

Mitochondrial DNA Diversity in Indigenous Populations of the Southern Extent of Siberia, and the Origins of Native American Haplogroups

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Summary

In search of the ancestors of Native American mitochondrial DNA (mtDNA) haplogroups, we analyzed the mtDNA of 531 individuals from nine indigenous populations in Siberia. All mtDNAs were subjected to high-resolution RFLP analysis, sequencing of the control-region hypervariable segment I (HVS-I), and surveyed for additional polymorphic markers in the coding region. Furthermore, the mtDNAs selected according to haplogroup/subhaplogroup status were completely sequenced. Phylogenetic analyses of the resulting data, combined with those from previously published Siberian arctic and sub-arctic populations, revealed that remnants of the ancient Siberian gene pool are still evident in Siberian populations, suggesting that the founding haplotypes of the Native American A-D branches originated in different parts of Siberia. Thus, lineage A complete sequences revealed in the Mansi of the Lower Ob and the Ket of the Lower Yenisei belong to A1, suggesting that A1 mtDNAs occasionally found in the remnants of hunting-gathering populations of northwestern and northern Siberia belonged to a common gene pool of the Siberian progenitors of Paleoindians. Moreover, lineage B1, which is the most closely related to the American B2, occurred in the Tubalar and Tuvan inhabiting the territory between the upper reaches of the Ob River in the west, to the Upper Yenisei region in the east. Finally, the sequence variants of haplogroups C and D, which are most similar to Native American C1 and D1, were detected in the Ulchi of the Lower Amur. Overall, our data suggest that the immediate ancestors of the Siberian/Beringian migrants who gave rise to ancient (pre-Clovis) Paleoindians have a common origin with aboriginal people of the area now designated the Altai-Sayan Upland, as well as the Lower Amur/Sea of Okhotsk region.

Keywords : mtDNA variation, native Siberians, Native Americans

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Introduction

The diffusion of the first modern humans in Siberia appears to have been restricted to the regions south of the 55°N parallel, where multiple Upper Paleolithic sites have been found and dated to 43,000–39,000 YBP (reviewed by Vasil'ev *et al.* 2002). Environmental conditions permitting human entry into the New World apparently existed shortly before and after 30,000 YBP

and 13,000 YBP, when many areas in Siberia/Beringia remained ice-free and may have been periodically connected to the North American interior through an ice-free corridor, which was repeatedly buried under continental glaciers (Wright, 1991; West, 1996; Goebel *et al.* 2003; Pitulko *et al.* 2004). In addition, the first Americans could have spread along the Pacific Coast among the islands and bays of Alaska and Canada, at a time when the North American interior was an inhospitable, ice-covered wasteland (Dalton, 2003)

The affinity of southern Siberians with Native Americans is supported by anthropological (Kozintsev *et al.* 1999), dental (Turner, 1994) and genetic evidence, including paternally inherited Y chromosome polymorphisms (Lell *et al.* 2002) and autosomal *HLA* class II gene(s) variation (Uinuk-ool *et al.* 2002; Volodko *et al.* 2003). Intriguingly, the *HLA* class II gene frequencies separate Siberian/Asian/Native American populations into two clusters, one of which encompasses nearly all of the Siberian and all of the Native American populations, while the other consists of central, eastern and southeastern Asians (Uinuk-ool *et al.* 2002). Thus, the debate continues about precisely where in Siberia the ancestors of Native Americans arose, and when and how they spread into the non-glaciated interior of Alaska, subsequently migrating southward into the American West.

Early studies of Native American mtDNA variation have shown that all Native American mtDNAs belong to haplogroups A, B, C, D and X, and that some of these haplogroups are also common along the northern Pacific Rim (Torroni *et al.* 1993a,b; Ward *et al.* 1993; Forster *et al.* 1996; Starikovskaya *et al.* 1998; Brown *et al.* 1998; Schurr *et al.* 1999). Specifically, the analysis of mtDNA diversity in the Chukchi and Siberian Eskimos of extreme northeastern Siberia revealed haplogroups A, C, and D. In contrast, the Koriak and Itelmen of the adjacent Kamchatka Peninsula, who speak a language from the same language phylum (Chukchi-Kamchatkan), harbour the east Eurasian haplogroups G, Z and Y, which are completely absent in Native Americans. Interestingly, the Aleuts of the Commander Islands, adjacent to the Kamchatka Peninsula, were founded by a single lineage of haplogroup D2 (Derbeneva *et al.* 2002 a). While haplogroup B is absent in aboriginal populations of northwestern and northern

Siberia (Derbeneva *et al.* 2002 b, c), it has been found in populations restricted to the south-western and south-central periphery of the subcontinent (Sukernik *et al.* 1996; Derenko *et al.* 2000, 2003;).

The headwaters of the Ob, Yenisei, Lena and Amur rivers, the largest rivers in Eurasia, served as major routes out of Inner Asia where migrations initiated northward to the northern and eastern perimeters of former Beringia. Hence, populations from these areas might be expected to harbour the founding mtDNA lineages present in Native Americans. The occurrence of haplogroups A, B, C and D in southern Siberia and adjacent areas of Mongolia (Kolman *et al.* 1996; Sukernik *et al.* 1996; Derenko *et al.* 2000, 2003; Keyser-Tracqui *et al.* 2003), and the traces of haplogroup X mtDNAs revealed in the mountainous Altai (Derenko *et al.* 2001), supports this hypothesis. To further clarify the relationship between Siberian and Native American mtDNA haplogroups and subhaplogroups, we have undertaken a broad survey of mtDNA variation by high-resolution RFLP analysis, sequencing of the HVS-I, and surveying some additional diagnostic markers, in nine distinct populations that have evolved in the southern extent of Siberia. Finally, mtDNAs selected according to haplogroup/subhaplogroup status were then completely sequenced. The data obtained were integrated with similar data sets representing Siberian arctic and sub-arctic populations: the Mansi of the Lower Ob River basin, the Ket of the Lower Yenisei and the Nganasan of the Taimir Peninsula (Derbeneva *et al.* 2002b,c), the Chukchi and Siberian Eskimos of Chukotka (Starikovskaya *et al.* 1998) and the Itelmen and Koriak of Kamchatka (Schurr *et al.* 1999). Together, these data encompass all of the linguistic groups of indigenous Siberian populations, and support a dual Siberian origin for Native Americans: a migration derived from the Altai-Sayan Upland and a migration from the Lower Amur River/Sea of Okhotsk region.

Subjects and Methods

Populations and Samples

Blood samples were collected from nine indigenous populations with appropriate informed consent during multiple field expeditions conducted by Rem I.



Figure 1 Approximate location of Siberian populations analysed for mtDNA variation.

Sukernik, Elena B. Starikovskaya, and Natalia V. Volodko (Figure 1). The individuals who participated in these studies were interviewed and verified their family histories prior to blood being drawn, only from those subjects who were unrelated through at least three generations and lacked non-native maternal ancestors. A brief description of each population follows. Note that the Nivkhi, the Udegey and most of the Evenki samples are those previously surveyed by Torroni *et al.* (1993a). In that study, RFLP analysis with 14 restriction endonucleases and sequence analysis of a few selected HVS-I regions representing haplogroups A, C and D were performed.

Tubalar – The present study includes 72 Tubalar, 27 individuals of whom were collected from the Tubalar admixed with Chelkan in the villages of Suronash, Tuloi and Artybash (Sukernik *et al.* 1996), while 45 are new samples drawn from the Tubalar in the villages of Pyzha, Tunzha, Paspaul, Salganda, Ynyrga, Kara-Koksha and Urlu-Aspak (Turochak and Choiski Districts, Altai Republic). The Tubalar and Chelkan are recent descen-

dants of small hunting-gathering bands who a century ago occupied the coniferous forest zone (taiga) of north-eastern Altai. They differ from numerous southern Altaians in culture, language, and physical appearance (Levin & Potapov, 1964).

Tuvan – The Tuvan are largely Turkic-speakers, and their total population size exceeds 100,000. A total of 95 Tuvan blood samples were collected across the Tuva Republic. Much of the Tuva Republic is situated in the south extreme of Siberia, with the taiga zone of Altai-Sayan ranges to the northwest, north and northeast, and the Mongolian arid sandy wastes to the south. Many families still perform nomadic/pastoral subsistence activities (Levin & Potapov, 1964).

Buryat – The Buryat are Mongolic-speakers who are regarded as a northern extension of the ethnic groups from Mongolia and former Manchuria (Levin & Potapov, 1964). At present, the total population size of the Russian Buryat is over 200,000. Geographically and culturally they are close to the Tuvan. Our sample consisted of 25 non-related individuals currently

residing in Kushun village (Nizhneudinsk District, Irkutsk Region), and represents the Buryat of the western Baikal Upland. The family history of a few of the individuals sampled indicated Tofalar ancestry through the maternal line.

Tofalar (former Karagas) – The Tofalar are a small tribe of hunters and reindeer breeders inhabiting the northern slopes of the Sayan mountain range. The current population, which totals less than four hundred individuals, is subdivided into two territorial groups, one residing in the western (the village of Upper Gutara) and the other in the eastern part of mountainous Sayan (the villages of Nerkha and Alygdzher). Originally, the Tofalar, as well as the Tubalar and Chelkan tribes of the northeastern Altai, spoke a Samoyed language of the Uralic language family, but later adopted a Turkic language (Levin & Potapov, 1964). Our sample included 46 individuals collected from Upper Gutara and Nerkha. Several current residents of Nerkha were born in the adjoining village of Alygdzher in the same Nizhneudinsk District, Irkutsk Region.

Evenki – The present study includes 71 Evenki mtDNAs, of which 53 were collected from the Evenki living in the villages of Poligus and Surinda, located in the middle of the Stony Tunguska River basin (Torroni *et al.* 1993a), and 18 are new samples obtained from the Evenki inhabiting the Sea of Okhotsk region (Lell *et al.* 2002; Uinuk-ool *et al.* 2002). Most of the Evenki are hunters and reindeer breeders, with a total population of approximately 20,000. They speak the northern Tungusic language of the Altaic linguistic family, and inhabit the vast expanses of boreal forest extending from the Lower Yenisei in the west to the coast of the Okhotsk Sea in the east (Levin & Potapov, 1964).

Negidal – Thirty-three Negidal samples were analysed. They were collected from the village of Vladimirovka (Polina Osipenko District, Khabarovsk Region), and from the villages of Takhta, Tir, and Mago (Nikolaevskiy District, Khabarovsk Region). Genetically, the Negidals may represent the last remnants in the southern Okhotsk region of a culture based on the hunting of small sea mammals, which was first described by Middendorf (1869). In the late 1800s, several hundred Negidal lived in small settlements spread along the course of the Amgun, the left tributary of the Lower Amur, and the adjacent Tugur, which flow into the Sea

of Okhotsk. Subsequently, they have been admixed and influenced by the expanding Tungusic Evenki and Even tribes, and now speak a dialect belonging to the Tungusic language group (Levin & Potapov, 1964).

Ulchi – We analysed 87 samples, obtained from elderly Ulchi residing in Old and New Bulava, two neighbouring villages (Ulchi District, Khabarovsk Region). Until recently, the Ulchi were a well-defined tribe of hunters and fishermen dispersed along the lakes and the reaches of the Lower Amur. They speak a language of the Tungusic-Manchurian group. The Ulchi sampled are representative of the few hundred individuals who are left from several small Ulchi villages (Levin & Potapov, 1964; Black, 1988) that are no longer in existence.

Nivkhi – The 56 Nivkhi were collected from the village of Nekrasovka and from the tiny fishing settlements located in the northwestern part of Sakhalin Island (Torroni *et al.* 1993a). In traditional times, the Nivkhi were small sea mammal hunters of the Lower Amur/Southern Okhotsk region and numbered several thousands. The Nivkhi is a language isolate with no known affiliation to existing language families (Levin & Potapov, 1964; Black, 1988). At present, this population is subdivided into two adjoining groups, one in the lower-most Amur River and Amur delta, and the other in the northern part of Sakhalin.

Udegey – We extended the analysis of 46 Udegey mtDNAs from the Gvasiugi village located in the central part of the Sikhote-Alin Range, Khabarovsk Region (Torroni *et al.* 1993a). This hunting-fishing group historically inhabited both slopes of the Sikhote-Alin Range, adjacent to the Sea of Japan, but are now on the brink of extinction. The Udegey language belongs to the southern branch of the Tungusic language group (Levin & Potapov, 1964; Krauss, 1988).

mtDNA Analysis

Genomic DNAs were extracted either from buffy coats or lymphoblast cell lines using standard procedures. To determine RFLP haplotypes, the entire mtDNA of each sample was amplified in ten overlapping PCR fragments (Torroni *et al.* 1993a; 1996). Each of the ten PCR segments was then digested with 14 restriction endonucleases (*AluI*, *AvaII*, *BamHI*, *DdeI*, *HaeII*, *HaeIII*, *HhaI*, *HincII*, *HinfI*, *HpaI*, *MboI*, *MspI*, *RsaI*,

and *TaqI*). In addition, all mtDNAs were screened for the presence/absence of the *BstNI* site at nucleotide position (np) 13704, the *AclI* sites at nps 14465 and 15254, the *BfaI* site at np 4914, the *NlaIII* sites at nps 4216 and 4577, and the *MseI* sites at nps 14766 and 16297. The polymorphism at np 12308 was also surveyed by using a mismatched primer that generates a *HinfI* site when the A12308G mutation is present (Torroni *et al.* 1996). RFLP analysis was supplemented by sequencing of HVS-I, which was performed as described in Starikovskaya *et al.* (1998). In addition, selected mtDNAs representing major lineages and sublineages in Siberia were completely sequenced using cycle sequencing, “Big Dye Terminators” (ABI/Perkin-Elmer Cetus) dideoxy nucleotide terminators and an ABI Prism 3100 DNA Analyzer. Trace files were analysed using the Sequencher (v.4.0.5 GeneCode Corp.) software. During the course of the present and related studies 67 mtDNAs collected throughout Siberia and the adjacent Commander Islands were subjected to complete sequencing (Derbeneva *et al.* 2002a; Mishmar *et al.* 2003). Twenty of these 67 mtDNA sequences are new, and have been submitted to GenBank under accession codes AY519484–AY519497 and AY570524–AY570526, AY615359–615361.

Categorization of Haplotypes and Haplogroups

The Siberian mtDNA types were categorized into haplogroups denoting monophyletic clusters of mitochondrial sequences that arose on the ancestral haplotype background (Torroni *et al.* 1993a, 1994, 1996; Forster *et al.* 1996; Macaulay *et al.* 1999; Finnila *et al.* 2001; Derbeneva *et al.* 2002 a, b, c; Herrnstadt *et al.* 2002; Mishmar *et al.* 2003). “Founder” or “ancestral” haplotypes (versus “derived” haplotypes) are those from which all other related haplotypes were “derived” through acquisition of new mutations. Haplogroups (subhaplogroups) are also referred to as mtDNA lineages of related mtDNA haplotypes. “Candidate” founders for Native American mtDNA haplogroups were identified by comparing the polymorphisms of similar haplotypes from Eurasia and the Americas. Relationships were confirmed and extended using complete mtDNA sequences.

Phylogenetic analysis

Phylogenetic relationships between 16 Siberian and two adjacent Native American populations of the North Pacific Rim (Aleut and Haida) were determined by the neighbor-joining algorithm using D_A distances:

$$D_A = 1 - \sum_i^m \sqrt{x_i y_i}$$

where m is the number of subhaplogroups across populations and x_i and y_i are frequencies of the i -th mtDNA subhaplogroups in populations X and Y , respectively. D_A distance is more efficient in obtaining a correct tree topology for a small number of loci (Takezaki & Nei, 1996).

Results

mtDNA Diversity

The mtDNAs of 531 maternally unrelated individuals from nine native Siberian populations were characterized by RFLP analysis and HVS-I sequencing (Table 1). The populations studied extend from the North-eastern Altai in the west to the Lower Amur/Sea of Okhotsk region in eastern Siberia. The overall distribution of mtDNA subhaplogroups across Siberia is shown in Table 2. Approximately, 66% of southern Siberian mtDNAs were found to belong to the “Asian” macro-haplogroup M (defined by presence of the 10394 *DdeI* and 10397 *AluI* sites) and its derivatives. The remaining mtDNAs belonged to the macro-haplogroup N, lacking both the 10394 *DdeI* and 10397 *AluI* sites, or alternatively having the 10394 *DdeI* site but lacking the 10397 *AluI* site.

Macro-haplogroup N mtDNAs

Haplogroup A is defined by the 663 *HaeIII* site and the HVS-I motif 16223–16290–16319–16362. It is rare but widespread, having been found in some Western and Middle Siberian populations such as the Mansi, Ket, Tubalar, Tuvan and Evenki (Torroni *et al.* 1993a; Sukernik *et al.* 1996; Derbeneva *et al.* 2002b,c; this study). Haplogroup A mtDNAs in these populations lack the mutation at np 16111 and are classified within

Table 1 mtDNA diversity in nine Siberian populations

Haplogroup	RFLPs	HVS-I (–16000)	Populations and number of subjects ^a									
			TB	TV	BR	TF	EV	NG	UL	NV	UD	Total
A1	+663e	192 223 290 319 362	8									8
	+663e	242 290 293 319		1								1
	+663e	086 223 290 293 319 362		1								1
	+663e +15606a	039 189 223 290 319 356 362					1					1
	+663e +15606a	039 189 218 223 290 319 356 362					1					1
B1	+663e –9751l/+15606a	039 189 223 290 319 356 362					1					1
	+663e –9751l/+15606a –15883e	039 189 223 290 319 356 362					1					1
	9bp del –6022a	086 136 189 217 519	1									1
	9bp del –6022a –9052n/+9612k	086 136 189 217 519	2	1								3
	9bp del +13940f	189 217 240 519			1							1
B3	9bp del	167 189 217 261 317T 519		4		2						6
	9bp del –14766u	129 189 217 261 356		1								1
B5	9bp del (+/–)	140 189 243 519		1								1
	9bp del (+/–) +14465s	140 148 189 218 266 243 519		1								1
	9bp del (+/–) +4895k (+/–) –10971g +15221j –15925l	111 129 140 189 234 243 244 463 519						4				4
F	+4732k –12406h/–12629b	189 232A 249 304 311 519	1	1			1					3
	+4732k –12406h/–12629b	172 189 232A 249 286 304 311			3							3
Y	+4732k –8391e –12406h/–12629b	172 189 232A 249 286 304 311			1							1
	–12406h/	189 304 327 519	3									3
	+545g –9052n/–12406h/–13268g	162 172 304 519							1			1
	+7933j –8391e (+/–)	126 231 266 304 519						1		1		2
	+7933j –8391e (+/–) –13608l	126 231 266 304 519						1				1
	–7641a +7933j –8391e (+/–)	126 231 266 304 380 519									1	1
	+7933j –8391e (+/–)	126 189 231 519								11		11
	+7933j –8391e (+/–)	126 189 231 266 519						5	18	10	2	35
	+7933j –8391e (+/–)	126 189 266 519							1			1
	+7933j –8391e (+/–)	126 189 231 292 519								3		3
	+7933j –8391e (+/–)	126 189 266 292 519								1		1
	+7933j –8391e (+/–)	126 189 231 266 292 519						2				2
	+7933j –8391e (+/–)	126 189 231 266 294 519						1	1			2
	+7933j –8391e (+/–)	126 140 189 231 266 399 519							1			1
	–322e +7933j –8391e (+/–)	126 189 231 266 519						5				5
	–322e +7933j –8391e (+/–)	093h 126 189 231 266 519						1				1
	–322e +7933j –8391e (+/–)	126 189 231 266 294 519							1			1

Table 1 continued.

Haplogroup	RFLPs	Populations and number of subjects ^a									
		HVS-I (–16000)	TB	TV	BR	TF	EV	NG	UL	NV	UD
H	–322e +7933j –8391e (+/–)	126 189 266 519							1		1
	+7933j –8391e (+/–) 4bp ins	126 189 231 519								4	4
	+7933j –8391e (+/–) +11900k	126 189 266 292 519								1	1
	+7933j –8391e (+/–) +11900k	126 189 231 266 292 519								4	4
	+7933j –8391e (+/–) +14168l	126 189 231 266 519						2		1	3
	–853c/+7933j –8391e (+/–) –15375g	126 189 231 519								1	1
V	–7025a –14766u	519	1								1
	–7025a –14766u	288 362	3								3
	–7025a –14766u	311 519		3							3
	–7025a –9380f –14766u	362 519		1							1
	–7025a –7598f +14249f –14766u	093 129 316 519			1						1
	–4577q –14766u +15904u	153 298			1						1
J	+4216q (+/–) –13704t	069 126 519			1						1
	+4216q (+/–) –13704t	069 126 145 172 261 278		1							1
	+4216q –4685a (+/–) +11001n/										
	+11414a –13704t –15254s	069 126 241		1							1
	+4643k +11329a +12308g	356 519	10	1							11
	+4643k +11329a +12308g	311 356 519	3								3
U4	+12308g	189 260 270 519		2							2
	+1718a +12308g +13634s	192 241 256 270 287 304 325 399	4								4
	+12308g	093 234 270			3						3
	+12308g	129C 189 214 258 362 519	2								2
	+8249b/ –8994e (12705)	104 223 284 519		2							2
	(–/–) +14465s (12705)	189 223 278 519	1								1
N9	(5417) (12705)	111 129 223 257A 261		3							3
	(5417) (12705)	223 248 257A 261 311 519	5								5
	(5417) –5742i (12705)	189 223							2		2
	(5417) –5742i (12705)	129 189 223 311							1		1
	(5417) (12705) –14258m/	223 519							3		17
	(–/–) –12282a	129 184i 190i 362 390 519	1								1
R6	(–/–)	145 192 243 309 390 519 527	1								1
R*	–7598f (+/+) –13259o/	223 298 325 327 356							1		1
C1a	(+/+) –13259o/	223 298 327 519	8	1							9
C2	–5983g/(+/+) –13259o/	223 298 327 519			2						2
	(+/+) –13259o/ +15606a	223 298 327 519				1					1

Table 1 continued.

Haplogroup	RFLPs	HVS-I (–16000)	Populations and number of subjects ^a									
			TB	TV	BR	TF	EV	NG	UL	NV	UD	Total
	(+/+) – 13259o/	129 223 298 327	1									1
	(+/+) – 13259o/	223 264 298 327 519	1									1
	(+/+) – 13259o/	223 298 311 327 519	2	2		3	5					12
	(+/+) – 13259o/	219 223 298 327 519					1					1
	(+/+) – 13259o/	223 294 298 311 327 519	1									1
	(+/+) – 13259o/	223 291 298 327 519	1		2							3
	(+/+) – 13259o/	223 261 298 519							1			1
	(+/+) – 12170g/ – 13259o/	223 249 291 295 298 327 519					2					2
	+8249b/(+/+) – 13259o/	223 298 327 519		10			5				5	20
	+8249b/(+/+) – 13259o/	093 223 298 327 519									1	1
	+8249b/(+/+) – 13259o/	093 129 223 298 327 519					1					1
	+8249b/(+/+) – 13259o/	190 223 232 298 327 519									1	1
	+8249b/(+/+) – 13259o/	223 259+A 298 327 519							2			2
	+8249b/(+/+) – 13259o/	223 262+C 291 298 327 519						2				2
	– 1715c (+/+) – 13259o/	150 223 298 327 519					1					1
	– 1715c (+/+) – 13259o/	129 150 223 298 327 519					1		1		1	3
	– 1715c (+/+) – 13259o/	093 129 223 298 327 519		1			1					2
	– 1715c (+/+) – 13259o/	129 223 298 327 519		1		2	1	1	1			6
	– 1715c (+/+) – 13259o/ + 15606a	093 129 223 298 327 519		8	3		7		1			19
	– 1715c (+/+) – 13259o/ + 15606a	093 129 223 235 298 327 390 519				10						10
	– 1715c – 6850i (+/+) – 13259o/ + 15606a	093 129 223 298 327 519			1							1
	– 1715c (+/+) – 13259o/ + 15606a	129 223 298 327 519					4					4
	– 1715c – 6850i (+/+) – 13259o/ + 15606a	129 223 298 327 519			2							2
	– 1004o – 1715c (+/+) – 13259o/	129 223 298 327 519					2					4
	(+/+) – 13259o/	223 298 327 344 357 519	1	2			2					5
	(+/+) – 13259o/	171 223 327 344 357 519					1					1
	(+/+) – 13259o/	171 223 298 327 344 357 519		1		2	3					6
	(+/+) – 13259o/	171 223 224 298 327 344 357 519			2	2						4
	(+/+) – 13259o/	171 209 223 298 327 344 357 519					1					1
	– 1715c (+/+) – 13259o/	093 129 167 223 298 327 357 519		1								1
	(+/+) – 13259o/	171 188 223 261 298 327 344 357 519					1					1
	(+/+) – 13259o/	167 223 298 327 344 357 519		1								1
	– 5983g/(+/+) – 13259o/	171 223 298 327 344 357 519		2		2						4
	(+/+) – 13259o/	093 171 223 298 327 344 357 519					1					1

Table 1 continued.

Haplogroup	RFLPs	HVS-I (–16000)	Populations and number of subjects ^a									
			TB	TV	BR	TF	EV	NG	UL	NV	UD	Total
C3	(+/+) –13259o/	129 140 171 223 298 327 344 357 519	1									1
	–1413l +8249b/(+/+) –13259o/	093 223 288 298 327 390 519	1									1
	–1413l (+/+) –13259o/	093 223 288 298 327 390 519					7					7
	(+/+) –13259o/	223 261 288 298 519	3						5			8
	(+/+) –13259o/	223 288 298 327 390 519					1					1
	(+/+) –13259o/	093 223 288 298 327 390 519						1				1
	(+/+) –13259o/(3460)	093 223 288 298 327 390 519	1									1
	(+/+) –13259o/	093 223 288 291 298 327 518T 519	2									2
	(+/+) –13259o/	093 223 288 298 327 390 500A 519						1				1
	–1715c +3397k (+/+) –13259o/9bp ins	148 223 288 298 327 519	1			3						4
	–1715c (+/+) –13259o/9bp ins	148 223 288 298 327 519	1			2						3
	+1718e –7297e (+/+) –13259o/	148 164 223 288 298 327 519					2					2
D	–5176a (+/+)	223 362	6	1								7
	–5176a (+/+)	223 291 362					1					4
	–5176a (+/+)	223 362 368					1		3			1
	–5176a (+/+)	174 223 362			4							4
	–5176a (+/+)	223 362 519										2
	–5176a (+/+)	189 223 362 519	2									2
	–5176a (+/+)	223 263 362 519	1									1
	–5176a (+/+)	145 311 362 368					4					4
	–5176a (+/+)	223 274 362 368					1					1
	–5176a (+/+)	223 245 311 362 368					2	1				3
	–5176a (+/+)	223 245 311 362 368	2									2
	–5176a (+/+)	093 223 232 290 362						1				1
	–5176a (+/+)	148 223 263 362 519					1	1				2
	–5176a (+/+)	223 362					1					1
	–5176a –5971f (+/+)	223 286 362	1									1
	+1973g –5176a (+/+)	223 245 362	1									1
	–5176a (+/+) +14583c	223 286 362	1									1
	–5176a –8074a (+/+)	223 362 519	1									1
	–5176a –9294e (+/+)	223 362 519	1									1
	–5176a (+/+) +14465s (14470)	223 362 519						1				1
	–5176a –7859j (+/+)	223 263 362 519					2					2
	–5176a (+/+) +10407k	129 223 362 519						2				2
	–5176a +7979e (+/+)	184 223 311 362							1			1
	–1715c –5176a (+/+)	042 223 362								1		1

Table 1 continued.

Haplogroup	RFLPs	HVS-I (–16000)	Populations and number of subjects ^a									
			TB	TV	BR	TF	EV	NG	UL	NV	UD	Total
D1a D3	–1715c –5176a (+/+)	042 214 223 362	1						10			11
	–1715c –5176a (+/+)	042 214 223 234 362							2			2
	–1715c –5176a +8865j (+/+)	042 172 223 362		1								1
	–5176a (+/+)	223 325 362 519						4				4
	–5176a –101801 (+/+)	223 319 362					1					1
D4	–951j –5176a –101801 (+/+) +15437e	223 319 362		2				2				2
	–5176a –101801 (+/+) +13717a +14923c +15437e	223 319 362										2
	–951j +4133q –5176a –5823a –101801 (+/+) +15437e	223 239 243 319 362		1								1
	–5176a –101801 (+/+) +15437e	093 172 173 215 223 319 362 519	1									1
	–5176a (+/+) +10646k	093 223 290 362							1			1
D5	–5176a (+/+) +10646k	093 223 232 290 362						3				3
	–5176a (+/+) +10646k –14015a	093 223 232 261 294 362						2				2
	–5176a (+/+) +10646k	093 223 232 290 362						2				2
	–5176a (+/+) +10646k	093 223 232 261 290 362						2				2
	–5176a (+/+) +10646k	183 223 274 290 319 362							1			1
G1	–5176a (+/+) +10646k	183 223 274 290 319 362 519							1			1
	–5176a (+/+) +10646k –13268g	176 183 223 274 290 319 342 362 519	1									1
	+4877a –5176a (–/–) +12026h/(12705)	092 172 189 223 266 362					1					1
	–5176a (–/–) +12026h/(12705)	092 126 164 189 223 266 362	3									3
	–5176a (–/–) +12026h/–13259o (12705)	092 164 172 189 223 266 362		1								1
G2	+4830n/+8198a (+/+)	017 129 223 519								1		1
	+4830n/+8198a (+/+)	017 093 129 223 519						2	2	1		5
	+4830n/+8198a (+/+)	017 093 129 223 303 519							1			1
	+4830n/+8198a (+/+)	017 051 093 207 223 399 519							1			1
	+4830n/+8198a +9253k (+/+)	017 093 129 223 311 519							1			1
G2	+4830n/+8198a (+/+)	017 129 223 519							1			1
	+4830n/+8198a (+/+)	017 093 129 207 223 519										2
	+4830n/+8198a (+/+)	017 093 207 223 244 399 519								1		1
	+4830n/+8198a +8957l (+/+) +15494c	017 051 093 207 223 399 519							1			1
	+4830n/+6629a +8198a (+/+)	017 129 223 519						1				1
G2	+4830n/+6629a +8198a (+/+)	017 093 129 223 519						5				5
	+4830n/+6629a –7658j +8198a (+/+)	017 093 129 223 313A 519						1				1
	+4830n/–7598f (7600) (+/+)	051 150 223 278 362	1									1

Table 1 continued.

Haplogroup	RFLPs	HVS-I (−16000)	Populations and number of subjects ^a										Total
			TB	TV	BR	TF	EV	NG	UL	NV	UD		
G3	+4830n/−7598f (7600) (+/+) +12752a	093 126 223 227 278 362	1									1	
	+4830n/−7598f (7600) (+/+) +12752a	223 227 234 278 362	1									1	
	+4830n/−7598f (7600) (+/+) +12752a	223 227 278 294 362	1									1	
	+4830n/−7598f (7600) +8249b/(+/+)	223 227 278 362 519	1									1	
	+4830n/−7598f (7600) −9438e (+/+) +12752a	220C 223 227 278 362		1								1	
	+4830n/−7598f (7600) (+/+) +12752a	093 129 223 271 278 362 399	1					1				2	
	+4830n/(+/+) −12406h	223 274 362 390	1									1	
	+4830n/(+/+) −12406h	093 223 274 362 390 519	1									1	
	+4830n/(+/+) −12406h	093 223 274 320 362 390 519			1							1	
	+4643k +4830n/(+/+) −12406h −15172e	093 223 274 362 390 519	1									1	
S	+4830n/−8858f (+/+) +12752a	215 223 274	1									1	
	(+/+) +14465s (14470)	184 223 293 298 319 519			1	7						8	
	(+/+) +14465s (14470)	184 189 223 293 298 319						1				1	
	(+/+) +14465s (14470)	184 223 298 319						2				2	
	(+/+) −13608l +14465s (14470)	184 223 298 319						1				1	
	+3744e (+/+) +14465s (14470) −15047e	086 223 298 319								7		7	
	(+/+) +14465s (14470)	185 223 260 298 360 519	1			5						6	
	(+/+) +14465s (14470)	129 185 223 224 260 298 519	1									1	
	−9052n/(+/+) +14465s (14470)	185 223 260 298 360 519			1							1	
	+9820g (+/+) +14465s (14470)	223 245 311 362 368							1			1	
M7	+9820g (+/+) +14465s (14470)	117 162delA 172 223 519		2								2	
	−4848e +9820g (+/+) +14465s (14470)	173 223 362 519		1								1	
	+4092e +9820g (+/+) +14465s (14470)	140 187 209 223 232 519								4		4	
	+4092e +9820g (+/+) +14465s (14470)	140 187 209 223 519								4		4	
	−1667c/+4092e +9820g (+/+) +14465s (14470)	140 187 209 223 232 519								1		1	
	+5351f −7853o +9820g (+/+) +14465s (14470)	129 152 179 189 223 362							1			1	
	+3391e (+/+) +14465s (14470)	223 234 316 362										1	
	Total		72	95	25	46	71	33	87	56	46	531	

Note: RFLP sites are numbered from the first nucleotide of the enzyme recognition sequence. A “+” indicates the presence of the restriction site, a “–” the absence. The restriction enzymes are given using the following single-letter code: a = *AluI*; b = *Avall*; c = *DdeI*; e = *HaeIII*; f = *HinfI*; g = *HinfI*; h = *HpaI*; i = *HpaII*; j = *TaqI*; m = *BamHI*; n = *HaeII*; o = *HincII*; q = *NlaIII*; r = *BfaI*; s = *AclI*; t = *BstNI*; u = *MseI*. A slash at the end of the restriction site implies the simultaneous presence/absence of a linked site due to the same nucleotide substitution. The presence/absence of the associated 10394 *DdeI*/10397 *AluI* sites is denoted through slash brackets (+/+), or (+/–), 9bp ins = 9bp COII/tRNA^{Lys}tripligation; 9bp del = 9bp COII/tRNA^{Lys}deletion; 4bp ins = 4bp COII/tRNA^{Lys} insertion; h = heteroplasmy. Additional mutations in the coding region verified through sequencing are shown in brackets. Founding RFLP/HVS-I haplotypes are shown in boldface. Only those nucleotide positions between 16013 and 16520 that differ from the revised Cambridge Reference Sequence (rCRS) (Andrews et al. 1999) are shown. Mutations are transitions, unless the base change is specified explicitly. The HVS-I mutations recognized also by RFLP analysis are shown only within the HVS-I sequence.

^a Population codes are as follows: TB = Tubalar; TV = Tuvan; BR = Buryat; EV = Evenki; NG = Negidal; UL = Ulchi; NV = Nivkhi; UD = Udegey.

Table 2 Frequencies of mtDNA haplogroups (%) in 16 Siberian populations

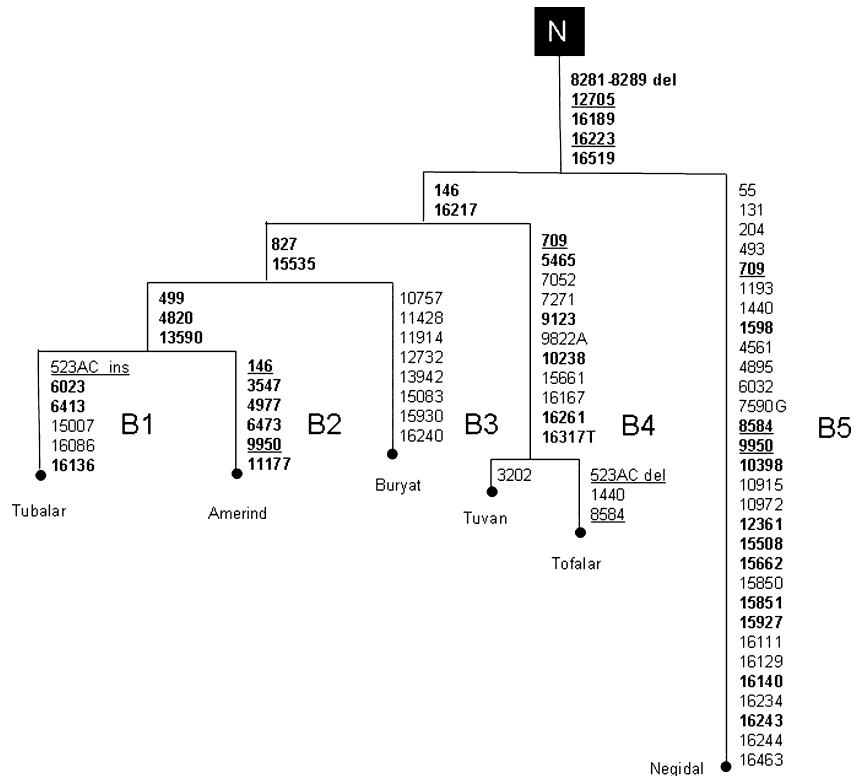
Haplogroup	MN (98)	KT (38)	NS (24)	TB (72)	TV (95)	BR (25)	TF (46)	EV (71)	NG (33)	UL (87)	NV (56)	UD (46)	IT (47)	KR (155)	CH (66)	ES (79)
A1	3.1	7.9	-	11.1	2.1	-	-	5.6	-	-	-	-	6.4	2.6	-	-
A2	-	-	-	-	-	-	-	-	-	-	-	-	-	2.6	68.2	77.2
B1	-	-	-	4.2	1.1	-	-	-	-	-	-	-	-	-	-	-
B3	-	-	-	-	-	4.0	-	-	-	-	-	-	-	-	-	-
B4	-	-	-	-	5.3	-	4.3	-	-	-	-	-	-	-	-	-
B5	-	-	-	-	2.1	-	-	-	12.1	-	-	-	-	-	-	-
F	1.0	23.7	-	1.4	4.2	-	8.7	1.4	-	1.1	-	-	-	-	-	-
Y	-	-	-	-	-	-	-	-	21.2	37.9	66.1	8.7	4.1	9.7	-	-
H	14.3	10.5	8.4	5.6	4.2	4.0	-	-	-	-	-	-	-	-	-	-
V	1.0	-	-	-	-	4.0	-	-	-	-	-	-	-	-	-	-
J	12.2	-	-	-	2.1	4.0	-	-	-	-	-	-	-	-	-	-
T	7.2	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
U	25.4	34.2	25.0	26.4	3.2	12.0	-	-	-	-	-	-	-	-	-	-
K	3.1	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
W	-	2.7	-	-	2.1	-	-	-	-	-	-	-	-	-	-	-
X	-	-	-	1.4	-	-	-	-	-	-	-	-	-	-	-	-
N9	-	-	-	6.9	3.2	-	-	-	-	6.9	-	30.4	-	-	-	-
R	-	-	-	2.8	-	-	-	-	-	-	-	-	-	-	-	-
C1a	-	-	-	-	-	-	-	-	-	1.1	-	-	-	-	-	-
C2	16.3	15.8	12.6	16.7	35.8	40.0	50.0	57.8	9.1	6.9	-	17.4	2.1	23.8	4.5	2.5
C3	1.0	-	20.8	2.8	7.4	-	10.9	14.1	6.1	5.7	-	-	12.8	12.3	6.1	-
D	6.2	2.6	8.3	12.5	9.5	16.0	-	18.3	18.2	4.6	23.2	-	-	-	-	-
D1a	-	-	-	-	-	-	-	-	-	4.6	-	-	-	-	-	-
D2	-	-	-	-	-	-	-	-	-	-	-	-	-	-	10.6	15.2
D3	1.0	-	16.7	1.4	3.2	-	-	1.4	-	2.3	-	-	-	1.3	1.5	5.1
D4	-	-	4.2	1.4	-	-	-	-	6.1	10.3	5.4	-	-	-	-	-
D5	1.0	-	-	4.2	1.1	-	-	1.4	-	-	-	-	-	-	-	-
G1	-	-	-	-	-	-	-	-	27.2	10.3	5.4	-	68.1	41.9	9.1	-
G2	6.1	-	-	-	6.3	4.0	-	-	-	1.1	-	-	-	-	-	-
G3	-	-	-	-	4.3	4.0	-	-	-	-	-	-	-	-	-	-
S	-	-	-	-	-	4.0	15.2	-	-	4.6	-	15.2	-	-	-	-
Z	-	2.6	4.2	1.4	1.6	4.0	10.9	-	-	-	-	-	6.4	5.8	-	-
M	1.0	-	-	-	1.6	-	-	-	-	2.3	-	28.3	-	-	-	-

Note: inferred from published data as follows: MN – Mansi (Derbeneva *et al.* 2002b); KT – Ket, NS – Nganasan (Derbeneva *et al.* 2002c); IT – Itelmen, KR – Koriak (Schurr *et al.* 1999); CH – Chukchi, ES – Eskimos (Starikovskaya *et al.* 1998). The double-letter code for populations in this study is given as in Table 1.

subhaplogroup A1 (Forster *et al.* 1996). In contrast, the Chukchi and Eskimo haplogroup A mtDNAs harbour the 16111 transition and belong to the same A subhaplogroup (A2) found in Native Americans (Starikovskaya *et al.* 1998). Thus, the homeland of A2 is most likely a Beringian refuge area, from where it subsequently expanded into the New World (Forster *et al.* 1996; Starikovskaya *et al.* 1998; Saillard *et al.* 2000). In the course of the present study, we completely sequenced two A1 mtDNAs (one Mansi from the Lower Ob and one Ket from the Lower Yenisei River) previously defined at the RFLP/HVS-I level (Derbeneva *et al.* 2002 b, c). While their control region harbored the CA deletion

at nps 522–523, the coding region showed five transitions at nps 663, 1736, 4248, 4824 and 8794. These mutations apparently predated the split of A2 from A1, since they are also found in A2 mtDNAs (Ingman *et al.* 2000; Mishmar *et al.* 2003). Interestingly, two haplogroup A mtDNA samples from China, which were completely sequenced by Kong *et al.* (2003), harboured the same ancestral motif characterizing the A1 mtDNA subgroup in the Ket and Mansi. However, haplogroup A2 mtDNAs, as noted in both the Chukchi and Native Americans, can be easily distinguished by the additional transitions at nps 8027 and 12007 (Ingman *et al.* 2000; Herrnstadt *et al.* 2002; Mishmar *et al.* 2003; Silva *et al.* 2003).

Figure 2 Schematic phylogeny of haplogroup B mtDNAs found in Native Siberians and Native Americans. The phylogeny is rooted from macrohaplogroup N as inferred from the complete sequences of Siberian/Asian haplogroup A, B, and Y (Ingman *et al.* 2000; Herrnstadt *et al.* 2002; Mishmar *et al.* 2003; this study). Mutation positions, relative to the revised Cambridge Reference Sequence (Andrews *et al.* 1999), are transitions unless the base change is specified. Underlining indicates recurrent mutations. Founding motifs are shown in bold type.



Haplogroup B mtDNAs are defined by the 9-bp deletion at nps 8281–8289 and the control-region motif 16189–16519. Several subhaplogroups of B have been delineated in Siberians (Table 1). B1 mtDNAs were found in a Tubalar and a Tuvan. They are distinguished by the 6022 *AluI* site loss and the control region motif 16086–16136–16189–16217–16519 (Table 1). A similar (at the HVS-I level) haplogroup B mtDNA has been detected in the skeletal remains exhumed from a 2000 year-old cemetery in northern Mongolia (Keyser-Tracqui *et al.* 2003). The complete sequence of the Tubalar mtDNA revealed five mutations (Figure 2) (499, 827, 4820, 13590, and 15535) which are shared with the Native American haplogroup B mtDNAs sequenced by Ingman *et al.* (2000) while two mutations (6023 and 6413) are shared with some, but not all, Asian haplogroup B mtDNAs (Ingman *et al.* 2000; this study). At the coding region level, the Native American branch (B2) of haplogroup B (Herrnstadt *et al.* 2002) differs from B1 for the additional transitions at nps 3547, 4977, 6473, 9950, 11177 and the lack of the mutations at nps 6023 and 6413. We also sequenced the haplogroup B mtDNA found in the Buryat (Table 1). It lacked three (499, 4820, 13590) of the transi-

tions shared by the Siberian B1 and the Native American B2. In addition, its coding region harboured seven transitions, and we classified this mtDNA within the novel subhaplogroup B3. An additional subset of B mtDNAs, which we designated B4, is defined by the HVS-I motif 16189–16217–16261 (Table 1) and was observed in five Tuvans and two Tofalars. This mtDNA subset seems to be widely dispersed in adjacent Mongolia and China (Kolman *et al.* 1996; Yao *et al.* 2000, 2002). We sequenced two B4 mtDNAs (one Tuvan and one Tofalar) and compared them with those obtained from Korean and Samoan haplogroup B mtDNAs (Ingman *et al.* 2000). The comparison showed that they all share the coding region variants 709, 5465, 9123, and 10238, suggesting that subhaplogroup B4 originated in either central or southeastern Asia. Finally, we observed two Tuvan and four Negidal mtDNAs belonging to subhaplogroup B5 (Kolman *et al.* 1996; Yao *et al.* 2002). This lineage, in contrast to the other subsets of B, harbours the 10394 *DdeI* site. The complete sequence of one Negidal B5 mtDNA revealed numerous mutations (Figure 2). A similar complete mitochondrial genome was reported in a Japanese cardiomyopathy patient (Shin *et al.* 2000). Lineage B5 is most likely widespread

in central/eastern Asia since it has been previously described at the RFLP/HVS-I level in a few Mongolians (Kolman *et al.* 1996), Chinese (Yao *et al.* 2000; 2002), and one Tuvan (Derenko *et al.* 2000). Overall, our analysis of haplogroup B mtDNAs from southern Siberia has revealed four distinct subhaplogroups (B1, B3, B4 and B5), but only one (B1) is closely related to B2, the branch of B found in Native Americans.

Haplogroup F mtDNAs characterized by the 4732 *RsaI* site, the lack of the 12406 *HpaI/HincII* and 12629 *AvaII* sites, and the HVS-I motif 16189-16232A-16249-16304-16311, were observed in the Tubalar, Tofalar, and Tuvan of the Altai-Sayan, the Mansi of the Lower Ob River, and the Ket and Evenki of the Middle Yenisei River (Derbeneva *et al.* 2002a,c, this study). A related mtDNA type was found in three Tuvan and one Ulchi but it lacked the 4732 *RsaI* site and was very different in HVS-I. Thus, this mtDNA type appears to be more closely related to the one which is common in Tibet (Torroni *et al.* 1994).

Haplogroup Y is characterized by the 7933 *MboI* and 10394 *DdeI* sites, the lack of the 8391 *HaeIII* site, the control-region motif 16126-16519, and was first localized in the Russian Far East (Torroni *et al.* 1993a; Schurr *et al.* 1999). The complete sequence of one Ulchi haplogroup Y mtDNA revealed five coding region transitions (3834, 7933, 8392, 14178, 14693) that distinguish haplogroup Y from other Asian haplogroups of the macro-cluster N (Mishmar *et al.* 2003). Haplogroup Y is restricted to the Lower Amur/Sea of Okhotsk region and adjacent Kamchatka. It reaches its maximum frequency in Sakhalin Island, where it represents 66% of the Nivkhi mtDNAs (Tables 1 and 2). Since Y has a limited geographic range and shows a relatively low level of intra-group diversity, it must be of recent origin, arising long after the migration of the progenitors of Native Americans to the New World.

A new unique "east" Eurasian haplogroup within macrohaplogroup N was found in three Tuvan, six Ulchi and 14 Udegey. It harbours transitions at nps 12705 and 16223 and is defined by the transition at np 5417. Several "west" Eurasian haplogroups, including H, V, J, U4, U5, W, and X, were also detected. Their confinement to the south-west part of Siberia might be indicative of an Upper Paleolithic dispersal from the Middle East/southeastern Europe, the traces of which

have not been erased by subsequent migrations and gene flow. Alternatively, a relatively recent gene flow mediated by women of European/West Asian ancestry could have occurred at the time of the expanding Mongolian Empire.

Macro-haplogroup M mtDNAs

Haplogroup C is defined by the 13262 *AluI* site and the HVS-I motif 16223-16298-16327. It is the most common and widespread throughout Siberia and can be divided into several subhaplogroups. One of these, which we termed C1 (Figure 3), is defined by the HVS-I transition at np 16325. It was found in one Ulchi of the Lower Amur, and has previously been reported in two Mongols and one Kirgiz (Kolman *et al.* 1996; Comas *et al.* 1998). The Ulchi C mtDNA with the HVS-I transition at np 16325 was completely sequenced and compared with Native American and Asian haplogroup C mtDNAs (this study). All were found to share the motif 3552A-9545-11914-13263-14318-16327, but the Ulchi mtDNA, in addition to the transition at np 16325, also harboured the deletion in HVS-II of nps 290-291 ubiquitously present in Amerindian haplogroup C mtDNAs (Ingman *et al.* 2000; Moraga *et al.* 2000; Malhi *et al.* 2001). Network analysis confirmed that the Ulchi mtDNA (C1a) and Native American mtDNA (C1b) are the most similar, while many of the Tuvan, Tofalar, Evenki, Nganasan, Udegey and Koriak complete sequences are distinguished by the HVS-I/coding region motif 2233 ins-6026-11969-15204 (Mishmar *et al.* 2003; this study), and are members of a different cluster that we termed C2. This lineage appears to be the most common and widespread in Siberia (Table 2). An additional lineage of haplogroup C has been discerned through complete sequencing of one of the Nganasan and Ulchi mtDNA samples (this study), and one Tuvan mtDNA with 3460A primary LHON mutation (unpublished data of Volodko *et al.*). We termed this lineage C3 as it lacked the coding region motif 2233 ins-6026-11969-15204 but has the control region variant marked by 16288. At the HVS-I level it has been observed not only in the populations of southern Siberia but also among the circumpolar groups, the Nganasan, Chukchi, Koriak and Itelmen (Starikovskaya *et al.* 1998; Schurr *et al.* 1999; Derbeneva *et al.* 2002b,

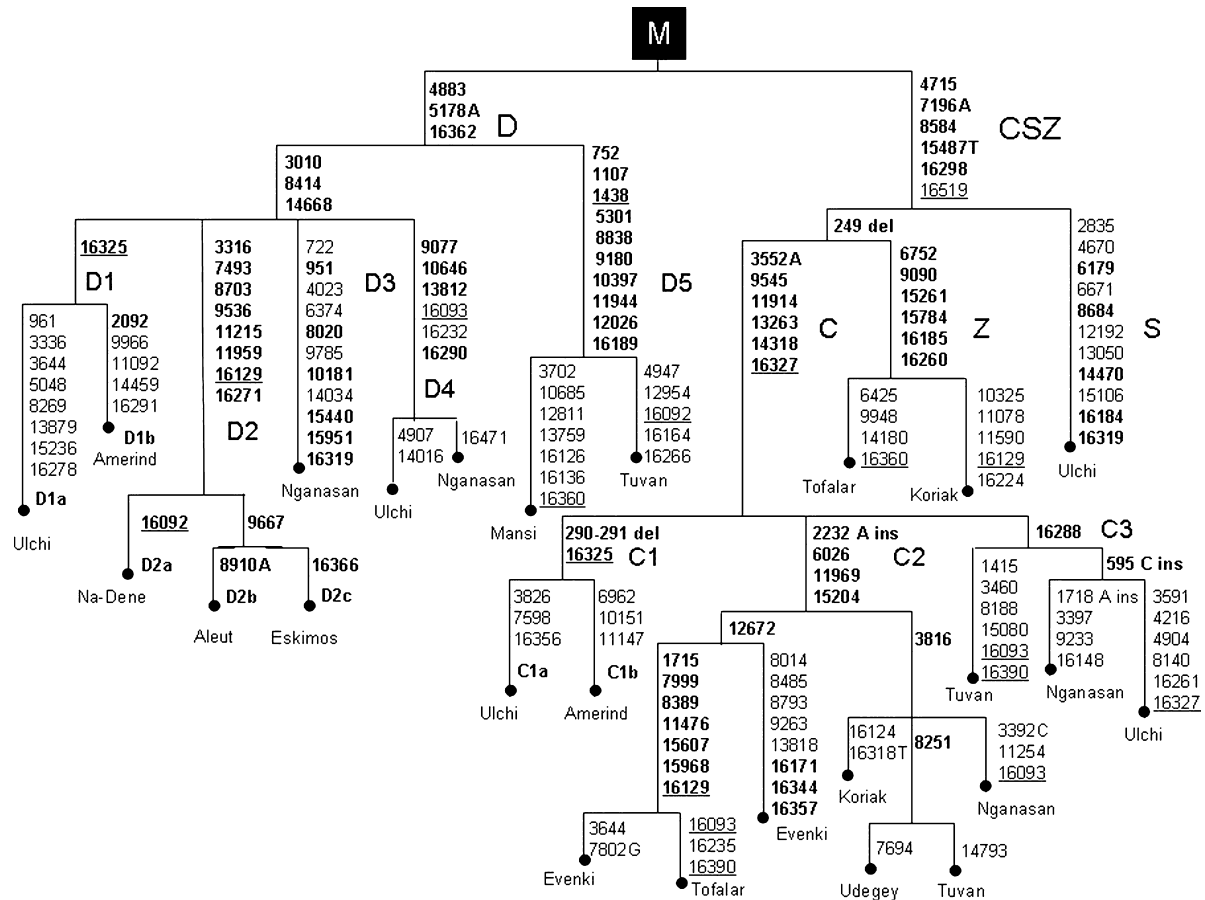


Figure 3 Schematic phylogeny of haplogroup C, D, S and Z mtDNAs found in Native Siberians in comparison with their Native American counterparts. The phylogeny is rooted from macrohaplogroup M. Note that a transition at np 13879 in the Amerindian D1b mtDNA was erroneously shown in the legend of Figure 4 by Derbeneva *et al.* (2002a).

c). The network analysis of haplogroup C2 and C3 complete sequences has revealed several distinct sub-lineages. Thus, the sublineage marked by 3816 (C2) is particularly widespread, being shared by the Buryat, Kirgiz, Koriak, Udegey, Tuvan and Nganasan haplogroup C mtDNA samples that were subjected to complete sequencing (Ingman *et al.* 2000; Mishmar *et al.* 2003; this study). It is worth noting the reversion in HVS-I of the 16327 mutation, which was found to characterize a C haplotype shared by three Tuvan and six Ulchi mtDNA samples (Table 1).

Haplogroup D is defined by the lack of the 5176 *AluI* site and the HVS-I motif 16223–16362. The majority of the haplogroup D haplotypes from southern Siberia did not exhibit sub-haplogroup-specific mutations, and thus they were attributed to a general haplogroup D category. However, we were able to identify at least five

distinct sub-clusters (D1–D5) branching off from the root of D. Lineage D1 is defined by the HVS-I transition at np 16325 and was found exclusively in four Ulchi of the Lower Amur. This transition is typically observed in Native American D mtDNAs (Torroni *et al.* 1993b; Moraga *et al.* 2000; Mahli *et al.* 2001, 2003) but has never been observed in central-eastern Asian populations (Kolman *et al.* 1996; Comas *et al.* 1998; Yao *et al.* 2002; Keyser-Tracqui *et al.* 2003). Thus, the sharing of the mutation at np 16325 in the Lower Amur and the Americas suggests a possible association between the Lower Amur and Native Americans.

Subhaplogroup D2 harbours the 8700 *AluI* site and lacks the 3315 *HaeIII* site in the presence of the HVS-I transition at np 16271. Complete sequence analyses show that overall it is defined by the motif 3316–7493–8703–9536–11215–11959–16129–16271. Lineage

D2 was most likely restricted to a relatively small area of Greater Beringia, since today it is concentrated in the Chukchi and Eskimo-Aleuts (Starikovskaya *et al.* 1998; Derbeneva *et al.* 2002a).

Subhaplogroup D3 at the RFLP/HVS-I level is distinguished by the presence of the 10180 *TaqI* and 15437 *HaeIII* sites, and the transition at np 16319. This lineage is uncommon but widespread in Siberia being represented in the Tubular, Mansi, Nganasan, Tuvan, Evenki, Ulchi, Chukchi, and Siberian Eskimos. One Nganasan D3 mtDNA, which was completely sequenced, exhibited the motif 951-8020-10181-15440-15951-16319 (Derbeneva *et al.* 2002a).

Subhaplogroup D4 is distinguished by the presence of the 10646 *RsaI* site linked with the transition at np 16290. It is common among the Negidal, Ulchi and Nivkhi of the Lower Amur, and has also been observed at low frequencies in the Tubular and Nganasan. The complete sequencing of two D4 mtDNA samples, one Ulchi of the Lower Amur and one Nganasan of the Taimir Peninsula, isolated by a distance of some 5,000 km, has revealed that they share the three coding region transitions, 9077, 10646, and 13812, thus delineating the D4 founding motif (Derbeneva *et al.* 2002a; this study).

Finally, there is subhaplogroup D5 which, in contrast to the other subsets of D, has back mutated at the 10394 *DdeI* and 10397 *AluI* sites. D5 mtDNAs have been observed in three Tubular, one Tuvan and one Evenki. The lack of the diagnostic 10394 *DdeI* and 10397 *AluI* sites has been observed also in one Mansi mtDNA, and the sequencing of the segment encompassing nps 10394-10397 revealed the expected macrohaplogroup M transitions at nps 10398 and 10400. However, a transition at np 10397 was also present that accounted for the simultaneous elimination of the adjacent restriction sites (Derbeneva *et al.* 2002b). The complete sequencing of both Mansi and Tuvan D5 mtDNAs, aside from the transition at np 10397, revealed in their coding regions seven additional transitions, namely at nps 752, 1107, 5301, 8838, 9180, 11944 and 12026 (this study). Furthermore, they differed from other derivatives of the M cluster by lacking the transition at np 1438 (Figure 3). Interestingly, a similar kind of variation is present in one of two Chinese haplogroup D5 mtDNAs that was subjected to complete sequencing by Kong *et al.* (2003). The lack of the 10394 *DdeI* and 10397 *AluI* sites in

haplogroup D mtDNAs was previously reported in the Taiwanese Han, Sabah aborigines (Ballinger *et al.* 1992), Tibetans (Torroni *et al.* 1994), and Mongolians (Kolman *et al.* 1996). A similar haplotype defined by the 10397, 10398 and 10400 variants has been reported among the Chinese Han (Yao *et al.* 2002), thus indicating a northward expansion of D5 from East Asia.

Haplogroup G is defined by the 4830/4831 *HaeII/HhaI* sites and is widespread in Siberian and adjacent Asian populations (Ballinger *et al.* 1992; Torroni *et al.* 1994; Yao *et al.* 2002; present study). It can be subdivided into three distinct subhaplogroups (Tables 1 and 2). Subhaplogroup G1 is distinguished by the 8198 *AluI* site and the transition at np 16017 in HVS-I. Geographically, G1 is restricted to the Sea of Okhotsk region, being the most prevalent in the Koriak and Itelmen of Kamchatka (Schurr *et al.* 1999). A variant, observed solely in four Chukchi of adjacent Chukotka, lacks the 8198 *AluI* site (Starikovskaya *et al.* 1998). Lineage G2 is characterized by the lack of the 7598 *HhaI* site, caused by a G-A transition at np 7600, and the HVS-I motif 16223-16278-16362. A Koriak G1 mtDNA and a Tuvan G2 mtDNA studied through complete sequencing were found to share the coding region motif 709-4833-5108-14569, thus forming the founding haplotype for haplogroup G (Mishmar *et al.* 2003; this study). In addition, the Koriak G1 showed the distinguishing motif 8200-12361-12972-15323-15497, while the Tuvan G2 harboured 1189-5601-7600-9377-9575-12372-13563-14198-15355. Subhaplogroup G2 appears to be widespread in central/eastern Asia as Korean haplogroup G sequences also harbour the motif 5601-7600-9377-9575-13563 (Snall *et al.* 2002). It has also been observed in the Mansi of Siberia's northwest (Derbeneva *et al.* 2002b), though it is rare or absent in the Russian Far East. Subhaplogroup G3 is distinguished from the root of G by transitions at nps 12408, 16274 and 16390 and has been seen in several Tuvan and one Buryat. The identity of this lineage was confirmed by sequencing the 12406 *HpaI-HincII* region which revealed that the distinguishing mutation is a transition at np 12408. However, one Tuvan G3 mtDNA was found to lack this transition, together with the 16390 variant (Table 1).

One haplogroup M lineage was characterized by the HVS-I motif 16223-16298-16319 and the 14465 *AccI*

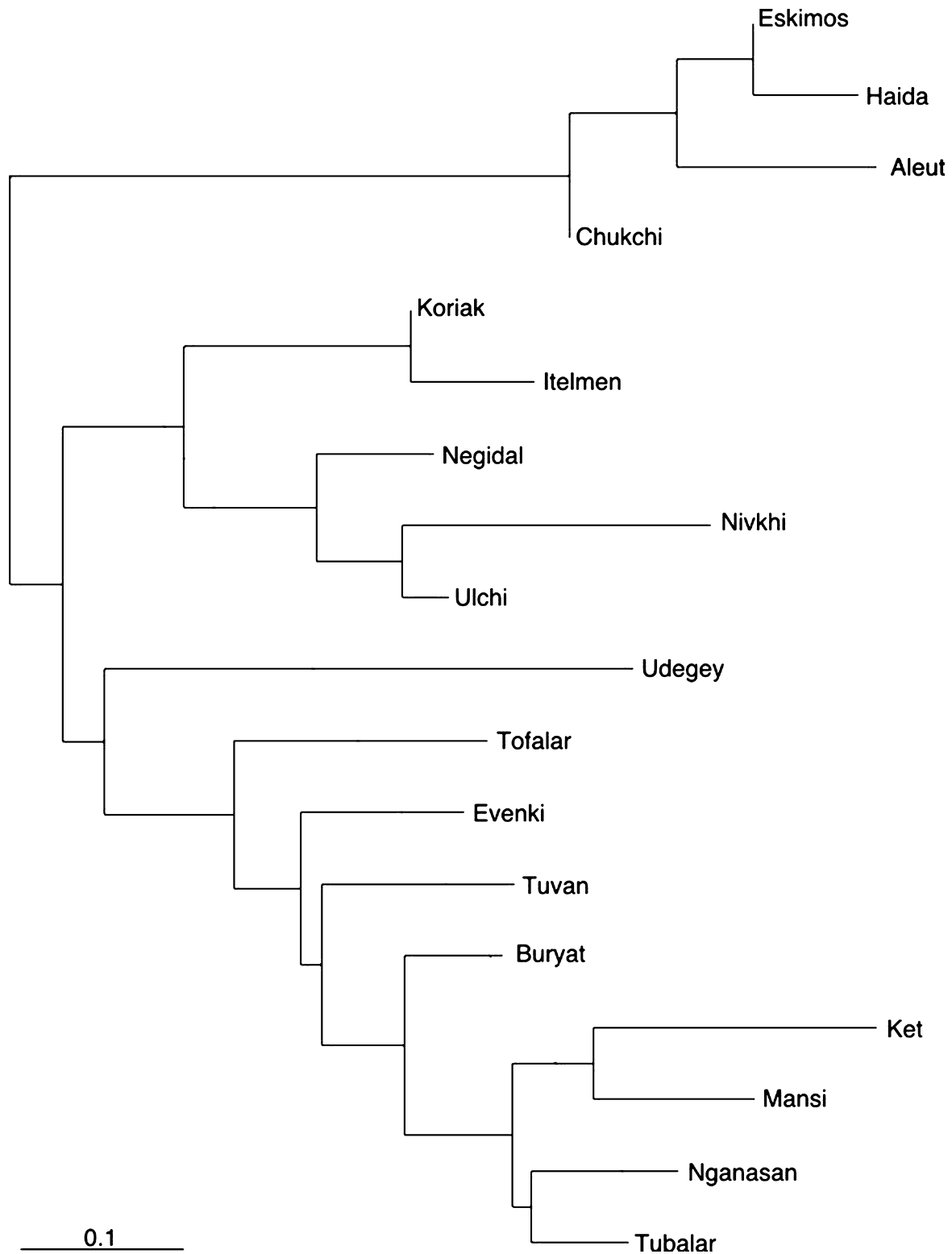


Figure 4 Phylogenetic relationships between 16 Siberian and two Native American populations of the North Pacific Rim (Aleut and Haida), determined by neighbor-joining using the frequencies of the 31 observed subhaplogroups. Haplogroup composition and frequencies in Haida and Aleut mtDNAs were adopted from Ward *et al.* (1993), and Derbeneva *et al.* (2002) and Rubicz *et al.* (2003) respectively.

site. This novel lineage, which we termed haplogroup S, was observed in one Buryat, seven Tofalar, four Ulchi and seven Udegey. A similar mtDNA type, designated M8a, has been recently described in China by Yao *et al.* (2002). One of the Ulchi haplogroup S mtDNA was completely sequenced. It shared with haplogroup C the coding region motif 4715–71976A–8584–15487T, revealing that these mutations predate the radiation of haplogroups C, S and Z. In addition, the Ulchi mtDNA also harboured the coding region variants 2835, 4670, 6179, 6671, 7196A, 8684, 12192, 13050, 14470 and 15106. Three of these (6179, 8684, and 14470) are shared with an Asian mtDNA described by Herrnstadt *et al.* (2002). The confinement of haplogroup S to southern and southeastern Siberia, inhabited by Mongolic-, Turkic-, and Tungusic-speaking populations which are members of the Altaic linguistic family, implies that it probably arose in one of the hunting groups ancestral to the Altaic-speakers. We presume that this lineage originated in the putative homeland (Mongolia/Manchuria) of the Altaic-speakers and was then dispersed by their geographical expansion.

Haplogroup Z is distinguished by the HVS-I motif 16185–16260 and was first identified in the Itelmen and Koriak of Kamchatka (Schurr *et al.* 1999). This haplogroup was also noted amongst the Entsi/Nganasan and Ket of the Lower Yenisei (Derbeneva *et al.* 2002c). In the southern part of Siberia, we found this haplogroup in one Northern Altaian (1.4%), one Tuvan (1.7%), one Buryat (4.0%), and five Tofalar (10.9%) mtDNAs. The sequence of one Tofalar and one Koriak haplogroup Z mtDNA revealed that they differ from the common root of C–Z for the 6752, 9090, 15261, 15784, 16185, 16260 variants (Mishmar *et al.* 2003; this study), mutations that they also share with the Finnish/Saami haplogroup Z mtDNAs (Finnila *et al.* 2001; Majamaa, personal communication). Thus, these six transitions are characteristic of the ancestral haplogroup Z haplotype. Based on the distribution of haplogroups C, S, Z and the number of variants encompasses, we postulate that south-central Siberia was the place of origin for the ancestral haplotype, and the populations bearing haplogroup Z must have spread into northern Eurasia after the ancestors of the Amerindians left the region.

The lineages of macro-haplogroup M were also found mainly in the populations of extreme south and south-

eastern Siberia (Table 1). One lineage marked by the 9820 *HinfI* site was initially detected in the Vietnamese and Taiwanese Han (Ballinger *et al.* 1992). Another lineage of M harbouring the 3391 *HaeIII* site (T3394C) in conjunction with the HVS-I motif 16223–16234–16316–16362 was found in four Udegey mtDNAs. A similar motif was recently observed in Chinese M mtDNAs, suggesting an East Asian origin for this lineage (Yao *et al.* 2002).

NJ analysis

The genetic distances between interior Siberian and north Pacific populations were computed from their subhaplogroup frequencies (Table 2), and analysed with the help of a dendrogram (Figure 4). The tree is drawn by mid-point rooting. The dendrogram supports the hypothesis that the Chukchi, Eskimos, Aleut and Haida, though quite divergent from one another, represent a detached population cluster isolated by sufficient time to be relatively distinct over the average genetic background of Siberia. In addition, the branching patterns of the Lower Amur and adjacent Okhotsk Sea coast/Kamchatkan populations, relative to those located in Western and Middle Siberia, may reflect relatively recent demic expansions which have brought haplogroups G, S, Y and Z into the Sea of Okhotsk/Kamchatka region. The observed geographical distribution of these haplogroups (Tables 1 and 2) would be expected from anthropological, archaeological and linguistic evidence indicating the expansion of the Altaic-speakers from the Lower Amur/Manchuria region during the Neolithic (Krauss, 1988; Turner, 1994; Janhunen, 1996; Kozintsev *et al.* 1999).

Discussion

A large variety of mtDNA haplogroups and subhaplogroups have been found in Siberia. However, only one or at most two subhaplogroups of A, B, C, and D appear to have contributed to the human colonization of the Americas, confirming that only a limited number of founder mtDNAs gave rise to the ancestral Native American populations (Wallace *et al.* 1985; Neel *et al.* 1994; Starikovskaya *et al.* 1998). Indeed, the evidence provided by the analysis of variation in the Y chromosome types, using almost the same set of Siberian DNAs,

is also consistent with Native Americans being founded by a limited number of individuals (Lell *et al.* 2002). Furthermore, many *HLA* class II genes (i.e., DRB1 allelic lineages) observed in the Siberian interior are absent in the New World (Uinuk-ool *et al.* 2002, 2004; Volodko *et al.* 2003).

Many of the mtDNA haplotypes found in Siberia are shared between culturally and linguistically distinct populations, indicating that extensive gene flow has occurred from the Ural Mountains to the Pacific Ocean, and from Mongolia/Manchuria/Southeastern Siberia (former Greater Manchuria) to the upper Arctic, since the early Holocene. However, still there exists a remarkable geographical and even tribal specificity for some of the Siberian subhaplogroups, and this provides major clues about the origin of Native Americans.

The trans-Beringian Relationships of Haplogroup A and B mtDNAs

Almost all Native American haplogroup A mtDNAs are characterized by the three distinguishing mutations, at nps 8027, 12007 and 16111, thus belonging to subhaplogroup A2 (Ingman *et al.* 2000; Silva *et al.* 2003; Mishmar *et al.* 2003). The phylogenetic relationship of A1 and A2 mtDNAs and their pattern of spatial distribution in Siberia/Asia (Starikovskaya *et al.* 1998; Schurr *et al.* 1999; Yao *et al.* 2002; Kong *et al.* 2003; this study) suggests that the A1 mtDNAs observed in the Mansi and Ket are part of an A1 northward dispersal that presumably originated in the Altai-Sayan region, and gave rise to A2 after expanding into eastern Beringia.

Haplogroup B is extremely diverse; several subsets of this haplogroup (B1, B3, B4, B5) that share a common root, but diverged early with very different subsequent evolutionary histories, have been found in southern Siberia. Subhaplogroup B1 is rare, but it is the only one that shares the coding region motif 827-4820-13590-15535 with B2, the Native American branch of haplogroup B. In our Siberian sample, B1 mtDNAs were found only in one Tuvan and three Tubalar, suggesting that the common ancestor of Siberian B1 and Amerind B2 could be traced back to the Altai-Sayan Upland. The failure to identify B1 mtDNAs north and northeast of the Altai-Sayan may be explained by their extinction

due to genetic drift and/or selection in small populations inhabiting Siberian arctic and sub-arctic prior to the last glacial maximum (>24,000 YBP).

Siberian Affinities of Native American Haplogroup C and D mtDNAs

A combined phylogenetic analysis of mtDNAs belonging to haplogroups C, D, S and Z revealed that, relative to the CRS, they all harbour the distinguishing M motif: 8701-9540-10398-10400-10873-12705-14783C-15043-15301-16223. However, while haplogroup D forms a cluster by itself within the macrogroup M, haplogroups C, S, and Z are sister groups sharing the motif 4715-7196A-8584-15487T-16298 (Figure 3). The origin of the Asian CSZ cluster appears to be rooted in central/eastern Asia during the Upper Paleolithic period, and to have later emerged as the progenitor of the Native American haplogroup C. However, since neither S nor Z mtDNAs have been reported in Native Americans mtDNAs, these haplogroups could have appeared in the Altai-Sayan-Baikal Upland and the Amur region after the immediate ancestors of Paleoindians had already migrated from the southern Siberia.

While lineages C2 and C3 are present at significant frequencies in Uralic, Ketic, Altaic, Chukchi/Koriak speakers, they are absent in Native Americans. The geographic distribution and phylogenetic relationships of C2 and C3 suggest that they arose in the Altai-Sayan-Baikal area. The subhaplogroup C1a mtDNA found in one Ulchi, similarly to Amerindian C1b mtDNAs, harbours the HVS-I marker 16325 and lack C2 and C3 markers. Moreover, C1a and C1b share the deletion of nps 290-291. This finding may imply that C1 and its distinguishing mutations (16325 and the deletion of nps 290-291) originated in the Amur River region and that from that region, the Native American branch C1b arose.

As was found in the course of this and our previous studies (Starikovskaya *et al.* 1998; Derbeneva *et al.* 2002 a, b, c), Siberian haplogroup D mtDNAs belong to a wide range of distinct subhaplogroups. Subhaplogroup D1a, which was observed in four Ulchi, is the only one which is more closely related to D1b, the Amerindian-specific branch of haplogroup D, than the other

sublineages of D in Siberia/Asia. However, Amerindian D1b mtDNAs are commonly associated with a transition at np 2092 (Ingman *et al.* 2000; Herrnstadt *et al.* 2002), which was not found in the Siberian D1a mtDNAs. The finding of both C1a and D1a only on the southwestern edge of former Beringia is intriguing, and indicates that the Amur River region could have been the main Siberian source for Amerindian haplogroups C and D. In contrast, subhaplogroup D2 is confined to Chukotka, Alaska and the Aleutian archipelago with a great geographical/tribal specificity; D2a in the Na-Dene, D2b in the Aleut and D2c in the Eskimos (Figure 3). The geographic specificity and phylogeny of haplogroup D complete sequences (Derbeneva *et al.* 2002a; this study) support the refugial hypothesis (Rogers *et al.* 1991), which proposes that the founding population of Eskimo-Aleut and Na-Dene Indians originated in the eastern Beringian/Alaskan refuge area during the early postglacial period. However, the genealogical relationships among the lineages of the exceptionally diverse haplogroup D cluster are not explicitly clear. Some intermediate haplotypes may be missing, suggesting a relatively long evolutionary history of Native American D.

Many factors must have influenced the success of human biological adaptation to the harsh glacial environment of Beringia and further movements into the New World. Late glacial and early postglacial climatic fluctuations have influenced population size and structure and, hence, the number of founder haplogroups. Natural selection might have either impoverished or enriched certain mtDNA haplogroups and subhaplogroups as people migrated into sub-arctic and then arctic regions (Mishmar *et al.* 2003; Ruiz-Pesini *et al.* 2004).

Conclusion

This study revealed that the remnants of ancient east Eurasian populations are still evident in Siberia due to the occurrence of lineages A1 and B1 in the Altai-Sayan area, and C1 and D1 in the Lower Amur. Assuming low effective population size, reduced diversity and a uniform distribution of mtDNA lineages prior to the last glacial maximum, the lineages A1, B1, C1 and D1 constituted the core genetic makeup of the Siberian progenitors of Paleoindians that spread northwards as a

uniform entity. However, it is more likely that the current mtDNA differences between Altai-Sayan Upland, where A1 and B1 occur, and the Amur region, where the traces of C1 and D1 are found, are due to genetic differentiation of the Upper Paleolithic inhabitants of Siberia. Hence, while subhaplogroups A1 and B1 could be a major part of the population dispersal from the Altai-Sayan area downstream via the Yenisei and Lena rivers and then toward eastern Beringia and Alaska, the migration originated in the Amur river basin brought the founders of C1 and D1 to the Americas along the Siberian Pacific. Indeed, a mosaic-like picture of different cultural variants (mammoth, bison and reindeer hunters, fishing and sea-mammal hunting populations) is evidenced by the diverse lithic industries that persisted in the river valleys and along the Siberian Arctic and Pacific regions (Dikov, 1994; Khlobystin, 1998). Moreover, the earliest traces of modern human occupation of northeastern Siberia (at 71°N) indicate that the ancestors of Paleoindians may have colonized the Lower Yenisei and Lena rivers and the now submerged Laptev, Eastern Siberian and Chukchi Sea coastal shelves during a relatively warm interstadial episode around 30,000–28,000 YBP (Vasil'ev, 2002; Pitulko *et al.* 2004). Since the glaciation of mountainous Chukotka hampered faunal and human migrations from the Lena river basin, this was the only practical route from the Middle Siberian Plateau to Alaska. Consequently, the bearers of the Amur microlithic tradition (Derev'anko & Volkov, 1997), lacking subhaplogroup B1, might have reached and occupied the maritime region of southwestern Beringia generally later. The phylogeny of haplogroup A, B, C and D lineages revealed through this study is consistent with the hypothesis that the direct ancestors of Native Americans were a hybrid product of different Siberian groups that had migrated to eastern Beringia at different times and following different routes (Kunz & Reanier, 1994).

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References

- Andrews, R. M., Kubacka, I., Chinnery, P. F., Lightowlers, R. N., Turnbull, D. M. & Howell, N. (1999) Reanalysis and revision of the Cambridge reference sequence for human mitochondrial DNA. *Nat Genet* **23**, 147.
- Ballinger, S. W., Schurr, T. G., Torroni, A., Gan, Y. Y., Hodge, J. A., Hassan, K., Chen, K.-H. & Wallace, D. C. (1992) Southeast Asian mitochondrial DNA analysis reveals genetic continuity of ancient mongoloid migrations. *Genetics* **130**, 139–152.
- Brown, M. D., Hosseini, S. H., Torroni, A., Bandelt, H.-J., Allen, J. C., Schurr, T. G., Scozzari, R., Cruciani, F. & Wallace, D. C. (1998) mtDNA haplogroup X: an ancient link between Europe/Western Asia and North America? *Am J Hum Genet* **63**, 1852–1861.
- Black, L. T. (1988) Peoples of the Amur and maritime regions. In: *Crossroads of continents: cultures of Siberia and Alaska* (eds W. W. Fitzhugh & A. Crowell), pp 24–31. Smithsonian Institution Press, Washington, DC.
- Comas, D., Calafell, F., Mateu, E., Perez-Lezaun, A., Bosch, E., Martinez-Arias, R., Clarimon, J., Facchini, E., Fiori, G., Luiselli, D., Pettener, D. & Bertranpetit J. (1998) Trading genes along the silk road: mtDNA sequences and the origin of central Asian populations. *Am J Hum Genet* **63**, 1824–38.
- Dalton, R. (2003) The coast road. *Nature* **422**, 10–12.
- Derbeneva, O. A., Sukernik, R. I., Volodko, N. V., Hosseini, S. H., Lott, M. T. & Wallace, D. C. (2002 a) Analysis of mitochondrial DNA diversity in the Aleuts of the Commander Islands and its implications for the genetic history of Beringia. *Am J Hum Genet* **71**, 415–421.
- Derbeneva, O. A., Starikovskaya, E. B., Wallace, D. C. & Sukernik, R. I. (2002 b) Traces of early Eurasians in the Mansi of northwest Siberia revealed by mitochondrial DNA analysis. *Am J Hum Genet* **70**, 1009–1014.
- Derbeneva, O. A., Starikovskaya, E. B., Volodko, N. V., Wallace, D. C. & Sukernik, R. I. (2002 c) Mitochondrial DNA variation in Kets and Nganasans and the early peopling of Northern Eurasia. *Genetika (Russian J. Genetics)* **38**, 1554–1560.
- Derenko, M. V., Malyarchuk, B. A., Dambueva, I. K., Shaikhaev, G. O., Dorzhu, C. M., Nimaev, D. D. & Zakharov, I. A. (2000) Mitochondrial DNA variation in two south Siberian aboriginal populations: implications for the genetic history of North Asia. *Hum Biol* **72**, 945–973.
- Derenko, M. V., Grzybowski, T., Malyarchuk, B. A., Czarny, J., Miscicka-Sliwka, D. & Zakharov, I. A. (2001) The presence of mitochondrial haplogroup X in Altayans from south Siberia. *Am J Hum Genet* **69**, 237–241.
- Derenko, M. V., Grzybowski, T., Malyarchuk, B. A., Dambueva, I. K., Denisova, G. A., Czarny, J., Dorzhu C. M., Kakpakov V. T., Miscicka-Sliwka, D. Wozniak M., & Zakharov, I. A. (2003) Diversity of mitochondrial DNA lineages in South Siberia. *Ann Hum Genet.* **67**, 391–411.
- Derev'anko, A. P. & Volkov, P. V. (1997) Evolution of the paleoeconomy of the ancient population of the Amur region (from the Upper Palaeolithic to Neolithic)//Suyanggae and her neighbors – Chungju, pp 35–44.
- Dikov, N. N. (1994) The Palaeolithic of Kamchatka and Chukotka and the problem of the peopling of America. In: *Crossroads of continents: cultures of Siberia and Alaska* (eds W.W. Fitzhugh & A. Crowell), pp 87–95. Smithsonian Institution Press, Washington, DC.
- Finnila, S., Lehtonen, M. S. & Majamaa, K. (2001) Phylogenetic network for European mtDNA. *Am J Hum Genet* **68**, 1475–1484.
- Forster, P., Harding, R., Torroni, A. & Bandelt, H.-J. (1996) Origin and evolution of Native American mtDNA variation: a reappraisal. *Am J Hum Genet* **59**, 935–945.
- Goebel T., Waters, M. R. & Dikova, M. (2003) The archeology of Ushki Lake, Kamchatka, and the Pleistocene peopling of the Americas. *Science* **301**, 501–505
- Herrnstadt, C., Elson, J. L., Fahy, E., Preston, G., Turnbull, D. M., Anderson, C., Ghosh, S. S., Olefsky, J. M., Beal, M. F., Davis, R. E. & Howell, N. (2002) Reduced-median-network analysis of complete mitochondrial DNA coding-region sequences for the major African, Asian, and European haplogroups. *Am J Hum Genet* **70**, 1152–1171.
- Ingman, M., Kaessmann, H., Paabo, S. & Gyllensten, U. (2000) Mitochondrial genome variation and the origin of modern humans. *Nature* **408**, 708–713.
- Janhunen J. Manchuria. An Ethnic History. Printed by the Finno-Ugrian Society at Vammalan Kirjapaino Oy, Helsinki (1996)
- Keyser-Tracqui, C., Crubezi, E. & Ludes, B. (2003) Nuclear and mitochondrial analysis of a 2,000-year-old necropolis

- in the Egyn Go Valley of Mongolia. *Am J Hum Genet* **73**, 247–260.
- Khlobystin, L. P. (1998) Ancient history of Taimyr and the formation of the North Eurasian cultures. Edited by V.V. Pitulko and V. Ya. Shumkin. Publisher - Dmitriy Bulanin. St. Petersburg (in Russian).
- Kolman, C. J., Sambuughin, N. & Bermingham, E. (1996) Mitochondrial DNA analysis of Mongolian populations and implications for the origin of New World founders. *Genetics* **142**, 1321–1334.
- Kong, Q.-P., Yao, Y.-G., Sun, C., Bandelt, H.-J., Zhu, C.-L. & Zhang, Y.-P. (2003) Phylogeny of East Asian mitochondrial DNA lineages inferred from complete sequences. *Am J Hum Genet* **73**, 671–676.
- Kozintsev, A. G., Gromov, A. V. & Moiseyev, V. G. (1999) Collateral relatives of American Indians among the Bronze Age populations of Siberia? *Am J Phys Anthropol* **108**, 193–204.
- Krauss, M. E. (1988) Many tongues-ancient tales. Peoples of the Amur and maritime regions. In: *Crossroads of continents: cultures of Siberia and Alaska* (eds W.W. Fitzhugh & A. Crowell), pp 145–150. Smithsonian Institution Press, Washington, DC.
- Kuntz, M. L. & Reanier, R. E. (1994) Paleoindians in Beringia: evidence from arctic Alaska. *Science* **263**, 660–662.
- Lell, J. T., Sukernik, R. I., Starikovskaya, Y. B., Su, B., Jin, L., Schurr, T. G., Underhill, P. A. & Wallace, D. C. (2002) The dual origin and Siberian affinities of Native American Y chromosomes. *Am J Hum Genet* **70**, 192–206.
- Levin, M. G. & Potapov, L. P. (1964) *The Peoples of Siberia*. The University of Chicago Press. Chicago and London.
- Macaulay, V., Richards, M., Hickey, E., Vega, E., Cruciani, F., Guida, V., Scozzari, R., Bonne-Tamir, B., Sykes, B. & Torroni, A. (1999) The emerging tree of west Eurasian mtDNAs: a synthesis of control-region sequences and RFLPs. *Am J Hum Genet* **64**, 232–249.
- Malhi, R. S., Schultz, B. A. & Smith, D. G. (2001) Distribution of mitochondrial DNA lineages among Native American tribes of northeastern North America. *Hum Biol* **73**, 17–55.
- Malhi, R. S., Mortensen, H. M., Eshleman, J. A., Kemp B. M., Lorenz J. G., Kaestle F. A., Johnson, J. R., Gorodezky, C. & Smith, D. G. (2003) Native American mtDNA prehistory in the American Southwest. *Am J Phys Anthropol* **120**, 108–124.
- Middendorf, A. (1869) *Travelling to the north and east of Siberia (Puteshestvi'e na Sever i vostok Sibiri)*. Part 2. S-Petersburg (In Russian). pp 707–711.
- Mishmar, D., Ruiz-Pesini, E., Golik, P., Macaulay, V., Clark, A. G., Hosseini, S., Brandon, M., Easley, K., Chen, E., Brown, M. D., Sukernik, R. I., Olckers, A. & Wallace, D. C. (2003) Natural selection shaped regional mtDNA variation in humans. *Proc Natl Acad Sci USA* **100**, 171–176.
- Moraga, M. L., Rocco, P., Miquel, J. F., Nervi, F., Llop, E., Chakraborty, R., Rothhammer, F. & Carvallo, P. (2000) Mitochondrial DNA polymorphisms in Chilean aboriginal populations: implications for the peopling of the southern cone of the continent. *Am J Phys Anthropol* **113**, 19–29.
- Neel, J. V., Biggar, R. J. & Sukernik, R. I. (1994) Virologic and genetic studies relate Amerind origins to the indigenous people of the Mongolia/Manchuria/southeastern Siberia region. *Proc Natl Acad Sci USA* **91**, 10737–10741.
- Pitulko, V. V., Nikolsky, P. A., Giry, E. Yu., Basilyan, A. E., Tumskey, V. E., Koulakov, S. A., Astakhov, S. N., Pavlova, E. Yu. & Anisimov, M. A. (2004) The Yana RHS site: humans in the Arctic before the Last Glacial Maximum. *Science* **303**, 52–56.
- Rogers, R. A., Rogers, L., Hoffmann, R. & Martin, L. D. (1991) Native American biological diversity and the biogeographic influence of Ice Age refugia. *J Biogeogr* **18**, 623–630.
- Rubicz, R., Schurr T. G., Babb P. L. & Crawford, M. H. (2003) Mitochondrial DNA variation and the origins of the Aleuts. *Hum Biol* **75**, 809–835.
- Ruiz-Pesini, E., Mishmar, D., Brandon, M., Procaccio V. & Wallace, D. C. (2004) Effects of purifying and adaptive selection on regional variation in human mtDNA. *Science* **303**, 223–226.
- Saillard, J., Forster, P., Lynnerup, N., Bandelt, H. J. & Norby, S. (2000) mtDNA variation among Greenland Eskimos: the edge of the Beringian expansion. *Am J Hum Genet* **67**, 718–726.
- Schurr, T. G., Sukernik, R. I., Starikovskaya, Y. B. & Wallace, D. C. (1999) Mitochondrial DNA variation in Koriaks and Itel'men: Population replacement in the Okhotsk Sea - Bering Sea region during the Neolithic. *Am J Phys Anthropol* **108**, 1–42.
- Shin, W. S., Tanaka, M., Suzuki, J., Hemmi, C. & Toyo-Oka, T. (2000) A novel homoplasmic mutation in mtDNA with a single evolutionary origin as a risk factor for cardiomyopathy. *Am J Hum Genet* **67**, 1617–1620.
- Snall, N., Savontaus, M.-L., Kares, S., Lee, M. S., Cho, E. K., Rinne, J. O. & Huoponen, K. (2002) A rare mitochondrial DNA haplotype observed in Koreans. *Hum Biol* **74**, 253–262.
- Silva, W. A. Jr., Bonatto, S. L., Holanda, A., Ribeiro-Dos-Santos, A. K., Paixao, B. M., Goldman, G. H., Abe-Sandes, K., Rodriguez-Delfin, L. A., Barbosa, M., Pako-Larson, M. L., Petzl-Erler, M. L., Valente, V., Santos, S. E. B. & Zago, M. (2003) Correction: mitochondrial DNA variation in Amerindians. *Am J Hum Genet* **72**, 1346–1348.
- Starikovskaya, Y. B., Sukernik, R. I., Schurr, T. G., Kogelnik, A. & Wallace, D. C. (1998) Mitochondrial DNA diversity in Chukchi and Siberian Eskimos: Implications for the

- genetic history of ancient Beringia and the peopling of the New World. *Am J Hum Genet* **63**, 1473–1491.
- Sukernik, R. I., Shurr, T. G., Starikovskaya, Y. B. & Wallace, D. C. (1996) Mitochondrial DNA variation in native inhabitants of Siberia, with reconstructions of the evolutionary history of American Indians. Restriction polymorphism. *Genetika (Russian J. Genetics)* **32**, 432–439.
- Takezaki, N. & Nei, M. (1996) Genetic distances and reconstruction of phylogenetic trees from microsatellite DNA. *Genetics* **144**, 389–399.
- Takezaki, N. NJBAFD: neighbor-joining tree constructions from allele frequency data.
- Torroni, A., Sukernik, R. I., Shurr, T. G., Starikovskaya, Y. B., Cabell, M. F., Crawford, M. H., Comuzzie, A. G. & Wallace, D. C. (1993 a) Analysis of mitochondrial DNA variation of indigenous Siberians confirms their genetic diversity and reveals distinctive affinity with Native Americans. *Am J Hum Genet* **53**, 591–608.
- Torroni, A., Schurr, T. G., Cabell, M. F., Brown, M. D., Neel, J. V., Larsen, M., Smith, D. G., Vullo, C. M. & Wallace, D. C. (1993 b). Asian affinities and continental radiation of the four founding Native American mtDNAs. *Am J Hum Genet* **53**, 563–590.
- Torroni, A., Miller, J. A., Moore, L. G., Zamudio, S., Zhuang, J. G., Droma, T. & Wallace, D. C. (1994) Mitochondrial DNA analysis in Tibet: implications for the origin of the Tibetan population and its adaptation to high altitude. *Am J Phys Anthropol* **93**, 189–199.
- Torroni, A., Huoponen, K., Francalacci, P., Petrozzi, M., Morelli, L., Scozzari, R., Obinu, D., Savontaus, M.-L. & Wallace, D. C. (1996) Classification of European mtDNAs from analysis of three European populations. *Genetics* **144**, 1835–1850.
- Torroni, A., Bandelt, H. J., D'Urbano, L., Lahermo, P., Moral, P., Sellitto, D., Rengo, C., Forster, P., Savontaus, M. L., Bonne-Tamir, B. & Scozzari, R. (1998) mtDNA analysis reveals a major late Palaeolithic population expansion from southwestern to northeastern Europe. *Am J Hum Genet* **62**, 1137–1152.
- Turner, C. G. II (1994) New dental anthropological observations relevant to the human population system of the Greater Beringian Realm. In: *Anthropology of the North Pacific Rim*. (eds W.W. Fitzhugh & V. Chassonet), pp 97–106. Smithsonian Institution Press. Washington and London.
- Uinuk-ool, T. S., Takezaki, N., Sukernik, R. I., Nagl, S. & Klein, J. (2002) Origin and affinities of indigenous Siberian populations as revealed by *HLA* class II gene frequencies. *Hum Genet* **110**, 209–226.
- Uinuk-ool, T. S., Takezaki, N., Derbeneva, O. A., Volodko N. V. & Sukernik (2004) Variation of *HLA* class II genes in the Nganasan and Ket, two aboriginal Siberian populations. *Eur. J. Immunogenet* **31**, 43–51.
- Vasil'ev, S. A., Kuzmin, Y. V., Orlova, L. A. & Dementiev, V. N. (2002) Radiocarbon-based chronology of the Palaeolithic in Siberia and its relevance to the peopling of the New World. *Radiocarbon* **44**, 503–530.
- Volodko, N. V., Derbeneva, O. A., Uinuk-ool, T. S. & Sukernik, R. I. (2003) Genetic history of Aleuts of the Commander Islands as revealed by *HLA* Class II gene frequencies. *Genetika (Russian J. Genetics)* **39**, 1710–1718 (in Russian).
- Wallace, D. C., Garrison, K. & Knowler, W. C. (1985) Dramatic founder effects in Amerindian mitochondrial DNAs. *Am J Phys Anthropol* **68**, 149–155.
- Ward, R. H., Redd, A., Valencia, D., Frazier, B. & Paabo, S. (1993) Genetic and linguistic differentiation in the Americas. *Proc Natl Acad Sci USA* **90**, 10663–10667.
- West, F. H. (1996) *American beginnings, the prehistory and paleoecology of Beringia*. University of Chicago Press, Chicago.
- Wright, H. E. Jr. (1991) Environmental condition for Paleoindian immigrations. In: *The first Americans: search and research* (eds T.D. Dillihay & D.J. Meltzer), pp 113–135. CRS, Boca Raton, FL.
- Yao, Y.-G., Watkins, W. S. & Zhang, Y.-P. (2000) Evolutionary history of the mtDNA 9-bp deletion in Chinese populations and its relevance to the peopling of East and Southeast Asia. *Hum Genet* **107**, 504–512.
- Yao, Y.-G., Kong, Q.-P., Bandelt, H.-J., Kivisild, T. & Zhang, Y.-P. (2002) Phylogeographic differentiation of mitochondrial DNA in Han Chinese. *Am J Hum Genet* **70**, 635–651.

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