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Terminal Pleistocene Alaskan genome reveals first founding population of Native Americans

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Despite broad agreement that the Americas were initially populated via Beringia, when and how this happened is debated 1–5. Key to this debate are human remains from Late Pleistocene Alaska. The first and only such remains were recovered at Upward Sun River (USR), and date to ~11.5 kya 6,7. We sequenced the USR1 genome to an average coverage of ~17X. We find USR1 is most closely related to Native Americans, but falls basal to all previously sequenced contemporary and ancient Native Americans 1,8,9. As such, USR1 represents a distinct Ancient Beringian (AB)
population. Using demographic modelling we infer the AB population and ancestors of other Native Americans descend from a single founding population that initially split from East Asians $\sim 36 \pm 1.5$ kya, with gene flow persisting until $\sim 25 \pm 1.1$ kya. Gene flow from ancient north Eurasians into all Native Americans took place 25-20 kya, with AB branching off $\sim 22-18.1$ kya. Our findings support long-term genetic structure in ancestral Native Americans, consistent with the Beringian Standstill Model $^{10}$. We find that the basal Northern (NNA) and Southern (SNA) branches, to which all other Native Americans belong, diverged $\sim 17.5-14.6$ kya, likely south of the North American ice sheets. After 11.5 kya, some NNA populations received gene flow from a Siberian population most closely related to Koryaks, but not Paleoeskimos$^1$, Inuit or Kets$^{11}$, and that Native American gene flow into Inuit was via NNA and not SNA groups$^1$. Our findings further suggest the far northern North American presence of NNA is from a back migration that replaced or absorbed the initial AB founding population.

The peopling of the Americas, and particularly the population history of Beringia, the land bridge that connected far northeast Asia to northwestern North America during the Pleistocene, remains unresolved $^{2,3}$. Humans were present in the Americas south of the continental ice sheets by $\sim 14.6$ kya $^{12}$, indicating they traversed Beringia earlier, possibly around the Last Glacial Maximum (LGM). Then, the region was marked by harsh climates and glacial barriers $^5$, which may have led to the isolation of populations for extended periods, and at times complicated dispersal across the region $^{13}$. Still controversial are questions of whether and how long Native American ancestors were isolated from Asian groups in Beringia prior to entering the Americas $^{2,10,14}$; if one or more early migrations gave rise to the founding population of Native Americans $^{1-4,8,15}$ (it is commonly agreed Paleoeskimos and Inuit represent separate and later migrations $^{1,16,17}$); and, when and where the basal split between SNA and NNA occurred. Unresolved too is whether the genetic affinity between some SNA groups and indigenous Australasians $^{2,3}$, reflects migration by non-Native Americans $^{3,4,15}$, early population structure within the first Americans $^3$, or later gene flow $^2$. Key to resolving these uncertainties is a better understanding of the population history of Beringia, the entryway for the Pleistocene peopling of the Americas.

Genomic insight into that population history has now become available with the recently recovered infant remains (USR1 and USR2) from the Upward Sun River site, Alaska (eastern Beringia), dated to $\sim 11.5$ kya $^{7,18}$. Mitochondrial DNA sequences (haplogroups C1 and B2, respectively) were previously acquired from these individuals $^{7,18}$ (SI 1,4,5). We have since obtained whole-genome sequence data, which provides a broader opportunity to investigate the number, source(s) and structure of the initial founding population(s), and the timing and location of their subsequent divergence. We sequenced the genome of USR1 to an average depth of $\sim 17X$, based on eight sequencing libraries from USER-treated extracts previously confirmed to contain DNA fragments with characteristic ancient DNA misincorporation patterns (SI 2-4). We estimated modern human contamination at $\sim 0.14\%$ based on the nuclear genome and $\sim 0.15\%$ based on
As expected, the error rate in the USER-treated sequencing data was low (0.09% errors per-base), and comparable to other high-coverage contemporary genomes, based on called genotypes (SI 4). While USR2 did not show sufficient endogenous DNA for high-coverage genome sequencing, we found both individuals were close relatives (SI 5), equally related to worldwide present-day populations (Figure S4g).

We assessed the genetic relationship between USR1, a set of ancient genomes \(^2,8,9,15,17\), and a panel of 167 worldwide populations genotyped for 199,285 SNPs \(^1,2,19\) (SI 6), using outgroup \(f_3\) statistics \(^20\), model-based clustering \(^21,22\) and multidimensional scaling (MDS) \(^23\) (SI 7-9). Outgroup \(f_3\) statistics of the form \(f_3(Yoruba; X, USR1)\) revealed that USR1 is more closely related to present-day Native Americans than to any other tested population, followed by Siberian and East Asian populations \(^1,2\) (Figure 1a). Pairwise comparisons of the \(f_3\)-statistics for USR1 and a set of ancient and contemporary Native American genomes \(^2,8,15\) (SI 6) showed that all are similarly related to Old World populations, though other Native American genomes (Aymara \(^2\), Athabascan1 \(^16\), 939 \(^2\), Anzick1 \(^8\) and Kennewick \(^15\) have a higher affinity for contemporary Native Americans than USR1 does (SI 9). MDS and ADMIXTURE analysis showed that the USR1 genome did not cluster with any specific Native American group (Figures 1d, S3b). These results imply that USR1 belonged to a previously unknown Native American population not represented in the reference dataset, herein identified as Ancient Beringians (SI 8.3).

To investigate if USR1 derived from the same source population that gave rise to contemporary Native Americans, we computed 11,322 allele frequency based-\(D\)-statistics \(^1,20\) of the form \(D(Native American, USR1; Siberian1/Han, Siberian2/Han)\) (SI 10.4). The resulting Z-score distribution corresponds qualitatively to the expected normal distribution under the null hypothesis that USR1 forms a clade with Native Americans to the exclusion of Siberians and East Asians – except for a set of Eskimo-Aleut, Athabascan and Northern Amerind-speaking populations for which recent Asian gene flow has been previously documented (Figures 1c, S5a, S6) \(^1,2,15,19\). Additionally, we found that present-day Native Americans and USR1 yield similar results for \(D(Native American/USR1, Han; Mal’Ta, Yoruba)\), suggesting they are equally related to the ancient north Eurasian population represented by the 24 kya Mal’ta individual \(^9\) (SI 10.5). These results confirm that USR1 and present-day Native Americans derived from the same ancestral source, which carried a mixture of East Asian and Mal’ta-related ancestry. We infer that descendants of this source represent the basal group that first migrated into the Americas.

To explore the relationship between USR1 and present-day Native Americans, we computed allele frequency-based and genome-wide \(D\)-statistics of the form \(D(Native American, Aymara; USR1, Yoruba)\). We could not reject the null hypothesis that USR1 is an outgroup to any pair of Native Americans, with the exception of a set of populations bearing recent Asian gene flow \(^1,2,15,19\) (Figures 1b, S7). We confirmed the phylogenetic placement of USR1 at a basal position in the Native American clade using TreeMix \(^24\) and two methods to estimate average genomic divergence and genetic drift, respectively (SI 14-16). These results support the branching of USR1 within the Native American
clade, but being equidistant to NNA and SNA. Below we discuss the potential geographic
locations of the USR1-NNA+SNA and the NNA-SNA splits (Figure 2) based on the
genetic results, the glacial geography of terminal Pleistocene North America\textsuperscript{25,26} and the
extant archaeological evidence (also SI 20).

Recent detection of an Australasian-derived genetic signature in some Native American
groups\textsuperscript{2,3} led us to explore whether USR1 bears that signal (SI 10.7, 11-13). Using
frequency-based and ‘enhanced’ D-statistics, we found no support for USR1 being closer
to Papuans (a proxy for Australasians) than other Native Americans.

We leveraged the position of USR1 on the Native American branch prior to the NNA-
SNA split to re-assess the origins of Athabascan and Eskimo populations by fitting
admixture graphs. We considered a whole-genome dataset including Siberian, East Asian,
Native American and Eskimo groups, as well as Mal’ta (SI 17). The heuristic approach in
TreeMix\textsuperscript{24} showed that the best proxies for the Asian component in Athabascans and
Greenlandic Inuit are Koryaks and the Saqqaq individual, respectively. We then followed
an incremental approach for fitting an $f$-statistic-based admixture graph\textsuperscript{20}, including the
Kets, previously suggested to share a linguistic and perhaps a genetic link with
Athabascans\textsuperscript{11,27}. This approach recapitulated the TreeMix results, and yielded a model
in which both Athabascans and Greenlandic Inuit derive from the NNA branch. However,
the Asian ancestry in Athabascans is most closely related to the Asian component in
Koryaks, while the Saqqaq genome is the best proxy for the Siberian component in the
Greenlandic Inuit (Figure 3). We infer the latter is a consequence of Palaeo- and Neo-
Eskimos having been derived from a similar Siberian population\textsuperscript{1,16}. This model appears
to be a good fit to the data, as the observed $f$-statistic that deviated the most from the
model prediction yielded $Z=3.27$. In SI 17.3 we tested the robustness of this model and
predictions by computing individual $D$ statistics, and re-fitting the model using alternative
datasets.

Lastly, we inferred the demographic history of USR1 with respect to Native Americans,
Siberians and East Asians, using two independent methods: diCal2\textsuperscript{28} and momi2\textsuperscript{29} (SI
18-19). diCal2 results indicate that the founding population of USR1, Native Americans,
and Siberians had a very weak structure from $\sim36$ kya up to $\sim24.5$ kya (Table S7), when
the ancestors of USR1 and Native Americans began to diverge substantially from
Siberians. USR1 diverged from other Native Americans around 20.9 kya, with a period
of ensuing moderate gene flow between them (Table S6 and S7), as indicated by a
simulation study that showed a significant increase in likelihood when comparing a 'clean
split' model to an 'isolation with migration' model (SI 18.4). Using momi2 and SMC++
we estimated a backbone demography where Karitiana and Athabascans split at $\sim15.7$
kya, while their ancestral population split from Koryaks $\sim23.3$ kya (Figure 4). With
momi2, we inferred the most likely branch (the population immediately ancestral to
NNA+SNA) and time ($\sim21$ kya) for the USR1 population to join the backbone
demography, while allowing for possible gene flow between USR and other populations
(SI 19, Figure 4b), results consistent with\textsuperscript{14} and the diCal2 inference.
These new findings, along with existing data, allow us to place Ancient Beringians (AB) within the broader context of the Pleistocene peopling of the Americas. The Native American founding population (comprised of both AB and NNA+SNA) began to diverge from ancestral Asians as early as ~36 kya, likely in northeast Asia, as there is no evidence of people in Beringia or northwest North America at this period. A high level of gene flow was maintained between them and other Asians until as late as ~25 kya. The subsequent isolation of the Native American founding population ~24 kya roughly corresponds with a decline in archaeological evidence for a human presence in Siberia. Both changes may result from the same underlying cause: the onset of harsh LGM climatic conditions. These findings, coupled with a divergence date of ~20.9 kya between USR1 and Native Americans, are in agreement with the Beringian Standstill Model (SI 21). The common ancestor of NNA+SNA and AB began to diverge ~20.9 kya, after which gene flow ensued, although whether it was with NNA+SNA, or the already differentiated NNA and SNA branches, cannot be determined owing to shallow divergence times among the groups.

These findings allow us to consider possible scenarios regarding where ancient Native American populations diverged (SI 20-21, Figure 2). Scenarios C-E require extended periods of strong population structure marking AB, NNA, and SNA as separate groups, for which we do not see compelling genetic evidence; hence these can be rejected. Scenarios A and B are compatible with our evidence of continuous gene flow among these groups, but differ as to the location of the AB versus NNA+SNA split at 20.9 kya, whether in northeast Asia (Scenario A) or eastern Beringia (Scenario B). Each has strengths and weaknesses relative to genetic and archaeological evidence: Scenario A best fits the archaeological and paleoecological evidence, as the earliest securely dated sites in Beringia are no older than ~15-14 kya, and the LGM cold period is unlikely to be associated with northward expanding populations. Scenario B is genetically most parsimonious, given evidence of continuous gene flow between the AB and NNA+SNA, suggesting their geographical proximity 20.9-11.5 kya, and that all three were isolated from Asian/Siberian groups after ~24 kya and form a clade.

Scenarios A and B are both consistent with the NNA-SNA split at ~15 kya having occurred in a region south of eastern Beringia. The ice sheets were then still a significant barrier to movement that would have helped maintain separation from the AB population. While members of the SNA branch have not been documented in regions that were once north of the glacial ice, NNA groups (including Athabascan-speakers) are present in Alaska today; thus, the latter are likely descendants of a population that moved north sometime after 11.5 kya.

The USR1 results provide the first direct genomic evidence that all Native Americans can be traced back to the same source population from a single Late Pleistocene founding event. Descendants of that population were present in eastern Beringia until at least 11.5 kya. By then, however, a separate branch of Native Americans had already established
itself in unglaciated North America, and diverged into the two basal groups that ultimately became the ancestors of most of the indigenous populations of the Americas.

**Data availability**

Sequence data was deposited in the ENA under accession: PRJEB20398.

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**Author Contributions**


**References**


27. Kari, J. M. & Potter, B. A. *The Dene-Yeniseian connection.* (University of
Figure 1. Genetic affinities between USR1, present-day Native Americans, and world-wide populations. a. $f_3$ statistics of the form $f_3(San; X, USR1)$, for each population in the genotype panel. Warmer colors represent greater shared drift between a population and USR1. b. $D$-statistics of the form $D(\text{Native American}, \text{Aymara}; USR1, \text{Yoruba})$ (points). The Andean Aymara were used to represent SNA. *: Native American populations with Asian admixture ($|Z|$ for $D(H1, \text{Aymara}; \text{Han}, \text{Yoruba})>3.3$) (Figure S5a). Error bars represent 1 and $\sim3.3$ standard errors ($p$-value $\sim0.001$). Native American populations were grouped by language family. c. Quantile-quantile plot comparing observed $Z$-scores to the expected normal distribution under the null hypothesis ($H_0$), for all possible $D(\text{Nat. Am.}, \text{USR1}; \text{Siberian1}, \text{Siberian2})$. Colors correspond to the $Z$-score obtained for $D(H1, \text{Aymara}; \text{Han}, \text{Yoruba})$. The expected normal distribution under the null hypothesis was computed for all groups jointly (SI Section 10.4). Thick and thin lines represent a $Z$-score of $\sim3.3$ ($p$-val $\sim0.001$) and a $Z$-score of $\sim4.91$ ($p$-val $\sim0.01$ after applying a Bonferroni correction for 11,322 tests). The bottom-right panel shows the expected tree under the null hypothesis. d. Admixture proportions estimated by ADMIXTURE assuming $K=20$ ancestral populations. Bars represent individuals, and colors represent admixture proportions from each ancestral component. Admixture proportions in ancient genomes (wider bars) were estimated using a genotype likelihood-based approach.

Figure 2. Possible geographic locations for the USR1 and NNA-SNA splits. We propose two possible locations for the split between USR1 and other Native Americans: the Old World (A, C, E) and Beringia (B, D); and three possible locations for the NNA-SNA split: the Old World (E), Beringia (C, D), and North America south of Beringia (A, B). Schematics show estimated glacial extent $\sim14.8$ kya. Dashed lines represent the Native American migration south of eastern Beringia, but they do not correspond to a specific migration route. Model discussion (SI Section 20) is based on extant archaeological evidence and inferred demographic parameters: a USR1-NNA+SNA split $\sim20$ kya with ensuing moderate gene flow and a NNA-SNA split $\sim15$ kya (SI 18-19).

Figure 3. A model for the formation of the different Native American populations. We fitted an admixture graph by sequentially adding admixed leaves to a 'seed' graph including the Yoruba, Han, Mal'ta, Ket, USR1, Anzick1 and Aymara genomes. For each 'non-seed' admixed group, we found the pair of edges that produced the best-fitting graph,
based on the fitting and maximum $|Z|$ scores (3.27 for this graph). Ellipse-shaped nodes: sampled populations; box-shaped nodes: metapopulations; *: single high-depth ancient genome. **: single low-depth genome. †: subgraphs whose structure we were unable to resolve due to sequencing and genotyping error in the Saqqaq genome (SI 17). Sample sizes and locations are shown at the top.

**Figure 4. USR1 demographic history in the context of East Asians, Siberians and other Native Americans.**

a. SMC++ inferred effective population sizes with respect to time for Athabascans (NNA), Karitiana (SNA), Han, Koryaks and USR1 (SI 19.1). We used these demographic histories as a basis for fitting a joint model for these populations.

b. A ‘backbone demography’ was fitted excluding USR1 using momi2, an SFS-based maximum likelihood approach (Figure S27), along with the most likely join-on point for USR1 onto the backbone demography (SI 19). We show the likelihood heatmap for the latter; warmer colors correspond to a higher likelihood of USR1 joining at a given point. These estimates agree with those obtained through diCal2, a method based on haplotype data (SI 18).