550 Indian mtDNA samples. We inferred a parsimonious phylogenetic tree from the data using the median network approach [10], which is particularly suitable for intraspecies analysis of mtDNA lineages and other highly variable data sets. Figure 1 is an outline of this Indian mtDNA tree within the background of the previously defined global mtDNA lineage clusters (haplogroups) [11–13]. Consistent with the recent out-of-Africa model of human origins [14], all of the Indian mtDNA lineages we inferred can be seen as deriving from the African mtDNA lineage cluster L3a, described in [15]. We found that more than 80% of the Indian mtDNA lineages belong to either Asian-specific haplogroup M (60.4%) or western-Eurasian-specific haplogroups H, I, J, K, U and W (20.5%), while the remaining 19.1% of lineages do not belong to any of the previously established mtDNA haplogroups (Table 1). We note that haplogroup K should now be considered a sub-cluster of haplogroup U [13].

The first and the most profound layer of overlap between the western-Eurasian and the Indian mtDNA lineages relates to haplogroup U, a complex mtDNA lineage cluster with an estimated age of 51,000–67,000 years [16]. Until now, this haplogroup has not been reported to occur in India nor east of India and was considered a western-Eurasian-specific haplogroup. Surprisingly, we found that haplogroup U is the second most frequent haplogroup in India as it is in Europe (Table 1). Nevertheless, the spread of haplogroup U subclusters in Europe and India differs profoundly (Figure 2). The dominant subcluster in India is U2. Although rare in Europe, the South-Asian form differs from the western-Eurasian one: western-Eurasian U2 includes a further characteristic transversion at nucleotide position (np) 16,129 [12], which is absent in Indian U2 varieties (Figure 2). We calculated the coalescence age essentially as described in [15,17] and estimate the split between the Indian and western-Eurasian U2 lineages as $53,000 \pm 4,000$ years before present (BP). We note that U5, the most frequent and ancient subcluster of haplogroup U

Figure 1

The skeleton network of Indian lineage clusters on the background of continent-specific mtDNA haplogroups. Red, Indians; green, western Eurasians; yellow, eastern Eurasians; blue, Africans. Haplogroup frequencies are proportional to node sizes. All Indian, eastern-Eurasian and western-Eurasian mtDNA lineages coalesce finally to the African node L3a. The former are shown magnified to account for higher mtDNA diversity in sub-Saharan Africans. The most likely root of the tree [15] is indicated within a pan-African cluster L1. The dashed line leading from the African external node L3a to the Eurasian mtDNA varieties identifies the position of L3a in the magnified part of the tree.

in Europe, has an almost identical coalescence age estimate [13]. Still, despite their equally deep time depth, the Indian U2 has not penetrated western Eurasia, and the European U5 has almost not reached India (Table 2).

Subcluster U7 (among U* in [12,13]) is another variety of haplogroup U present in India (Figure 2). Unlike the Indian U2, it has been sampled, albeit rarely, in southern Europe, the Near East [12,13] and (according to HVR I sequence identification only) also in Central Asia [18]. We calculated the coalescence age of this subcluster in India as $32,000 \pm 5,500$ years: still deep in late Pleistocene but considerably younger than that for U2. Table 2 compares the frequency of varieties of haplogroup U in India, in the Trans-Caucasus populations and in Europe.

Typical western-Eurasian mtDNA lineages found in India belong to haplogroups H, I, J, T, X and to subclusters U1,

Table 1

MtDNA haplogroup frequencies (%) among some Indian and Eurasian populations.

Haplogroup affiliation	Population or group of populations						
	India			Western and eastern Eurasia			
	1	2	3	4	5	6	7
African*	0	0	0	0	0	0	0
Eastern Eurasian	53.3	65.7	60.4	1.5	0.7	61	94.5
Western Eurasian	29.3	14.5	20.5	80.9	95.4	30.5	0
H	3	1.2	1.8	24.8	41.1	14	0
I	2	0	0.7	1.8	2.6	1	0
J	0	0.4	0.5	6.7	10.3	2.5	0
K	0	0.4	0.2	8.2	4.4	0.5	0
T	1	1.7	1.8	11.8	10.1	3.5	0
U	23.3	10.3	13.1	21.2	20.8	8	0
V	0	0	0	0	3.1	0	0
W	0	0.4	2.2	0.9	1.6	1	0
X	0	0	0.2	5.5	1.4	0	0
Others†	17.4	19.8	19.1	17.6	3.9	8.5	5.5

The numbers in italics represent the following populations: 1, North India (Uttar Pradesh, $n = 103$, this study); 2, South India (Andhra Pradesh Telugus, $n = 250$, this study); 3, India total ($n = 550$, this study); 4, The Caucasus – Armenians ($n = 192$, this study), Georgians ($n = 138$, this study); 5, Europe – Slovaks ($n = 129$, this study), Russians ($n = 100$, this study), Czechs ($n = 95$, this study), Estonians ($n = 100$, this study), Italians ($n = 99$ [27]), Finns ($n = 49$ [16]); 6, Central Asia – Kirghiz ($n = 95$, deduced from [18]), Kazakhs ($n = 55$, deduced from [18]), Uighurs ($n = 55$, deduced from [18]); 7, Tibet ($n = 54$ [26]). *L1 and L2 defined by +3592 HpaI. †Lineages that do not belong to any of the previously established haplogroups.

U4, U5 and K of haplogroup U (Figure 1; Tables 1,2). Frequencies of these lineages in Indian populations are more than an order of magnitude lower than in Europe: 5.2% versus 70%, respectively (normalised from Table 1). This finding might be explained by gene flow, as suggested previously [2]. Nevertheless, we note that the frequency of these mtDNA haplogroups reveals neither a strong north–south, nor language-based gradient: they are found both among Hindi speakers from Uttar Pradesh (6%) and Dravidians of Andhra Pradesh (4%). Assuming that they are largely of western-Eurasian origin, we may ask when their spread in India started. To assign a tentative date for their introduction, we calculated the averaged minimal distance of the corresponding mtDNA hypervariable region sequences in Indians from the branches shared with western Eurasians. We obtained a value for the statistic p (see Materials and methods) equal to 0.46, consistent with a

Indians. Furthermore, this specific distribution of mtDNA varieties in India compared with the distribution observed among Mongoloids and the Caucasoid populations of western Eurasia (Figure 1) is, at present, best explained by two separate late Pleistocene migrations of modern humans to India. One of them, possibly arriving by the southern route, brought to India an ancestral population carrying haplogroup M and was spread further eastward. The second migration brought the ancestors of haplogroup U. Although the admixture of these major waves started perhaps very early — explaining the spread of these major mtDNA varieties all over the subcontinent — it is likely that it happened after the carriers of haplogroup M found their way further east, explaining the absence of haplogroup U lineages among Mongoloid populations studied so far.

Materials and methods

Samples from 86 Lambadi, 62 Lobana (Lamani speakers; Indo-Aryan languages), 12 Tharu and 18 Buksa (Indo-Aryan languages), 122 predominantly Indo-Aryan language speakers from Uttar Pradesh (GenBank accession numbers AJ234902–AJ235201) and a set of 250 Telugu samples (Dravidic speakers) were sequenced for hypervariable region I of mtDNA and typed for the presence of major continent-specific markers, described in [11,16,26]. The HVR I polymorphic sites of all 550 Indian mtDNAs sequenced by us are provided in the Supplementary material. The phylogenetic analysis also included 101 published HVR I sequences from south-western India [6].

Phylogenetic analysis was performed by reduced median networks [10], applied here using parsimony analysis of the data. The median network analysis allows one to reveal simultaneously multiple parallel, equally probable, phylogenetic pathways in the form of reticulations induced by highly variable markers. The distinct mtDNA lineage clusters are referred to here as haplogroups. The time to the most recent common ancestor of a cluster of lineages (haplogroup) or, where appropriate, a sub-cluster inside a particular haplogroup, was calculated as described [17], using an estimator ρ , which is the average transitional distance from the founder haplotype sequence.

Supplementary material

Supplementary material including a table listing the mtDNA HVR I sequence polymorphisms in different Indian populations and a more detailed description of the materials and methods is available at <http://current-biology.com/supmat/supmatin.htm>.

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Supplementary material

Deep common ancestry of Indian and western-Eurasian mitochondrial DNA lineages

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Supplementary materials and methods

Samples from 86 Lambadi, 62 Lobana (Lamani speakers; Indo-Aryan languages), 12 Tharu and 18 Buksa (Indo-Aryan languages) were collected as part of the ongoing genetic studies of the populations of the Indian subcontinent at the Division of Human Genetics of the University of Newcastle-upon-Tyne. In addition, 122 samples with mixed-caste status, predominantly Indo-Aryan language speakers from Uttar Pradesh and Kashmir, were included in the analyses (GenBank accession numbers AJ234902–AJ235201). The Uttar Pradesh sequences are from our independent Gypsy study (K.K., F. Calafell, T.K., M.J.B., J.P., E.M. *et al.*, unpublished observations). The set of 250 Telugu samples that was used to represent Dravidic speakers will be published elsewhere. Altogether, 550 samples from the Indian peninsula were sequenced for hypervariable region I of mtDNA and typed for the presence of major continent-specific markers, described in [S1–S3].

Phylogenetic analysis was performed by reduced median networks [S4], applied here using parsimony analysis of the data. In general, parsimony methods for inferring phylogenies operate by selecting trees that minimize the total tree length [S5]. In particular, the median networks approach allows one to reveal simultaneously multiple parallel, equally probable, phylogenetic pathways in the form of reticulations induced by highly variable markers. These reticulations reflect either parallel mutations or, more often, ambiguities in the branching pattern of a phylogenetic tree. Compared with any 'single tree' method, the network approach does not increase the phylogenetic resolution artificially. Nevertheless, considerable reduction of the network towards a tree can be achieved by giving higher weight to conservative markers versus hypervariable ones. Here, the reduced median networks were constructed using RFLP-typed conservative markers from the mtDNA coding region, with haplogroups specified according to the nomenclature proposed in [S1–S3,S6] and were further refined using sequence data from HVR I of the D-loop of the mtDNA genome. Every cluster of mtDNA lineages thus inferred should, in theory, constitute a monophyletic clade in the human mtDNA pool. These distinct clusters are referred to here as mtDNA haplogroups. The time to the most recent common ancestor of a cluster of lineages (haplogroup) or, where appropriate, of a sub-cluster inside of a particular haplogroup, was calculated as described [S7], using an estimator ρ , which is the average transitional distance from a founder haplotype sequence. We considered only transitions between nucleotide positions 16,090–16,365 in the HVR I of mtDNA and one substitution per 20,180 years was taken as an average distance from a specified founder [S8]. The phylogenetic analyses also included 101 published D-loop sequences from southwestern India [S9]. Western Eurasian samples that were used as a comparator included our unpublished sequences and RFLP data on the Caucasus area ($n = 330$), Slavic populations ($n = 324$) and approximately 2000 sequences retrieved from data banks [S10,S11] and recent publications [S12,S13].

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Table S1

The mtDNA HVR I sequence polymorphisms in different Indian populations.

Sample number	Origin	HVS-I sequence polymorphisms*	Haplogroup	RFLP polymorphisms*
Ind625	UP	51 145 179 234 240AC 242CG 353 362	U2i	+12308 <i>Hinf</i> I; -11465 <i>Tru</i> I
Ind873	UP	86 126 223	M1	+10394 <i>Ddel</i> ; +10397 <i>Alu</i> I
Ind872	UP	234	H	-00073 <i>Alw</i> 44I; -7025 <i>Alu</i> I
Ind624	UP	126 223 311	M1	+10394 <i>Ddel</i> ; +10397 <i>Alu</i> I
Ind874	UP	223 319	M2	+10394 <i>Ddel</i> ; +10397 <i>Alu</i> I
Ind623	UP	223 290 319 362	A	+663 <i>Hae</i> III
Ind868	UP	51 129 223 362	M4	+10394 <i>Ddel</i> ; +10397 <i>Alu</i> I
Ind869	UP	126 362	pTJ	-00073 <i>Alw</i> 44I
Ind621	UP	refer	R*	
Ind871	UP	172 304 362	R*	
Ind620	UP	51 209 239 352 353	U2i	+12308 <i>Hinf</i> I; -11465 <i>Tru</i> I
Ind616	UP	51 93TA 154 206AC 230 311	U2i	+12308 <i>Hinf</i> I; -11465 <i>Tru</i> I
Ind622	UP	184 189 223 300	M*	+10394 <i>Ddel</i> ; +10397 <i>Alu</i> I
Ind618	UP	189 223 294	L3a	
Ind612	UP	51 209 239 352 353	U2i	+12308 <i>Hinf</i> I; -11465 <i>Tru</i> I
Ind614	UP	51 92 168	U2i	+12308 <i>Hinf</i> I; -11465 <i>Tru</i> I
Ind613	UP	256 309 318AT	U7	+12308 <i>Hinf</i> I; -11465 <i>Tru</i> I
Ind619	UP	51 209 239 352 353	U2i	+12308 <i>Hinf</i> I; -11465 <i>Tru</i> I
Ind617	UP	93 223 266 304	R1	
Ind615	UP	51 207 227	U2i	+12308 <i>Hinf</i> I; -11465 <i>Tru</i> I
Ind604	UP	126 223	M1	+10394 <i>Ddel</i> ; +10397 <i>Alu</i> I
Ind605	UP	51 206AC 242 291 311	U2i	+12308 <i>Hinf</i> I; -11465 <i>Tru</i> I
Ind606	UP	71	R*	
Ind607	UP	86 223 335	M*	+10394 <i>Ddel</i> ; +10397 <i>Alu</i> I
Ind608	UP	93 129 223	M4	+10394 <i>Ddel</i> ; +10397 <i>Alu</i> I
Ind388	Guj	93 188 223 231 318	M5	+10394 <i>Ddel</i> ; +10397 <i>Alu</i> I
Ind610	UP	92 126 223	M1	+10394 <i>Ddel</i> ; +10397 <i>Alu</i> I
Ind538	UP	93 172 304 362	R*	
Ind429	Mah	153	R*	
Ind404	Pun	179 227 245 266 278 362	R*	
Ind600	UP	129 192 213 223	M4	+10394 <i>Ddel</i> ; +10397 <i>Alu</i> I
Ind601	UP	51 172 209 239 352 353	U2i	+12308 <i>Hinf</i> I; -11465 <i>Tru</i> I
Ind602	UP	111 192 223 275	M*	+10394 <i>Ddel</i> ; +10397 <i>Alu</i> I
Ind603	UP	189 223 254 270 311	M*	+10394 <i>Ddel</i> ; +10397 <i>Alu</i> I
Ind609	UP	189 304	F	-12704 <i>Hinc</i> II
Ind438	Kash	223 278	M*	+10394 <i>Ddel</i> ; +10397 <i>Alu</i> I
Ind611	UP	266 304 311 355 356	R1	
Ind508	UP	51 206AC 230 304 311	U2i	+12308 <i>Hinf</i> I; -11465 <i>Tru</i> I
Ind537	UP	316	H	-00073 <i>Alw</i> 44I; -7025 <i>Alu</i> I
Ind590	UP	51 209 239 352 353	U2i	+12308 <i>Hinf</i> I; -11465 <i>Tru</i> I
Ind498	UP	51 206AC 230 261 304 311	U2i	+12308 <i>Hinf</i> I; -11465 <i>Tru</i> I
Ind499	UP	129 213 249	M4	+10394 <i>Ddel</i> ; +10397 <i>Alu</i> I
Ind509	UP	51 82 92 189 325	R*	
Ind502	UP	51 209 239 352 353	U2i	+12308 <i>Hinf</i> I; -11465 <i>Tru</i> I
Ind503	UP	134 356	U4	+12308 <i>Hinf</i> I; -11465 <i>Tru</i> I
Ind506	UP	223	M*	+10394 <i>Ddel</i> ; +10397 <i>Alu</i> I
Ind540	UP	189 192 223 299	M*	+10394 <i>Ddel</i> ; +10397 <i>Alu</i> I
Ind539	UP	126 192 223	M1	+10394 <i>Ddel</i> ; +10397 <i>Alu</i> I
Ind541	UP	51 206AC 242 291 311	U2i	+12308 <i>Hinf</i> I; -11465 <i>Tru</i> I
Ind500	UP	48 93 129 218 223 243	M4	+10394 <i>Ddel</i> ; +10397 <i>Alu</i> I
Ind591	UP	223 286	M*	+10394 <i>Ddel</i> ; +10397 <i>Alu</i> I
Ind592	UP	223	M*	+10394 <i>Ddel</i> ; +10397 <i>Alu</i> I
Ind593	UP	223 318AT	M*	+10394 <i>Ddel</i> ; +10397 <i>Alu</i> I
Ind596	UP	126 189 223 344	M1	+10394 <i>Ddel</i> ; +10397 <i>Alu</i> I
Ind594	UP	134 356	U4	+12308 <i>Hinf</i> I; -11465 <i>Tru</i> I
Ind595	UP	51 206AC 230 304 311	U2i	+12308 <i>Hinf</i> I; -11465 <i>Tru</i> I
Ind597	UP	129 189 260 362	F	-12704 <i>Hinc</i> II
Ind598	UP	51 93 223 304	M*	+10394 <i>Ddel</i> ; +10397 <i>Alu</i> I
Ind599	UP	187 223	M*	+10394 <i>Ddel</i> ; +10397 <i>Alu</i> I

Table S1 cont.

Sample number	Origin	HVS-I sequence polymorphisms*	Haplogroup	RFLP polymorphisms*
Ind505	UP	214	H	-00073Alw44I; -7025Alul
Pak454	Pak	126 154 223 239	M1	+10394Ddel; +10397Alul
Ind455	Ben	129 140 223 271	M4	+10394Ddel; +10397Alul
Ind456	Ori	51 169 234 278	U2i	+12308 HinfI; -11465 Tru1I
Ind458	UP	129 169 223	I	+10237HphI; +10394Ddel; +8249Avall; -1715Ddel
Ind459	Tamil	189 223 278 362	M*	+10394Ddel; +10397Alul
Ind462	UP	184 223 241 311	M3	+10394Ddel; +10397Alul
Ind464	Bih	126 223 247	M1	+10394Ddel; +10397Alul
Ind465	Bang	129 209 223	M4	+10394Ddel; +10397Alul
Ind466	Mah	51 209 239 244 352 353	U2i	+12308 HinfI; -11465 Tru1I
Ind467	AP	129 362	R*	
Ind489	Mah	172 278	R*	
Ind491	UP	129 362 365	R*	
Ind487	Raj	71 293	R*	
Ind488	Guj	223	M*	+10394Ddel; +10397Alul
Ind492	UP	126 192 223	M1	+10394Ddel; +10397Alul
Ind493	UP	129 169 223	I	+10237HphI; +10394Ddel; +8249Avall; -1715Ddel
Ind494	UP	129 223	M4	+10394Ddel; +10397Alul
Ind495	UP	223 319	M2	+10394Ddel; +10397Alul
Ind496	UP	51 168 172 192 243 287	U2i	+12308 HinfI; -11465 Tru1I
Ind497	UP	223	M*	+10394Ddel; +10397Alul
Ind264	Guj	51 145 189 271 300	R*	
Ind431	AP	refer	R*	
Ind426	UP	145 223 261 311	M3	+10394Ddel; +10397Alul
Pak260	Pak	refer	H	-00073Alw44I; -7025Alul
Ind423	Bih	223 368	M*	+10394Ddel; +10397Alul
Ind428	Tamil	217 243	R*	
Pak267	Pak	223 362	M*	+10394Ddel; +10397Alul
Ind507	UP	126 181 209	pTJ	
Pak451	Pak	129 223 264	M4	+10394Ddel; +10397Alul
Pak403	Pak	92	Ö	-00073Alw44I
Ind504	UP	129 266 290 318 320 362	M4	+10394Ddel; +10397Alul
Pak453	Pak	51 154 206AC 230 311	U2i	+12308 HinfI; -11465 Tru1I
Pak261	Pak	318AT	U7	+12308 HinfI; -11465 Tru1I
Ind439	Ori	189 223 260 294 295 325	M	+10394Ddel; +10397Alul
Ind510	UP	309 318AT	U7	+12308 HinfI; -11465 Tru1I
Ind387	Guj	266 304	R1	
Ind433	Mah	189 249	U1	+12308 HinfI; -11465 Tru1I
Pak425	Hary	51 93TA 154 178 206AC 230 261 311	U2i	+12308 HinfI; -11465 Tru1I
Pak262	Pak	129 223 298 311 327	M-C	+10394Ddel; +10397Alul; +13262Alul
Ind501	UP	189 223 274 319 320	M2b	+10394Ddel; +10397Alul
Lam1	AP	189 223 274 311 319	M2b	+10394Ddel; +10397Alul
Lam2	AP	129 223	M4	+10394Ddel; +10397Alul
Lam3	AP	223 311	M3	+10394Ddel; +10397Alul
Lam4	AP	189 233 304 325 362	R*	
Lam5	AP	223	M*	+10394Ddel; +10397Alul
Lam6	AP	223 270 311 319 352	M2a	+10394Ddel; +10397Alul
Lam7	AP	189 223 278	X	-1715Ddel; +14465Accl
Lam8	AP	129 223	M4	+10394Ddel; +10397Alul
Lam9	AP	223 270 274 319 352	M2b	+10394Ddel; +10397Alul
Lam10	AP	292	R*	
Lam11	AP	223 270 274 319 352	M2b	+10394Ddel; +10397Alul
Lam12	AP	126 275 294 296 325	T	+13366BamHI; +15606Alul
Lam13	AP	93 223	M*	+10394Ddel; +10397Alul
Lam14	AP	93 192 223 311	M3	+10394Ddel; +10397Alul
Lam15	AP	167 172 318AT	U7	+12308 HinfI; -11465 Tru1I
Lam16	AP	51 172 206AC 286	U2i	+12308 HinfI; -11465 Tru1I
Lam17	AP	126 129 223	M1	+10394Ddel; +10397Alul
Lam18	AP	126 344	M1	+10394Ddel; +10397Alul
Lam19	AP	93 192 223 311	M3	+10394Ddel; +10397Alul
Lam20	AP	129 144TA 223 362	M4	+10394Ddel; +10397Alul

Table S1 cont.

Sample number	Origin	HVS-I sequence polymorphisms*	Haplogroup	RFLP polymorphisms*
Lam21	AP	260 261 319 362	R*	
Lam22	AP	129 144TA 223 362	M4	+10394Ddel; +10397Alul
Lam23	AP	86 129 223 249 259 311	M3	+10394Ddel; +10397Alul
Lam24	AP	129 144TA 223 362	M4	+10394Ddel; +10397Alul
Lam25	AP	166 223 311 359	M3	+10394Ddel; +10397Alul
Lam26	AP	188 189 223 231 356 362	M5	+10394Ddel; +10397Alul
Lam27	AP	188 223 231 362	M5	+10394Ddel; +10397Alul
Lam28	AP	223 311 316 355	M3	+10394Ddel; +10397Alul
Lam29	AP	129 144TA 223 362	M4	+10394Ddel; +10397Alul
Lam30	AP	189 192 223 292	W	+8249Avall; -8994HaeIII
Lam31	AP	260 261 319 362	R*	
Lam32	AP	223	M*	+10394Ddel; +10397Alul
Lam33	AP	69 126 145 222 261	J	-13704BstNI; +10394Ddel
Lam34	AP	129 144TA 223 362	M4	+10394Ddel; +10397Alul
Lam35	AP	223 270 319	M2a	+10394Ddel; +10397Alul
Lam36	AP	188 223 231 362	M5	+10394Ddel; +10397Alul
Lam37	AP	51 179 234 247 278 /240 /1	U2i	+12308 HinfI; -11465 Tru1I
Lam38	AP	188 223 231 362	M5	+10394Ddel; +10397Alul
Lam39	AP	75 92 189 223	M*	+10394Ddel; +10397Alul
Lam40	AP	223 318AT	M*	+10394Ddel; +10397Alul
Lam41	AP	309 318AT	U7	+12308 HinfI; -11465 Tru1I
Lam42	AP	260 261 319 362	R*	
Lam43	AP	260 261 319 362	R*	
Lam44	AP	51 172 206AC 286	U2i	+12308 HinfI; -11465 Tru1I
Lam45	AP	223 324 357	M*	+10394Ddel; +10397Alul
Lam46	AP	213 223 231 278 356 362	M5	+10394Ddel; +10397Alul
Lam47	AP	213 223 231 278 356 362	M5	+10394Ddel; +10397Alul
Lam48	AP	223 318AT	M*	+10394Ddel; +10397Alul
Lam49	AP	93 126 145 223	M1	+10394Ddel; +10397Alul
Lam50	AP	213 223 231 278 318 324 356 362	M5	+10394Ddel; +10397Alul
Lam51	AP	51 206AC 230 311	U2i	+12308 HinfI; -11465 Tru1I
Lam52	AP	188 223 231 362	M5	+10394Ddel; +10397Alul
Lam53	AP	223 270 274 292 319 352	M2b	+10394Ddel; +10397Alul
Lam54	AP	223 304 311	M3	+10394Ddel; +10397Alul
Lam55	AP	223 311 316 355	M3	+10394Ddel; +10397Alul
Lam56	AP	51 189 218 292	R	
Lam57	AP	51 172 206AC 286	U2i	+12308 HinfI; -11465 Tru1I
Lam58	AP	260 261 294 319 362	R*	
Lam59	AP	75 92 189 223	M*	+10394Ddel; +10397Alul
Lam60	AP	51 172 206AC 286	U2i	+12308 HinfI; -11465 Tru1I
Lam61	AP	260 270	U5	+12308 HinfI; -11465 Tru1I
Lam62	AP	126 169 223	M1	+10394Ddel; +10397Alul
Lam63	AP	223 318AT	M*	+10394Ddel; +10397Alul
Lam64	AP	184 274	H	-00073Alw44I; -7025Alul
Lam65	AP	223	M*	+10394Ddel; +10397Alul
Lam66	AP	129 223 304 311	I	+10237HphI; +10394Ddel; +8249Avall; -1715Ddel
Lam67	AP	51 234	U2i	+12308 HinfI; -11465 Tru1I
Lam68	AP	154 221	R*	
Lam69	AP	185 223 270 274 319 352	M2b	+10394Ddel; +10397Alul
Lam70	AP	178 223 288	M*	+10394Ddel; +10397Alul
Lam71	AP	129 223 335	M4	+10394Ddel; +10397Alul
Lam72	AP	188 223 231 362	M5	+10394Ddel; +10397Alul
Lam73	AP	129 144TA 223 362	M4	+10394Ddel; +10397Alul
Lam74	AP	129 223 304 311	I	+10237HphI; +10394Ddel; +8249Avall; -1715Ddel
Lam75	AP	304 311	R*	
Lam76	AP	185 189 223 270 274 319	M2b	+10394Ddel; +10397Alul
Lam77	AP	223 292	W	+8249Avall; -8994HaeIII
Lam78	AP	111 189 223 327 330	M*	+10394Ddel; +10397Alul
Lam79	AP	221	R*	
Lam80	AP	129 242 356	Ö	-00073Alw44I
Lam81	AP	223 311	M3	+10394Ddel; +10397Alul
Lam82	AP	147CA 172 223 248 294 320 355	O	+10394Ddel; +10237HphI; -1715Ddel

Table S1 cont.

Sample number	Origin	HVS-I sequence polymorphisms*	Haplogroup	RFLP polymorphisms*
Lam83	AP	93 223	M*	+10394Ddel; +10397Alul
Lam84	AP	148 189 223 270 274 319	M2b	+10394Ddel; +10397Alul
Lam85	AP	92 223 362	M*	+10394Ddel; +10397Alul
Lam86	AP	129 223 304 311	M*	+10394Ddel; +10397Alul
Lob2	Pun	189 192 223 260 291 292 325 355	W	+8249Avall; -8994HaeIII
Lob3	Pun	185 223 289 311 362	M*	+10394Ddel; +10397Alul
Lob4	Pun	93 223	M*	+10394Ddel; +10397Alul
Lob5	Pun	223 263	M*	+10394Ddel; +10397Alul
Lob6	Pun	51 234 247 304	U2i	+12308 HinfI; -11465 Tru1I
Lob7	Pun	189 283 304	F	-12704HincII
Lob8	Pun	188 223 231 362	M5	+10394Ddel; +10397Alul
Lob9	Pun	145 189 223 292 320	W	+8249Avall; -8994HaeIII
Lob10	Pun	223 289	M*	+10394Ddel; +10397Alul
Lob11	Pun	223 234 304	M*	+10394Ddel; +10397Alul
Lob12	Pun	223 263	M*	+10394Ddel; +10397Alul
Lob14	Pun	223 293	M*	+10394Ddel; +10397Alul
Lob17	Pun	223 292	W	+8249Avall; -8994HaeIII
Lob19	Pun	92 189 298 299	F	-12704HincII
Lob21	Pun	189 249	U1	+12308 HinfI; -11465 Tru1I
Lob24	Pun	292	R*	
Lob26	Pun	126 294 296 304	T	+13366BamHI; +15606Alul
Lob27	Pun	172 223 362	M-D	+10394Ddel; +10397Alul
Lob28	Pun	185 223 289 311 362	M*	+10394Ddel; +10397Alul
Lob30	Pun	126 223 261 344	M1	+10394Ddel; +10397Alul
Lob33	Pun	129 223 304	M4	+10394Ddel; +10397Alul
Lob35	Pun	223 270 319 352	M2a	+10394Ddel; +10397Alul
Lob36	Pun	223	M*	+10394Ddel; +10397Alul
Lob38	Pun	126 223 261 344	M1	+10394Ddel; +10397Alul
Lob40	Pun	126 223 261 344	M1	+10394Ddel; +10397Alul
Lob41	Pun	223 290 292	W	+8249Avall; -8994HaeIII
Lob42	Pun	223	M*	+10394Ddel; +10397Alul
Lob43	Pun	223 293	M*	+10394Ddel; +10397Alul
Lob45	Pun	223	M*	+10394Ddel; +10397Alul
Lob46	Pun	223	M*	+10394Ddel; +10397Alul
Lob47	Pun	292	R*	
Lob52	Pun	223 231 234 311 356 362	M5	+10394Ddel; +10397Alul
Lob54	Pun	223 231 311 356 362	M5	+10394Ddel; +10397Alul
Lob55	Pun	292	R*	
Lob56	Pun	223 231 311 356 362	M5	+10394Ddel; +10397Alul
Lob57	Pun	51 206AC	U2i	+12308 HinfI; -11465 Tru1I
Lob58	Pun	223 231 311 356 362	M5	+10394Ddel; +10397Alul
Lob59	Pun	223 256	M*	+10394Ddel; +10397Alul
Lob60	Pun	129 189 223	M4	+10394Ddel; +10397Alul
Lob62	Pun	292	R*	
Lob63	Pun	217	R*	
Lob64	Pun	48 129 218 223	M4	+10394Ddel; +10397Alul
Lob65	Pun	223 290 292	W	+8249Avall; -8994HaeIII
Lob67	Pun	126 294 296 304	T	+13366BamHI; +15606Alul
Lob68	Pun	223 289	M*	+10394Ddel; +10397Alul
Lob69	Pun	145 189 223 292 320	W	+8249Avall; -8994HaeIII
Lob70	Pun	92 189 298 299	F	-12704HincII
Lob71	Pun	223 292	W	+8249Avall; -8994HaeIII
Lob72	Pun	129 223 311	M4	+10394Ddel; +10397Alul
Lob74	Pun	71 189 278	R*	
Lob77	Pun	126 294 296 304	T	+13366BamHI; +15606Alul
Lob78	Pun	217	R*	
Lob79	Pun	92 189 298 299	F	-12704HincII
Lob81	Pun	92 189 298 299	F	-12704HincII
Lob86	Pun	292	R*	
Lob89	Pun	223 290 292	W	+8249Avall; -8994HaeIII
Lob91	Pun	71 93	R*	
Lob92	Pun	223 234 311	M*	+10394Ddel; +10397Alul

Table S1 cont.

Sample number	Origin	HVS-I sequence polymorphisms*	Haplogroup	RFLP polymorphisms*
Lob94	Pun	223 270 319	M2a	+10394Ddel; +10397Alul
Lob95	Pun	223 270 319	M2a	+10394Ddel; +10397Alul
Lob97	Pun	111 184 189 223 274 295	M*	+10394Ddel; +10397Alul
Lob101	Pun	223 318AT	M*	+10394Ddel; +10397Alul
Bog1	UP	124 179 189 223 249 294	M*	+10394Ddel; +10397Alul
Bog2	UP	223 241	M*	+10394Ddel; +10397Alul
Bog3	UP	223 318AT	M*	+10394Ddel; +10397Alul
Bog4	UP	223	M*	+10394Ddel; +10397Alul
Bog5	UP	129 223 311	M4	+10394Ddel; +10397Alul
Bog6	UP	93 223	M*	+10394Ddel; +10397Alul
Bog8	UP	95 223 249 359	M*	+10394Ddel; +10397Alul
Bog10	UP	129 362	R*	
Bog12	UP	223 270 319 352	M2a	+10394Ddel; +10397Alul
Bog14	UP	111CA 223	M*	+10394Ddel; +10397Alul
Bog15	UP	refer	R*	
Bog17	UP	126 223 311	M1	+10394Ddel; +10397Alul
Bog20	UP	129 189 223 325	M4	+10394Ddel; +10397Alul
Bog21	UP	95 223 249 359	M*	+10394Ddel; +10397Alul
Bog22	UP	140 189 223 293 311	M*	+10394Ddel; +10397Alul
Bog23	UP	51 93TA 154 206AC 230 311	U2i	+12308 HinfI; -11465 Tru1I
Bog29	UP	126 176 181 209	pTJ	
Bog99	UP	92 126 223 286	M1	+10394Ddel; +10397Alul
UP106	UP	175 223 234	M*	+10394Ddel; +10397Alul
UP111	UP	51 129 209 239 291 325 352 353	U2i	+12308 HinfI; -11465 Tru1I
UP183	UP	223	M*	+10394Ddel; +10397Alul
Tha2	UP	51 223 298 327	M-C	+10394Ddel; +10397Alul; +13262Alul
Tha4	UP	114 223 294 318 362	M-D	+10394Ddel; +10397Alul; -5176Alul
Tha5	UP	223 362	M-D	+10394Ddel; +10397Alul; -5176Alul
Tha15	UP	refer	R*	
Tha25	UP	93 126 163 186 189 294	T1	+13366BamHI; +15606Alul
Tha36	UP	71	R*	
Tha46	UP	223 302	M*	+10394Ddel; +10397Alul
Tha47	UP	356	U4	+12308 HinfI; -11465 Tru1I
Tha48	UP	129 189 241 266 304	R1	
Tha49	UP	126 223 368	M1	+10394Ddel; +10397Alul
Tha50	UP	93 129 223	M4	+10394Ddel; +10397Alul
Tha51	UP	51 209 239 352 353	U2i	+12308 HinfI; -11465 Tru1I
Ksh1	Kash	189 304	F	-12704HincII
Ksh2	Kash	309 325	Ö	-00073Alw44I
Ksh3	Kash	refer	Ö	-00073Alw44I
Ksh4	Kash	189 304	F	-12704HincII
Ksh5	Kash	129 213 362	R*	
Ksh6	Kash	69 126 145 261	J	-13704BstNI; +10394Ddel
Ksh7	Kash	189 304	F	-12704HincII
Ksh8	Kash	189 304	F	-12704HincII
Ksh9	Kash	309 325 362	Ö	-00073Alw44I
Ksh10	Kash	148 175 223 292	W	+8249Avall; -8994HaeIII
Ksh11	Kash	refer	H	-00073Alw44I; -7025Alul
Ksh12	Kash	51 206AC 291	U2i	+12308 HinfI; -11465 Tru1I
Ksh13	Kash	145 176 223 311	M3	+10394Ddel; +10397Alul
Ksh14	Kash	188 223 231 362	M5	+10394Ddel; +10397Alul
Ksh15	Kash	145 239 241 325	R*	
Ksh16	Kash	51 209 239 352 353	U2i	+12308 HinfI; -11465 Tru1I
Ksh17	Kash	192 223 300 316	M*	+10394Ddel; +10397Alul
Ksh18	Kash	188 223 231 362	M5	+10394Ddel; +10397Alul
APx	AP	223 274 311 319	M2b	+10394Ddel; +10397Alul
BV10	AP	146 311	H	-7025Alul
BV12	AP	179 227 245 266 278 362	R*	
BV14	AP	51 234		+10394Ddel; +10397Alul
BV15	AP	223 257	M*	+10394Ddel; +10397Alul
BV16	AP	126 223 311		+12308 HinfI
BV19	AP	126 223 271	M1	+10394Ddel; +10397Alul

Table S1 cont.

Sample number	Origin	HVS-I sequence polymorphisms*	Haplogroup	RFLP polymorphisms*
BV2	AP	51 172 286 291 206AC	U2i	+12308 <i>Hinf</i> I
BN22	AP	189 197CA 223 287 327 330	M*	+10394Ddel; +10397Alul
BN24	AP	129 223 311	M4	+10394Ddel; +10397Alul
BV25	AP	93 224 311	U-K	P+12308 <i>Hinf</i> I; +10394Ddel; -9025HaeII
BN26	AP	126 266 304 309 325 356	R	
BN28	AP	129 223 286 291	M4	+10394Ddel; +10397Alul
BV29	AP	69 274 280 318AT	U7	+12308 <i>Hinf</i> I
BN30	AP	223	M*	+10394Ddel; +10397Alul
BN31	AP	172 223 243 270 319 352	M2a	+10394Ddel; +10397Alul
BV32	AP	223 324 362	M*	+10394Ddel; +10397Alul
BV33	AP	223 274	M*	+10394Ddel; +10397Alul
BV34	AP	129 223	M4	+10394Ddel; +10397Alul
BV35	AP	126 223 311	M1	+10394Ddel; +10397Alul
BV36	AP	104 189 223 243 319 362	M*	+10394Ddel; +10397Alul
BV38	AP	178 223 256	M*	+10394Ddel; +10397Alul
BV4	AP	223	M*	+10394Ddel; +10397Alul
BV40	AP	51 172 206AC	U2i	+12308 <i>Hinf</i> I
BV42	AP	145 176 223 261 311	M3	+10394Ddel; +10397Alul
BV43	AP	304 311	R*	
BV44	AP	207 309 318AT	U7	+12308 <i>Hinf</i> I
BV45	AP	93 223	M*	+10394Ddel; +10397Alul
BV47	AP	223 240AC 274 311 319	M2b	+10394Ddel; +10397Alul
BV48	AP	37 51 234 325	U2i	+12308 <i>Hinf</i> I
BV49	AP	207 309 318AT 352	U7	+12308 <i>Hinf</i> I
BV5	AP	126 223 271	M1	+10394Ddel; +10397Alul
BN50	AP	184 214 357	R*	
BN51	AP	223 274 319	M2b	+10394Ddel; +10397Alul
BN52	AP	172 278		+10394Ddel; +10397Alul
BN53	AP	126 185 223		
BN54	AP	311 320		+10394Ddel; +10397Alul
BN55	AP	54AT 223 325		-7025Alul
BN56	AP	223 325	M*	+10394Ddel; +10397Alul
BN57	AP	93 223 278	M*	+10394Ddel; +10397Alul
BN9	AP	223 304	M*	+10394Ddel; +10397Alul
J11	AP	223 256 294	M*	+10394Ddel; +10397Alul
J13	AP	51 86 209 239 354 362	U2i	+12308 <i>Hinf</i> I
J15	AP	92 145 223 261 311	M3	+10394Ddel; +10397Alul
J21	AP	92 145 223 261 311	M3	+10394Ddel; +10397Alul
J24	AP	223	M*	+10394Ddel; +10397Alul
J25	AP	93 104 234 243 244	M*	+10394Ddel; +10397Alul
J26	AP	93 104 108 234 243 244	M*	+10394Ddel; +10397Alul
KT1	AP	93 223	M*	+10394Ddel; +10397Alul
KT10	AP	126 294 296 325	T	
KT11	AP	184 189 223 288TA 300	M*	+10394Ddel; +10397Alul
KT12	AP	223 274 301	M*	+10394Ddel; +10397Alul
KT13	AP	51 93TA 154 206AC 230 311	U2i	+12308 <i>Hinf</i> I
KT14	AP	223 270 274 319 352	M2b	+10394Ddel; +10397Alul
KT15	AP	189 223 270 278	M*	+10394Ddel; +10397Alul
KT16	AP	126 223 309	M1	+10394Ddel; +10397Alul
KT17	AP	188 189 223 231 355 362	M5	+10394Ddel; +10397Alul
KT2	AP	129 213 319 362	R*	
KTK22	AP	42 51 93 179 234 240AC	U2i	+12308 <i>Hinf</i> I
KTK23	AP	69 126 145 172 222 261	J	
KTK24	AP	refer	U*	+12308 <i>Hinf</i> I
KTK25	AP	93 223	M*	+10394Ddel; +10397Alul
KTK27	AP	266 304 311 356	R1	
KTK28	AP	223 318AT	M*	+10394Ddel; +10397Alul
KTK30	AP	51 93TA 154 206AC 230 311	U2i	+12308 <i>Hinf</i> I
KT32	AP	223 263	M*	+10394Ddel; +10397Alul
KTK33	AP	223 270 274 292 319 352	M2b	+10394Ddel; +10397Alul
KTK34	AP	refer	H	-7025Alul
KTK35	AP	292	R*	

Table S1 cont.

Sample number	Origin	HVS-I sequence polymorphisms*	Haplogroup	RFLP polymorphisms*
KTK36	AP	129 223	M4	+10394Ddel; +10397AluI
KT38	AP	51 189 223 274 319 320 362	M2b	+10394Ddel; +10397AluI
KT39	AP	172 278 344CA	R*	
KTK4	AP	223 304	M*	+10394Ddel; +10397AluI
KT40	AP	189 223 300	M*	+10394Ddel; +10397AluI
KT41	AP	179 223 294 319 356	M*	+10394Ddel; +10397AluI
KTK42	AP	51 230 311 206AC	U2i	+12308 HinfI
KT43	AP	93 223 258AC 274	M*	+10394Ddel; +10397AluI
KT44	AP	129 223 270 274 319 352	M2b	+10394Ddel; +10397AluI
KT45	AP	129 223 264 265AC	M4	+10394Ddel; +10397AluI
KTK46	AP	223	M*	+10394Ddel; +10397AluI
KT47	AP	86 221 278	R*	
KT48	AP	188 223 231 362	M5	+10394Ddel; +10397AluI
KT49	AP	223 234	M*	+10394Ddel; +10397AluI
KTK5	AP	93 189 223 232 270 311	M*	+10394Ddel; +10397AluI
KT50	AP	37 51 154 206AC 230 311	U2i	+12308 HinfI
KT51	AP	126 223 312	M1	+10394Ddel; +10397AluI
KTK52	AP	189 192 223 300	M*	+10394Ddel; +10397AluI
KT54	AP	223 234	M*	+10394Ddel; +10397AluI
KT55	AP	223	M*	+10394Ddel; +10397AluI
KTK57	AP	223 318AT	M*	+10394Ddel; +10397AluI
KT59	AP	119 193 223 303GT	M*	+10394Ddel; +10397AluI
KTK6	AP	93 223	M*	+10394Ddel; +10397AluI
KTK60	AP	260 261 319 362	R*	
KTK64	AP	111 223	M*	+10394Ddel; +10397AluI
KTK66	AP	111 223	M*	+10394Ddel; +10397AluI
KT67	AP	274	R*	
KT68	AP	92TA 145 223 227 245 290CA 291		
KTK69	AP	75 223 270 274 319 352	M2b	+10394Ddel; +10397AluI
KT8	AP	304 311	R*	
KT9	AP	169 172 223	M*	+10394Ddel; +10397AluI
KS1	AP	223	M*	+10394Ddel; +10397AluI
KS10	AP	179 223 294	M*	+10394Ddel; +10397AluI
KS11	AP	126 223 265	M1	+10394Ddel; +10397AluI
KS2	AP	270	R*	
KS3	AP	270	R*	
KS4	AP	270	R*	
KS7	AP	223	M*	+10394Ddel; +10397AluI
KS8	AP	179 223 294	M*	+10394Ddel; +10397AluI
KS9	AP	129 223 311	M4	+10394Ddel; +10397AluI
M11	AP	126 181 209 362	R*	
M12	AP	188 189 223 256 311	M*	+10394Ddel; +10397AluI
M13	AP	104 223 234 243 244	M*	+10394Ddel; +10397AluI
M14	AP	129 223 291	M4	+10394Ddel; +10397AluI
M15	AP	129 223 291	M4	+10394Ddel; +10397AluI
M16	AP	172 223 270 274 319 352	M2b	+10394Ddel; +10397AluI
M17	AP	172 278	R*	
M18	AP	129 223 291	M4	+10394Ddel; +10397AluI
M19	AP	189 223 248 274 291 319 320	M2b	+10394Ddel; +10397AluI
M2	AP	223 318AT	M*	+10394Ddel; +10397AluI
M21	AP	223 301	M*	+10394Ddel; +10397AluI
M22	AP	223 231 362	M5	+10394Ddel; +10397AluI
M23	AP	93 145 189 223 290 312 355 381	M*	+10394Ddel; +10397AluI
M24	AP	129 223 264 265AC	M4	+10394Ddel; +10397AluI
M25	AP	129 223 264 265AC	M4	+10394Ddel; +10397AluI
M26	AP	221	R*	
M27	AP	126 223	M1	+10394Ddel; +10397AluI
M29	AP	75 92 189 223 270	M*	+10394Ddel; +10397AluI
M3	AP	37 51 234 283AC	U2i	+12308 HinfI
M30	AP	266 289 304	R1	
M31	AP	75 92 189 223 270	M*	+10394Ddel; +10397AluI
M32	AP	223	M*	+10394Ddel; +10397AluI

Table S1 cont.

Sample number	Origin	HVS-I sequence polymorphisms*	Haplogroup	RFLP polymorphisms*
M4	AP	304 311	R*	
M7	AP	145 189 192 266 304 309 325 356	R1	
M8	AP	223 318AT	M*	+10394Ddel; +10397Alul
M9	AP	126 163 186 189 294	T1	
ML1	AP	124 179 209 223 294 311 319 356	M*	+10394Ddel; +10397Alul
ML12	AP	223 324	M*	+10394Ddel; +10397Alul
ML13	AP	184 223	M*	+10394Ddel; +10397Alul
ML14	AP	93 134 223 318AT	M*	+10394Ddel; +10397Alul
ML15	AP	193 223 256	M*	+10394Ddel; +10397Alul
ML16	AP	129 179 189 223 292 294 355	M*	+10394Ddel; +10397Alul
ML17	AP	129 223	M4	+10394Ddel; +10397Alul
ML18	AP	309 318AT	U7	+12308 HinfI
ML19	AP	172 223	M*	+10394Ddel; +10397Alul
ML2	AP	189 223 327 330	M*	+10394Ddel; +10397Alul
ML20	AP	136 248 266 304 325 356	R1	
ML21	AP	223 234 318AT	M*	+10394Ddel; +10397Alul
ML23	AP	223 304	M*	+10394Ddel; +10397Alul
ML24	AP	51 114 193 278 357	M*	+12308 HinfI
ML25	AP	223 231 356 362	M5	+10394Ddel; +10397Alul
ML26	AP	223	M*	+10394Ddel; +10397Alul
ML27	AP	116AC 169 234 283 317AT 351AT	R*	
ML3	AP	256 266 304 356	R1	
ML4	AP	145 185 189 239 325	R*	
ML5	AP	136 248 266 304 325 356	R1	
ML6	AP	188 223 231 362	M5	+10394Ddel; +10397Alul
ML7	AP	266 304 311	R1	
ML8	AP	223	M*	+10394Ddel; +10397Alul
ML9	AP	189 223 270 287 296 318AT	M*	+10394Ddel; +10397Alul
R1	AP	189 218 223 274 319 320	M2b	+10394Ddel; +10397Alul
R10	AP	223 318AT	M*	+10394Ddel; +10397Alul
R11	AP	189 223 228CG 242 274 319 320 355	M2b	+10394Ddel; +10397Alul
R12	AP	189 218 223 274 319 320	M2b	+10394Ddel; +10397Alul
R13	AP	147 189 223 243 278 355 362	M*	+10394Ddel; +10397Alul
R15	AP	93 292	R*	
R16	AP	178 223 288	M*	+10394Ddel; +10397Alul
R17	AP	178 223 288	M*	+10394Ddel; +10397Alul
R18	AP	93 172 223 327	M*	+10394Ddel; +10397Alul
R19	AP	223 274 319 357	M2b	+10394Ddel; +10397Alul
R2	AP	93 301 317AT	R*	
R20	AP	93 292	R*	
R21	AP	178 223 288	M*	
R3	AP	209 223 156GT	M*	+10394Ddel; +10397Alul
R4	AP	223 318AT	M*	+10394Ddel; +10397Alul
R5	AP	189 218 223 274 319 320 228CG	M2b	+10394Ddel; +10397Alul
R6	AP	51 247 254 362	U2i	+12308 HinfI
R7	AP	156GT 184 189 223 316AT	M*	+10394Ddel; +10397Alul
R8	AP	156GT 223 318AT	M*	+10394Ddel; +10397Alul
R9	AP	48 63 129 223 362	M4	+10394Ddel; +10397Alul
VS1	AP	223 292	W	
VS10	AP	117 126 223 278	M1	+10394Ddel; +10397Alul
VS2	AP	126 223 278	M1	+10394Ddel; +10397Alul
VS3	AP	51	U2i	+12308 HinfI
VS4	AP	292	R*	
VS5	AP	51	U2i	+12308 HinfI
VS6	AP	126 223 278	M1	+10394Ddel; +10397Alul
VS7	AP	126 223 278	M1	+10394Ddel; +10397Alul
VS8	AP	51	U2i	+12308 HinfI
VS9	AP	126 223 278	M1	+10394Ddel; +10397Alul
WB15	AP	172 304	R*	
WB17	AP	104 223 234 243 244	M*	+10394Ddel; +10397Alul
WB18	AP	223	M*	
WB19	AP	92 145 223 261 311	M3	+10394Ddel; +10397Alul

Table S1 cont.

Sample number	Origin	HVS-I sequence polymorphisms*	Haplogroup	RFLP polymorphisms*
WB20	AP	266 304 311 355 356		+10394Ddel; +10397Alul
WB21	AP	155AT 356	R*	
WB22	AP	104 223 234 243 244		
WB23	AP	104 223 234 243 244	M*	+10394Ddel; +10397Alul
Y1	AP	184 223 311	M3	+10394Ddel; +10397Alul
Y11	AP	192 223 231 356 362	M5	+10394Ddel; +10397Alul
Y15	AP	126 275 294 296 325	T	
Y17	AP	51 172 209 224 239 352 353	U2i	+12308 HinfI
Y18	AP	126 223 311	M1	+10394Ddel; +10397Alul
Y19	AP	129 242 356	R*	
Y2	AP	129 193 223	M4	+10394Ddel; +10397Alul
Y20	AP	189 222 223 256 274 319 320	M2b	+10394Ddel; +10397Alul
Y21	AP	223	M*	+10394Ddel; +10397Alul
Y22	AP	129 193 223	M4	+10394Ddel; +10397Alul
Y23	AP	223	M*	+10394Ddel; +10397Alul
Y24	AP	193 223	M*	+10394Ddel; +10397Alul
Y25	AP	129 242 356	R*	
Y26	AP	93 223	M*	+10394Ddel; +10397Alul
Y27	AP	86 172 189 223 227 278 362	M-E	+10394Ddel; +10397Alul; -7598Hhal
Y28	AP	223 304	M*	+10394Ddel; +10397Alul
Y29	AP	223	M*	+10394Ddel; +10397Alul
Y3	AP	223 304	M*	+10394Ddel; +10397Alul
Y30	AP	189 231 270 319 362	R*	
Y31	AP	129 242 356	R*	
Y32	AP	104 223 234 243 244 354	M*	+10394Ddel; +10397Alul
Y33	AP	172 189 228CG 270 278 296 355	R*	
Y34	AP	126 294 296 304	T	
Y35	AP	309 318 AT	U7	+12308 HinfI
Y36	AP	51 234	U2i	+12308 HinfI
Y37	AP	193 223	M*	+10394Ddel; +10397Alul
Y38	AP	188 223 231 362	M5	+10394Ddel; +10397Alul
Y39	AP	145 221 260 261 311 319 343 362	R*	
Y4	AP	223	M*	+10394Ddel; +10397Alul
Y40	AP	213 223 231 356 362	M5	+10394Ddel; +10397Alul
Y41	AP	304 311	R*	
Y42	AP	223 270 362 381	M*	+10394Ddel; +10397Alul
Y43	AP	266 304 310 311 356	R1	
Y44	AP	221	R*	
Y45	AP	172 278	R*	
Y46	AP	126 223	M1	+10394Ddel; +10397Alul
Y47	AP	184 223 311	M3	+10394Ddel; +10397Alul
Y49	AP	129 193 223	M4	+10394Ddel; +10397Alul
Y51	AP	51 126 179 227 234 240AC	U2i	+12308 HinfI
Y52	AP	223 270 319 352	M2a	+10394Ddel; +10397Alul
Y53	AP	93 223 243	M*	+10394Ddel; +10397Alul
Y54	AP	189 212 223 228CG 270 327 330	M*	+10394Ddel; +10397Alul
Y55	AP	86 172 189 223 227 278 362	M-E	+10394Ddel; +10397Alul; -7598Hhal
Y56	AP	126 275 294 296 325 357	T	
Y6	AP	129 193 223	M4	+10394Ddel; +10397Alul
Y7	AP	129 213 320 362	R*	
Y9	AP	250 260 261 293 311 319	R*	

All mtDNAs were compared to the R node which equals [S14] in HVS-I sequence but differs from it by the presence of an *Alw44I* site at nucleotide position (np) 00073 and the presence of an *Alul* site at np 7025. Origin of the samples: AP, Andhra Pradesh; Bang, Bangladesh; Ben, Western Bengal; Bih, Bihar; Guj, Gujarat; Hary, Haryana; Kash, Kashmir; Mah, Maharashtra; Ori, Orissa; Pak, Pakistan; Pun, Punjab; Raj, Rajasthan; Tamil, Tamil-Nadu; UP, Uttar Pradesh. For those data sets, published by others, with which the Indian data were compared, see Supplementary material and methods section.