

1 HIGH RESOLUTION MITOCHONDRIAL DNA ANALYSIS SHEDS LIGHT ON HUMAN
2 DIVERSITY, CULTURAL INTERACTIONS AND POPULATION MOBILITY IN NORTHWESTERN
3 AMAZONIA

4 Leonardo Arias¹, Chiara Barbieri², Guillermo Barreto³, Mark Stoneking¹, Brigitte Pakendorf⁴

5 ¹ Department of Evolutionary Genetics, Max Planck Institute for Evolutionary Anthropology, D-04103,
6 Leipzig, Germany

7 ² Department of Linguistic and Cultural Evolution, Max Planck Institute for the Science of Human
8 History D-07745 Jena, Germany

9 ³ Laboratorio de Genética Molecular Humana, Universidad del Valle, Cali, Colombia,

10 ⁴ Dynamique du Langage, UMR5596, CNRS & Université Lyon 2, 69363 Lyon Cedex 07, France

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13 Leonardo Arias, Max-Planck-Institute for Evolutionary Anthropology, Deutscher Platz 6, D-04103,
14 Leipzig, Germany. Telephone number: +49 341 3550 505, Fax: +49 341 3550 555. E-mail:
15 leoarias2@gmail.com

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22 **ABSTRACT**

23 **Objectives**

24 Northwestern Amazonia (NWA) is a center of high linguistic and cultural diversity. Several language
25 families and linguistic isolates occur in this region, as well as different subsistence patterns: some groups
26 are foragers while others are agriculturalists. In addition, speakers of Eastern Tukanoan languages are
27 known for practicing linguistic exogamy, a marriage system in which partners must come from different
28 language groups. In this study, we use high resolution mitochondrial DNA sequencing to investigate the
29 impact of this linguistic and cultural diversity on the genetic relationships and structure of NWA groups.

30 **Methods**

31 We collected saliva samples from individuals representing 40 different NWA ethnolinguistic groups and
32 sequenced 439 complete mitochondrial genomes to an average coverage of 1030x.

33 **Results**

34 The mtDNA data revealed that NWA populations have high genetic diversity with extensive sharing of
35 haplotypes among groups. Moreover, groups who practice linguistic exogamy have higher mtDNA
36 diversity, while the foraging Nukak have lower diversity. We also find that rivers play a more important
37 role than either geography or language affiliation in structuring the genetic relationships of populations.

38 **Discussion**

39 Contrary to the view of NWA as a pristine area inhabited by small human populations living in isolation,
40 our data support a view of high diversity and contact among different ethnolinguistic groups; movement
41 along rivers has probably facilitated this contact. Additionally, we provide evidence for the impact of
42 cultural practices, such as linguistic exogamy, on patterns of genetic variation. Overall, this study
43 provides new data and insights into a remote and little-studied region of the world.

44

45

46 Northwestern Amazonia (NWA) contains tremendous biological, linguistic, and cultural diversity, which
47 likely reflects the heterogeneity of the landscape, especially the complex and extensive network of rivers
48 found in this area. The region (Figure 1) extends from the Andean foothills in the west to the area
49 between the Orinoco River and the Rio Negro in the east, and extends south until the confluence between
50 the Rio Negro and the Amazon River. The northern border is defined by the Eastern Andean Cordillera
51 and the Colombian-Venezuelan llanos, and in the south by the full length of the Putumayo River (Eriksen
52 2011).

53 In terms of linguistic diversity, NWA harbors ethnolinguistic groups belonging to the main South
54 American language families accepted by specialists (Campbell 1997; Chacon 2014; Dixon and
55 Aikhenvald 1999), namely Arawakan, Carib, Tupi, and Quechua. Additionally, several local families are
56 also present, such as Tukanoan, Guahiban, Huitotoan, Boran, Peba-Yaguan, Piaroa-Saliban and Maku-
57 Puinave, as well as various isolate languages like Tikuna, Cofan, and Kamentsa (Landaburu 2000).
58 Furthermore, several indigenous groups live in voluntary isolation, and almost nothing is known about
59 their linguistic affiliation (Franco 2002). The area has been proposed as the place of origin of the
60 Arawakan family, since it contains the highest linguistic diversity within the family (Aikhenvald 1999;
61 Heckenberger 2002; Zucchi 2002). In addition, all 20 languages of the Tukanoan family are found in the
62 area; these are classified into two branches: The Western Tukanoan branch (WT) distributed along the
63 Putumayo, Caquetá, and Napo rivers, and the Eastern Tukanoan branch (ET) along the Vaupés, Rio
64 Negro, and Apaporis rivers and their tributaries (Chacon 2014). The language families Carib, Tupi, and
65 Quechua are probably recent immigrants in NWA, since only one language per family is present in the
66 area. In addition, the Tupi language Nheengatú or Lingua Geral is found in the region; however, this is a
67 very recent introduction spread by missionaries during the 17th and 18th centuries and by traders during
68 the rubber boom in the 19th century, when it was used as a trade language (Sorensen 1967; Stenzel 2005).
69 [Figure 1 here]

70 In terms of cultural diversity, while NWA has often been viewed as a pristine area inhabited only by
71 small, isolated, seminomadic tribes with an economy based on hunting and gathering (Denevan 1992;
72 Meggers 1954), in fact there is considerable variation in subsistence and marriage practices. While some
73 groups are traditional foragers, others engage in agriculture, and instead of isolated groups, archaeological
74 and anthropological evidence now shows that NWA was indeed part of a continent-wide network of
75 exchange and trade. Complex societies organized in chiefdoms and multiethnic confederations arose in
76 the region, and multilingualism and extensive interactions among ethnolinguistic groups were the norm
77 (Heckenberger 2002; Hornborg 2005; Santos-Granero 2002; Vidal 1997). In particular, the groups
78 speaking Eastern Tukanoan languages and some of their Arawakan neighbors living in the basin of the
79 Vaupés River and Rio Negro engage in an exceptional marital practice known as linguistic exogamy
80 (Aikhenvald 1996; Chacon and Cayón 2013; Sorensen 1967; Stenzel 2005). According to this cultural
81 norm, individuals are required to marry someone from a different language group, with each individual's
82 linguistic affiliation determined by the language of the father. Linguistic exogamy thus creates a situation
83 of multilingualism and movement of people (especially women, since it is accompanied by patrilocality
84 and patrilineality) among the groups participating in the system (Sorensen 1967).

85 Historical linguistics, cultural anthropology, and archaeology are the main disciplines that have
86 traditionally addressed questions regarding the origins, prehistory, and genetic relationships among the
87 NWA ethnolinguistic groups (Campbell 1997; Chacon 2014; Heckenberger 2008; Lathrap 1970; Meggers
88 1948; Nettle 1999). However, due to the scarcity of the archaeological record, the time-depth limitations
89 of linguistic methods based on lexical cognates to establish deep relationships (Dediu and Levinson 2012;
90 Hock and Joseph 2009), and the insufficiency of documentation and description of a large number of the
91 NWA societies, many of these questions remain to be fully answered. The oldest archaeological evidence
92 of human occupation in NWA comes from a single site on the middle Caquetá River dated between 9250
93 and 8100 BP, containing a great variety of stone artifacts, carbonized seeds and other botanical remains
94 from different palm species, as well as phytoliths of bottle gourd, leren, and pumpkin (Aceituno et al.

95 2013; Gnecco and Mora 1997), indicating that these early human groups relied on vegetable resources
96 that are still exploited by contemporary societies in NWA.

97 One hypothesis about the peopling of NWA was proposed by Nimuendajú (1950), who suggested that the
98 region was first inhabited by hunter-gatherer populations, perhaps the ancestors of the Maku-Puinave
99 groups, most of whom still practice a foraging lifestyle. Proto-Arawakan groups then started expanding in
100 the region from their place of origin located between the Orinoco river and the Rio Negro (Heckenberger
101 2002; Lathrap 1970), and finally, the Tukanoans are assumed to have arrived in the area and displaced
102 peoples speaking Arawakan and Maku-Puinave languages from the Vaupés (the Tukanoans probably
103 came from the Napo-Putumayo, where Western Tukanoans still live). However, this scenario does not
104 account for the presence of groups belonging to the Carib, Guahiban, Huitotoan, and Boran language
105 families and the various language isolates in the region.

106 Genetic studies can provide insights into population history, and indeed studies of mitochondrial DNA
107 (mtDNA) genetic variation in Native American populations have contributed greatly to our knowledge
108 about the peopling of the Americas. Early studies using restriction fragment length polymorphisms
109 (RFLP) and sequencing of the hypervariable region one (HVS-I) identified five founder lineages or
110 haplogroups, designated as A to D and X (Bailliet et al. 1994; Barbieri et al. 2011; Gaya-Vidal et al.
111 2011; Keyeux et al. 2002; Lewis et al. 2007; Schurr 2004; Torroni et al. 1993). Whereas haplogroups A-D
112 are widely distributed in the Americas, haplogroup X is restricted to North America (Bolnick and Smith
113 2003; Malhi et al. 2001). The analysis of HVS-I in several Native American populations showed that
114 haplogroups A-D exhibit similar levels of diversity (Bonatto and Salzano 1997), supporting the
115 hypothesis of a single origin of all Native American populations from a Northeast Asian source.
116 Additionally, HVS-I data have been used to determine the genetic relationships among indigenous
117 populations in South America and to test hypotheses concerning how genetic variation is structured at the
118 regional and continental levels (Barbieri et al. 2011; Gaya-Vidal et al. 2011; Lewis et al. 2007; Marrero et
119 al. 2007; Melton et al. 2007). These studies revealed that Andean (or western) populations show higher

120 levels of diversity and low genetic distances in contrast to the Eastern populations, who show the opposite
121 pattern. However, in previous studies NWA populations have been generally underrepresented, and hence
122 the inferences about the genetic structure of the entire Amazonian region are based on a small number of
123 populations.

124 Recent developments in sequencing technology allow the determination of complete mtDNA genomes at
125 the population level and thus enable unbiased insights into the maternal history of human populations
126 (Delfin et al. 2014; Gunnarsdottir et al. 2011; Kivisild 2015). At present, no such studies are reported for
127 South American indigenous populations. Available studies of complete mtDNA genomes from Native
128 Americans have been restricted to a limited number of individuals carrying particular haplogroups,
129 usually selected based on their HVS-I sequences (Achilli et al. 2013; Bodner et al. 2012; de Saint Pierre et
130 al. 2012; Fagundes et al. 2008; Lee and Merriwether 2015; Perego et al. 2009; Perego et al. 2010), or to
131 archaeological remains from different time periods (Fehren-Schmitz et al. 2015; Llamas et al. 2016).

132 These studies have primarily focused on inferences about the peopling of the continent, the number of
133 migrations, the divergence times, and changes in the effective population size through time. Nevertheless,
134 several problems and biases are associated with this sampling strategy. First, the overall diversity might
135 be underestimated, since individuals carrying the same HVS-I sequence can exhibit considerable variation
136 in the coding region (Gunnarsdottir et al. 2011). Secondly, the reconstruction of demographic trends can
137 be skewed, since the estimation of effective population sizes through time using Bayesian coalescent
138 methods (i.e. Bayesian skyline plots in BEAST) can generate spurious signals of population growth when
139 based on samples selected by haplogroup (Gunnarsdottir et al. 2011). Lastly, the histories and origins of
140 specific populations cannot be investigated, since the coalescent age of a particular lineage does not
141 correspond to the age of the population, especially when the diversity within each lineage is unknown
142 (Schurr 2004). In this study, we use complete mtDNA sequencing in a large and representative sample of
143 populations covering the extant ethnolinguistic diversity from NWA to reconstruct their maternal history,
144 as well as to determine their genetic diversity and to make inferences about the origins of this diversity.

145 Finally, we aim to investigate the impact of prehistoric population dynamics and cultural interactions on
146 the structure of the genetic variation observed among present-day NWA populations.

147

148 **MATERIALS AND METHODS**

149 **Sample collection**

150 Samples from unrelated individuals belonging to 40 ethnolinguistic groups were collected during several
151 expeditions carried out by one of the authors (L.A.) in five departments (administrative divisions) of
152 NWA, namely: Amazonas, Guainía, Guaviare, Meta and Putumayo (Table 1, Figure 1). The samples
153 consisted of either saliva, collected as 3 mL of saliva in 3 mL of lysis buffer (Quinque et al. 2006), or
154 blood samples stabilized with EDTA. Written informed consent was obtained from each participant, and
155 from the community leader and/or local/regional indigenous organizations, after giving a full description
156 of the aims of the study. Local translators and fieldwork assistants helped to explain and translate into the
157 local languages when individuals or communities were not proficient in Spanish. Additionally, each
158 participant answered a short questionnaire soliciting information regarding their birthplace, language,
159 ethnic affiliation and that of their parents and grandparents. The study was approved by the ethics
160 committee of the Universidad del Valle in Cali, Colombia and the Ethics Commission of the University of
161 Leipzig Medical Faculty. All procedures were undertaken in accordance with the Declaration of Helsinki
162 on ethical principles and an export permit was issued by the Colombian Ministry of Health and Social
163 Protection.

164 [Table 1 here]

165 **DNA sequencing and sequence processing**

166 The DNA was extracted from blood samples with the “salting out” method (Miller et al. 1988) and from
167 the saliva samples with the QIAamp DNA Midi kit (Qiagen), starting from 2.0 mL of the saliva/buffer

168 mixture. The concentration of DNA was quantified with the NanoDrop 8000 spectrophotometer (Thermo
169 Scientific). We prepared genomic libraries with double indices and enriched for full mtDNA genomes
170 using a hybridization-capture method described previously (Kircher et al. 2012; Maricic et al. 2010).
171 From the enriched libraries, paired-end sequences of 100 bp length were generated on the Illumina HiSeq
172 2500 platform. Base-calling was performed using freeIbis (Renaud et al. 2013), and Illumina adapters
173 were trimmed and completely overlapping paired sequences were merged using leeHOM (Renaud et al.
174 2014a). The sequencing data were de-multiplexed using deML (Renaud et al. 2014b) and the sequences
175 aligned against the human reference genome 19 using BWA's *aln* algorithm (Li and Durbin 2009). After
176 duplicate removal using PicardTools v2.1.1 (<https://github.com/broadinstitute/picard>), we performed an
177 iterative alignment for each library individually to obtain mtDNA consensus sequences. In the first step,
178 we extracted all sequencing reads of a library that aligned either to the mitochondrial genome or to a list
179 of nuclear copies of mtDNA (NUMTs) (Li et al. 2012). We subsequently aligned these reads to the
180 revised Cambridge Reference Sequence (rCRS; Andrews et al. 1999) using BowTie2's *very-sensitive*
181 algorithm (Langmead and Salzberg 2012) and called a consensus sequence. In the second step, the reads
182 were re-aligned to the library's respective consensus sequence generated in the first step, using the same
183 BowTie2 settings. After the second alignment step, we called a final consensus sequence that was used
184 throughout the rest of the analysis. Final sequences in fasta format were aligned to the Revised
185 Cambridge Reference Sequence (rCRS (Andrews et al. 1999)) with the multiple sequence alignment
186 software Mafft (Katoh and Standley 2013), and manually inspected for alignment errors with Bioedit ver.
187 7.2.5 (Hall 1999). The two poly-C regions (np 303–315 and 16,183–16,194) were excluded from the
188 subsequent analysis; although one position (16,189) diagnostic for haplogroup B2 is therefore not
189 considered in the haplogroup-calling analysis, the additional substitutions defining this haplogroup that
190 occur elsewhere in the mitochondrial genome enable unambiguous assignment of sequences to lineage
191 B2.

192

193 **Data analysis**

194 We considered populations with a sample size of 10 individuals or more, and merged populations with
195 sample sizes smaller than 10 based on linguistic criteria when our initial analyses did not show significant
196 genetic differences, as follows (Table 1). The Arawakan groups Achagua (n=6) and Piapoco (n=18) were
197 merged into a single population, since their indigenous reservations are adjacent and individuals often
198 intermarry (data available on request); one Bare (n=1) individual was added to the Curripaco (n=16)
199 sample among whom he was living when sampled on the Atabapo River; Yucuna (n=31) and Matapi
200 (n=8) were merged into a single population, since they both speak Yucuna, live along the same river, and
201 intermarry (data available on request); and Murui (n=18) and Uitoto (n=8) were merged, as these two
202 groups belong to the same language family, which is composed of several dialects that are mutually
203 intelligible (<http://glottolog.org/resource/languoid/id/huit1251>, accessed on 31.05.2017). Finally,
204 following the latest classification of the Tukanoan family (Chacon 2014), the Eastern Tukanoan groups
205 Piratapuyo (n=8) and Wanano (n=5) were merged as Pira-Wanano; Tukano (n=8) and Tatuyo (n=2) were
206 merged as Tuka-Tatuyo; and Tuyuca (n=7), Yuruti (n=1), Pisamira (n=1), and Karapana (n=1) were
207 merged as Other-ET. The only group with a sample size smaller than 10 that we retained as a separate
208 group in the analyses were the Carijona (n=8), since this is the only Carib-speaking group living in NWA.
209 Moreover, they are at risk of disappearing both physically and culturally, with less than 30 active speakers
210 of Carijona scattered in two communities, and they occupy an important place in the ethno-history of the
211 region (Franco 2002). We excluded Barasano (n=4), Kubeo (n=5), Cofan (n=6), Cabiari (n=1),
212 Guambiano (n=1), and Nasa (n=1) from all the analyses except the haplotype networks, since this analysis
213 represents the evolutionary relationships among individual sequences. We furthermore excluded nine
214 individuals with maternal ancestry outside of NWA (labeled 'Mestizo' in Table 1) from all analyses.
215 After merging and filtering as described above, 412 sequences from 24 groups were kept in the
216 population-based analyses.

217 Based on information from D-PLACE (Kirby et al. 2016) and HG database
218 (<https://huntergatherer.la.utexas.edu/home>, accessed on 06.06.2017), we divided the populations into
219 agriculturalists (AG) and hunter-gatherers (HGP). In the latter category we included the Nukak, who
220 currently still practice a foraging way of life, as well as the Puinave, Sikuani, and Guayabero, who have
221 all adopted agriculture only recently (Kondo 2002; Uribe Tobón and Instituto Colombiano de Cultura
222 1992).

223 The haplogroup affiliation of the individual sequences was determined with Haplogrep (Kloss-
224 Brandstatter et al. 2011), based on Phylotree build 16 (van Oven and Kayser 2009). Haplogroup
225 frequencies by population were estimated by simple counting, and a correspondence analysis (CA) based
226 on the frequency of sub-haplogroups (e.g. A2a) was performed and visualized with the R-packages
227 FactoMineR (Le et al. 2008) and factoextra (Kassambara and Mundt 2016), respectively.

228 Population-based statistical analyses were performed with Arlequin v3.5 (Excoffier and Lischer 2010);
229 these include the analysis of molecular variance (AMOVA), estimation of molecular diversity indices, the
230 estimation of pairwise genetic distances based on Φ_{ST} , and Tajima's D test of selective neutrality. A
231 Multidimensional Scaling analysis (MDS) was performed on the matrix of pairwise Φ_{ST} values to
232 visualize the distances between populations. Additionally, we performed a Mantel test to evaluate the
233 correlations between genetic distances and geographic distances. The matrix of geographic distances was
234 built using the geographic coordinates of the location where the majority of samples for each
235 ethnolinguistic group were collected and then calculating the great circle distances between locations via
236 the R packages ade4 and geosphere (Dray and Dufour 2007; Hijmans 2016). Furthermore, a multiple
237 regression analysis on distance matrices (MRM) (Goslee and Urban 2007) with the form:
238 $MRM(as.dist(gen.dist) \sim as.dist(geo.dist) + as.dist(rivers.dist))$ was performed; the analysis takes into
239 consideration a matrix of geographic distances and a matrix of proximity along rivers as predictor
240 variables of the genetic distances (pairwise Φ_{ST} values) between populations (Pugach et al. 2016;
241 Yunusbayev et al. 2012). For the matrix of river distances a value of zero was given to populations living

242 along the same river or on rivers that are closely connected, and a value of one was given to populations
243 living on different rivers.

244 The sharing of haplotypes between populations was estimated with in-house R scripts as the proportion of
245 pairs of identical sequences shared between populations. Additionally, networks of haplotypes were
246 constructed with the software Network ver. 4.6.1.3 and visualized with Network Publisher ver. 2.0.0.1
247 (<http://www.fluxus-engineering.com>). Finally, Bayesian skyline plots (BSP) were constructed with
248 BEAST ver. 1.8.2 (Drummond et al. 2012). For this analysis, the best substitution model was estimated
249 with jModeltest 2.1.7 (Darriba et al. 2012), and BEAST was used to estimate whether a strict or a relaxed
250 clock model best fits the data. This analysis was performed on both the complete sequences and the
251 sequences partitioned into coding (577-16023) and non-coding (16024-576) regions, applying the
252 corresponding substitution rates reported previously (Soares et al. 2009).

253

254 **RESULTS**

255 We generated 439 complete mitochondrial sequences to an average coverage per sample of 1030x, which
256 were deposited in GenBank with accession numbers: XXXXXXXXX-XXXXXXXXX and XXXXXXXXX-
257 XXXXXXXXX. All sequences belonged to one of the main Native American haplogroups, namely A2, B2,
258 C1 and D1. Haplogroups A2 and C1 were the most frequent lineages in the NWA populations (excluding
259 the so-called ‘Mestizos’), with more than half of all sequences belonging to A2 (90 haplotypes in 138
260 sequences) and C1 (95 haplotypes in 181 sequences) together; in contrast, B2 (49 haplotypes in 73
261 sequences) and D1 (32 haplotypes in 38 sequences) were less frequent. Table 2 provides a breakdown of
262 the haplogroup frequencies for the ethnolinguistic groups included in the population analyses.

263 [Table 2 here]

264 A Correspondence Analysis (CA) (Figure 2) shows the clustering of populations based on the frequency
265 of sub-haplogroups. We observed differences among populations without a clear clustering by language

266 family, with the exception of the Eastern Tukanoan groups Siriano, Desano, Pira-Wanano, Tuka-Tatuyo
267 and Other-ET, which were near one another in the left side of the plot. However, Tanimuka, who also
268 speak an Eastern Tukanoan language, were far apart from their linguistic relatives. Additionally,
269 Guayabero and Sikuaní (who speak languages belonging to the Guahiban family) were located close to
270 each other in the lower left pane of the plot. In addition to language affiliation, a few cases of proximity in
271 the CA plot could be attributed to geographic proximity, as in the case of Kamentsá, Pasto and Inga, who
272 all live close to one another in the Andean foothills. In other cases, the relatively close proximity of
273 populations could be attributed to their being settled along the same river or on rivers that are part of the
274 same basin (Supporting information Figure S1), as in the case of Curripaco and Puinave, who live on the
275 Inírida and Atabapo rivers.

276 [Figure 2 here]

277 **Molecular diversity indices**

278 The genetic variation in these communities was assessed through different molecular diversity indices
279 (Figure 3, Supporting information Table S1). On average, the gene diversity in these groups was high
280 (0.9), but there were differences amongst them. For example, Eastern Tukanoan groups showed
281 consistently high values of gene diversity, with the exception of the Tanimuka, who had one of the lowest
282 values (0.73). The Western Tukanoan groups Coreguaje (0.92) and Siona (0.82) showed lower values
283 than Eastern Tukanoan groups. Among Arawakans, the Ach-Piapoco had the lowest value (0.77). The
284 hunter-gatherer group Nukak showed the lowest gene diversity of all groups (0.64): only four haplotypes
285 were observed among the 16 individuals analyzed. Additionally, we observed that agriculturalist groups
286 tended to have higher gene diversities (average = 0.92) than hunter-gatherer groups (average = 0.80)
287 (Mann-Whitney U test, P-value = 0.03).

288 [Figure 3 here]

289 The mean number of pairwise differences (MPD) per population showed less variation, with an average
290 of 41.07 +/- 17.86 differences. The smallest values were found in Sikurangi (24.08 +/- 11.18) and Nukak
291 (31.07 +/- 14.33), and the largest values were observed in Cocama (44.64 +/- 20.37), Carijona (42.71 +/-
292 20.8), and Siriano (41.38 +/- 19.66) The D values of Tajima's test of neutrality (Tajima 1989) ranged
293 from -0.735 to 2.318. Under neutrality, Tajima's D is expected to be equal to zero and significant
294 departures are interpreted as a result of selection or changes in population size. Although none of the D
295 values were significant (all P-values > 0.2), positive D values >1.2 were obtained for Guayabero, Nukak,
296 Sikurangi and Tanimuka, which may reflect recent reductions in the size of these populations. This
297 hypothesis was supported both by the distribution of pairwise differences by population (Supporting
298 information Figure S2), which showed increased frequencies for the category of small differences (0 and
299 1 differences) and for the category of large differences (50 or more), as well as by the Bayesian
300 reconstruction of population size changes through time (BSP plots, Supporting information Figure S3 and
301 below). Furthermore, the Tanimuka and Nukak had the lowest gene diversity values.

302

303 **Shared haplotypes**

304 A total of 216 different haplotypes were observed among the 412 sequences included in this analysis,
305 pointing to a considerable number of shared sequences. Of these, 146 were unique haplotypes and 70
306 haplotypes were shared among 266 sequences: 52 within populations, 31 between populations, and 13
307 both within and between populations. The shared haplotypes accounted for 64.6% of all the sequences
308 analyzed. This amount of haplotype sharing between populations is considerably high when compared to
309 other population-based studies of complete mitochondrial genomes (Table 3). In other studies, the
310 majority of shared haplotypes were generally observed within populations, with the exception of two
311 African datasets from Burkina Faso and Zambia (Barbieri et al. 2013; Barbieri et al. 2012), which showed
312 low levels of sharing both within and between populations. The highest level of sharing between
313 populations was observed for Siberian populations spread over a large geographic area (Duggan et al.

314 2013); the NWA populations analyzed in this study showed the second highest value of sharing between
315 populations.

316 [Table 3 here]

317 Figure 4 shows the proportion of pairs of sequences shared between and within NWA populations.
318 Siriano, Other-ET, and Pasto were the only groups without shared haplotypes within the populations. The
319 majority of between-group haplotype sharing involved Arawakan and Eastern Tukanoan groups. The
320 Arawakan groups share mostly with groups living in close proximity (Figure S4), e.g. Yucu-Matapi
321 shared with Tanimuka; Curripaco with Puinave and Nukak, and Ach-Piapoco with Saliba and with the
322 Guahiban groups Sikuani and Guayabero. In contrast, most Eastern Tukanoan groups, who practice
323 linguistic exogamy, shared haplotypes among each other (except for Tanimuka, who shared only with
324 Yucu-Matapi). In contrast, the Western Tukanoan groups Siona and Coreguaje shared primarily within
325 their populations and did not share haplotypes with the Eastern Tukanoan groups.

326 The groups from the Andean foothills--Inga, Kamentsa, and Pasto--showed different patterns of shared
327 haplotypes, despite the fact that they live in close geographic proximity. The Pasto, a group that has lost
328 its native language and that is highly incorporated into the admixed local population, shared no
329 haplotypes with any population. The Kamentsa shared haplotypes only with the Inga, while the Inga also
330 shared haplotypes with three other groups located further inside the Amazonian area: Carijona, Coreguaje
331 and Mur-Uitoto. Finally, of the three groups living on the banks of the Amazon River close to the town of
332 Leticia, the Cocama shared with both the Yagua and Tikuna, whereas the latter two groups lacked
333 common haplotypes.

334 [Figure 4 here]

335 **Haplotype networks**

336 The networks of haplotypes (Supporting information Figure S5 A-D) complement the patterns of
337 sequence sharing, but in addition allow us to discern clusters of related (not just identical) haplotypes. We

338 observed that some of these clusters were common among different language families and others were
339 restricted to specific language families or to groups living in close geographic proximity; these are
340 highlighted in Figures S5 (A-D). For instance, Arawakan and Eastern Tukanoan groups exhibited several
341 haplotypes within haplogroups A2 (Cluster I, Figure S5A), B2 (Cluster I and II, Figure S5B), and C1
342 (Cluster III, IV, V and VI, Figure S5C) that were either shared or separated by only a few mutational
343 steps. Notably, several of these clusters also included individuals speaking Maku-Puinave languages (cf.
344 Cluster I and II, Figure S5B, and Cluster III, IV, and V, Figure S5C). Clusters of haplotypes restricted to
345 specific groups are represented by clusters I and II in Figure S5D, exclusive to Eastern Tukanoan and
346 Huitotoan populations, respectively. Furthermore, the haplotypes of the Inga (Quechuan) and the
347 Kamentsa, who live in close proximity in the Andean foothills, were either shared between them or
348 closely related (e.g. cluster II in Figure S5C). Finally, the haplotypes of the Guayabero and Sikuani
349 (Guahiban) were mostly differentiated from those of other populations and generally shared by several
350 individuals within the family (clusters II and III, Figure S5A; cluster I, Figure S5C). The sequences
351 belonging to cluster I in haplogroup C (cluster I, Figure S5C) lack the diagnostic mutation A13263G for
352 haplogroup C, but contain other diagnostic mutations that allow unambiguous assignment to haplogroup
353 C. MtDNAs with this variant were previously identified in eastern Colombia by RFLP typing (Torres et
354 al. 2006), where they occurred at high frequency in Guahibo, Piapoco, and Saliba groups. Given their
355 high frequencies in the Guahiban groups, these haplotypes appear to belong to an autochthonous lineage
356 that has then diffused into other groups living in the Orinoco basin.

357

358 **Genetic structure and genetic distances**

359 The AMOVA analysis (Table 4) allows us to test different hypotheses concerning how genetic variation
360 is structured in NWA. We defined groups *a priori* based on language affiliation, geographic proximity,
361 and distribution along major rivers or their tributaries to evaluate how much of the observed variation is
362 explained by each grouping strategy. We observed that of the three grouping strategies, grouping

363 populations by their distribution along rivers resulted in the largest among-group component of the
364 genetic variance. In contrast, language was a poor predictor of the genetic structure, showing negative and
365 nonsignificant values for the component of variance due to differences among groups, indicating larger
366 genetic differences among groups of populations speaking related languages than among linguistically
367 different groups. Finally, geographic proximity was also a poor predictor; although the among-group
368 component was larger than for language, it was not significantly different from zero. An important aspect
369 to note is that although grouping by rivers performed better than grouping by geography or language, it
370 still did not provide a very good description of the genetic structure, since the percentage of variance due
371 to differences among populations within groups was still higher than the among groups component,
372 suggesting the existence of other levels of substructure within populations.

373 [Table 4 here]

374 The matrix of genetic distances between populations based on pairwise Φ_{ST} values (Supporting
375 information Figure S6) was used to construct an MDS plot (Figure 5). The populations do not form any
376 clear clustering: the majority of populations are grouped together in the center of the plot (indicated by
377 the inner circle in Figure 5) with an average pairwise $\Phi_{ST} = 0.03$, while around the main cluster a second
378 group of populations showed higher differentiation (external circle, average $\Phi_{ST} = 0.07$). Finally, Sikuani,
379 Siona, and the hunter-gatherer Nukak appeared as outliers with high genetic differentiation (average Φ_{ST}
380 = 0.22). This picture did not change after adding an additional dimension to the MDS plot (Supporting
381 information Figure S7). Particularly striking were the small genetic distances between the Eastern
382 Tukanoan groups, who clustered together in the center of the MDS plot. Although Tanimuka appeared
383 more distant from the main cluster of Eastern Tukanoan groups, their pairwise Φ_{ST} values were not
384 significantly different (Supporting information Figure S6) and the average Φ_{ST} (0.02) indicated low
385 genetic differentiation among all Eastern Tukanoan groups. In contrast, the Coreguaje and the Siona, who
386 speak languages of the Western Tukanoan branch, showed larger genetic distances, both with the Eastern
387 Tukanoan groups and with each other. Populations from each of the other language families did not form

388 clusters with their linguistic relatives. For example, Arawakan groups occupied different positions in the
389 plot and their Φ_{ST} values were significantly different.

390 [Figure 5 here]

391 The results of the Mantel test showed a lack of significant correlation between geographic distances,
392 estimated as great-circle distances, and the matrix of pairwise Φ_{ST} values ($r = 0.07$, p -value = 0.28).
393 However, since rivers emerged as an important factor explaining the structure of genetic variation in the
394 AMOVA results (Table 4), we also performed a multiple regression analysis on distance matrices
395 (MRM), where we added rivers as an additional predictor variable. Adding rivers to the regression model
396 resulted in an increase in the amount of variation explained by the model (Table 5), with rivers being a
397 significant predictor (p -value = 0.01). We then jack-knifed over populations (Pugach et al. 2016;
398 Ramachandran et al. 2005) and identified three populations as outliers: Sikuani, Siona, and Nukak, who
399 appeared as outliers in the MDS plot as well (Figure 5). We performed the multiple regression analysis
400 excluding the outliers; this resulted in an increase of 3.4 % in the R square value, a better correlation
401 between genetic and geographic distances, and geography becoming a significant predictor factor (p -
402 value < 0.05) (Table 5, supporting information Figure S8), although rivers were no longer a significant
403 predictor of genetic subdivision.

404 [Table 5 here]

405 **Bayesian demographic reconstruction**

406 Bayesian skyline plots (BSP) were generated by haplogroup (A2, B2, C1 and D1) and by population. All
407 four haplogroups showed an increase in effective population size between 17,500 – 25,000 years before
408 present. This signal was more evident for haplogroups A2 and C1, which have the highest number of
409 sequences (Supporting information Figure S9). In contrast, the BSP plots by population showed different
410 outcomes. We observed four main trajectories (Supporting information Figure S3): first, a signal of
411 population size increase shown by Yucu-Matapi, Curripaco, Desano, Siriano, Inga, Pasto, Mur-Uitoto,

412 Tikuna, and Cocama (exemplified by Yucu-Matapi in Figure S3A); second, population stability through
413 time shown by Ach-Piapoco, Tanimuka, Coreguaje, Siona, Kamentsa, Puinave, and Yagua (exemplified
414 by Coreguaje in Figure S3B); and third, population contraction shown by Sikuani, Guayabero, and Nukak
415 (exemplified by Nukak in Figure S3C), which is particularly striking for Sikuani (Supporting information
416 Figure S3D). These differences in the effective population size through time suggest that these
417 populations have followed independent demographic histories.

418

419 **DISCUSSION**

420 We have investigated the genetic diversity of ethnolinguistic groups from NWA at the level of complete
421 mitochondrial genomes. This area is underrepresented in previous studies, and our data contribute to fill a
422 gap in our knowledge about the genetic diversity of modern human populations. We have found that
423 NWA harbors a considerable amount of genetic diversity, with evidence for contact among different
424 ethnolinguistic groups, contrary to the common picture of Amazonian populations as small and isolated
425 with low genetic diversity (Fuselli et al. 2003; Wang et al. 2007). NWA populations show values of
426 nucleotide diversity as high as or higher than those observed in most other non-African populations
427 (Supporting information Figure S10), and they display the second-highest amount of sequence sharing in
428 a world-wide comparison (Table 3). The complete mitochondrial genome is the maximum level of
429 resolution one can achieve to differentiate individuals and populations at the maternal level, so the
430 presence of identical sequences among populations living in distant geographic areas indicates recent
431 contact and/or common ancestry.

432

433 **Lack of genetic structure along linguistic lines**

434 Although our dataset includes populations speaking languages belonging to different language families,
435 we found that linguistic affiliation is a poor predictor of genetic structure, as shown by the AMOVA

436 analysis (Table 4) and by the Correspondence Analysis based on sub-haplogroup frequencies (Figure 2).
437 This indicates that language does not constitute a barrier to gene flow, and that groups have been
438 interacting with other neighboring groups, especially along rivers, which in our analyses performed better
439 in explaining the patterns of genetic diversity. Archaeological and linguistic evidence demonstrates that
440 NWA has been an area of intense contact and movement of peoples of different cultural traditions,
441 evidenced by the diffusion of ceramic styles (Heckenberger 2002; Lathrap 1970; Zucchi 2002) and shared
442 subsistence strategies, by the existence of language areas and contact-induced linguistic change
443 (Aikhenvald 1999), and the generalized multilingualism among groups (Sorensen 1967; Stenzel 2005).
444 Likewise, cultural anthropology provides additional evidence of contact among groups. For example, both
445 Arawakans and Eastern Tukanoans share a ceremonial complex for male initiation known as Yurupari, in
446 which sacred flutes and trumpets are only played by males, as well as sharing myths concerning the hero
447 Kúwai (Hugh-Jones 1979; Jackson 1983; Vidal 2002). In addition, the Eastern Tukanoan groups from the
448 Pira-Parana and Apaporis rivers (Barasano, Makuna, and Tanimuka) reveal Arawakan influence, since
449 they also practice dances with masks during the season of high abundance of the palm tree fruit pupunha
450 (*Bactris gasipaes*) (Hugh-Jones 1979). The genetic distances among populations provide additional
451 evidence in this regard: although the global Φ_{ST} value of 0.11 indicates moderate differentiation (Hartl
452 and Clark 2007), this value is driven by three populations, namely the Siona, Sikuani, and Nukak. These
453 are highly differentiated from the other populations, likely reflecting the effects of genetic drift due to
454 bottlenecks, as indicated by the positive Tajima's D values (Figure 3) and the distribution of pairwise
455 differences (Supporting information Figure S2). When we exclude these populations, we observe an
456 average pairwise Φ_{ST} of 0.07, and populations appear close together in the MDS plot (Figure 5),
457 indicating low genetic differentiation among NWA populations.

458 In this general picture the Eastern Tukanoan groups stand apart, since they cluster together in the CA and
459 MDS plots (Figure 2 and Figure 5), and their pairwise genetic distances are small and non-significant
460 (Figure S6). Linguists have proposed a time depth for the Tukanoan family of 2000-2500 years, based on

461 a comparison of the diversity in Tukanoan languages with the diversity in Romance and Germanic
462 languages (Chacon 2014). The time depth of the Eastern Tukanoan branch (and thus the time to the most
463 recent common ancestor of the Eastern Tukanoan languages) would be even more recent, which might
464 indicate that the peoples speaking these languages share recent common genetic ancestry as well (at least
465 on the maternal side). However, the Eastern Tukanoan groups practice linguistic exogamy, and the close
466 genetic relationships among these populations might be the result of this marital system in which women
467 move among different ethnolinguistic groups. The consequences of the linguistic exogamy are also
468 evident in the gene diversity values and the patterns of shared haplotypes. Eastern Tukanoans are the
469 groups with the highest values of gene diversity, and they share more haplotypes among themselves than
470 with other non-Eastern Tukanoan groups. In addition, their haplotypes tend to be closely related, as can
471 be seen in the phylogenetic networks (Supporting information Figure S5). Analyses of the Y-chromosome
472 as well as nuclear markers will help to disentangle the effects of linguistic exogamy vs. recent common
473 ancestry on the patterns of genetic variation among Eastern Tukanoan groups.

474 The Tanimuka stand apart from the other Eastern Tukanoan groups in the analyses, which may reflect
475 their settlement further south, along the Apaporis and Mirití-Paraná rivers. Moreover, they do not
476 participate in the exogamic system with other Eastern Tukanoan groups, but interact mainly with the
477 Arawakan groups Yucuna and Matapi. This is reflected in the patterns of haplotype sharing (Figure 4) as
478 well as in their language, which shows evidence of Arawakan influence (Barnes 1999; Chacon 2014).

479

480 **The role of rivers in structuring genetic variation**

481 Besides language, geography is another important factor in structuring the patterns of genetic variation in
482 human populations (Ramachandran et al. 2005; Schonberg et al. 2011; Wang et al. 2007). One of the most
483 salient characteristics of the physical landscape of NWA is the high density of rivers that drain the area,
484 and their importance for human populations was earlier recognized by explorers and ethnographers that

485 traveled through the region (Koch-Grünberg 1995; Wallace 1853). We found that the distribution along
486 rivers is an additional important factor influencing the genetic structure of NWA populations: our
487 AMOVA analyses (Table 4) show that clustering populations according to the rivers where they are
488 distributed explains more of the genetic variation that is due to differences among groups than does
489 grouping them by linguistic affiliation, i.e. populations living on the same river basin or in closely
490 connected rivers are genetically more similar than those living on different rivers. This pattern is also
491 observed in the distribution of sub-haplogroups among populations (Figure S1). For example, Curripaco
492 and Puinave, who live on the Inírida and Atabapo rivers, are located close together in the plot, and the
493 presence of Coreguaje, Yucu-Matapi, and Mur-Uitoto in the center of the plot could reflect their presence
494 in a region where the Putumayo and Caquetá rivers are separated by their shortest distance, therefore
495 facilitating contact among people inhabiting the basins and tributaries of these two rivers. Indeed, one
496 Murui individual was sampled in a Coreguaje community, and two Uitoto individuals were sampled in
497 the Mirití-Parana region, thus providing evidence for the movement of people among these groups. The
498 results of the MRM analysis provide additional evidence in this regard: even though no correlation
499 between genetic distances and geographic distances was observed via the Mantel test, we observed that
500 adding river distances as an additional predictor variable resulted in an increase of around 3% of the R-
501 square value (Table 5), indicating that rivers contribute to explaining a slightly higher percentage of the
502 variation observed in the genetic distances.

503 Rivers in Amazonia serve a double function in providing a means of communication as well as
504 subsistence, and the wide distribution of certain cultural traits (e.g., the production of Saladoid-
505 Barranoid ceramics and circular plaza village settlement patterns), has been associated with the
506 expansion of Arawakan-speaking populations along the extensive system of NWA waterways
507 (Heckenberger 2002; Hornborg 2005; Lathrap 1970; Lowie 1948). They also mark a distinction in
508 subsistence strategies between the more numerous “river people” that build canoes, settle along rivers,
509 and rely on horticulture and fishing, and the “forest people” that inhabit the interfluvial areas, settle away

510 from the major rivers, and base their subsistence on foraging (Epps and Stenzel 2013). Additionally, the
511 rivers have profound meanings and are embedded in the cosmogonies of several NWA indigenous
512 groups. The Eastern Tukanoan creation myths describe the journeys that the ancestors of the people made
513 to settle this world on board an anaconda canoe that travelled up the Vaupes River; from the anaconda's
514 body all the Tukanoan siblings emerged (Chernela 2010; Jackson 1983). Arawakan groups also describe a
515 series of ever returning voyages from the sacred center of the world and the place of emergence of the
516 first ancestors at the rapids of Hípana on the Aiary River, covering the major arteries of the Rio Negro,
517 Orinoco, and Amazon (Wright 2002; Zucchi 2002). Therefore, our findings about the role of rivers in
518 structuring the genetic variation are in keeping with the pivotal role that rivers play for NWA populations.

519 The lack of fit between genetic and simple geographic distances may be the result of relatively recent
520 movements and the displacement of ethnolinguistic groups from their traditional territories. Population
521 dynamics and population sizes were drastically altered during the last five centuries, starting with early
522 colonial times (16th and 17th centuries), when many groups were decimated by newly introduced
523 epidemics and moved away from the accessible margins of the major rivers to avoid the slave raids of the
524 Spanish, Portuguese, and Dutch colonizers. Similar perturbations happened during the time of the
525 Christian missions in the 18th century, when many groups were forced to relocate to multiethnic mission
526 settlements, and finally during the rubber boom between the 19th and beginning of the 20th centuries,
527 when the groups who managed to escape the mercenaries exploiting the rubber fields resettled in remote
528 areas in the headwaters of small rivers (Dixon and Aikhenvald 1999; Hill and Santos-Granero 2002;
529 Stenzel 2005). The inferred reduction in population size of the Tanimuka, Sikuani, Guayabero, and
530 Nukak, as indicated by their low diversity values, the positive Tajima's D values (Figure 3), the
531 distribution of pairwise differences (Figure S2), and the reconstruction of effective population sizes
532 (Figure S3C, D), might be a result of these social upheavals.

533

534

535 **The impact of subsistence strategies on the genetic diversity**

536 NWA contains groups with different subsistence strategies, with manioc (*Manihot esculenta*) as the main
537 staple among horticulturalist groups, who are best described as riverine horticultural societies, given their
538 close association with rivers. The Nukak, in contrast, are traditionally foragers, who still rely on hunting
539 and gathering and move throughout the extensive area between the Guaviare and Inírida rivers.
540 Furthermore, the Guayabero, Sikuani, and Puinave are traditional foragers who have only recently
541 undergone the transition to agriculture, and are therefore considered as HGP together with Nukak in our
542 analyses (Table 1). Our data show that agricultural societies (AG) have higher levels of diversity on
543 average than forager groups (HGP) as indicated by the Mann-Whitney U test (P-value = 0.03), while the
544 HGP groups have larger values of Tajima's D statistic (Figure 3) and do not show signals of population
545 expansion (Figure S3). These findings agree with patterns reported for other hunter-gatherer populations
546 around the world (Aime et al. 2013; Excoffier and Schneider 1999; Oota et al. 2005) and contrast with the
547 genetic signature of an agricultural way of life, namely higher effective population size (Patin et al. 2014),
548 higher levels of diversity, and significantly negative values of Tajima's D test (Aime et al. 2013).

549 However, subsistence strategies are flexible and diverse among NWA populations. Horticulturalists
550 complement their diet with occasional hunting and/or gathering of several kinds of palm fruit, and
551 extensive exchanges between AG and HGP groups have been reported. In this system, HGP populations
552 usually provide meat and several products from the forest, such as the poison curare for the tips of darts
553 and arrows, in exchange for different cultivated products, such as manioc and other trade goods (Epps and
554 Stenzel 2013; Jackson 1983; Milton 1984). Nonetheless, this exchange seems to be exclusively restricted
555 to goods and labor, with little or no intermarriage documented between AG and HGP groups (Aikhenvald
556 1996). In contrast, we observed shared haplotypes between AG and HGP groups, which likely reflects
557 intermarriage or recent common ancestry. For example, the most frequent haplotype in the Arawakan AG
558 group Curripaco (Haplotype H_84 in Figure S4) is observed at high frequency in the HGP Nukak (and in
559 the Eastern Tukanoan AG group Siriano). Moreover, the HGP Puinave share several haplotypes with the

560 AG group Curripaco (H_219, H_161, H_117 in Figure S4), a likely result of intermarriage between these
561 groups, since there are communities on the Inírida River where one finds individuals from both groups.
562 Similarly, the Guahiban HGP groups Sikuni and Guayabero exhibit a haplotype at high frequency (H_43
563 in Figure S4) that is shared with the AG Ach-Piapoco as well as further haplotypes related to haplotypes
564 found in AG Arawakan groups (cluster I Figure S5C and cluster II S5B). This may reflect contact among
565 them, since there are Piapoco communities on the lower Guaviare River as well as Sikuni communities
566 on the Meta River, places where these groups overlap. However, it is difficult to determine the direction
567 of the gene flow or to distinguish between contact and common ancestry as explanations for shared
568 mtDNA haplotypes. Nevertheless, it is plausible that where haplotypes are shared the source population is
569 the one in which the haplotype is present at higher frequency. For instance, the shared haplotype between
570 the HGP Puinave and the AG Curripaco (H_219 in Figure S4) has a likely origin in Puinave, because of
571 its higher frequency and the presence of related haplotypes in Puinave (cluster I Figure S5B). The source
572 of the shared haplotype among the HGP Nukak and the AG Curripaco and Siriano (H_84 in Figure S4) is
573 more difficult to infer, since its frequency is similar in the Nukak and in the Curripaco; furthermore, three
574 other haplotypes present in the HGP Nukak and Guayabero are only one mutation apart from it (cluster II
575 Figure S5B). Therefore, it is likely that this haplotype, too, moved from the HGP populations into the AG
576 Curripaco. A similar explanation could be given for H_43 in Figure S4, which is part of the cluster I in
577 Figure S5C, moving from the HGP Guayabero and Sikuni into the AG Ach-Piapoco. Thus, these
578 observations seem to fit a scenario of asymmetric gene flow in which women move from HGP to AG, a
579 pattern that has been reported for populations in Central and Southern Africa (Barbieri et al. 2014;
580 Destro-Bisol et al. 2004; Verdu et al. 2013). However, this scenario will be further refined by analyses of
581 Y-chromosome and genome-wide data, which will allow us to determine whether the gene flow among
582 groups was sex-biased (i.e. involving the movement of only females or only males among groups) and to
583 make inferences about the time and magnitude of these events.

584 In conclusion, this study provides new data from this remote and little-studied part of the world, which
585 allow insights into the impact of cultural practices on the patterns of genetic variation and on the
586 population dynamics of NWA groups. Although our current data do not allow us to distinguish whether
587 the population movements took place prior to European contact or only later, analyses of Y-chromosome
588 variation and genome-wide data will shed further light on the genetic history of NWA. Furthermore,
589 historical genetic studies will benefit from more archaeological work in NWA, since huge areas remain
590 completely unexplored.

591

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606

607

608 **LITERATURE CITED**

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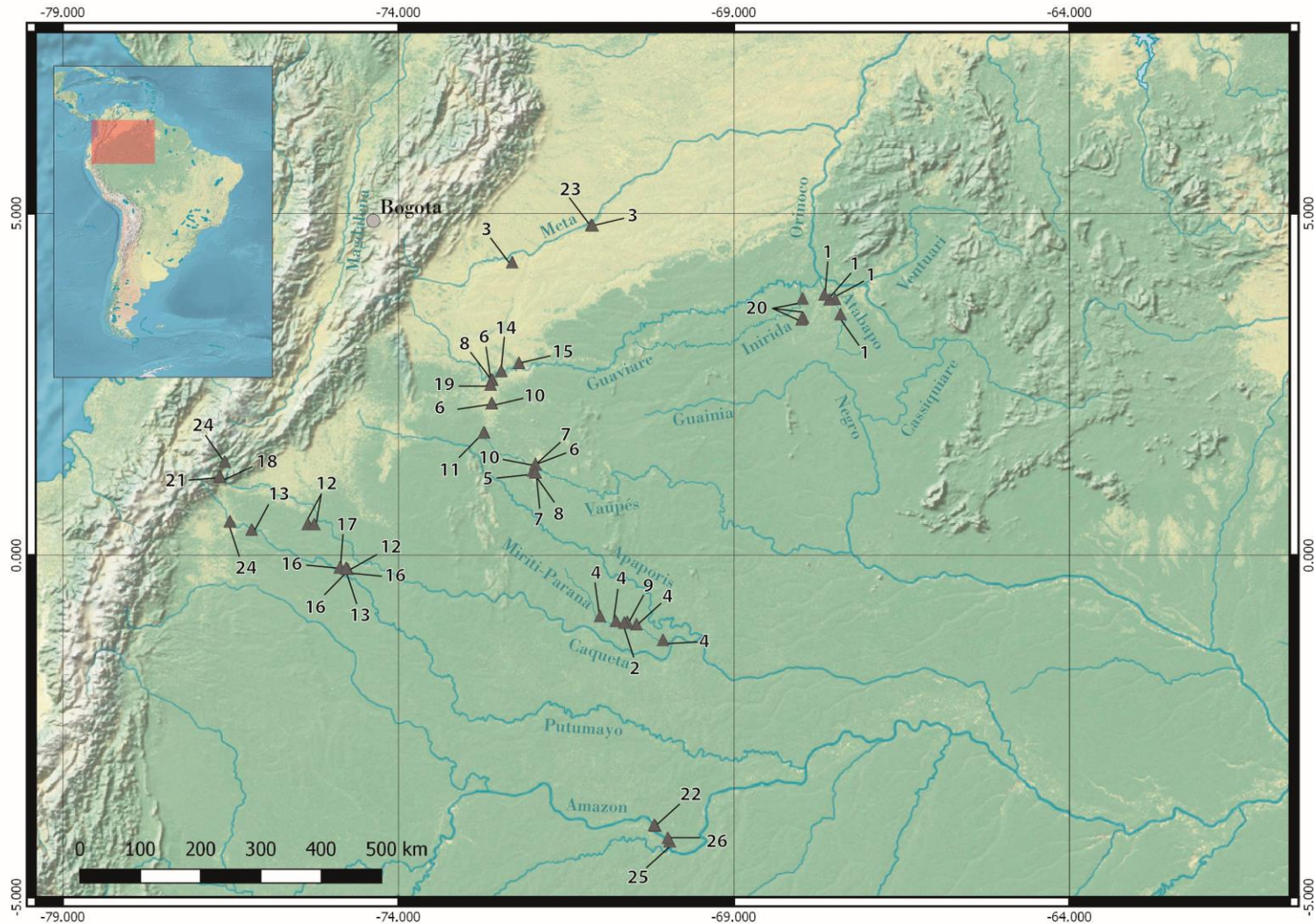
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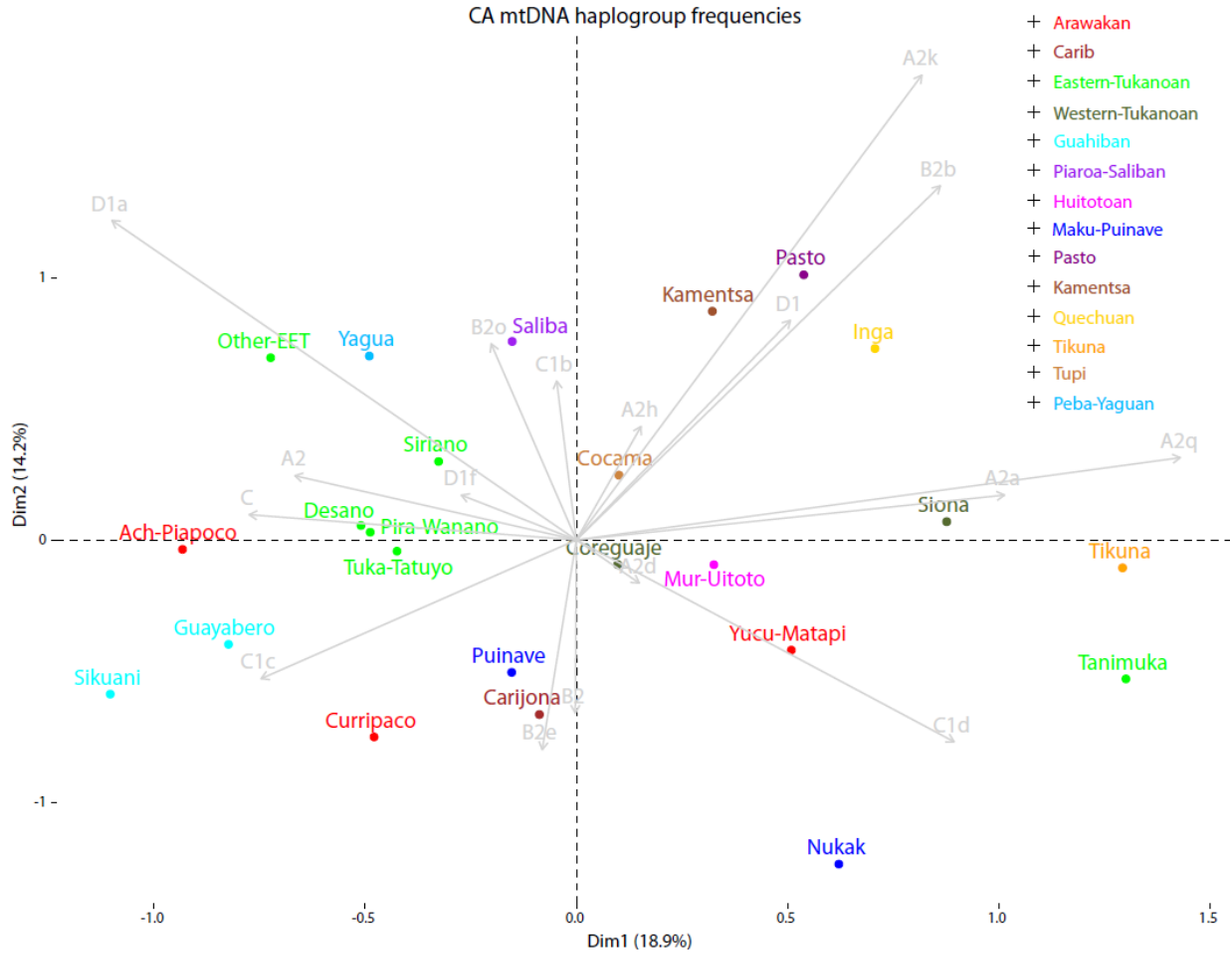
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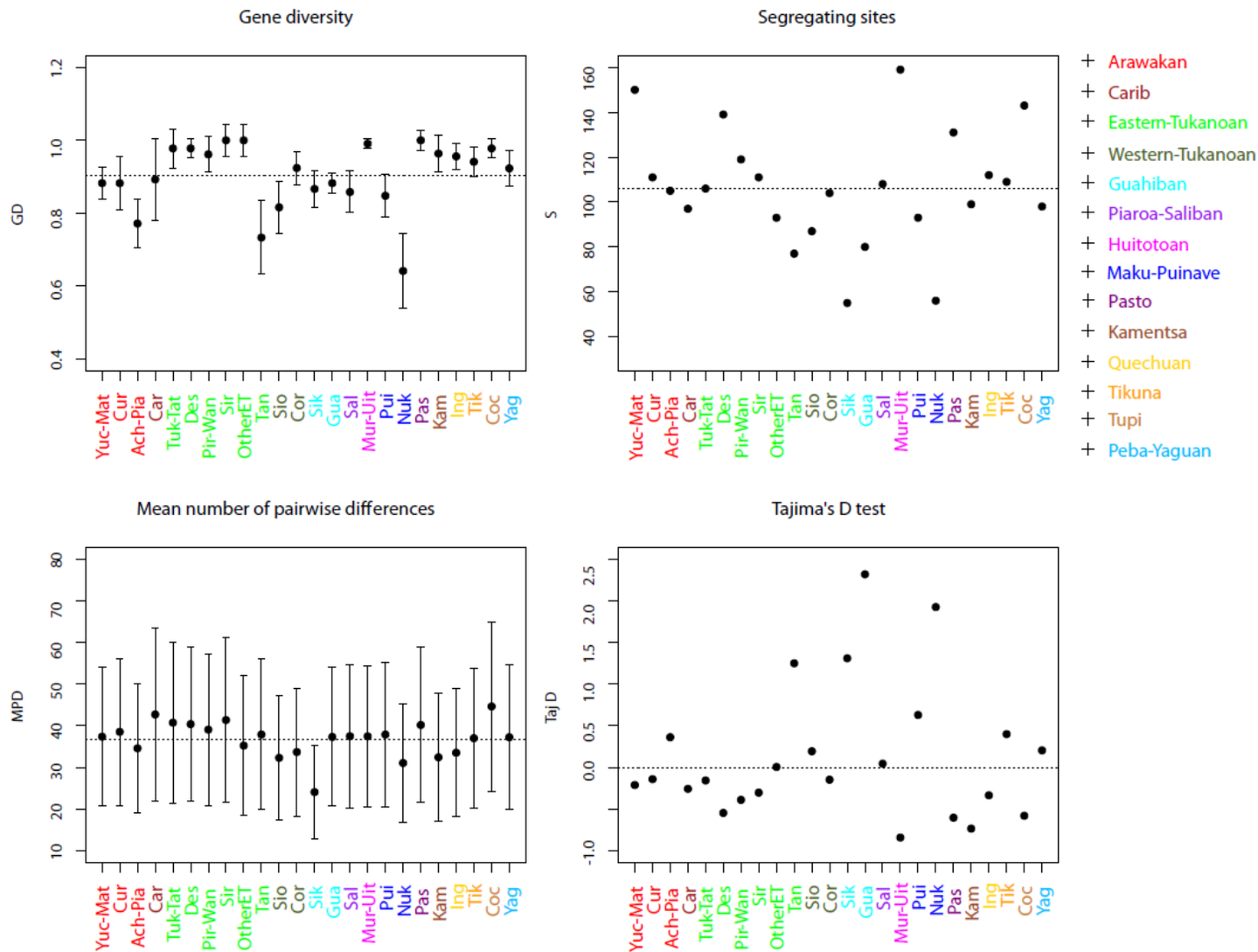
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885 **Figure 1.** Geographic location of the sampling sites. Every triangle corresponds to a single community, which may contain more than one
 886 ethnolinguistic group. 1. Curripaco and Bare, 2. Matapi, 3. Ach-Piapoco, 4. Yucuna, 5. Carijona, 6. Desano, Yuruti, Pisamira, and
 887 Karapana, 7. Pira-Wanano, 8. Siriano, 9. Tanimuka, 10. Tukano, 11. Tuyuca and Tatuyo, 12. Coreguaje, 13. Siona, 14. Guayabero, 15.
 888 Sikuani, 16. Murui, 17. Uitoto, 18. Kamentsa, 19. Nukak, 20. Puinave, 21. Pasto, 22. Yagua, 23. Saliba, 24. Inga, 25. Tikuna, 26. Cocama.



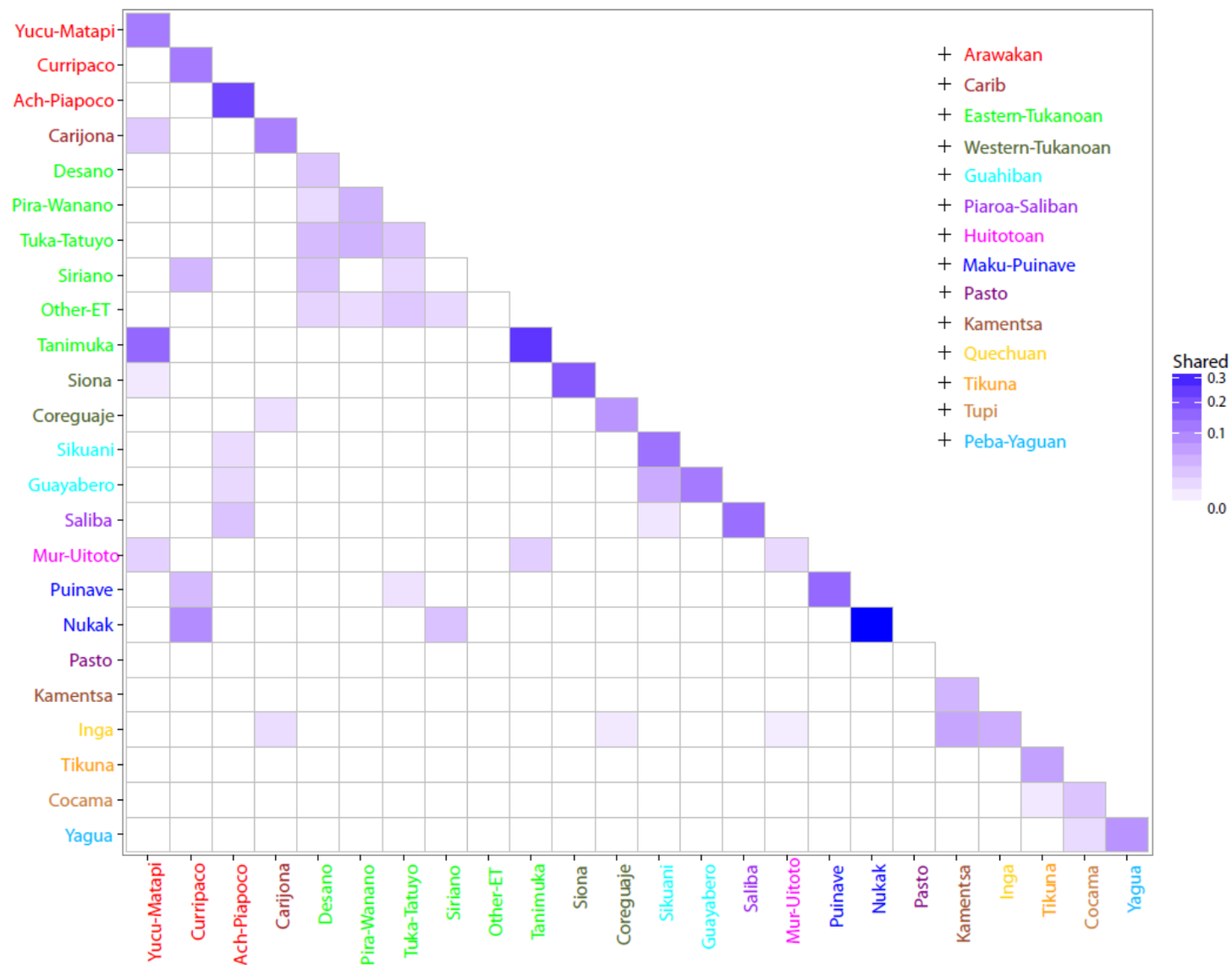
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890 **Figure 2.** Correspondence analysis based on the sub-haplogroup frequencies by population. Populations
 891 are color-coded by linguistic affiliation.



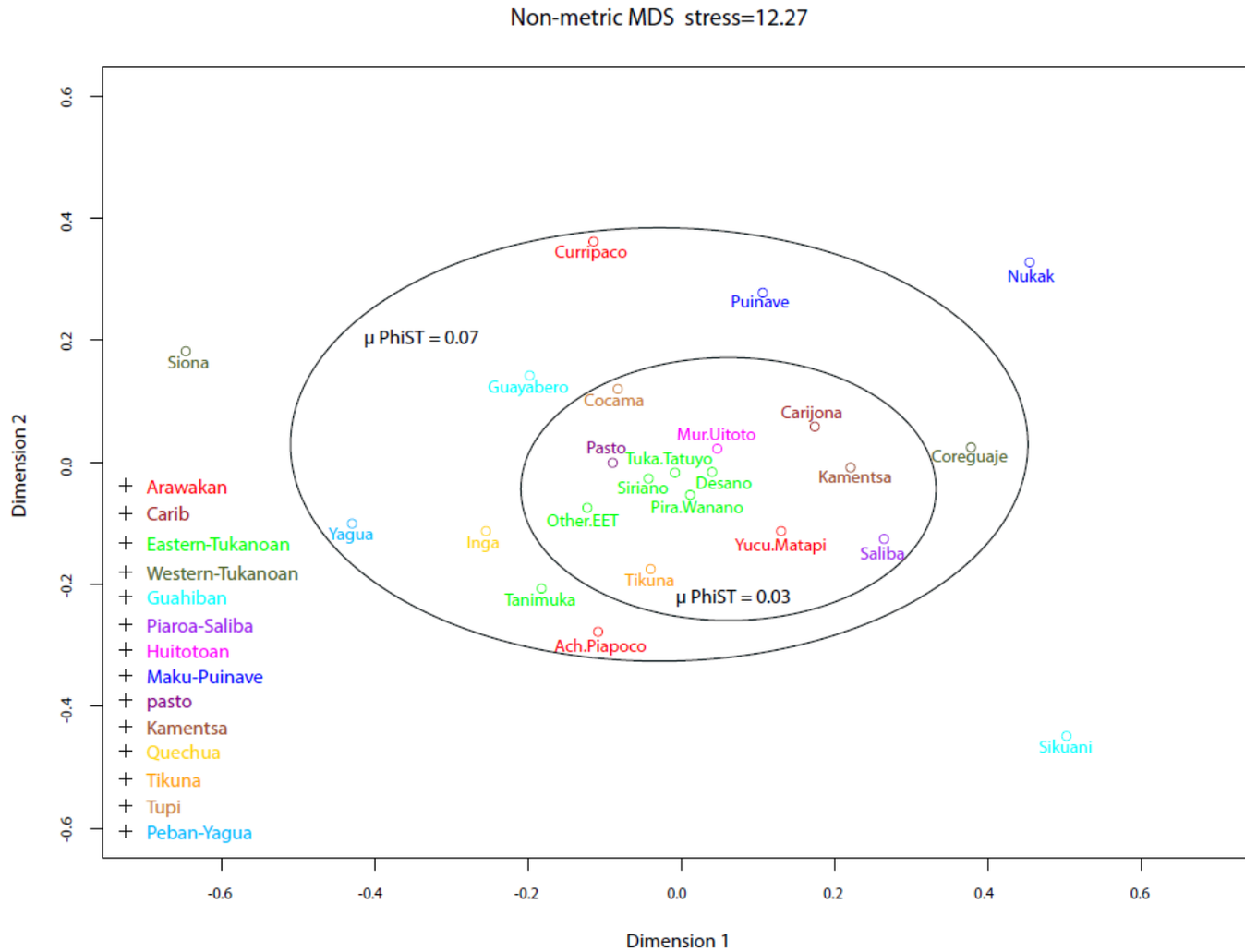
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893 **Figure 3.** Molecular diversity indices by population. Dashed lines correspond to average values, except for Tajima's D test which corresponds to
 894 zero. Populations are color-coded by linguistic affiliation as in Figure 2.



895

896 **Figure 4.** Matrix of shared haplotypes between populations. The color scale indicates the proportion of the total haplotypes that are shared within
 897 (on the diagonal) or between (below the diagonal) populations.



898

899 **Figure 5.** Multidimensional Scaling plot based on Φ_{ST} genetic distances. Stress value is given in percentage. The inner circle indicates populations
 900 with low genetic differentiation and the outer circle indicates populations with moderate differentiation. μ PhiST is the average pairwise Φ_{ST} value
 901 within each circle.

Table 1. Sampled ethnolinguistic groups with information on merged groups (see Material & Methods: Data Analysis) given below the compound names.

Population	Label in figure 1	N	Census Sizea	Language family	Subsistence Strategyb	River/place of residence
Yucu-Matapi		39		Arawakan	AG	Mirití-Paraná
Yucuna	4	31	550	Arawakan	AG	Mirití-Paraná
Matapi	2	8	220	Arawakan	AG	Mirití-Paraná
Curripaco		17		Arawakan	AG	Atabapo
Curripaco	1	16	7827	Arawakan	AG	Atabapo
Bare	1	1	NAC	Arawakan	AG	Atabapo
Ach-Piapoco		24		Arawakan	AG	Meta
Achagua	3	6	283	Arawakan	AG	Meta
Piapoco	3	18	4926	Arawakan	AG	Meta
<i>Cabiyarid</i>		1	311	Arawakan	AG	Mirití-Paraná
Carijona	5	8	307	Carib	AG	Upper-Vaupés
<i>Cofan</i>		6	877	Cofan	AG	Guamúez
<i>Barasano</i>		4	2008	Eastern Tukanoan	AG	Upper-Vaupés
Desano	6	17	2457	Eastern Tukanoan	AG	Upper-Vaupés
<i>Kubeo</i>		5	6647	Eastern Tukanoan	AG	Upper-Vaupés
Other-EET		10		Eastern Tukanoan	AG	Upper-Vaupés
Tuyuca	11	7	642	Eastern Tukanoan	AG	Upper-Vaupés
Yuruti	6	1	687	Eastern Tukanoan	AG	Upper-Vaupés
Pisamira	6	1	61	Eastern Tukanoan	AG	Upper-Vaupés
Karapana	6	1	464	Eastern Tukanoan	AG	Upper-Vaupés
Pira-Wanano		13		Eastern Tukanoan	AG	Upper-Vaupés
Piratapuyo	7	8	697	Eastern Tukanoan	AG	Upper-Vaupés
Wanano	7	5	1395	Eastern Tukanoan	AG	Upper-Vaupés
Siriano	8	10	749	Eastern Tukanoan	AG	Upper-Vaupés
Tanimuka	9	10	1247	Eastern Tukanoan	AG	Mirití-Paraná
Tuka-Tatuyo		10		Eastern Tukanoan	AG	Upper-Vaupés
Tukano	10	8	6996	Eastern Tukanoan	AG	Upper-Vaupés
Tatuyo	11	2	331	Eastern Tukanoan	AG	Upper-Vaupés
Siona	13	17	734	Western Tukanoan	AG	Putumayo
Coreguaje	12	19	2212	Western Tukanoan	AG	Caquetá
Sikuani	15	16	23006	Guahiban	HGP	Guaviare
Guayabero	14	35	1118	Guahiban	HGP	Guaviare
Saliba	23	16	1929	Piaroa-Saliban	AG	Meta
Mur-Uitotoe		26	7343	Huitotoan	AG	Putumayo
Murui	16	18		Huitotoan	AG	Putumayo
Uitoto	17	8		Huitotoan	AG	Putumayo
Puinave	20	19	6604	Maku-Puinave	HGP	Inirida
Nukak	19	16	1483	Maku-Puinave	HGP	Interfluvial
Pasto	21	14	69789	Pasto	AG	Andean

Kamentsa	18	11	4773	Kamentsa	AG	Andean
Inga	24	17	19079	Quechuan	AG	Andean
Tikuna	25	18	7102	Tikuna	AG	Amazonas
Cocama	26	17	792	Tupi	AG	Amazonas
Yagua	22	13	297	Peba-Yaguan	AG	Amazonas
<i>Guambiano</i>		1	23462	Barbacoan	AG	Andean
<i>Nasa</i>		1	138501	Nasa	AG	Andean
<i>Mestizo</i>		9	NA	Mestizo	NA	NA
Total		439				

a. Data from: (Arango and Sánchez 2004). b. AG: agriculturalist; HGP: Hunter-gatherer populations, data from D-PLACE (Kirby et al 2016).and HG (<https://huntergatherer.la.utexas.edu/home>, accessed on 06.06.2017). d. Populations with label in italics were not considered in the population-based analyses. e. Census data reports the population size including groups that speak five dialectal varieties.

Table 2. Frequency of haplogroups for the 24 NWA ethnolinguistic groups included in the population analyses

Population	N	A2	B2	C1	D1
Yucu-Matapi	39	0.28	0.10	0.56	0.05
Curripaco	17	0.18	0.53	0.24	0.06
Ach-Piapoco	24	0.54	0.04	0.42	0.00
Carijona	8	0.13	0.25	0.63	0.00
Desano	17	0.29	0.12	0.41	0.18
Other-ET	10	0.50	0.00	0.30	0.20
Pira-Wanano	13	0.31	0.08	0.38	0.23
Siriano	10	0.40	0.10	0.40	0.10
Tanimuka	10	0.50	0.10	0.40	0.00
Tuka-Tatuyo	10	0.30	0.10	0.30	0.30
Siona	17	0.59	0.35	0.00	0.06
Coreguaje	19	0.11	0.16	0.74	0.00
Sikuani	16	0.25	0.00	0.75	0.00
Guayabero	35	0.43	0.23	0.34	0.00
Saliba	16	0.19	0.06	0.56	0.19
Mur-Uitoto	26	0.23	0.12	0.31	0.35
Puinave	19	0.11	0.42	0.47	0.00
Nukak	16	0.00	0.31	0.69	0.00
Pasto	14	0.36	0.14	0.21	0.29
Kamentsa	11	0.18	0.09	0.64	0.09
Inga	17	0.59	0.06	0.29	0.06
Tikuna	18	0.44	0.00	0.44	0.11
Cocama	17	0.29	0.35	0.29	0.06
Yagua	13	0.62	0.15	0.23	0.00

Table 3. Shared haplotypes in a worldwide sample of complete mitochondrial sequences sampled at the population level

Geographic region	#Sequences	#Haplotypes	%Unique haplotypes	Shared Within population	Shared Between populations	Source
NW Amazonia	412	216	0.676	0.241	0.144	Present study
Burkina Faso	335	332	0.991	0.006	0.003	(Barbieri et al. 2012)
SW Zambia	169	146	0.897	0.048	0.055	(Barbieri et al. 2013)
Botswana/Namibia	218	128	0.75	0.188	0.133	(Barbieri et al. 2014)
Philippines	365	233	0.734	0.227	0.077	(Delfin et al. 2014)
Sumatra	72	48	0.771	0.229	0.021	(Gunnarsdottir et al. 2011)
Taiwan	549	299	0.669	0.308	0.084	(Ko et al. 2014)
Oceania	1331	650	0.689	0.277	0.106	(Duggan et al. 2014)
Siberia	525	244	0.574	0.336	0.217	(Duggan et al. 2013)
Mexico ^a	113	90	0.867	0.133	0	(Mizuno et al. 2014)

Note: The proportions do not sum up to 1 since some haplotypes are shared both within and between populations. a. The individuals from Mexico are all of Amerindian origin.

Table 4. Analysis of molecular variance (AMOVA)

	# groups	Among groups	Within groups	Within populations	Global FST
One group	1		11.12**	88.88	0.1112
Language ^a	14	-1.33	12.37**	88.96**	0.1104
Geography ^b	6	1.04	10.24**	88.72**	0.1128
Rivers ^c	11	5.42**	5.99**	88.59**	0.1141

** Significant at 0.01 level

^a. 1.Arawak: Yucu-Matapi, Curripaco, Ach-Piapoco; 2.Carib: Carijona; 3.Eastern-Tukanoan: Desano, Pira-Wanano, Siriano, Tuka-Tatuyo, Other-ET, Tanimuka; 4.Western-Tukanoan: Coreguaje, Siona; 5.Guahiban: Sikuani, Guayabero; 6.Huitoto: Mur-Uitoto; 7.Maku-Puinave: Puinave, Nukak; 8.Kamentsa; 9.Pasto; 10.Piaroa-Saliba: Saliba; 11.Peba-Yaguan: Yagua; 12.Quechua: Inga; 13.Tikuna; 14.Tupi: Cocama.

^b. 1. Saliba, Ach-Piapoco; 2. Sikuani, Guayabero, Nukak, Desano, Pira-Wanano, Siriano, Tuka-Tatuyo, Other-ET, Carijona; 3. Coreguaje, Siona, Mur-Uitoto, Inga, Kamentsa, Pasto; 4. Curripaco, Puinave; 5. Yucu-Matapi, Tanimuka; 6. Cocama, Tikuna, Yagua.

^c. 1.Meta: Saliba, Ach-Piapoco; 2.Vaupés: Desano, Pira-Wanano, Siriano, Tuka-Tatuyo, Other-ET, Carijona; 3.Guaviare: Guayabero, Sikuani; 4.Interfluve: Nukak; 5.Atabapo-Inirida: Curripaco, Puinave; 6.High-Putumayo: Inga, Kamentsa, Pasto; 7.Middle-Putumayo: Siona; 8. Lower-Putumayo: Mur-Uitoto; 9. Middle-Caqueta: Coreguaje; 10.Mirití-Parana: Yucu-Matapi, Tanimuka; 11.Amazon: Cocama, Tikuna, Yagua.

Table 5. Multiple regression analysis on distance matrices

		gen.dist all populations			
		Reg.coefficient	P-val	R.square	P-val
Simple regression	geo.dist	3.28×10^{-8}	0.444	0.009	0.444
Multiple regression	geo.dist	9.64×10^{-10}	0.983	0.036	0.126
	rivers.dist	5.71×10^{-2}	0.011		
		gen.dist without outliers			
Simple regression	geo.dist.no.outliers	6.70×10^{-8}	0.026	0.067	0.026
Multiple regression	geo.dist.no.outliers	5.76×10^{-8}	0.112	0.070	0.040
	rivers.dist.no.outliers	1.38×10^{-2}	0.458		

Supporting information Table S1. Molecular diversity indices by population

Population	N	No.hap	S	GD	GD.SD	ND	ND.SD	MPD	MPD.SD	TajD	Pval
Yucu-Matapi	39	19	150	0.88	0.045	0.0023	0.0011	37.38	16.61	-0.211	0.440
Curripaco	17	11	111	0.88	0.072	0.0023	0.0012	38.54	17.64	-0.141	0.456
Ach-Piapoco	24	8	105	0.77	0.065	0.0021	0.0011	34.60	15.62	0.363	0.719
Carijona	8	6	97	0.89	0.111	0.0026	0.0014	42.71	20.80	-0.257	0.405
Tuka-Tatuyo	10	9	106	0.98	0.054	0.0025	0.0013	40.76	19.37	-0.156	0.467
Desano	17	14	139	0.98	0.027	0.0024	0.0012	40.40	18.47	-0.546	0.322
Pira-Wanano	13	11	119	0.96	0.050	0.0024	0.0012	39.08	18.18	-0.389	0.368
Siriano	10	10	111	1.00	0.045	0.0025	0.0013	41.38	19.66	-0.303	0.388
Other.ET	10	10	93	1.00	0.045	0.0021	0.0011	35.24	16.80	0.006	0.533
Tanimuka	10	4	77	0.73	0.101	0.0023	0.0012	37.91	18.04	1.250	0.928
Siona	17	8	87	0.82	0.071	0.0020	0.0010	32.31	14.84	0.194	0.613
Coreguaje	19	12	104	0.92	0.046	0.0020	0.0010	33.70	15.37	-0.147	0.508
Sikuani	16	7	55	0.87	0.050	0.0015	0.0008	24.08	11.18	1.310	0.941
Guayabero	35	11	80	0.88	0.028	0.0023	0.0011	37.31	16.62	2.318	0.992
Saliba	16	7	108	0.86	0.057	0.0023	0.0012	37.49	17.22	0.046	0.577
Mur-Uitoto	26	23	159	0.99	0.013	0.0023	0.0011	37.46	16.84	-0.841	0.207
Puinave	19	9	93	0.85	0.059	0.0023	0.0012	37.89	17.25	0.630	0.788
Nukak	16	4	56	0.64	0.103	0.0019	0.0010	31.07	14.33	1.925	0.989
Pasto	14	14	131	1.00	0.027	0.0024	0.0013	40.19	18.59	-0.603	0.281
Kamentsa	11	9	99	0.96	0.051	0.0020	0.0010	32.44	15.35	-0.735	0.227
Inga	17	13	112	0.96	0.037	0.0020	0.0010	33.53	15.38	-0.334	0.427
Tikuna	18	13	109	0.94	0.042	0.0022	0.0011	37.02	16.90	0.399	0.722
Cocama	17	14	143	0.98	0.027	0.0027	0.0014	44.64	20.37	-0.580	0.274
Yagua	13	8	98	0.92	0.050	0.0023	0.0012	37.26	17.35	0.205	0.646

No.hap: Number of haplotypes; S: Segregating sites; GD: Gene diversity; GD.SD: GD standard deviation; ND: Nucleotide diversity; ND.SD: ND standard deviation; MPD: Mean number of pairwise differences; MPD.SD: MPD standard deviation; TajD: Tajima's D values; Pval: P values of Tajima's D.