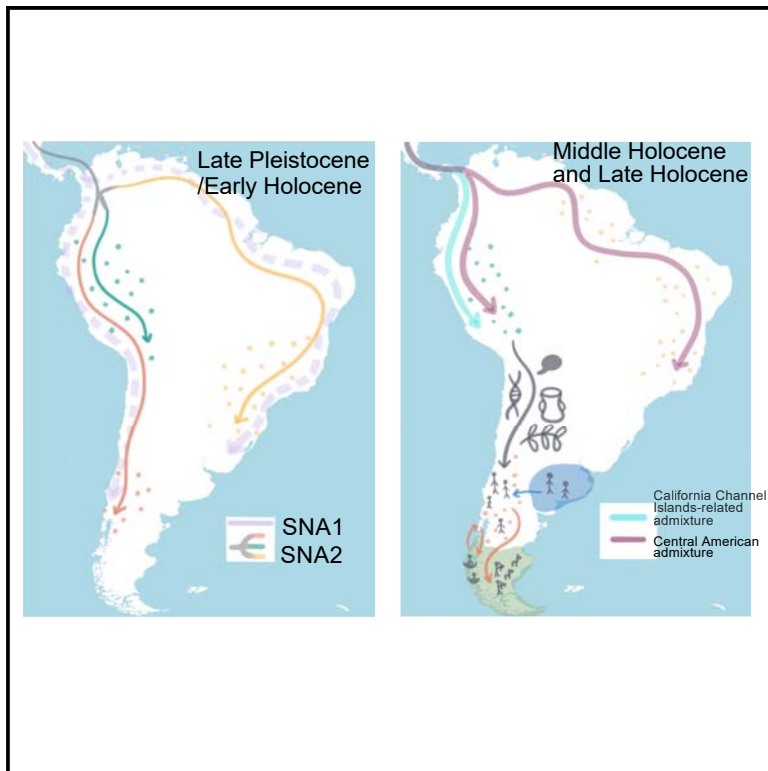


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The genetic history of the Southern Andes from present-day Mapuche ancestry

Graphical abstract



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In brief

Arango-Isaza et al. provide a genetic analysis of the South American Indigenous Mapuche ancestry that reveals local origin up to the Middle Holocene, relatively high isolation, and connections with the Central Andes. The study connects present-time Indigenous roots to the ancient history of southern South America.

Highlights

- Mapuche's genetic profile belongs to the Southern Cone broad genetic ancestry
- Their ancestors exchanged genes, words, crops, and ceramics with the Central Andes
- They are connected to ancient individuals in Chile up to 5,100 years ago
- They present relatively high isolation from other South American groups

Article

The genetic history of the Southern Andes from present-day Mapuche ancestry

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SUMMARY

The southernmost regions of South America harbor some of the earliest evidence of human presence in the Americas. However, connections with the rest of the continent and the contextualization of present-day indigenous ancestries remain poorly resolved. In this study, we analyze the genetic ancestry of one of the largest indigenous groups in South America: the Mapuche. We generate genome-wide data from 64 participants from three Mapuche populations in Southern Chile: Pehuenche, Lafkenche, and Huilliche. Broadly, we describe three main ancestry blocks with a common origin, which characterize the Southern Cone, the Central Andes, and Amazonia. Within the Southern Cone, ancestors of the Mapuche lineages differentiated from those of the Far South during the Middle Holocene and did not experience further migration waves from the north. We find that the deep genetic split between the Central and Southern Andes is followed by instances of gene flow, which may have accompanied the southward spread of cultural traits from the Central Andes, including crops and loanwords from Quechua into Mapudungun (the language of the Mapuche). Finally, we report close genetic relatedness between the three populations analyzed, with the Huilliche characterized additionally by intense recent exchanges with the Far South. Our findings add new perspectives on the genetic (pre)history of South America, from the first settlement through to the present-day indigenous presence. Follow-up fieldwork took these results back to the indigenous communities to contextualize the genetic narrative alongside indigenous knowledge and perspectives.

INTRODUCTION

The peopling of the Americas represents the last of the major human migrations, beginning—according to the latest interpretations of the genetic data—no earlier than ~23 thousand years ago (kya).¹ Historical admixture with European and African individuals and the devastating decline of indigenous populations caused by contact with Europeans hamper our ability to reconstruct ancient demographic history in the Americas. The patterns and timing of the initial migrations remain debated in population genetics and other disciplines.²

From a genetic perspective, all non-Arctic Native American groups descend from an ancestral population that split into a northern and a southern branch while still within North America.^{3–6} Within the southern branch, an early wave related to a 12,600 BP individual associated with the Clovis culture (Anzick-1)⁷ and labeled SNA1 spread rapidly southward during the Late Pleistocene (before 13,000 BP).^{8–10} A second wave, associated with a North American sample from 10,100 BP (Spirit Cave) and labeled SNA2, also entered South America possibly as early as the Late Pleistocene.^{9,10}

The genetic profile of present-day indigenous populations of South America stems predominantly from this second migration wave, which differentiated into three main ancestries characteristic of three broadly defined ecogeographic regions: one primarily represented in the Andes, one in the Amazonian lowlands, and one in the Southern Cone.^{4,11} During the Holocene, further migration waves from North and Central Americas reached the Andes and Amazonia.^{8,12} Genetic studies in South America have so far focused on Amazonia or the Central Andes. In contrast, only a few recent genome-wide studies have addressed populations of the remaining macro-region, the Southern Cone.^{11,13,14} In comparison, a rich literature based on uniparental markers revealed the presence of characteristic early diverging lineages in the Southern Cone.^{15–21} The lack of genome-wide data leaves open questions on how the earliest human migrations reached the south, which routes they took, and how much they interacted with subsequent migration waves. With our study, we focus in particular on southernmost South America, i.e., the Southern Cone more narrowly defined as modern Chile (excluding the northernmost regions), Argentina, and Uruguay.

Archaeological evidence points to the southern regions of South America as having been inhabited from the earliest stages of human settlement of South America. The site of Monte Verde in southern Chile, dated to at least ~14,500 cal BP, bears the earliest widely accepted traces of human presence in the continent.²² The migration routes taken by the first settlers are debated: some argue that routes along the Pacific or Atlantic coasts are equally likely,¹⁵ whereas others strongly support the Pacific coast as the predominant route.^{16,17} The eastern Southern Cone could have been settled from trans-Andean,¹⁶ Atlantic, or inland routes.¹⁷ Ancient DNA (aDNA) data from southern Patagonia show a genetic continuity from 6,600 BP onward, as well as differentiation between sea nomads along the Pacific coast, who relied on marine resources, and foot nomads in eastern Patagonia, who relied on hunting wild guanaco.^{13,23} In the Late Holocene (~3,000 BP), more complex resource management and the first settlements appeared.^{24,25} More work is needed to link the archaeological and genetic evidence through the Holocene and to recent indigenous history in the southern regions of South America.

Today, the Mapuche represent one of the main indigenous groups of the Southern Cone, with a major presence in southern Chile and small parts of Argentina. The archaeological record suggests population continuity from the first centuries CE to the groups encountered by Europeans in the 16th century.²⁶ Central Chile was conquered by the Inca Empire in the 15th century and then came under Spanish control from 1541. Further south, a conflict known as the Arauco War affected the indigenous populations. By 1641, a frontier was established, south of which the region known as “Araucanía” remained largely independent until Chilean and Argentinean conquest from the 1860s to 1880s.

Geographic and ecological factors draw the boundaries between several self-identified Mapuche territorial identities such as Lafkenche (people of the sea) along the Pacific coast, Pehuenche (people of the araucaria pine) in the Andean mountains, and Huilliche (people of the south) south of the Toltén river and into the Chiloé archipelago. Some groups in southern Chile, in particular, self-identify as just Huilliche, rather than as a Huilliche

subgroup of the Mapuche people, to mark a separation from the broad Mapuche group. Mapudungun is considered a single-broad language with various regional varieties (“geolects”).²⁷ Previous population genetics studies^{4,7} have followed an ethnolinguistic categorization of the Americas that placed Mapudungun in a supposed “Andean” macro-group.²⁸ This framework has been rejected within mainstream linguistics, however, for its lack of sound methodological foundation.^{29,30} In standard classifications, Mapudungun is a language isolate, with no demonstrable shared origin to any other language.^{31,32} Nonetheless, a small number of words (e.g., for “hundred,” “thousand,” “fish,” and perhaps also “sun”) that appear to be borrowings do imply that Mapudungun was once in sporadic contact with Quechua and Aymara from the Central Andes. The borrowings remain phonologically very similar to the source words, suggesting that these contacts do not predate the Inca period by long, if at all.^{33–35} Further exchanges can also be traced with crops such as potato, quinoa, beans, squash, and maize,^{24,25} domesticated in the Central Andes and then introduced to these southern regions. Gene-flow events may have accompanied the entry of loanwords and crops from the Central Andes: such demographic contacts could be dated with genetic analysis and provide a time frame for the exchanges.

Despite the distinct role played by the Mapuche in the indigenous history of South America, their internal population structure and relationships with other indigenous groups remain poorly understood. The origins of the Mapuche have been explained variously as (1) a local trajectory in continuity since the earliest occupation,²⁶ (2) migration from the Central Andes,³⁶ or (3) migration from the Amazonian rainforest or the Gran Chaco region.³⁷ Published genome-wide data from one Mapuche and one Huilliche population suggest that they are closely related to other ancient and modern Patagonian populations.¹¹ Overall, it remains unclear which major migration wave the Mapuche stem from, or whether they were affected by the population movements of the Holocene, with potentially major implications for our understanding of the peopling of the Americas.

To reconstruct the origins of Mapuche genetic ancestry and trace its trajectory within South America, we generated genome-wide SNP-chip data for 64 participants of Mapuche descent from two regions of Araucanía and from the island of Chiloé. We analyze their genetic makeup, compare it with published data from Native American groups and aDNA, and set our findings alongside linguistic and archaeological evidence. Our results reveal in more detail the nature of the genetic connections between ancient and living American populations, describe the recent demographic effect of European contact, and clarify the genetic relationships between different Mapuche groups.

RESULTS

Overall genetic patterns in the Americas

We generated genome-wide data with the Axiom Human Origins SNP array³⁸ from the following three populations: Pehuenche from the mountains of Araucanía; Lafkenche from the coast of Araucanía (both groups who recognize their ancestry as Mapuche); and Huilliche-Chiloé, a population from the island of Chiloé, which, in part, recognizes its ancestry as Huilliche. The label

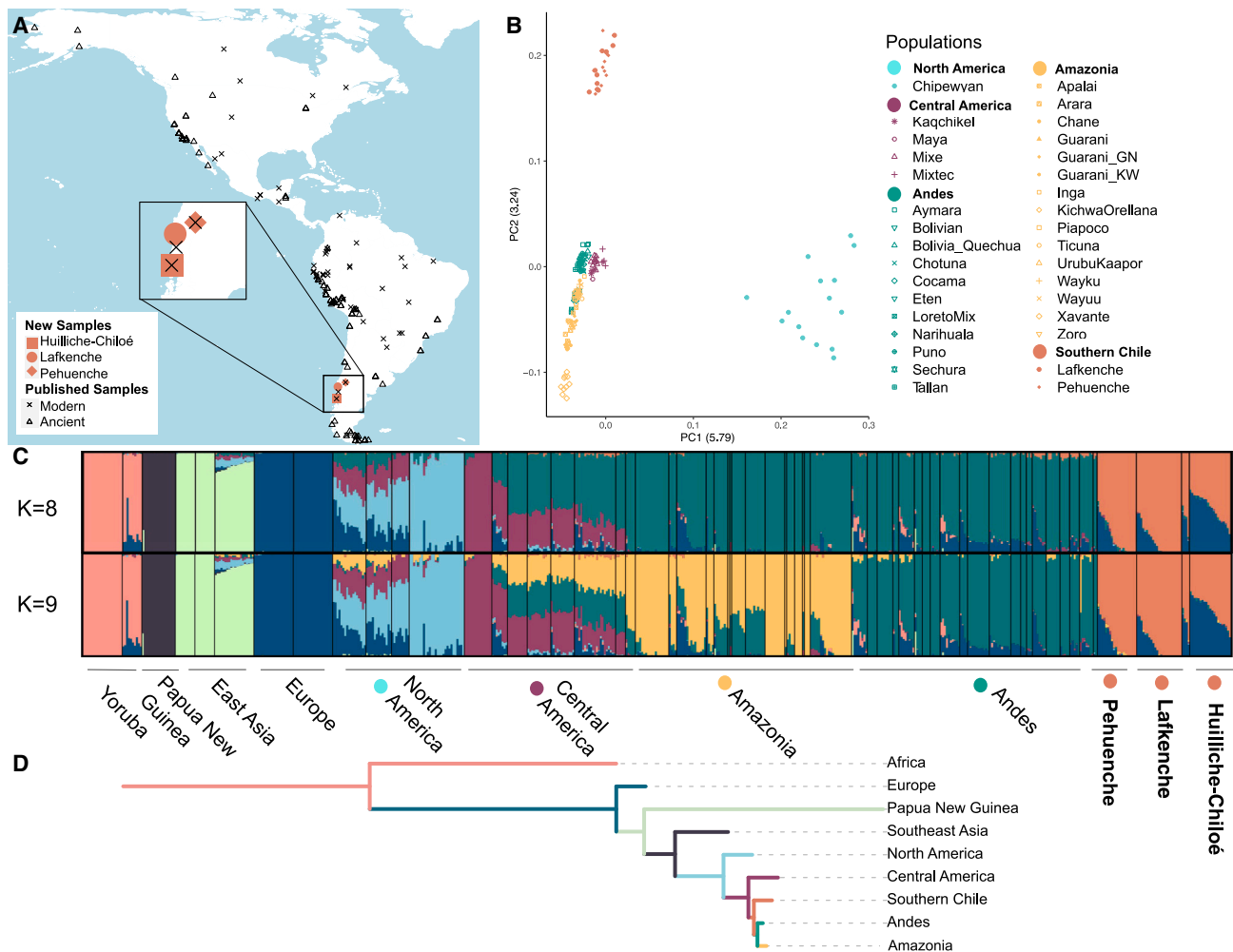


Figure 1. Sampling locations of newly genotyped individuals, reference individuals, and overall genetic patterns in the Americas

(A) Map of the Americas showing the approximate location of the newly generated samples and the selection of individuals from published literature, both modern (crosses) and ancient (triangles).

(B) PCA including only unadmixed individuals from the Americas, defined as having at least 99.9% of Native American component, as inferred by ADMIXTURE at $K = 8$ (dataset 1.3; Figure S1B). Only 20 individuals from the Lafkenche and Pehuenche populations passed this filter and no individuals from the Huilliche-Chiloé population. Color coding corresponds to broad macro-regions of the Americas (Figure S1A).

(C) ADMIXTURE run for the global dataset 1 (Figure S1B) for $K = 8$ and $K = 9$. Runs from $K = 2$ to $K = 15$ are available in Figure S2. For the ADMIXTURE run with dataset 1.2. See Figure S3.

(D) NJ tree of covariance-derived distances among ancestry components for $K = 9$ (dataset 1), computed with OHANA based on the global ADMIXTURE run with the highest likelihood score.

Huilliche-Chiloé is to distinguish them from Huilliche groups on the mainland (Figure 1A). We merged the genotypes of these individuals with modern publicly available data from relevant populations (datasets 1, 2, and 3)^{38–41} and with aDNA data (dataset 3.3; see Figure S1B for a schematic description of the different datasets used).^{6–8,11–13,38–43} To understand the global pattern of genetic relatedness among modern samples (dataset 1), we used ADMIXTURE.⁴⁴ The ADMIXTURE analysis showed the lowest cross-validation (CV) errors for $K = 8$ and $K = 9$ (Figure S2B). Major ancestry components specific to the Americas (Native American ancestry) start to differentiate from $K = 6$ (Figure S2A). At $K = 7$, a component emerges that is prevalent in our Mapuche sample and that we refer to here as the Southern Chile (SC) component (orange in Figure 1C). This SC component is

also sporadically present in the Andes. The Mapuche individuals sampled have a variable percentage of European admixture (dark blue in Figure 1C), ranging from averages of 9.2% in the Lafkenche and 13.3% in the Pehuenche to 43.4% in the Huilliche-Chiloé (at $K = 8$). Dataset 1 includes four individuals from a previous publication labeled as Chilote (i.e., from the island of Chiloé),³⁹ which shows a similar admixture profile to our Huilliche-Chiloé sample. We ran an analogous ADMIXTURE analysis with a dataset that included the modern samples from the study of De la Fuente et al.¹¹ but with fewer overlapping SNPs (dataset 1.2; Figure S1B). The results are consistent with Figure 1; our Mapuche samples are genetically similar to the Pehuenche sample from the study of De la Fuente et al. (Figure S3A). Populations in this dataset from the Far South

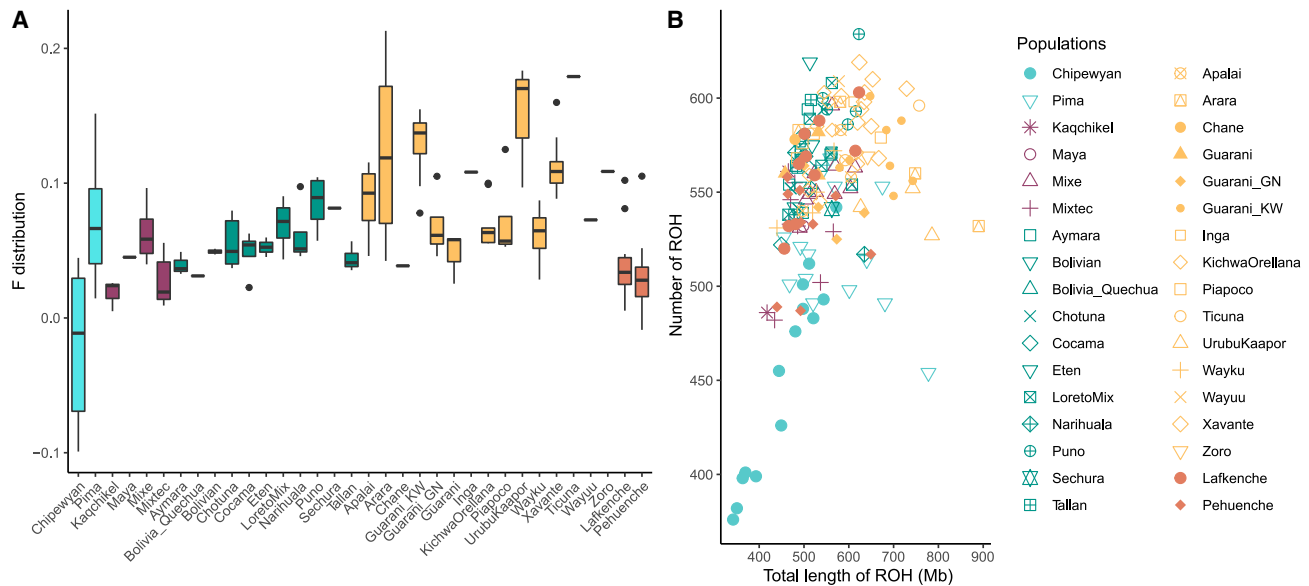


Figure 2. Within-population diversity

(A) Individual values of consanguinity (F) averaged for each population from South America. (B) Distribution of ROH fragments. Both analyses are computed on the unadmixed individuals (Dataset 1.3).

(FS, defined here as south of $\sim 50^\circ\text{S}$), namely Yámana and Kawéskar, display a characteristic component at $K = 12$. The relationship between the admixture components is visualized with a neighbor-joining (NJ) tree, which supports all South American ancestries branching from each other closely in quick succession (Figure 1D).

To exclude historical gene flow from Europe, we retained only “unadmixed” individuals (here defined as having 99.9% Native American ancestry as computed by ADMIXTURE at $K = 8$, dataset 1.3) and performed a PCA on this subset (Figure 1B). Here, the first component separates North and South American groups, and the second component separates SC and Amazonian groups, whereas the Central American and Andean populations remain close to the Amazonian ones.

Both ADMIXTURE and PCA suggested that the SC populations are genetically distinct from the rest of South America. The marked differentiation of the SC component could imply an early divergence in the population structure of South America but could alternatively be due to an overrepresentation of Mapuche individuals in the dataset or to a recent bottleneck associated with strong genetic drift in small, isolated SC groups.⁴⁵ An analysis of the degree of consanguinity and the distribution of runs of homozygosity (ROHs) (Figure 2) shows that our SC individuals are not especially high in homozygosity compared with other populations of South America, ruling out recent strong drift as the predominant explanation for this genetic divergence. A mixed scenario with early divergence and moderate drift is also plausible.

Recent demography and connectivity

To further investigate the structure of the Mapuche populations in relation to the rest of the continent, we looked at recent demography (~ 3 ka) and gene-flow patterns using shared identity by descent (IBD) blocks, inherited from the same common ancestor. Due to recombination, the length of IBD blocks shared

by two populations decays with time since these populations split,^{46,47} but admixture may (re)introduce shared IBD blocks.⁴⁸ The individuals from the Southern Cone are connected to each other by a dense network of IBD sharing (Figure 3A). The Mapuche populations, and, in particular, the two Pehuenche populations from this and a previous study,¹¹ show a high shared ancestry with each other, whereas the Huilliche-Chiloé population shares more blocks with Chilote (from Lazaridis et al.³⁹) and with the distant Kawéskar and Yámana (from de la Fuente et al.¹¹) than with the neighboring Lafkenche and Pehuenche. Across South America as a whole, three broad networks of shared ancestors roughly correspond to the regions where the three main ancestries from our ADMIXTURE analysis are represented: Andes, Amazonia, and Southern Cone. The three regions share a significant number of blocks, especially between the Andes and Amazonia. In contrast, the southern regions are less integrated into this network of shared IBD blocks, which suggests a higher degree of isolation. We found only one persistent link between our SC samples and the Northern Andes, and a similar link to the Gran Chaco region (Figures 3A, 3B, S4A, and S4B). This analysis is also performed with fragments of Native American descent, identified with the masking process that filtered out variants of possible African and European origins (Figures S4D and S4E). The overall pattern is consistent with Figure 3B, confirming a connection between Mapuche populations and the Andes for fragments smaller than 10 cM.

The length of the shared IBD blocks has been correlated with the number of generations back to shared ancestors in previous studies of European,⁴⁶ Asian,⁴⁹ and Native American populations.⁵⁰ In these studies, fragments between 5 and 10 cM have been associated with sharing events occurring 500–1,500 years ago. In a recent study, Ioannidis et al.⁵¹ dated a gene-flow event from indigenous Americans into Polynesians at around 1200 CE and independently matched it with IBD blocks longer than 7 cM.

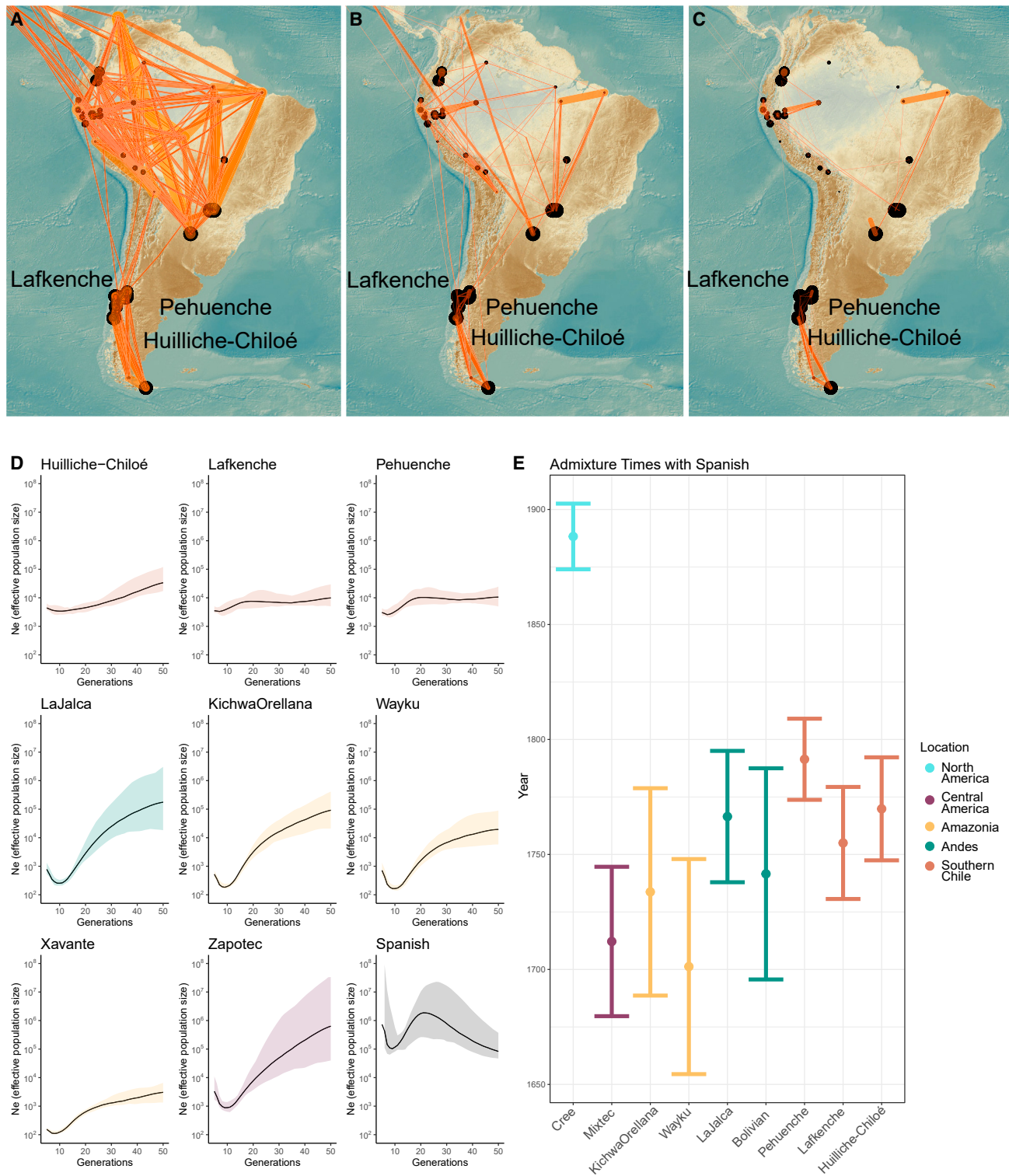


Figure 3. Recent demography and connectivity

IBD sharing probability network among South Americans (dataset 2.2). The network represents the probability of a pair of individuals from populations A and B sharing an IBD fragment, adjusted by population size. Thicker width and lighter orange color of the lines correspond to higher exchange between populations. The size of the black circles is proportional to sample size.

(A) Shared fragments from 4 to 7 cM. For visualization purposes, only population pairs with a probability of sharing higher than 10% are considered.

(B) Shared fragments from 7 to 10 cM. For visualization purposes, only population pairs with a probability of sharing higher than 2% are considered.

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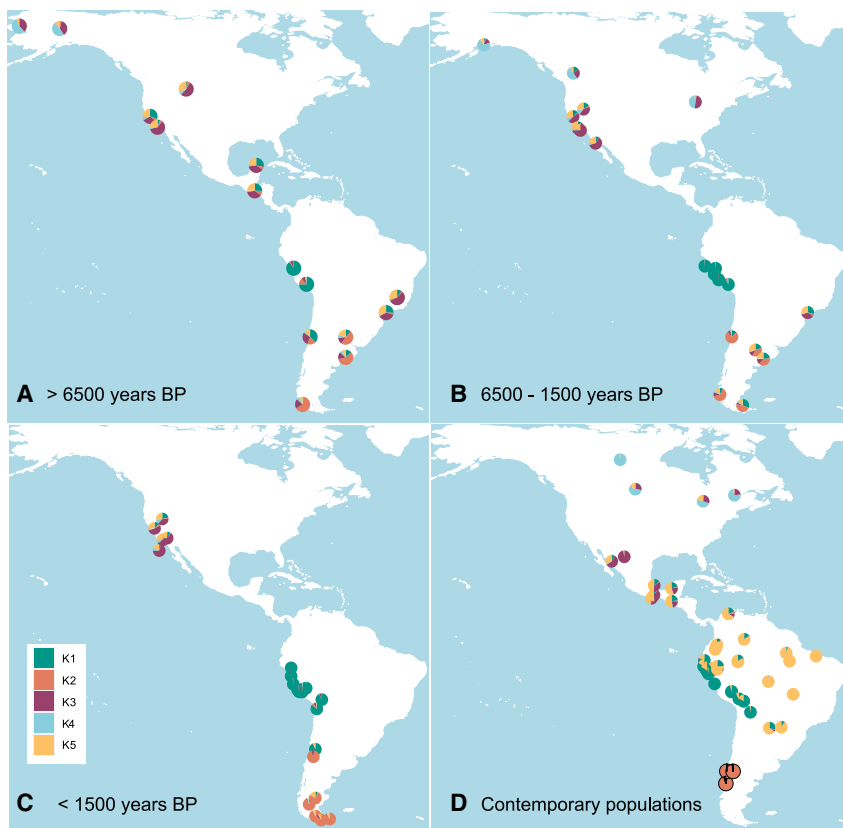


Figure 4. The ancient population structure of South America

ADMIXTURE run with ancient and masked individuals (dataset 3.3) for $K = 5$. Modern samples are grouped by population (in D), and ancient samples by proximity in geographical space and time.

(A) Up to 6,500 BP.

(B) 6,500–1,500 BP.

(C) After 1,500 BP but not contemporary.

(D) Contemporary. For details about the masking procedure, see [Figure S5](#).

disequilibrium and is most sensitive to recent large admixture events, ignoring minor admixture episodes and multiple admixture pulses that could have occurred previously.⁵⁴ The inferred admixture times vary across populations. The Wayku in lowland Peru show the earliest estimate (late 16th century), whereas the Cree in North America show the most recent estimate (late 19th century). The Mapuche populations show an intermediate admixture time to the mid-18th century ([Figure 3E](#)).

The ancient population structure of South America

To focus on indigenous ancestry, we performed masking to filter out variants associated with European or African descent.

To check the performance of the masking process, we ran f_4 -tests and PCA ([Figures S5A–S5C](#); https://github.com/epifaniarango/popgen_with_epi/tree/Local-ancestry-and-masking). We then merged masked individuals with a selection of ancient samples from the Americas into a new dataset (dataset 3.3; [STAR Methods](#); [Data S1A](#)). On this dataset, we ran ADMIXTURE⁴⁴ from $K = 2$ to $K = 10$. $K = 5–7$ were associated with the lowest CV errors ([Figures S5D and S5E](#)), and $K = 5$ had the least variance between runs. At $K = 3$, a distinct component emerges that is present primarily in the ancient and modern SC samples ([Figure S5E](#)).

We take the results of the ADMIXTURE analysis at $K = 5$ and explore the genetic relationships across geographic locations and timescales. Most samples older than 6,500 BP (i.e., from the Late Pleistocene to the start of the Middle Holocene) harbor all five ancestry components, but at varying proportions ([Figure 4A](#)). Only the two Andean samples have a single-predominant component, which persists at high frequency in the Andes through all later periods. Between 6,500 and 1,500 BP, we observe an increasing differentiation between the Central Andes and the Southern Cone ([Figure 4B](#)). The samples from Central Chile and the Far South are structured and non-homogeneous, with the SC local component present at varying proportions. In

However, it is important to note that patterns of IBD sharing can also be affected by population-specific histories. With this caveat, and considering the data from other studies as an indicative reference, the IBD sharing between SC and the Central Andes may date back more than ~ 500 years ago, as we do not find shared fragments longer than 10 cM.

The analysis of IBD fragments among individuals within a population can also give insights into variations in the effective population size (N_e).⁵² We inferred the demography of populations of the Americas represented by large sample sizes (min 7 individuals) and of the Spanish population as a reference from outside the Americas ([Figure 3D](#)). Before ~ 30 generations ago (~ 840 years), the three Mapuche populations are characterized by a relatively small and constant N_e . Starting around 15–20 generations ago, all Native American populations underwent a severe bottleneck that corresponds to the European colonial impacts (including pathogens) and the ensuing historically documented population decline to its lowest point at 10 generations ago. This bottleneck is not strongly visible in the three populations of Mapuche ancestry, contrary to reports by other studies based on simulation methods.⁵³

Finally, we used ALDER to estimate the date of admixture with Europeans ([Figure 3E](#)). This software is based on linkage-

(C) Shared fragments longer than 10 cM.

(D) Variation in effective population size for selected Native American and Spanish populations over the last 50 generations, calculated with IBDNe.

(E) Estimated admixture times of selected Native American populations with a Spanish source, calculated using ALDER. The error bars represent a 95% confidence interval (generation time: 28 years, only the most recent admixture pulse is reconstructed with this method). Matrix visualization of IBD sharing is available in [Figure S4A](#). IBD sharing with Native American ancestry specific markers is available in [Figure S4B](#).

the most recent period represented by aDNA (1,500–100 BP), the three major ecogeographical regions appear more differentiated, although Amazonia is poorly represented (Figure 4C).

Populations in North and Central America and the most ancient individuals of the dataset show the highest number in different ancestry components. We do not interpret this strictly as an admixture event between distinct ancestries but as a characteristic of the initially undifferentiated gene pool, which drifted as the populations were migrating southward. Finally, among the present-day populations, the Andes, Amazonia, and Southern Cone are clearly distinguished from each other, with various degrees of admixture in the contact zone between Amazonia and the Andes (Figure 4D).

An alternative perspective on the genetic prehistory of the Americas can be gained from outgroup f_3 -statistics,⁵ which estimate shared genetic drift among two populations relative to an outgroup. We selected commonly used transformations of f_3 -statistics (NJ and MDS) to visualize the shared genetic history (Figure 5).

The NJ tree suggests an early split of the North and Central American samples (Figure 5A), which is in line with the North-to-South population expansion through the Americas.^{1,4,6–8,12} The next groups to branch off independently are the Amazon and the Andean clades. A further clade includes the most ancient individuals of the dataset, which have been associated with SNA1^{8,12} (here defined as “Anzick-related”). The Southern Cone samples then divide into three sub-branches: ancient Argentinean samples from the Pampas; ancient CC and modern SC; and ancient FS. These three SC populations are closely related and share drift mostly with ancient samples from the Pampas and with CC (*Conchali_700BP* and *LosRieles_5100BP*) and overlap with them in the MDS plot (Figure 5B).

Population history of the Southern Cone

We used f_4 -statistics to study changes in the genetic structure over time. F_4 -statistics measures the shared genetic drift between a set of four populations/individuals to provide insights into population contact. We explored allele sharing within Mapuche groups using the configuration $f_4(\text{Mbuti}, X; Y, Z)$, where X is any ancient or modern population, and Y and Z are two populations taken from Lafkenche, Pehuenche, and Huilliche-Chiloé. As expected, our SC samples show a high and symmetrical allele sharing among each other (Z score ~ 0 ; Data S1C). Of all the populations/individuals tested in position X , only the *Conchali_700BP* samples and, to a lesser extent, the *ArroyoSeco2_7700BP* sample showed a shared drift with Lafkenche (Figure 6A), when Lafkenche is paired together with Huilliche-Chiloé in position Z (for details, see https://github.com/epifaniarango/popgen_with_epi/tree/Dstats-plots).

We further explored the relationship between SC and Conchalí by computing f_4 -statistics of the form $f_4(\text{Mbuti}, [\text{targeted ancient or modern sample}]; \text{Conchali}_700BP, \text{Modern SC})$. All three SC Mapuche populations consistently show a greater affinity to *Conchali_700BP* than to any other ancient or contemporary group (Z score $< |3|$; Data S1D), with the exception of *LapaDoSanto_9600BP* which shows affinity to Conchalí, but only when all SNPs are used (Z score < -3.3).

A previous study related *Conchali_700BP* to Late Holocene FS samples using the test $f_4(\text{Mbuti}, \text{Conchali}_700BP; \text{Middle}$

Holocene FS, and Late Holocene FS).¹³ We repeated this test, also with our Mapuche SC populations in the position of Conchalí. We obtained higher f_4 values with our SC Mapuche populations than with Conchalí, in most combinations tested (Figure 6B; Data S1E). We consistently found significant Z scores (>3.3) in those combinations that involved more recent Late Holocene FS individuals, i.e., those dated between 400 and 200 BP. This result suggests that the genetic ancestors of the Mapuche were involved in contact with the Late Holocene FS, either through the same contact event described in the literature for Conchalí or possibly with a further, more recent contact event between recent ancestors of Mapuche and FS populations. We also searched for evidence of Late Holocene gene flow between SC and other areas of South America (Argentinean Pampas and Central Andes) using $f_4(\text{Mbuti}, \text{SC/Conchali}_700BP; [\text{Middle Holocene Argentinean Pampas/Central Andes}], [\text{Late Holocene Argentinean Pampas/Central Andes}])$ but found none (Data S1F and S1G).

With the configuration $f_4(\text{Mbuti}, X; \text{SouthernCone 1 and SouthernCone 2})$, where X is any ancient or modern sample outside the Southern Cone, we did not find evidence of a significant shared drift from other regions (Z scores $< |3|$), suggesting a single origin for the Southern Cone populations. Only a few configurations involving the Middle Holocene Pampas as one of the Southern Cone populations are significant (Data S1H; see https://github.com/epifaniarango/popgen_with_epi/tree/Dstats-plots). The Middle Holocene Pampas individuals display a more distinctive genetic profile within the Southern Cone, which could be tentatively related to other substrates.

We also explored the formation of ancestries and admixture events affecting the Southern Cone using admixture graph modeling with qpGraph. We created an initial scaffold using individuals representing the three Southern Cone clades, plus *LapaDoSanto_9600BP* (associated with SNA1), and two outgroups (African Mbuti and Ancient Beringian USR1). The best-fitting topology showed a split among the main three clades: Central-Southern Chile (*LosRieles_5100BP*), Pampas (*ArroyoSeco2_7700BP*), and FS, with FS comprising the two lineages associated with the sea nomads (*Ayayema_4700BP*) and foot nomads (*LaArcillosa2_5800BP*) (Figure S7A). We then added Late Holocene samples from CC and FS (Figures S7C and S7D). The FS sampled is modeled by a strong contribution from the lineages of Central-Southern Chile, confirming the f_4 results (Data S1E). In the last step, we added our modern SC Mapuche samples. Their genetic profile stems from a common ancestor with *Conchali_700BP* (Figure S7E).

The qpGraph scaffolding in some configurations requires admixture edges that are not clearly supported by direct f_4 -tests. For example, to model the genetic profile of *Conchali_700BP* and *LosRieles_5100BP*, an admixture edge is required from an ancestral population close to *LapaDoSanto_9600BP*. To model the Mapuche, an admixture edge is required from a population related to *ArroyoSeco2_7700BP*, which is supported in the f_4 described above with a Z score of 3.174 (Figures 6A, S7E, and S7G). (See https://github.com/epifaniarango/popgen_with_epi/tree/Dstats-plots for further discussion on the f_4 -tests.) When adding the Huilliche-Chiloé population, the best configuration requires a substantial (10%) admixture edge from an ancestral population at the root of the South American lineages (Figure S7H). This effect could

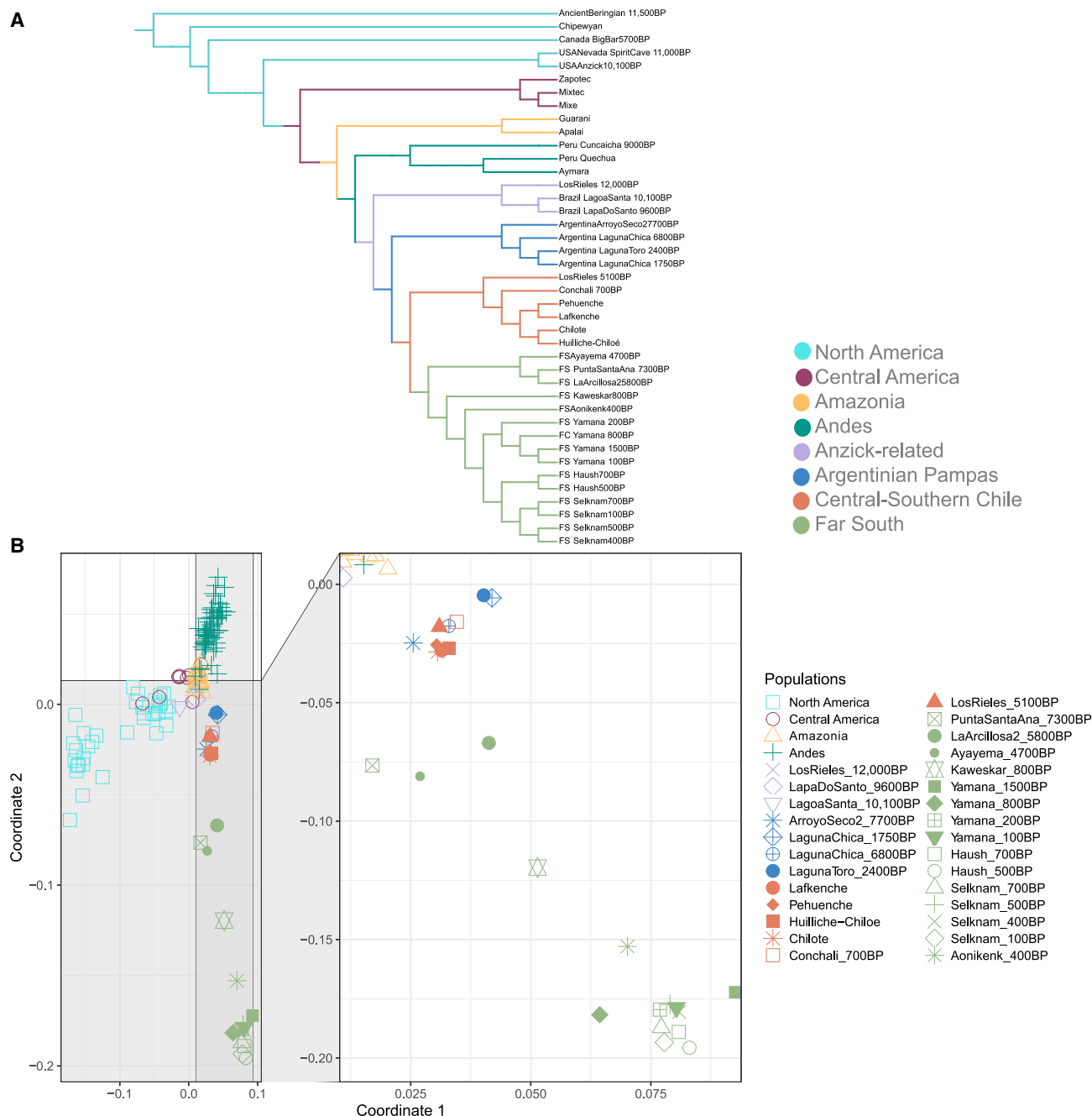


Figure 5. F_3 -statistics analyses

(A) NJ tree of the matrix of inverted outgroup f_3 -statistics ($1/f_3(\text{Mbuti}; X, Y)$) using Ancient Beringian as an outgroup. Ancient samples are filtered for a minimum of 100,000 SNPs. The tree is a graphic simplification, which does not include all populations/samples and displays the cladogram without information on branch lengths. The original tree with all samples and branch length to scale can be found in [Figure S6](#). Color code corresponds to broad regions and time transects. (B) MDS plot of the matrix of $1-f_3(\text{Mbuti}, X, Y)$. The blow-up to the right zooms in on the Far South, Argentinian, and Central-Southern Chile samples.

be due to the Huilliche-Chiloé having a higher proportion of European ancestry than the other Mapuche populations, which results in higher missingness after the masking. Alternatively, some European ancestries could have eluded the masking, resulting in a signal of gene flow from Eurasia. Nevertheless, configurations where the admixture edge comes from nodes upstream to the entry into the Americas return a worse fit, favoring a Native

American source ([Figure S7H](#)). Possible sources of bias in these discrepancies between qpGraph and f_4 -statistics include having modern and ancient samples modeled together, the small number of SNPs available when merging ancient samples with masked modern samples, and the presence of ancient samples sequenced with a different technology (shotgun sequencing versus SNP capture).

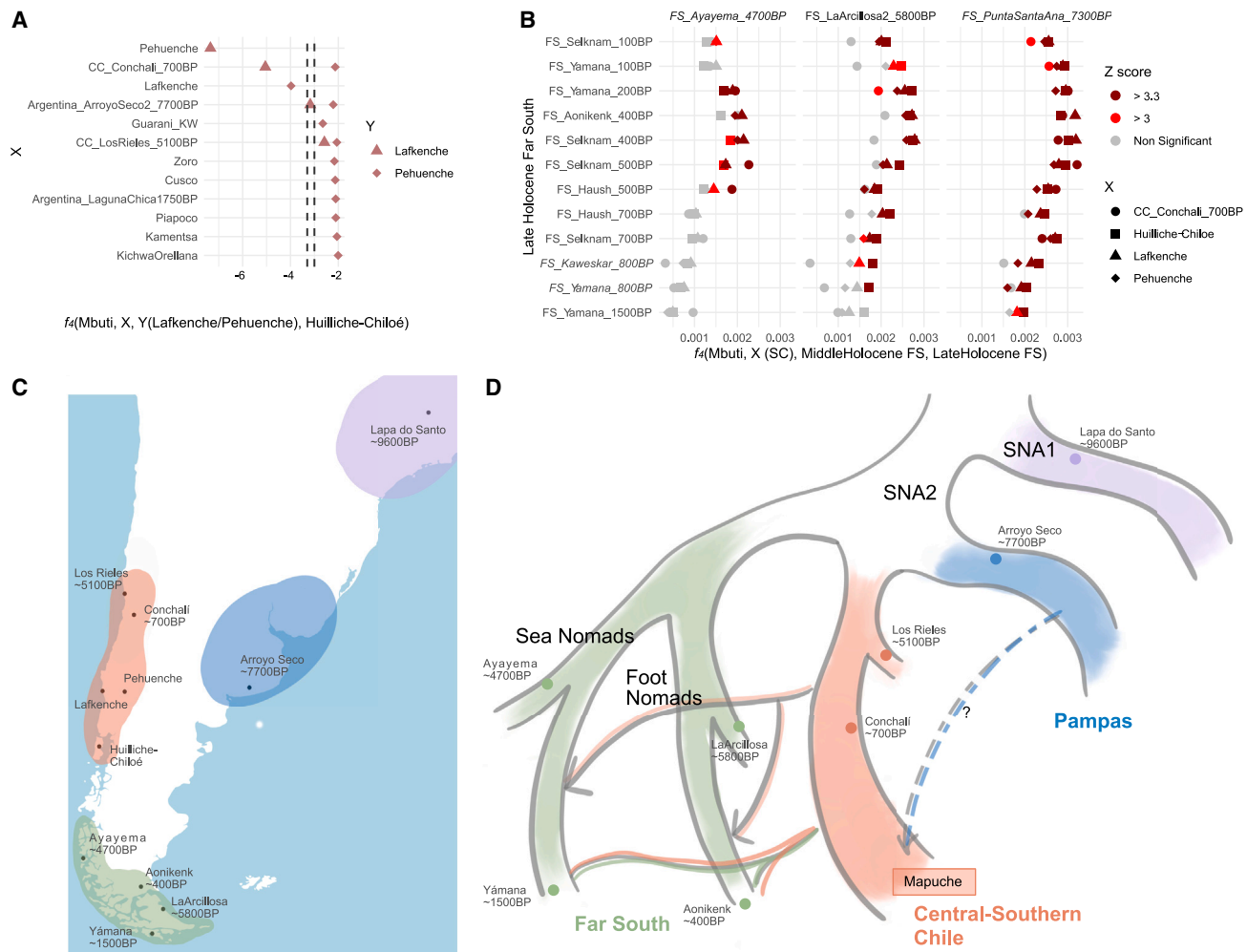


Figure 6. Population history of the Southern Cone

(A) f_4 -tests exploring allele sharing between South American samples (ancient and modern) and present-time SC Mapuche. In the y axis, only the individuals/populations associated with a Z score < -2 are displayed. Two vertical dashed line mark the significance thresholds of -3 and -3.3 .

(B) f_4 -tests exploring connections between Central-Southern Chile ancient and modern individuals and Late Holocene FS. F_4 -values are plotted in the x axis. Plot marks are colored by the significance of the test based on Z scores. Individual names in italics distinguish those samples sequenced with a shotgun technology from the rest of the samples genotyped with a capture approach.

(C) Geographical location of key samples and possible geographic distribution of main genetic lineages of the Southern Cone.

(D) Schematic representation of the most important regional lineages shaping the genetic landscape of the Southern Cone, summarized from the f_3 , f_4 , and qpGraph results in Figure S7. Admixture edges between Late Holocene FS and Central-Southern Chile refer to the f_4 in (B). The connection between the Pampas and Mapuche, marked with a question mark, refers to the f_3 -results in Figure 5B and is partially supported in the f_4 configuration in (A).

In the scheme of Figures 6C and 6D, we reconstruct a possible scenario for the formation of ancestries in the Southern Cone, considering possible ancestry divergence and admixture pulses as reconstructed from f_3 , f_4 , and qpGraph analysis.

DISCUSSION

Southern ancestry formed through isolation from the rest of South America

Our findings allow us to trace the formation of the Mapuche ancestry in the wider context of genetic diversity across South America. As previously noted,¹¹ Mapuche ancestry belongs to a genetic cluster characteristic of southern South America. We

observe that this ancestry cluster is equally distinct from the two other main clusters, which characterize the Central Andes and Amazonia, respectively (Figures 1B and 1D). The split between these ancestry clusters could trace back to the early Holocene: simulation studies dated the split between Mapuche and Andean Aymara at 8,750 BP.⁵³ Our IBD sharing profiles between modern samples suggest that the Southern Cone has been developing in relative isolation from other regions (Figures 3A–3C). Small, isolated groups could have persisted until $\sim 1,000$ years ago; hence, their relatively small N_e (Figure 3D). Homozygosity and ROH data suggest that this isolation pattern is not confounded by high levels of recent consanguinity within populations (Figure 2).

A recent publication has proposed that genomic variation within South America during the Pleistocene derives principally from two ancestries, labeled SNA1 and SNA2, and that the genetic ancestry of South America emerges mostly from SNA2.¹⁰ Between the Middle/Late Holocene, the Central Andes and Amazonia admixed with further migration waves from the California Channel and/or Central America.^{5,8,12} Based on an f_4 analysis of the single ancient sample from *Ayayema_4700BP*, Moreno-Mayar et al. proposed that these waves did not reach the Southern Cone.¹² We can now extend this finding to the whole Southern Cone, consistently over most of the Holocene and into present-day populations (Data S1H and S1I).

Our results suggest that the ancestry of the Mapuche, and of the rest of SC and the FS, originates in local continuity from an early migration wave (SNA2), followed by relative isolation. This consistent genome-wide pattern matches other evidence of relative isolation, such as the presence of early diverging lineages coming from uniparental markers.^{15–17} This finding has important repercussions for our understanding of the cultural and demographic background of these populations. First, it excludes any extensive pan-Andean regional development, as assumed in the putative “Andean” population group claimed in previous analyses.^{4,7} Second, it excludes a major external population source for changes associated with the adoptions of pottery and crops in Central-Southern Chile during the Late Holocene.^{36,37} Third, it corresponds with the status of Mapudungun and the extinct Chono language as language isolates.³² Mapudungun has long attracted speculation that it could be related to other languages in the Americas, not only those further north in the (Central) Andes but also alternatively to Arawak, Tupí, and even Mayan; however, there is no accepted linguistic support for any of those claims.⁵⁵ Our finding of the genetic isolation of the Mapuche is compatible with the standard linguistic view that Mapudungun is indeed a language isolate. Finally, the lack of any evident serial founder effect cannot straightforwardly be associated with either a Pacific/Andean or an Amazonian route for the first settlement. This does not directly match the scenario of a single major route along the Pacific coast, as hypothesized from archaeology and mtDNA.^{16,22}

Interaction between lineages of the Southern Cone

In the finer-scale structure of the Southern Cone through time (Figure 5), our f_3 results distinguish three main lineages: Argentinean Pampas (*ArroyoSeco2_7700BP*); Central-Southern Chile (*LosRieles_5100BP*), including the modern SC samples; and FS (*LaArcillosa2_5800BP* and *Ayayema_4700BP*) (Figures 5A, 6C, 6D, and S6). The appearance of population structure in the region could correspond with the warming climatic conditions of the Holocene, which peaked around 8.5 kya.^{56,57} The ancient CC individuals (Conchalí in particular) confirm the presence of Mapuche ancestry in regions further north than those where most Mapuche live today.

In the FS, we confirmed the presence of two sub-lineages associated with the sea nomads and foot nomads.¹³ The first lineage, in the western archipelagos, was genetically related to *Ayayema_4700BP* and is associated with groups such as the Chono, Kawéskar, and Yámana. The second lineage, in eastern Patagonia, was genetically related to *LaArcillosa2_5800BP* and is associated with other groups such as the Selk’nam, Haush (also

known as the Manek’enk), and Tehuelche, who spoke languages of the Chonan family (note, not related to Chono in the west). Both the sea nomads and foot nomads experienced admixture from the CC ancestry described here, to which present-day Mapuche are closely related (Figure 6B).

Differences between Mapuche groups: Connections with the Central Andes and with the FS

Multiple lines of evidence suggest past connections between the Central Andes and Central-Southern Chile: (1) cultivated plants that appeared in Central-Southern Chile during the Late Holocene,^{24,25} (2) evidence from Quechua loanwords into Mapudungun,^{33–35} and (3) historical contact with the southernmost expansion of the Inca Empire. We investigated demographic connections with f_4 -statistics and IBD block sharing. The f_4 -statistics do not support preferential allele sharing with the Central Andes, in contrast to the results of another study on a present-day Huilliche sample.⁵⁸ Nevertheless, our IBD analysis (Figures 3 and S4) does show a subtle but robust signal of shared ancestry, but which does not persist into the most recent time-frame considered (IBD fragments over 10 cM). This is compatible with contacts that may have predated the Inca period and may have brought not only crops and loanwords southward but also a faint genetic legacy too.¹⁶

The three SC populations analyzed are genetically closely related to each other. This is in line with Mapuche territorial identities being shaped essentially by geographic residence rather than by different demographic histories, and with their limited linguistic divergence. The fact that together, regional varieties of Mapudungun still form a coherent single language²⁷ is consistent with a relatively recent common origin, followed by geographical expansion and divergence over a timescale of the order of many centuries, but not millennia. However, earlier population structure could have been altered by Mapuche relocating southward first because of the Arauco War and then the so-called “Pacification” of Araucanía,⁵⁹ as well as by long-term internal migration driven by forced population transfers, economic necessity, imposed restructuring of land ownership, and other factors.⁶⁰ In the face of such pressures, local communities also formed alliances that led to the absorption of previously unrelated groups.^{61,62} This fusion of genetically stratified groups would have increased the population diversity of the current Mapuche groups (Figure 2), mirroring the effect of relatively constant effective population size for the SC populations (Figure 3D). A similar effect of recent ethnogenesis through the fusion of structured populations has been suggested to account for the IBDNe profiles of Mexican populations.⁶³ Our result contrasts with the decimation of indigenous groups reported in historical sources^{61,64} and with the results obtained by Lindo et al.⁵³ The different demographic trends obtained by Lindo et al.⁵³ could result from the different methods employed and could be further explored with simulations on a high-resolution dataset.

The IBD analysis shows a high level of shared ancestry between our Pehuenche and Lafkenche samples, suggesting that the two groups differentiated only recently or have continued in close contact for generations. A slightly different genetic profile emerges for Huilliche–Chiloé, distinguished by its high level of IBD sharing with FS populations, indicatively dated at 500–1,000 BP (Figures 3A–3C). Historical sources report strong migration

waves from Chiloé into southern Patagonia for economic reasons, during the 19th century, but these recent migrations cannot explain the sharing of shorter fragments (<7 cM), which date back to an earlier time frame.⁶⁵ A study of mtDNA haplogroups in Chiloé found a composition more similar to ancestries in southern Patagonia than among the Pehuenche and Lafkenche, suggesting a connection to the Chono people, who occupied the Chonos Archipelago, southernmost Chiloé, and the coast around the Gulf of Corcovado,^{66,67} and are associated with the sea nomads. The putative Chono toponymy further north through Chiloé suggests the Chono and Huilliche could have come into contact there.

Integrating genetic results with indigenous Mapuche perspectives

Our genetic results reconstruct the genetic trajectory of Mapuche ancestors back to the first peopling of the Americas. Set into their archaeological and linguistic contexts, our findings enrich and complement the historical records and local narratives of the indigenous populations of southern Chile. As recent scholarship points out (see Tsoie et al.,⁶⁸ Malhi,⁶⁹ Hudson et al.,⁷⁰ and Muller and Dortch⁷¹), best practice in anthropological and genetic research to fill in the gaps in a region's history entails direct indigenous participation. We involved the local community through the process of data collection in 2019 and with a return visit to discuss our results with different members from the various locations in 2022 (STAR Methods). These conversations contributed to the drafting our manuscript. Returning results represents a natural extension of the scientific work. It creates trust and accountability between communities, participants, stakeholders, and the scientific community. From our conversations with local partners and participants, we note that the research focus on the pre-Hispanic period was frequently commented on and positively received. We also note that personal involvement of local partners can be culturally charged, as it requires participants to overcome the stigma long associated with indigenous descent, given the historical contexts of long-standing abuse and exploitation by representatives of non-Mapuche cultures. With this work, we aim to contribute to an emerging scientific framework that takes more into account local cultural codes and complex, dynamic social contexts, and that challenges old models of scientific knowledge production.

STAR★METHODS

Detailed methods are provided in the online version of this paper and include the following:

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at <https://doi.org/10.1016/j.cub.2023.05.013>.

A video abstract is available at <https://doi.org/10.1016/j.cub.2023.05.013#mmc4>.

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AUTHOR CONTRIBUTIONS

C.B. conceived the study with the support of S.S., P.H., and F.I.M. E.A.-I. performed the genetic analysis. C.B. and H.B. performed laboratory analyses. C.B. and K.K.S. supervised the study. C.B. and M.J.A. organized the fieldwork expedition and collected samples. C.B., M.J.A., and E.A.-I. organized the trip to return the genetic results to the communities. M.R.C. provided guidance and expertise with qpGraph and masking analysis. S.A. provided expertise with admixture analysis. C.P. provided expertise with f_4 analysis. C.B., M.R.C., C.P., and A.A. provided expertise with the genetic contextualization within South America. S.S., P.H., R.C., and F.I.M. provided linguistic, historical, and archaeological contextualization. M.J.A. curated community engagement and data interpretation. E.A.-I. wrote the first draft of the manuscript. E.A.-I. and C.B. wrote the final manuscript with major contributions from P.H., S.S., M.R.C., S.A., M.J.A., R.C., and K.K.S.

DECLARATION OF INTERESTS

The authors declare no competing interests.

INCLUSION AND DIVERSITY

We support inclusive, diverse, and equitable conduct of research.

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STAR★METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Biological samples		
Saliva samples from Chile	This study	N/A
Chemicals, peptides, and recombinant proteins		
Proteinase K	Sigma Aldrich	Cat# P6556
Water	Sigma Aldrich	Cat# W4502
1M Tris-HCl pH 8.0	Sigma Aldrich	Cat# AM9856
Ethanol	Sigma Aldrich	Cat# E7023
3M Sodium Acetate (pH 5.2)	Sigma Aldrich	Cat# S7899
EDTA (0.5M) pH 8.0	Thermo Fisher Scientific	Cat# AM9261
UltraPure Agarose	Thermo Fisher Scientific	Cat# 15581044
TBE-Buffer 10x	Thermo Fisher Scientific	Cat# AM9261
6x Loading Dye	Thermo Fisher Scientific	Cat# R0611
SYBR Safe DNA Gel Stain	Thermo Fisher Scientific	Cat# S33102
Critical commercial assays		
Oragene Kit, OG-500	DNA Genotek	Cat# OG-500
Axiom Genome-Wide Human Origins 1 Array	Thermo Fisher Scientific	Cat# 901853
Qubit dsDNA BR Essay Kit	Thermo Fisher Scientific	Cat#Q32853
Deposited data		
Modern DNA Southern Chile individuals	This study	EGAS00001007200
Ancient samples	Allen Ancient DNA Resource	https://reich.hms.harvard.edu/allen-ancient-dna-resource-aadr-downloadable-genotypes-present-day-and-ancient-dna-data
Software and algorithms		
PLINK	Chang et al. ⁷²	https://www.cog-genomics.org/plink/1.9/
ADMIXTURE	Alexander et al. ⁴⁴	http://dalexander.github.io/admixture/index.html
OHANA	Cheng et al. ⁷³	https://github.com/jade-cheng/ohana
Itol	Letunic et al. ⁷⁴	https://itol.embl.de/
BEAGLE	Browning et al. ⁷⁵	https://faculty.washington.edu/browning/beagle/b5_1.html
Refined IBD	Browning et al. ⁷⁶	https://faculty.washington.edu/browning/refined-ibd.html
ALDER	Loh et al. ⁵⁴	http://cb.csail.mit.edu/cb/alder/
IBDNe	Browning et al. ⁵²	http://faculty.washington.edu/browning/ibdne.html
RFMIX	Maples et al. ⁷⁷	https://sites.google.com/site/rfmixlocalancestryinference/
ADMIXTOOLS	Patterson et al. ³⁸	https://github.com/DReichLab/AdmixTools

RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources should be directed to and will be fulfilled by the lead contact, Chiara Barbieri (barbieri.chiara@gmail.com).

Materials availability

This study did not generate new unique reagents.

Data and code availability

- Modern genotype data have been deposited in the European Genome-phenome Archive (EGA; <https://ega-archive.org/>) with accession number EGA: EGAS00001007200 (see [key resources table](#)). Given the sensitive nature of the human genetic data

generated in this study, these will not be made publicly available, but access to the data will be granted by a Data Access Committee upon agreeing the conditions on the Data Access Agreement Form available upon request.

- Main scripts to reproduce the analyses are available on GitHub (https://github.com/epifaniarango/popgen_with_epi).
- Any additional information required to reanalyze the data reported in this paper, together with other scripts used for analysis and plots, is available from the corresponding authors upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Sample collection

The study involved individuals either of self-declared Mapuche ancestry and/or who lived in regions where Mapuche presence was historically attested. Sampling was conducted in early 2019 in Chile's Araucanía region and on the island of Chiloé. Local authorities such as municipalities, cultural centers and figures such as *lonkos* (traditional leaders of Mapuche communities) were consulted prior to and/or during the sampling. In Araucanía, residents of rural inland Andean regions are grouped as individuals of putative Pe-huenche ancestry, while those living near the coast are grouped as individuals of putative Lafkenche ancestry. Individuals of putative Huilliche ancestry were recruited in towns across Chiloé. This sampling underrepresents the documented differences between rural communities in various parts of the island. Exact sampling locations are not disclosed to protect participants' privacy. Participants of both sexes and all ages were recruited. The composition of age and sexes does not influence our analysis of genetic history of the region. Participants agreed to participate after the project's aims had been explained to them extensively, and all signed consent forms. Cultural indicators like grandparents' places of birth, local surnames (often of Mapuche origin), and the native language of parents and grandparents were also noted. The biological sample consisted of ~2 ml of saliva, collected in Oragene tubes (DNAgenotek), and stored with an anonymous code. The research project and sample collection were approved by the Unidad de Ética y Seguridad de Investigación of the Pontificia Universidad Católica de Chile's IRB (project #171009001, decree #1520863561038). All project's steps were performed in compliance with the Declaration of Helsinki. The samples analyzed in this study represent only a small fraction of the population living in the target regions of Araucanía and Chiloé and are only partially representative of these regions' populations and their complex demographic histories.

Ethics and Community Engagement

After finalizing the data analysis, we organized a return expedition trip in early 2022 with the goal of making our results accessible to the participants and the local population. To achieve this, we translated the scientific results into a language accessible to the general public (in Spanish). The presented material was printed in a large format to be displayed conveniently without the aid of screens or video projectors. Additionally, we engaged with local schools in each study area to share our findings with lectures for students and teachers. Our contacts with local stakeholders were crucial for framing of research questions, displaying our results appropriately, and, more generally, being aware of the cultural and social context. The return expedition was conducted before the writing of the present manuscript to incorporate participants' suggestions.

METHOD DETAILS

DNA extraction and genotyping

Before lab processing, a second anonymization step assigned a new random code to each sample to ensure that sample numbers did not follow the sampling chronological order. DNA was extracted from the Oragene kit, following the manufacturer's protocols, in the molecular biology laboratories of the Max Planck Institute for the Science of Human History in Jena, Germany. DNA samples were screened and quantified with a Nanodrop spectrophotometer and Qubit fluorometer and visually assessed by gel electrophoresis. Samples were genotyped at the ATLAS Biolab in Berlin, on the Axiom Human Origins array.³⁸ Genotyping data were processed using Affymetrix Genotyping Console v4.2.0.26. In total, 64 samples were genotyped for 629,443 SNPs. PLINK v1.90b5.2⁷² was used to calculate the missing genotype rate with the command '`--missing`'. The proportion of missing calls per sample is <0.005. A small fraction of sites on the array are potentially triallelic in specific populations and are included by reporting the SNP several times (either 2 or 3 times) with a different name. Most population genetic analyses are designed based on biallelic sites, so the triallelic SNPs were removed from the final dataset. PLINK was then used to calculate the consanguinity coefficient F (i.e., $[\text{observed hom. count} - \text{expected count}] / [\text{total observations} - \text{expected count}]$) and P_i values (degree of relatedness as Proportion of IBD, i.e., $P[\text{IBD} = 2] + 0.5 \cdot P[\text{IBD} = 1]$) between pairs of individuals, filtering for minimum allele frequencies of 0.05. All pairs of individuals have P_i values below 0.2, which excludes the presence of first- or second-degree relatives in the dataset.

The final data set comprises 597,167 SNPs.

QUANTIFICATION AND STATISTICAL ANALYSIS

Comparative datasets

For different sets of analysis, we assembled different datasets. Dataset 1 and variations of it were used to study global relationships; samples from the Americas^{38–41} were merged with a selection of reference populations for each continent (Africa represented by

Yoruba,³⁸ Asia represented by Han Chinese,³⁸ Southeast Asia and Oceania represented by Ami,³⁹ Atayal,³⁹ and Papuan,³⁸ and Europe by French³⁸ and Spanish³⁹) genotyped with the Axiom Human Origins array.³⁸ This dataset contains 584 individuals and an average missing call rate of 0.99642. Dataset 1.2 includes the 61 samples from De la Fuente et al. 2018,¹¹ which were genotyped with a different SNP array (Axiom LAT1 platform, Affymetrix). Dataset 1.2 thus consists of 645 individuals and 96,492 filtered SNPs that overlap in the two genotyping platforms. Dataset 1.3 retains only those individuals from Dataset 1 whose Native American ancestry component is higher than 99.9%. Datasets 2 and 2.2 include a selection of individuals from Datasets 1 and 1.2, respectively, for phasing and identity-by-descent analysis. Finally, Dataset 3 was used to study relationships with aDNA. It combines the modern data with a selection of relevant ancient samples^{6–8,11–13,38–43} downloaded from the Allen Ancient DNA Resource⁷⁸ that are compatible with the Human Origins SNP array format. All the modern individuals are masked to focus on indigenous American history and exclude European and African ancestries. Details on each dataset and the analysis for which it is used can be found in the [Data S1A](#) and [Figure S1B](#).

Population structure analysis

ADMIXTURE⁴⁴ was used on the modern global dataset (Dataset 1) to estimate the proportions of ancestry components for each individual. Before the analysis, variants were pruned to limit pairwise linkage disequilibrium (LD) r^2 to at most 0.4 among neighboring SNPs, in sliding windows of 200 SNPs (step size: 25) using PLINK (*-indep-pairwise 200 25 0.4*), which left 218,339 SNPs for the analysis. We performed 10 replicates from $K = 2$ to $K = 15$. Results were visualized using Pong version 1.4.7.⁷⁹ The analysis was also performed on Dataset 1.2, which includes more populations from the Southern Cone ([Figure S3](#)). With this dataset, 73,212 SNPs were left after LD pruning.

We used OHANA⁷³ to create a covariance matrix of the ancestry components for each value of K from 2 to 15, from the average of the allele-frequency matrices generated by ADMIXTURE (selecting the run with the highest likelihood) from Dataset 1 (P.matrix). We converted the covariance matrix into a distance matrix and made an NJ tree following the OHANA protocol. We used Ito⁷⁴ to rearrange the branches with the Yoruba population as an outgroup and visualize the tree.

To inspect genetic relationships among Native Americans, we selected Native American individuals with no European or African ancestry, according to the previous ADMIXTURE analysis at $K = 8$ (Dataset 1.3). PCA was performed on this dataset using the PLINK option (*-pca*) with LD-pruning.

We also used PLINK on Dataset 1.3 to check for inbreeding coefficients (*-het*) and infer ROHs and the distribution of their lengths (with the setting *-homozyg -homozyg-density 50 -homozyg-gap 100 -homozyg-kb 500 -homozyg-snp 50 -homozyg-window-het 1 -homozyg-window-snp 50 -homozyg-window-threshold 0.05*).

Analysis of Identity-by-Descent fragments

To infer blocks of identity by descent (IBD) shared among populations, we first phased all individuals using Beagle 5.1⁷⁵ without a reference panel and with the following options: *window=20 trim=0.3*. Refined IBD software⁷⁶ was used to identify the IBD fragments with the same options as above: *window=20 trim=0.3*. We used three replicates of phasing and IBD analysis to remove breaks and gaps in IBD segments. Afterward, all replicates were merged using the *merge-ibd-segments* tool using a gap of 0.5 cM (all software versions are available at <https://faculty.washington.edu/browning/refined-ibd.html>). In the Americas, most populations share fragments over 4 or 5 cM, as also verified in other studies.^{41,80} For this reason, we apply a high cutoff and consider only fragments above 4 cM. We excluded pairwise comparisons between two Guaraní populations as this very high proportion of shared fragments would have obscured the remaining continental patterns. We then binned the fragments in three categories: 4–7 cM, 7–10 cM, >10 cM. For each bin, we then calculated the probability of an individual from population A sharing an IBD fragment with an individual from population B. These probabilities were calculated by dividing the number of pairs of individuals from populations A and B who do share fragments by the total number of possible combinations of pairs of individuals from A and B (which is obtained by multiplying the number of individuals in population A by the number of individuals in population B) (following Ioannidis et al.⁵¹). The same analysis is applied to the same set of IBD fragments, filtering for fragments with more than 80% presence of SNPs of Native American ancestry, as defined during the masking step (see below) – therefore excluding fragments of African, European, or “unassigned” descent. The probability was projected as a network onto a map in which populations are nodes and edges between them are scaled in width by the probability of IBD sharing. Only probabilities above 0.005 are displayed. The raster file for creating the map was downloaded from Natural Earth.⁸¹

Dating and demographic analysis

We used ALDER 1.03⁵⁴ to reconstruct admixture times from linkage disequilibrium patterns. To infer the admixture time with Europeans (primarily Spanish) in each target population, we chose two source populations: one proxy for the Spanish population and one for Native American ancestry. For the Spanish parental population, all Spanish individuals were merged. The best representative for Native American population depends on which of the various ancestries of the Americas is most represented in the target population. We therefore ran all possible combinations of source populations, and then selected the runs with the best Z-score and p-value. We only considered populations with a high population size (>7 individuals).

To estimate variation in effective population size over time, we used IBDNe⁵² with the default settings. We used only fragments over 2cM and reconstructed population sizes only for the past 50 generations, as this method is not able to reconstruct older ancestor relationships reliably from SNP array data.⁵² A generation time of 28 years was used to convert generations to years.

Local ancestry analysis and masking

Modern samples can be used for generating high-quality genome-wide data with less effort than ancient samples. On the other hand, modern samples from the Americas harbor a range of ancestries, in particular the highly divergent ancestries brought by European impacts since 1492, which act as confounds for our research focus on the prehistory of the native populations of the Americas. Masking is the process of filtering out variants associated with other components, in our case those from outside the Americas (https://github.com/epifaniarango/popgen_with_epi/).

This regional ancestry analysis is a semi-supervised approach, requiring a reference panel for each ancestry of interest, performed with the software RFMIX v1.5.4.⁷⁷ This analysis uses the same phased haplotypes as the IBD analysis. The reference panels for African and European admixture were constituted by Yoruba and Spanish individuals respectively. The Native American Reference panel was built as follows. We first selected individuals previously identified as “unadmixed” in ADMIXTURE. This set was further filtered with an f_4 -statistic of the form f_4 (Unadmixed Native American Population, Target individual; Han, San)⁴ designed to detect more subtle European and African admixture in each individual. The selected Unadmixed Native American populations for the f_4 were Karitiana, Mixe, and Xavante. The individuals who passed this filter with a non-significant f_4 -statistic were included in the Native American Reference panel (116).

We ran RFMIX with 2 expectation-maximization iterations ($-e\ 2$) that also screens the reference panel in the ongoing analysis, as recommended by the authors.⁷⁷ Parameters settings were: window size = 0.2 cM, spacing = CFR, node size = 5, and number of generations since admixture = 11, according to the analysis from Homburger et al.⁸² and our ALDER results ($-G\ 11\ -n\ 5\ -forward-backward\ -use-reference-panels-in-EM\ -e\ 2\ -w\ 0.2$). The threshold for local ancestry assignment was a probability level of 0.9. To check the consistency of the method, we compared the global Native American ancestry proportions estimated with RFMIX, using a weighted mean by chromosomes, with the proportions calculated by ADMIXTURE. The correlation is almost linear (>0.9 after performing a Spearman’s correlation test), except for the North American samples, for which the RFMIX estimates of Native American ancestry is sometimes smaller (see https://github.com/epifaniarango/popgen_with_epi/blob/Local-ancestry-and-masking/README.md).

African and European ancestries were then “masked” for each sample. Following the previous analysis of local ancestry, we kept only those SNPs assigned to Native American ancestry above a threshold of >0.9 probability. The remaining SNPs were coded as missing data, and individuals were separated into the two-phased haplotypes in a pseudo-haploidization process. We evaluated various masking strategies using quality checks and confirmed that the commonly used pseudo-haploid masking^{10,51} performs well and retains more SNPs for the analysis.

We then removed individuals with $<30\%$ SNPs typed and SNPs with $>50\%$ missing genotypes ($-mind\ 0.7\ -geno\ 0.5$ with PLINK 1.9). To check the performance of the masking protocol, we again computed an individual-based f_4 -statistic of the form f_4 (Unadmixed Native American, Test (Admixed Native American); Han, San) following Reich et al.,⁴ and compared the results before and after masking (for details, see https://github.com/epifaniarango/popgen_with_epi/). The selected unadmixed populations were Kaqchikel, Karitina, Mixe, and Xavante. The results were consistent with a positive f_4 after masking. We also performed PCA visualization to confirm the absence of outliers and no attraction towards European and African individuals (Figures S5A–S5C).

After these quality checks, the masked dataset was merged with the Native American Reference panel and the ancient samples (which do not bear traces of European and African admixture), retaining only individuals with more than 100,000 SNPs (Dataset 3.3).

Dataset 3 was used to perform another ADMIXTURE analysis, following the same protocol described above (Figures S5D and S5E). We summarized the results at $K = 5$ as pie-charts on a map, distinguished by time period (Figure 4). Ancestry proportions were averaged among populations. Ancient samples were grouped as populations if they belong to the same archaeological site and time period.

Marker-frequency-based statistics and ancestry modeling

We compute D-statistics with ADMIXTOOLS³⁸ to analyze fine-scale population dynamics between ancient and modern samples. Genetic affinity in terms of shared genetic drift was quantified with the outgroup f_3 -statistic, with the Mbuti as the outgroup, i.e. f_3 (Mbuti; Pop1, Pop2), using qp3Pop.³⁸ For Pop1 and Pop2 we used all possible combinations of individuals or populations from Dataset 3 and then created an f_3 -distance matrix. Higher f_3 -values imply higher genetic affinity (more shared genetic drift) between Pop1 and Pop2. The converted dissimilarity matrix $1-f_3$ was used to generate an MDS plot using R, and the matrix $1/f_3$ to generate an NJ tree with the R package “ape⁸³”, using Ancient Beringian as the outgroup. The tree was displayed using Itol.⁷⁴

f_4 -statistics were designed to search for an excess of allele sharing between populations, and were computed with qpDstats using the default parameters: “f4mode: YES”, and block jackknife over 5-Mb. Most statistics were computed in the form f_4 (Mbuti, Target; X, Y). X and Y are paired only if their data were generated through the same sequencing technology (SNPchip vs. Shotgun sequencing), to minimize bias and attraction effects. In cases where this was not possible, we also compared various configurations of the f_4 , in order to exclude possible attraction effects. For robustness, the tests were computed with all available SNPs and verified with transversions to confirm that the signal was not biased by aDNA degradation (Dataset 3.3 and 3.4). Transitions in aDNA data often result from miscoding lesions; selecting for “transversions only” allows us to avoid those errors.

To model the relationships between the various Southern Cone populations, we used qpGraph,³⁸ considering only transversions and using the default settings. The qpGraph combines f_2 , f_3 , and f_4 -statistics to check the robustness of the tree topologies that we provide. To reduce bias, we used only those ancient samples genotyped with a capture method close to the SNP chip data used for the modern samples, except for *USR1* and *Ayayema_4700BP*, which were genotyped with shotgun sequencing. We focused on

samples from the Southern Cone, and contextualized them with Mbuti as the basal outgroup, and *Ancient_Beringian* and *Brazil_LapaDoSanto_9600BP* as non-SNA2 references. Our basal tree was built with *Mbuti*, *Ancient_Beringian*, *Brazil_LapaDoSanto_9600BP*, *ArroyoSeco2_7700BP*, *LosRieles_5100BP*, *LaArcillosa2_5800BP*, and *Ayayema_4700BP* (Figure S7A). We used the simplest tree topology (without the basal admixture in the ancestors of *ArroyoSeco2_7700BP* and *LosRieles_5100BP*) for building the following topologies. We tested different configurations without *Ayayema_4700BP* to test for biased in the sequencing method (Figure S7B). We successively added populations in various configurations, keeping only graphs with $|Z| < 3.5$ (following Posth et al.⁸) (Figures S7B–S7D). We explore topologies without admixture edges from *LosRieles_5100BP* and *ArroyoSeco2_7700BP* which are not fully supported in f_4 configurations (Figures S7F and S7G). When incorporating the modern samples, we merged Pehuenche and Lafkenche individuals into a single larger population labeled “Mapuche” (Figures S7E).

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Supplemental Information

**The genetic history of the Southern Andes
from present-day Mapuche ancestry**

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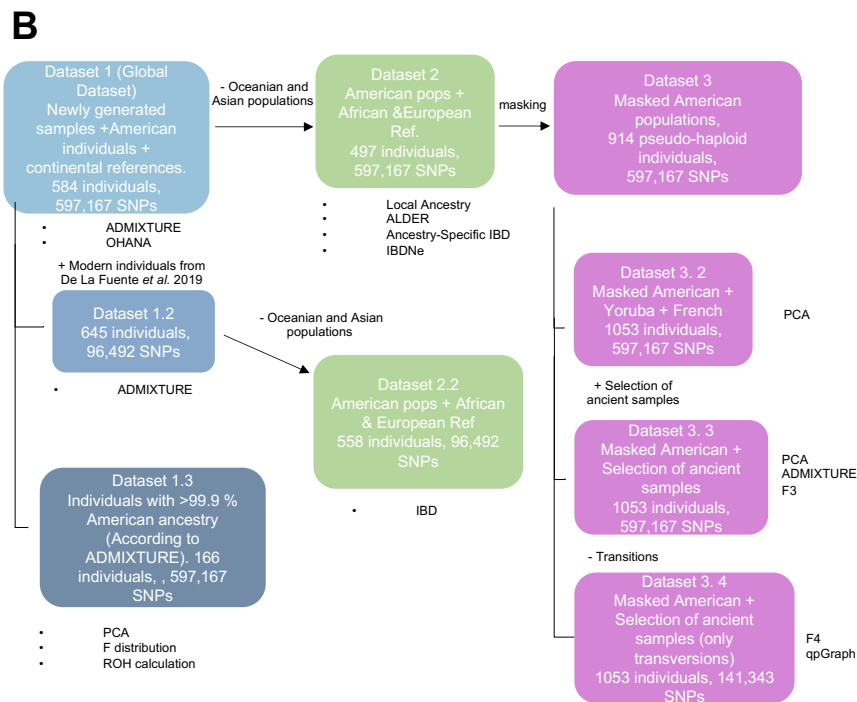
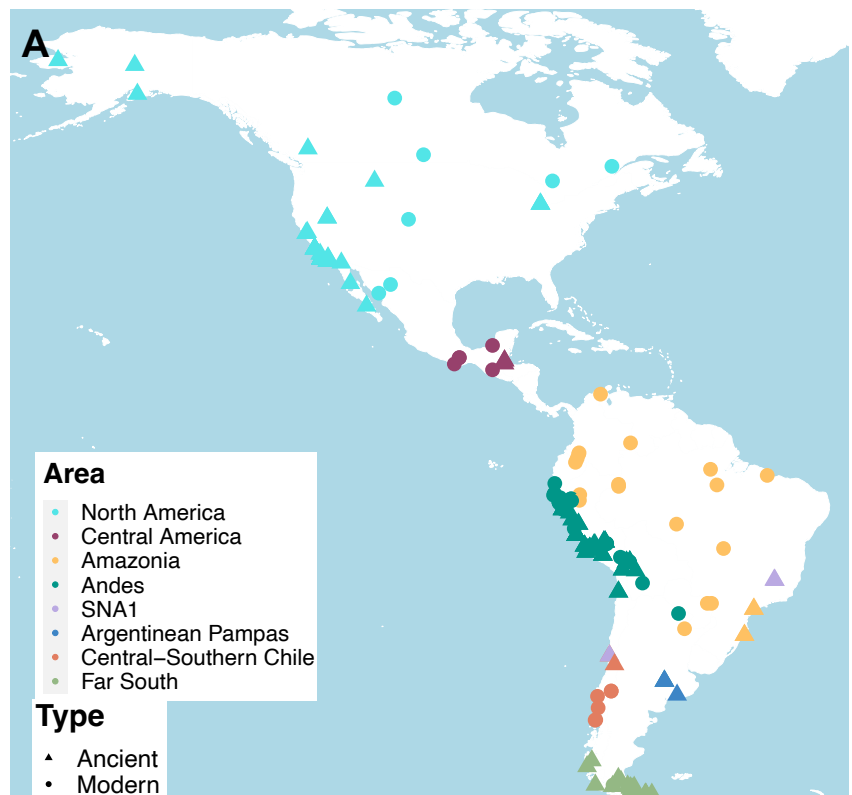


Figure S1: New and published samples used for the analysis, related to Figure 1. (A) Modern populations and ancient samples from the Americas merged with our new dataset in this study. Modern samples are color-coded according to the major ancestry component characterizing them, at $K = 9$ (See Figure S2). Labels correspond to broad ecogeographic regions. Ancient samples are color coded according to major branches from the f_3 outgroup statistic NJ tree (see Figure S6). Labels correspond to broad geographic regions and specific ancestry related to an early wave in the continent (SNA1). **(B)** Schematic description of the datasets, and the analysis for which they are used. For further detail about the individuals on the datasets, refer to the Supplementary Table.

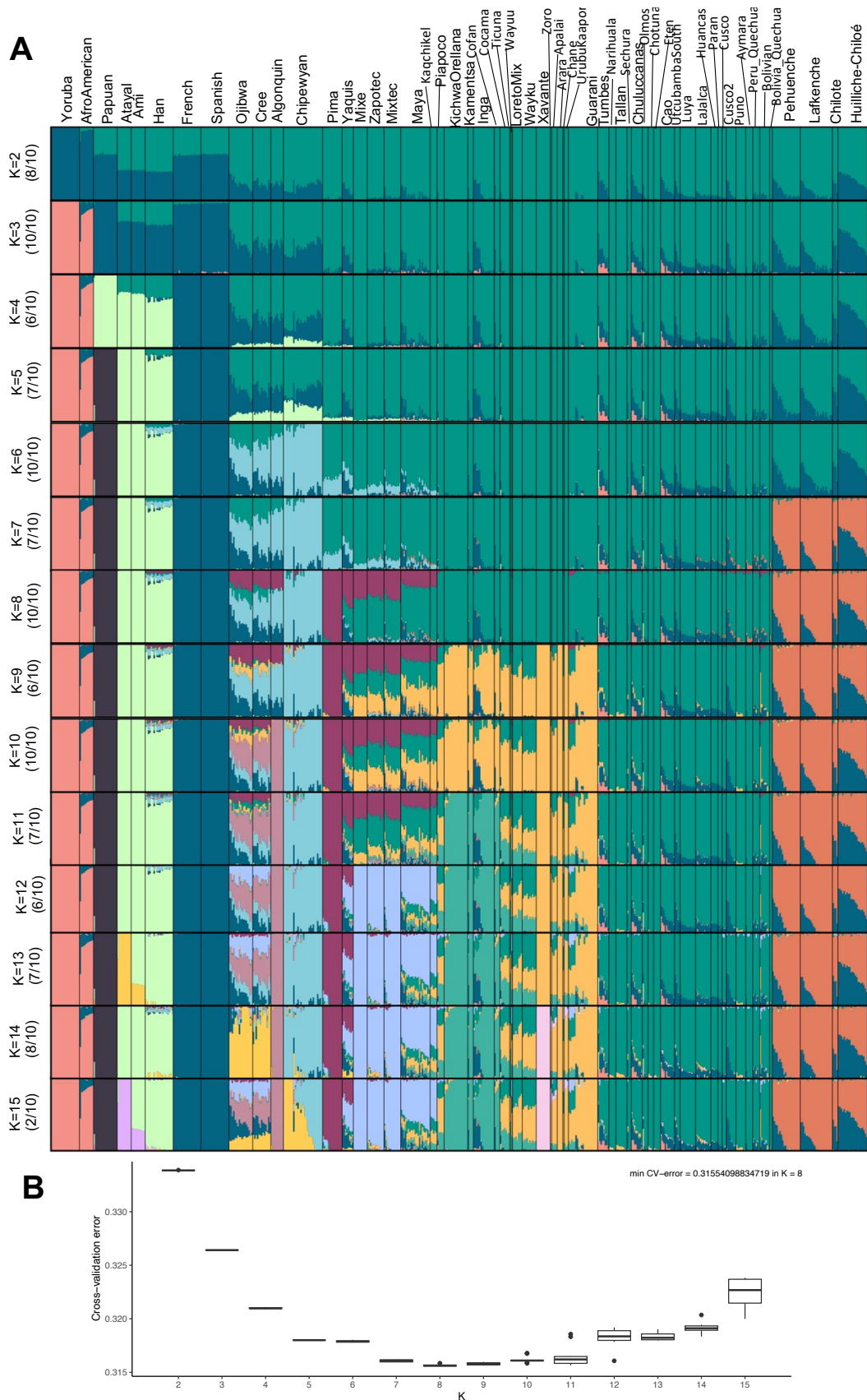


Figure S2: ADMIXTURE Analysis of Global Dataset (Dataset 1) for K = 2-15, related to Figure 1. (A) ADMIXTURE run with the global dataset (Dataset 1). Values in brackets represent the proportion of runs supporting the configuration shown. **(B)** Cross-Validation error associated to each K in the global ADMIXTURE run.

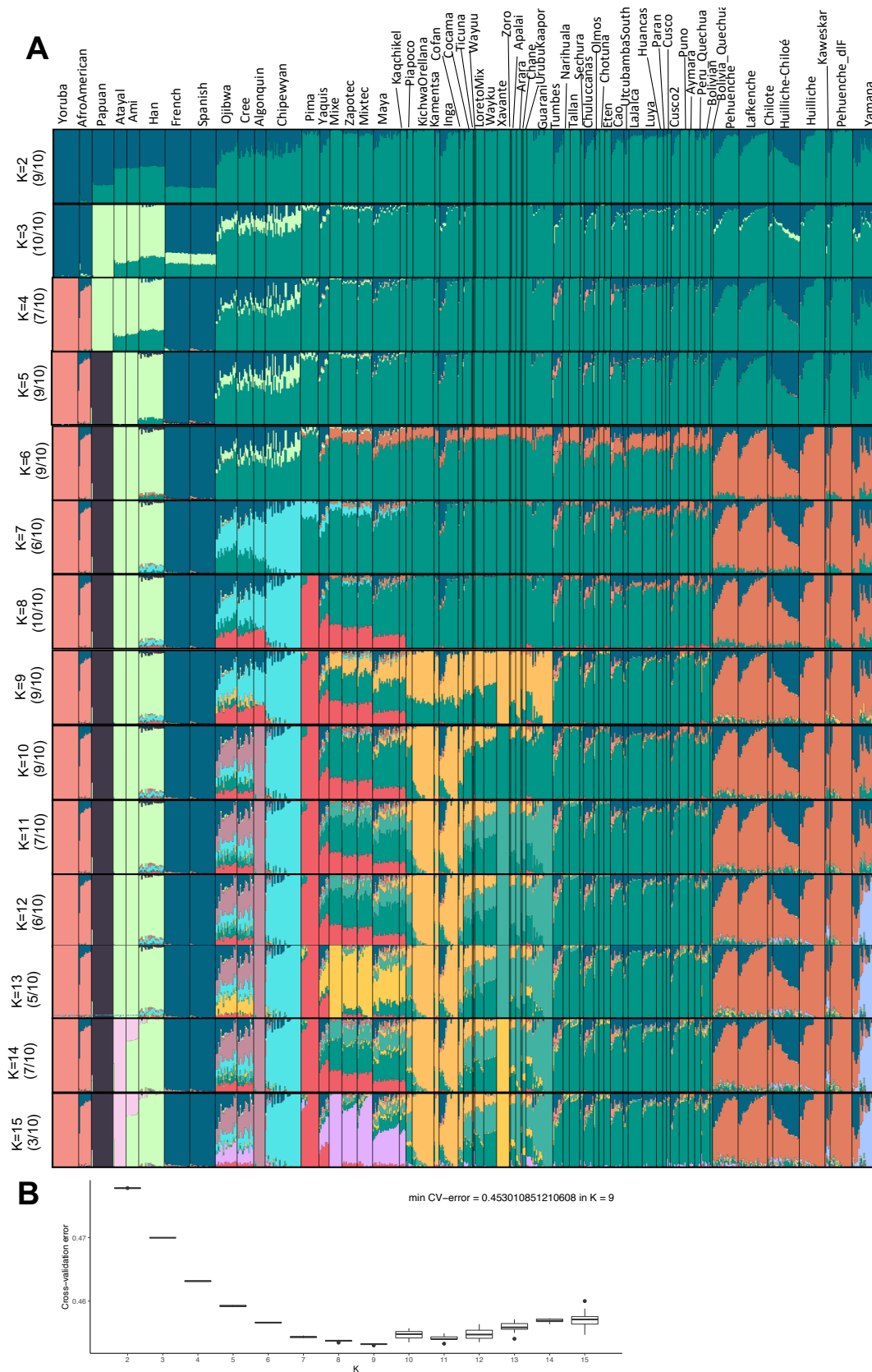


Figure S3: ADMIXTURE Analysis with samples from De La Fuente et al. 2019 (Dataset 1.2) for K = 2-15, related to Figure 1. (A) ADMIXTURE run with Dataset 1.2, including Mapuche, Yamana, and Kaweskar samples from de La Fuente *et al.* (^{S1}) (the last 4 populations on the right). Values in brackets represent the proportion of runs supporting the configuration shown. **(B)** Cross-Validation error associated to each K with Dataset 1.2

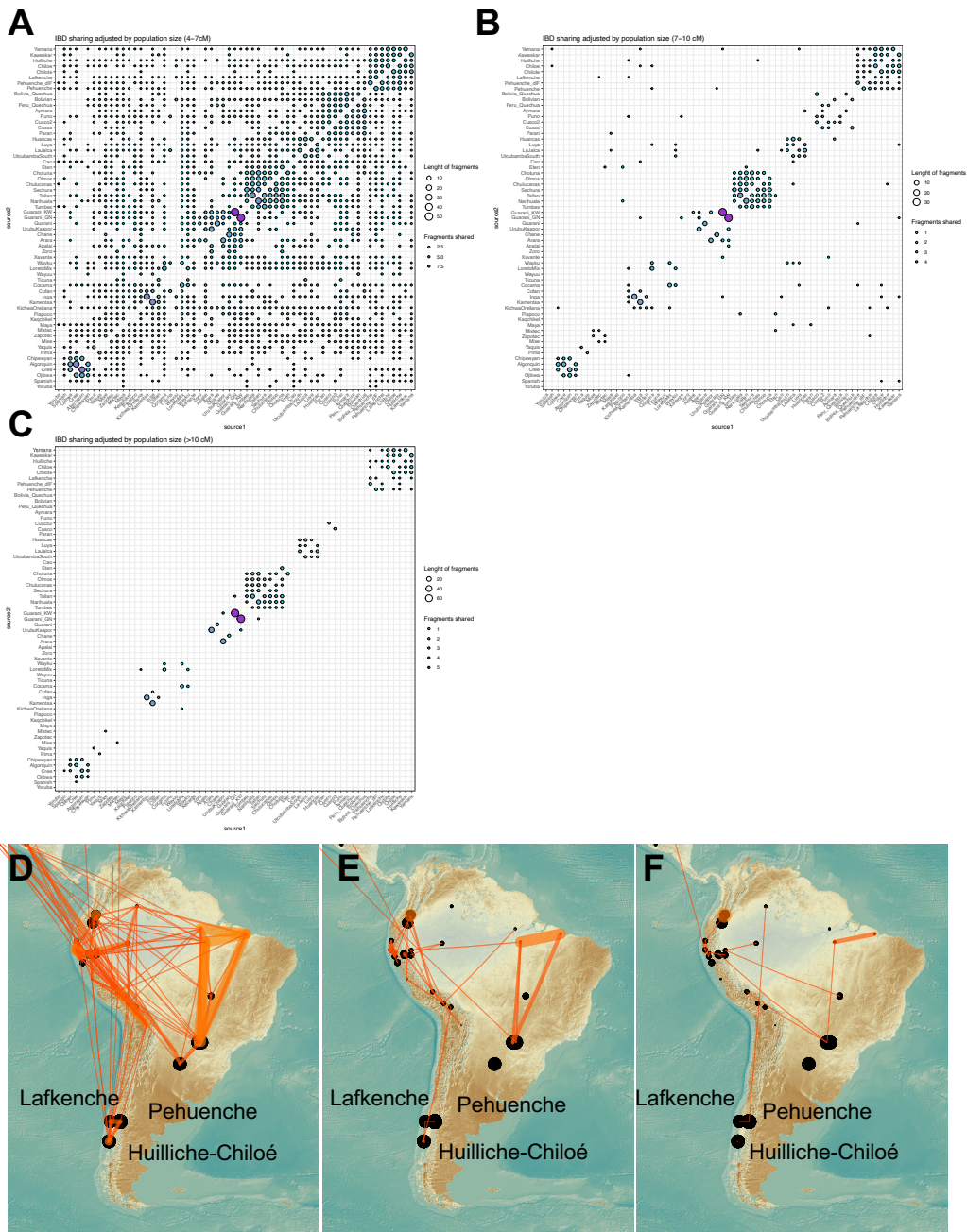


Figure S4: Matrices of Length-Corrected IBD Sharing Among Populations (Dataset 2.2) and IBD Sharing Network of Native American Fragments (Dataset 2), related to Figure 3. The symmetrical matrices of pairwise IBD blocks sharing using Dataset 2.2 show the total length and the number of fragments, adjusted for sample size by dividing by the product of the number of individuals in the pairwise comparison of populations 1 and 2. **(A)** Shared fragments from 4 to 7 cM. As the sharing for this range is almost ubiquitous, we counted only where a pair of populations share more than one fragment. **(B)** Shared fragments from 7 to 10 cM. **(C)** Shared fragments longer than 10 cM. In the maps, IBD sharing probability network among South Americans (Dataset 2). The networks represent the probability of a pair of individuals from populations A and B sharing an IBD fragment, adjusted by population size. The thicker width and lighter orange color of the lines correspond to higher exchange between populations. The size of the black circles is proportional to the sample size. **(D)** Shared fragments from 4 to 7 cM, as sharing within this section is common; we only plotted probabilities higher than 2%. **(E)** Shared fragments from 7 to 10 cM. **(F)** Shared fragments longer than 10 cM.

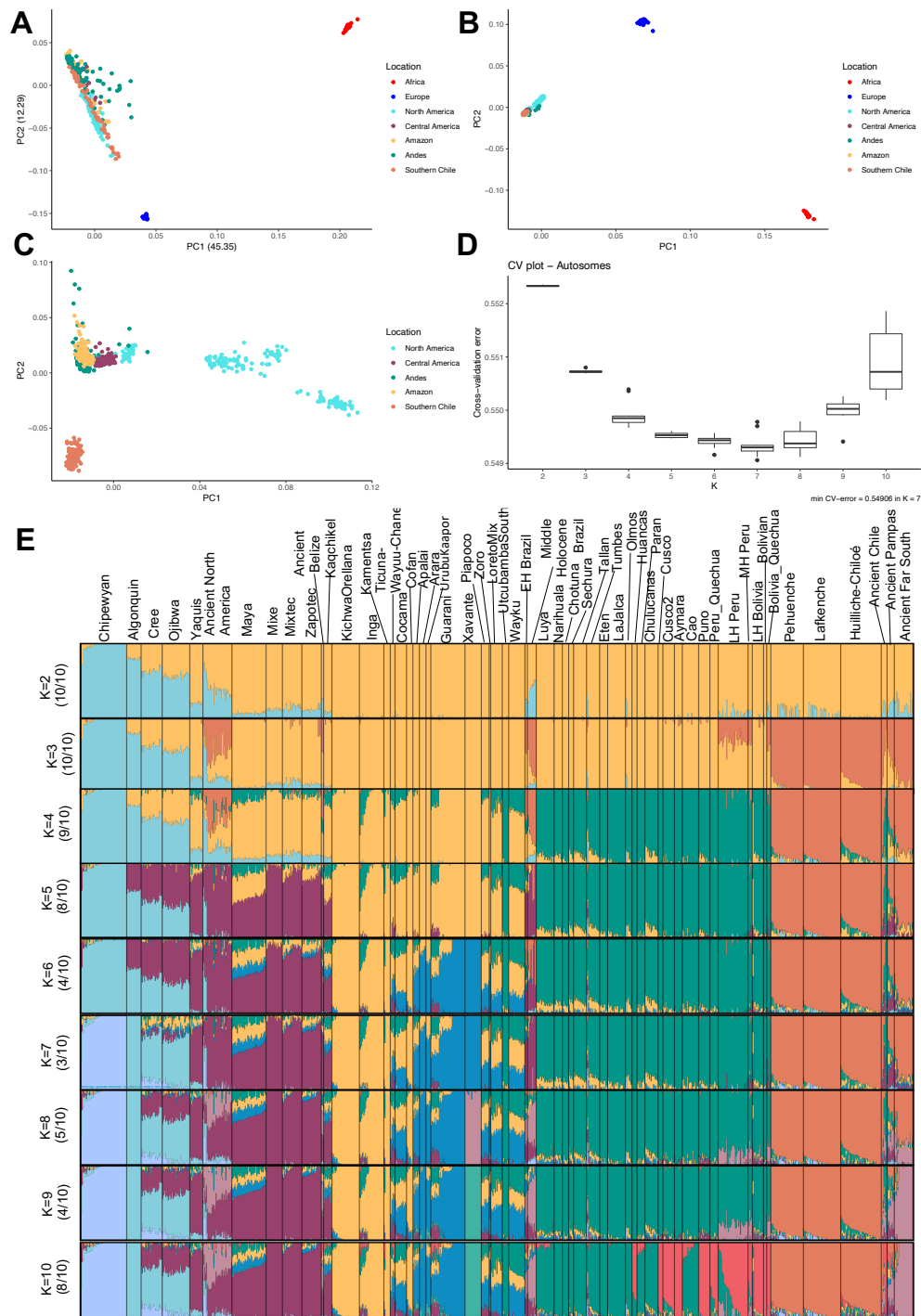


Figure S5: Genetic diversity of masked dataset (Dataset 3.3) illustrated with PCA plots (before and after masking) and ADMIXTURE analysis, related to Figures 4. (A) Samples from Dataset 2 (before masking). Admixed individuals from the Americas are placed in a gradient towards European individuals. **(B)** Masked Native American samples with African and European references (Dataset 3.1). The gradient towards the European individuals is not visible anymore. **(C)** Masked Native American samples (Dataset 3). **(D)** Cross-Validation error associated to each K in the ADMIXTURE run with Dataset 3.3. **(E)** ADMIXTURE run with ancient and masked individuals (Dataset 3.3). Ancient samples are grouped based on geographical proximity and compatible time frame. Values in brackets represent the proportion of runs supporting the configuration shown.

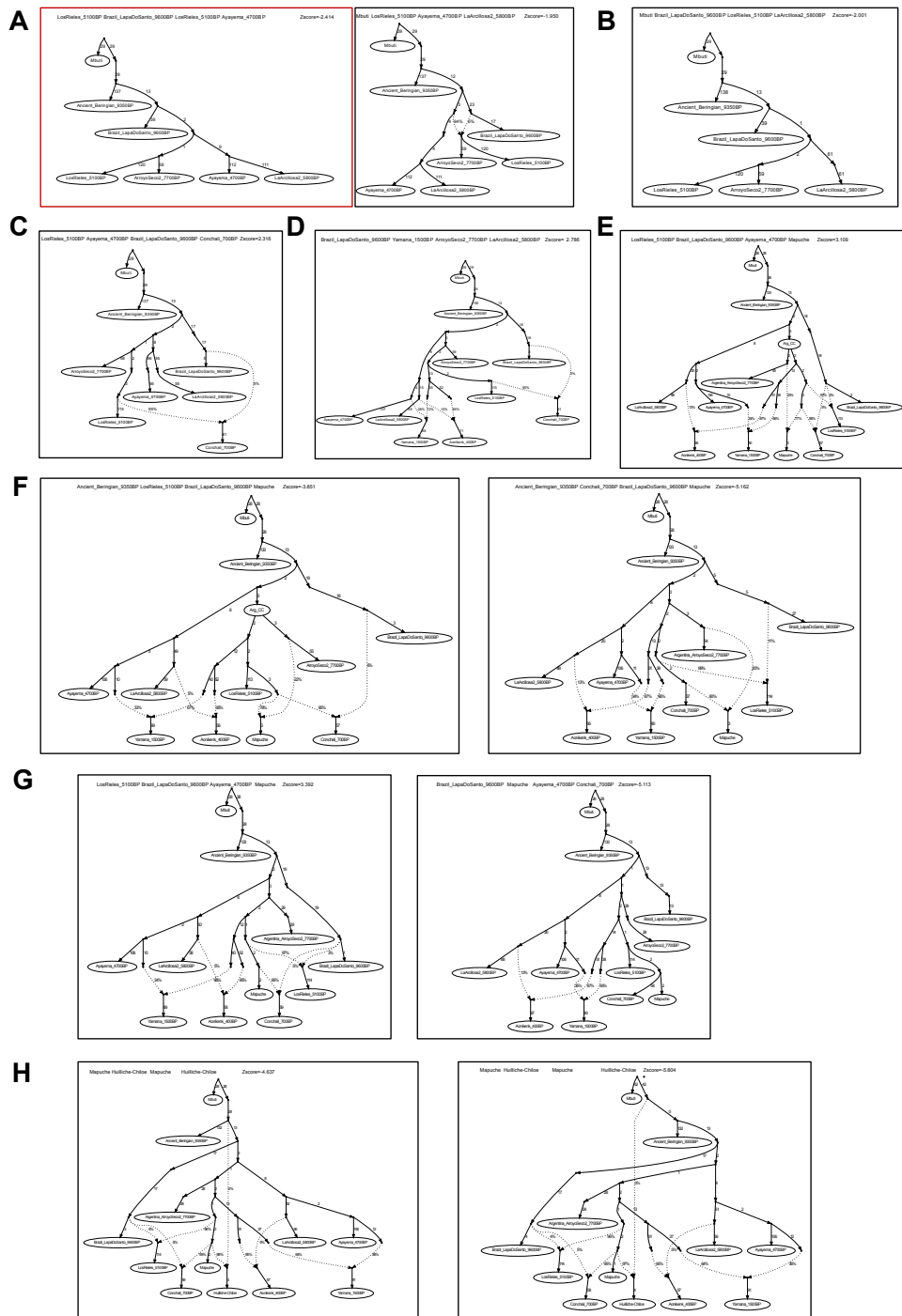


Figure S7: Scaffolds and configurations considered in qpGraph analysis, related to Figure 6D. Numbers on solid lines correspond to genetic drift with units of $F_{ST} \times 1000$. Z-scores were determined from standard errors obtained from jackknife resampling. **(A)** Basal qpGraph trees with Middle Holocene samples from the Southern Cone. The best fitting topology on the left suggests an admixture edge from LapaDoSanto which is not backed up by direct f_4 comparisons. **(B)** Tree topology without the sample of Ayayema, which is sequenced with a different technology (Shotgun). **(C-D)**: Best fitting topologies after adding the ancient samples representative of the CC and FS: **(C)** *Conchali_700BP* and **(D)** *Yamana_1500BP* and *Aonikenk_400BP*. **(E)** Best fitting topology after adding the modern samples of SC Mapuche (which comprises Lafkenche and Pehuenche individuals). **(F)** Alternative tree topologies to (E), without the admixture edges coming from the *Brazil_LapaDoSanto_9600BP* into *LosRieles_5100BP* and *Conchali_700BP*. **(G)** Alternative tree topologies to (E) without the admixture edges coming from the *Argentina_ArroyoSeco2_7700BP* into Mapuche, and without all the admixture edges from *Brazil_LapaDoSanto_9600BP* and *Argentina_ArroyoSeco2_7700BP*. **(H)** Possible tree topologies after adding the modern samples from Huilliche-Chiloé.

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