Discrepancy between Cranial and DNA Data of Early Americans: Implications for American Peopling

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Abstract

Currently, one of the major debates about the American peopling focuses on the number of populations that originated the biological diversity found in the continent during the Holocene. The studies of craniofacial variation in American human remains dating from that period have shown morphological differences between the earliest settlers of the continent and some of the later Amerindian populations. This led some investigators to suggest that these groups—known as Paleoelectronics and Amerindians respectively—may have arisen from two biologically different populations. On the other hand, most DNA studies performed over extant and ancient populations suggest a single migration of a population from Northeast Asia. Comparing craniofacial and mtDNA data of diachronic samples from East Central Argentina dated from 8,000 to 400 years BP, we show here that even when the oldest individuals display traits attributable to Paleoamerican crania, they present the same mtDNA haplogroups as later populations with Amerindian morphology. A possible explanation for these results could be that the craniofacial differentiation was a local phenomenon resulting from random (i.e. genetic drift) and non-random factors (e.g. selection and plasticity). Local processes of morphological differentiation in America are a probable scenario if we take into consideration the rapid peopling and the great ecological diversity of this continent; nevertheless we will discuss alternative explanations as well.

Introduction

The biological diversity of South American human populations has been the focus of extensive research for more than a hundred years (see review in [1]). These investigations have been associated with intense interdisciplinary studies regarding the peopling of the Americas. The great interest in this subject is partially due to the fact that America was the latest continent colonized by modern humans (ca. 11,000–13,000 years B.P.; [2]) and also due to the high levels of morphological variation found in Native American populations. In this context, two main hypotheses have been proposed to account for this biological variation: a) the migratory hypothesis, which suggests that the biological variation among South American groups was the result of a variable number of migratory waves [3,4]; and b) the local diversification hypothesis, i.e. that all South American groups descend from the same ancestral population or from populations related to each other, with local random (i.e. genetic drift) and non-random factors (i.e. selection and phenotypic plasticity) as the main causes of the diversification [5–7].

In recent years, the migratory hypothesis that postulates different biological origins for South American populations has received increased attention by researchers working with craniofacial evidence [8–10]. This hypothesis, known as two main biological components, asserts that the morphological diversity of American human populations results from two successive migratory events. The first component, named Palaeoamericans, derived from Pleistocene Southeast Asian populations which expanded into America around 14,000 years BP. Morphologically they were characterized by long and narrow cranial vault (i.e. dolichocephalic morphology) and a narrow face. The second component, named Amerindians, from which most of modern American groups derive, corresponds to a migration of populations from Northeast Asia which occurred during the Early Holocene (ca. 8,000 years BP; [8–11]). These populations exhibited short and wide cranial vault, along with wide faces (i.e. brachycephalic morphology). In addition, it was pointed out that this Amerindian morphology corresponds with a mongoloid pattern of craniofacial shape. The presence of this cranial shape in America has been explained as the result of a “fixation” of the mongoloid morphology in North Asia, previous to the Amerindian migration.

In contrast, the molecular evidence available to date (i.e. mtDNA and nuclear DNA information) supports a single origin in Northeast Asia ca. 15,000 years BP for almost all American populations, followed by local diversification—probably with the exception of the Eskimo and Na-Dene groups [12–14]. Particularly, mtDNA studies have detected four major pan-American founding haplotypes (A2, B2, C1, D1), which are also frequent in Asia. In addition, other founding mtDNA haplotypes occur in the Americas, such as X2a, D2, and D3, which are found nearly exclusively in North America [12,13]. The haplogroup distribution, together with the similar coalescence time for these haplotypes, has been used to support a single origin for extant
American populations, as well as a swift pioneering process of the initial north to south migration [12]. In addition, coalescent analyses suggest an initial differentiation of the Northeast Asia populations, a bottleneck in Beringia ca. 20,000 years BP, ended with a population expansion in America ca. 15,000 years BP [12,13].

The discrepancies between craniometric and molecular data, as well as the hypotheses supported by each kind of evidence, could be related either to the properties of both types of data, which provide different types of genealogical information, or to differences between the samples studied in each case. Particularly, quantitative traits and mtDNA differ in their respective mechanisms of inheritance (uniparental in mtDNA and biparental in quantitative traits), rate of change and degree of environmental influence [15–17]. On the other hand, the molecular data have been mainly obtained from extant or recent populations, whereas craniofacial variation has been assessed using skeletal samples from Early and Late Holocene populations. Hence, researchers who proposed the hypothesis of two main biological components assert that if the Paleoamericans did not survive or if their contribution to the biological variation of modern American populations was very small [7–9], the variation found among Later Late Holocene groups would not be relevant to discuss the early peopling.

One way to approach this problem is by analyzing the cranial morphology of diachronic samples, ranging from Early to Late Holocene, for which ancient mtDNA data are also available. The few areas able to provide human remains dated as Early Holocene on the basis of 14C dates of human bones are [18]: East Central Brazil (Lagoa Santa, ca. 9,000–5,000 yr 14C BP; [10,19]), the Bogotá savannah, Colombia (Tequendama, ca. 7,500–5,800 yr 14C BP; [20]) and the East Central Argentina (Arroyo Seco 2, ca. 7,800–6,300 yr 14C BP; [21]). However, East Central Argentina is the only region with a diachronic sequence ranging from 8,000 to 200 years BP [21,22] for which both mtDNA and craniometric data are available. Even though this region holds important evidences, it has not yet been included in the discussion about the biological diversity of South American populations from a diachronic perspective. In this study we present the first analysis of a skeletal sample from East Central Argentina including both craniometric and molecular data. The goal of this work is to compare the pattern of temporal and spatial variation in both types of data and to discuss them in light of the current hypotheses about the peopling of America. The analysis of these data allows for a renewed approach to the problem of the biological diversity and peopling of this continent.

Materials and Methods

Samples

We studied the early site from East Central Argentina (i.e. Southeast of Pampa and Northeast of Patagonia, Fig. 1, Table 1) known as Arroyo Seco 2, dated between Late Pleistocene and Early/Middle Holocene—the human remains that were used here are dated on ca. 7,800-6,300 yr 14C BP [21,23]—plus four samples of human remains corresponding to Middle and Earlier Late Holocene and four samples corresponding to Late Late Holocene from the same region. In addition, seven Late Holocene samples from neighbour regions were also analyzed (Table 1). All these samples include adult individuals of both sexes from hunter-gatherer groups, with presence of pottery in the Later Late Holocene.

The Arroyo Seco 2 archaeological site presents exceptional evidence to study the early peopling of America [23]. This multi-component open-air site is dated from 12,500 14C yr BP to the XIX Century [24] and nowadays is located at about 50 km north from the Atlantic Coast in the Buenos Aires Province of Argentina (38°21’ lat S. and 60°14’ lon W). Arroyo Seco 2 has an early component containing a lithic assemblage of unifacial, marginally retouched tools associated with bone remains of guanaco (camelid), Pampian deer, and nine extinct megafauna: *Paleolama*, *Equus*, *Hippeion*, *Toxodon*, *Megatherium*, *Esuttus*, *Glossotherium*, *Macrauchenia*, and *Clyptodon* [23]. Apart from this early component, the site contains one of the best records of human remains for the Early/Middle Holocene transition in South America. To date, 45 human skeletons have been uncovered and there are 21 dates from ca. 7,800 to 4,500 14C yr BP related to them [23]. The span of dates from the primary and secondary burials of Arroyo Seco 2, suggests the use of the site—not continuously but redundantly—for inhumations purposes, for more than 3,000 years during the Early and Middle Holocene.

Middle and Earlier Late Holocene samples from East Central Argentina contain individuals of different sites from Laguna del Juncal archaeological locality (Laguna del Juncal, Rio Negro Valley 1 and 2; see Table 1), placed south from Viedma city in the Rio Negro Province of Argentina (40°48’ lat S. and 62°58’ lon W), and one sample from Southeast Pampa (Table 1). The samples from East Central Argentina dated on Later Late Holocene come from various archaeological sites from Rio Negro, placed near Laguna del Juncal and Peninsula San Blas (40°33’ lat S. and 62°13’ lon W), and the Buenos Aires Province (Table 1).

Specimens are housed at División Antropología of the Museo de La Plata, Museo Etnográfico ‘J. B. Ambrosetti’ in Buenos Aires and INCUAPA in Olavarria, Argentina.

Preliminary analyses

Because most samples are sex balanced, males and females were pooled in the analyses to obtain a greater sample size. In order to control some sources of variation related to sex, we analyzed size standardized adult individuals of both sexes. The observational error was controlled using the experimental design introduced by Perez [1]. The results showed that photographing and digitalization of landmarks and semilandmarks procedures did not generate significant observational error [1].

Morphometric analyses

The craniofacial variation was analyzed with geometric morphometrics techniques [25,26] employing an arrangement of two-dimensional coordinates of biologically definable landmarks and semilandmarks (Fig. 2). Most comparisons were done on the facial skeleton, which is not affected by the cranial deformation present in these samples. We also performed an analysis of vault morphology in non-deformed skulls. Specimens were photographed with an Olympus SP 350 digital camera with the skull positioned according to the Frankfurt plane. For facial images, the camera lens was located in the coronal plane [27] and digital images were obtained from the crania in frontal view. Facial images were taken at 250 mm from the prosthion point. Eight landmarks and seventy-four semilandmarks (Fig. 2A) were obtained from the facial skeleton. For vault skeleton, digital images were obtained from the crania in lateral (left side) view. Lateral view images were taken at 300 mm from the Euryon. Coordinates for two landmarks and seventy-eight semilandmarks were recorded on the lateral view of the crania (Fig. 2B). The landmarks were located following the definitions of Buikstra and Ubelaker [27]. The application MakeFanb [28], which places alignment ‘fans’ at equal angular displacements along a curve, was used to ensure consistent placement of the craniofacial semilandmark coordinates. Both landmarks and semilandmarks were afterwards digitized by one of us (SIP) using tpsDIG 1.40 software [29].
In geometric morphometrics, shape variation can be defined as the information that remains in the coordinates of landmarks and semilandmarks after the differences due to location, scale and orientation (i.e. non-shape differences) have been removed [25]. To eliminate non-shape variation in such coordinates—by overlaying them according to a least-square optimization criteri-
on—the superimposition method known as Generalized Procrustes Analysis was used [25,26]. At the start the coordinates of any single individual are centered at the origin (0,0) by subtracting the centroid or mean location of all landmarks and semilandmarks. After that, the centroid size of the configuration (the square root of the summed square distance of all landmarks from the centroid) is set to 1 dividing the coordinates by the initial centroid size of the individual. An iterative procedure is used to determine the mean form onto which all individuals are aligned. To do so, all individuals are first aligned as a single individual, and their mean shape is calculated. All individuals are then rotated to minimize the added squared differences of point coordinates between each one of them and the estimated mean shape or reference form. This procedure is repeated until the mean shape does not change substantially after iteration of the orientation procedure. At this point, the individuals are in partial Procrustes superimposition onto the reference form [25,26]. When outlines are digitized as points, the individuals are in partial Procrustes superimposition, but the Generalized Procrustes Analysis used in geometric morphometrics is the simplest approach (see above; [31]).

A permutation procedure (10,000 permutations) was used afterwards to assess the statistical significance of the Procrustean fit. PROTEST analysis was performed using vegan 1.8–8 package for R 2.6.1 [32]. There are several strategies for superimposition, but the Generalized Procrustes Analysis used in geometric morphometrics is the simplest approach (see above; [31]). A permutation procedure (10,000 permutations) was used afterwards to assess the statistical significance of the Procrustean fit. PROTEST analysis was performed using vegan 1.8–8 package for R 2.6.1 [32].

Molecular data

mtDNA haplogroups for some of the East Central Argentina samples have previously been obtained in different works [33–36]. aDNA analytical methods used by Lalueza et al. [33] and Figueiro and Sans [36; Figueiro personal communication] are similar. Teeth and well-preserved bone pieces were handled under stringent precautionary measures to prevent extraneous contaminations. Teeth were sequentially soaked in 15% or 20% HCl for 15 min to remove dirt and carbonate deposits (in addition, Lalueza et al. [33] employed 70% ethanol for 10 min and rinsed the teeth in sterile double-distilled water for 30 min). Subsequently, teeth and bones were irradiated with UV lamp for 15 min. Next, the external surface of the samples was removed using a sand-blast to eliminate both soil and exogenous DNA contaminants. Samples

### Table 1. Sample composition, abbreviations, age, gender distribution and sample sizes.

| Samples                  | Abbrev. | Region                | Age*               | F  | M  | Total |
|--------------------------|---------|-----------------------|--------------------|----|----|-------|
| Southeast Pampa          | SEP-emH | Southeast Pampa       | Early/Middle Holocene (ca. 7,800–6,300 years BP) | 3  | 3  | 6     |
|                          | SEP-eH  | Southeast Pampa       | Earlier Late Holocene (ca. 2,500–1,500 years BP) | 2  | 7  | 9     |
|                          | SEP-IH  | Southeast Pampa       | Later Late Holocene (ca. 1,500–200 years BP) | 4  | 7  | 11    |
| Isla Gama                | IG-IH   | Northeast Patagonia    | Later Late Holocene (ca. 1,500–200 years BP) | 7  | 5  | 12    |
|                          | SB-IH   | Northeast Patagonia    | Later Late Holocene (ca. 1,500–200 years BP) | 15 | 18 | 33    |
| Laguna del Juncal        | LJ-eIH  | Northeast Patagonia    | Earlier Late Holocene (ca. 3,500–2,500 years BP) | 12 | 19 | 31    |
| Negro River Valley       | RN-eH1  | Northeast Patagonia    | Earlier Late Holocene (ca. 3,500–2,500 years BP) | 13 | 10 | 23    |
|                          | RN-eH2  | Northeast Patagonia    | Earlier Late Holocene (ca. 2,500–1,500 years BP) | 2  | 8  | 10    |
|                          | RN-IH   | Northeast Patagonia    | Later Late Holocene (ca. 1,500–200 years BP) | 9  | 12 | 21    |
| San Antonio Este         | SAE-IH  | Centre Patagonia       | Later Late Holocene (ca. 1,500–200 years BP) | 3  | 5  | 8     |
| Chubut River Valley      | ChV-eH  | Centre Patagonia       | Earlier Late Holocene (ca. 2,500–1,500 years BP) | 6  | 10 | 16    |
|                          | ChV-IH  | Centre Patagonia       | Earlier Late Holocene (ca. 1,500–200 years BP) | 18 | 20 | 38    |
| Southwest Chubut         | SWCh-IH | Centre Patagonia       | Later Late Holocene (ca. 1,500–200 years BP) | 7  | 7  | 14    |
| South Mendoza            | SM-eH   | Northwest Patagonia    | Earlier Late Holocene (ca. 2,500–1,500 years BP) | 8  | 15 | 23    |
|                          | SM-IH   | Northwest Patagonia    | Later Late Holocene (ca. 1,500–200 years BP) | 6  | 9  | 15    |
| Delta of Parana          | Del-IH  | Northeast Pampa       | Later Late Holocene (ca. 1,500–200 years BP) | 5  | 8  | 13    |
| **Total**                |         |                       |                    | 283|    |       |

*Approximate sample ages according to radiocarbon dating obtained from human bones and contextual information.

**Table 1. Sample composition, abbreviations, age, gender distribution and sample sizes.**

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|---------|-----------------------|--------------------|----|----|-------|
| SEP-emH | Southeast Pampa       | Early/Middle Holocene (ca. 7,800–6,300 years BP) | 3  | 3  | 6     |
| SEP-eH  | Southeast Pampa       | Earlier Late Holocene (ca. 2,500–1,500 years BP) | 2  | 7  | 9     |
| SEP-IH  | Southeast Pampa       | Later Late Holocene (ca. 1,500–200 years BP) | 4  | 7  | 11    |
| IG-IH   | Northeast Patagonia    | Later Late Holocene (ca. 1,500–200 years BP) | 7  | 5  | 12    |
| SB-IH   | Northeast Patagonia    | Later Late Holocene (ca. 1,500–200 years BP) | 15 | 18 | 33    |
| LJ-eIH  | Northeast Patagonia    | Earlier Late Holocene (ca. 3,500–2,500 years BP) | 12 | 19 | 31    |
| RN-eH1  | Northeast Patagonia    | Earlier Late Holocene (ca. 3,500–2,500 years BP) | 13 | 10 | 23    |
| RN-eH2  | Northeast Patagonia    | Earlier Late Holocene (ca. 2,500–1,500 years BP) | 2  | 8  | 10    |
| RN-IH   | Northeast Patagonia    | Later Late Holocene (ca. 1,500–200 years BP) | 9  | 12 | 21    |
| SAE-IH  | Centre Patagonia       | Later Late Holocene (ca. 1,500–200 years BP) | 3  | 5  | 8     |
| ChV-eH  | Centre Patagonia       | Earlier Late Holocene (ca. 2,500–1,500 years BP) | 6  | 10 | 16    |
| ChV-IH  | Centre Patagonia       | Earlier Late Holocene (ca. 1,500–200 years BP) | 18 | 20 | 38    |
| SWCh-IH | Centre Patagonia       | Later Late Holocene (ca. 1,500–200 years BP) | 7  | 7  | 14    |
| SM-eH   | Northwest Patagonia    | Earlier Late Holocene (ca. 2,500–1,500 years BP) | 8  | 15 | 23    |
| SM-IH   | Northwest Patagonia    | Later Late Holocene (ca. 1,500–200 years BP) | 6  | 9  | 15    |
| Del-IH  | Northeast Pampa       | Later Late Holocene (ca. 1,500–200 years BP) | 5  | 8  | 13    |
| **Total** |                       |                    | 283|    |       |
Figure 2. Allocated geometric coordinates are displayed with different symbols. Landmarks are represented as squares (■), whereas semilandmarks are represented as circles (●) on face (A) and vault (B) views. The numbers correspond to the following landmarks: nasion (1); nasospinale (2); prosthion (3); alare (4); ectoconchion (5); frontotemporale (6); frontomalare temporale (7); ectomolare (8); post-mastoid (9).

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were powdered under liquid nitrogen in a Spex freezer mill fitted with UV-sterilized tubes and impactors. The obtained powder was used to DNA extraction, employing a standard, high-volume phenol/chloroform protocol [33]. Several strategies were strictly followed with the object to demonstrate authenticity of the obtained data. All analyses were performed in laboratories exclusively dedicated to ancient DNA manipulation. To trace possible contamination, mtDNA sequences from the authors and other laboratory members who had manipulated the bones were obtained. To characterize the mtDNA lineages, DNA purified from bone and teeth was amplified by PCR using specific primers [33,35,36]. After amplification, the mtDNA products were classified with the specific endonucleases defining each Amerindian haplogroup and then electrophoresed on agarose gels. In addition, several samples that yielded significant PCR amplification products for the HVRI mtDNA region were used for further mtDNA sequencing characterization [33,35].

Results

For the facial skeleton, the relative warp 1 shows that the samples from Arroyo Seco 2 and the four samples corresponding to the Middle and Earlier Late Holocene from East Central Argentina separate themselves from the four Later Late Holocene samples of the same region (Fig. 3A). Almost all samples of the neighbour regions—with exception of the Delta of Parana sample—have a similar shape to the Later Late Holocene samples from East Central Argentina (Fig. 3A). Figures 3B and 3C display the deformation grids for these data, showing that the main differences along the first axis are located in the orbital and zygomatic shape, as well as in the relative size of the orbit. Particularly, the Later Late Holocene samples show the widest facial skeleton, with wider malar bones, and relatively smaller orbits. The Procrustes analysis confirm this diachronic pattern of differences (Fig. 4), showing a significant association between facial shape and temporal plus geographic dimension (m1,2 = 0.447, P = 0.016), being both, temporal and geographic variation, important to explain facial shape differences (temporal variation m1,2 = 0.446, P = 0.016; geographic variation m2,2 = 0.489, P = 0.012). The relative warp 1 of volt variation indicates that the individuals from Arroyo Seco 2 and those of the Middle and Earlier Late Holocene from East Central Argentina (mainly the sample from Laguna del Juncal) are different from the Later Late Holocene individuals of the same region (Fig. 5A). Figures 5B and 5C display the deformation grids for these data, showing that the earlier samples have longer, or dolichocephalic, cranial vault. This cranometric variation, particularly in the relative cranial length and facial width, seems to be consistent with the pattern of morphological variation interpreted as differences between Paleoamerican and Amerindian groups [9].

The ancient mtDNA analyses of individuals from Arroyo Seco 2 and Laguna del Juncal samples, however, show the presence of native American haplogroups B, C and D [33,35,36]. This agrees with the main incidence of these three haplogroups in mtDNA sequences of recent populations from the same region [33,35,36,37], related to our Later Late Holocene samples. Figueiro and Sans [36] successfully recover DNA from 8 individuals from a total of 25 individuals studied in Arroyo Seco 2 site. The haplogroups B (n = 3; 37.5%), C (n = 4; 50%) and D (n = 1; 12.5%) were found [36]. In Laguna del Juncal haplogroups C (n = 4; 26.7%) and D (n = 11; 73.3%) are present [33]; but note that this sample sizes represent the haplogroups from Aonikenk plus Laguna del Juncal site. In addition, several samples of haplogroups C and D were successfully cloned and sequenced by Lalouea et al. [33] and Garcia-Bour et al. [35], verifying the results provided by the analysis of restriction site polymorphisms. The haplogroup frequencies from Arroyo Seco 2 and Laguna del Juncal are very similar to the haplogroup frequencies from recent groups of Central Argentina. Particularly, a recent population from Pampa, the Mapuche, has values of A = 6.14%, B = 35.96%, C = 23.9% and D = 34% [38] and a recent population from Northwest Patagonia, the Pewenche, has values of A = 2%, B = 9%, C = 37% and D = 52% [39]. Such values match with the expected clinal change of haplogroups frequencies from north to south observed in South America [34-37,39].

Discussion

The results obtained show that morphological variation in East Central Argentina does not correlate with mtDNA differences. The oldest samples from the region under study, dated on ca. 8,000–2,000 years BP, present more elongated crania than the Later Late Holocene samples, but both groups have the same mtDNA haplogroups (and even haplotypes). It was pointed out that mtDNA variation in modern Native Americans support a single expansion into America of groups from Northeast or Central Asia [12–14,37]. Conversely, the same morphological differences were also observed in other regions of South America and have been used as evidence of different migratory waves, according with the two main biological components hypothesis, with a major population replacement taking place around 8,000–3,000 years BP [8–10].

This hypothesis asserts, in particular, that Amerindians have a mongoloid craniofacial shape, while Paleoamericans have a generalized morphological. Although mongoloid phenotypic pattern encompasses highly variable groups, most populations of North East Asia share two phenotypic traits: facial flatness [40] and the synodont dental pattern [41,42]. However, despite the two main biological components affirmation, facial flatness is absent among Late Holocene South American groups [40,43]. In addition, several studies that analyze craniofacial similarities between American groups and other worldwide populations demonstrate that modern aborigines do not present the typical morphology of North East Asia [44,45]. Specifically, “the American groups are more apt to join Europeans than Asians” [44], suggesting that the Late Holocene South American groups have not specific mongoloid craniofacial traits.

The lack of concordance between molecular data and craniofacial morphology has also been observed when studying the groups who inhabited the southernmost part of America during historic times [1]. The Fueguian groups have been classified as Paleoamericans by cranial shape (i.e. high levels of dolicocephaly and robusticity), and differ morphologically from other South American groups with brachycephalic morphology (i.e. Amerindian morphology sensu Neves and co-workers) [1]. However, the molecular studies show that they carried Native American mtDNA haplotypes C and D [34–35]. In addition, the study of Y-STRs sequences shows similar results [35], suggesting that the Fueguians are close to Amerindian populations from South Central Chile and Argentina.

Different hypotheses can be suggested to explain the discrepancy between mtDNA and craniofacial variation in South America. Because mtDNA is essentially a single locus, it could have been subject to considerable genetic drift, even more than morphological traits [7], during the Pleistocene-Early Holocene. Particularly, some North American aDNA studies suggest that the founding migrants exhibited greater molecular diversity than what has been previously recognized, showing that during the Early-
Middle Holocene there were more than five founding mtDNA lineages [46,47]. If the hypothetical Paleoamerican component had a particular mtDNA variation, it could have been modified during the initial South American peopling by founder effect. As Paleoamericans moved south, new territories were colonized by small groups carrying a subsample of haplotypes from the ancestral populations. Therefore, some Paleoamerican specific haplotypes could have gotten lost in Southern South America as the consequence of this process [34,37,39].

An alternative hypothesis is the existence of a selective sweep in mtDNA variation. Some investigators have pointed out that mtDNA variation has been influenced by climatic selection related to heat generation [48]. According to these authors some haplogroups, that cause lowered coupling efficiency generating less ATP and more heat, were positively selected during the radiation of modern humans into colder climates. Hence, natural selection might have favored certain mtDNA haplogroups when the Asian groups migrated into colder climates in Northeast Asia and peopled America through Beringia. Assuming that the haplogroups found in America were positively selected, their presence in Paleoamericans and Amerindians could be explained by convergent evolution. However, several recent papers found

![Figure 3. Relative Warp analysis of the face. A) Ordination of the 16 East Central samples in the space of the first two relative warps, based on the partial warp and uniform component variables calculated for the face. The circles (•) represent the consensus individual or mean shape for each Later Late Holocene sample (llH), stars (★) represent the consensus individual for each Earlier Late Holocene sample (elH), and number sign (#) represents the consensus individual for the Early Holocene sample from Arroyo Seco 2 (eH). B and C) Facial shape changes implied by variation along the first relative warp axis is shown as deformation grids. Grids show shape changes for negative (B) and positive (C) deviations from the mean for RW1. doi:10.1371/journal.pone.0005746.g003](image-url)
results contrary to what this hypothesis predicted, and support random genetic drift as the main factor in shaping mtDNA variation [49,50]. Alternatively, if we consider the hypothesis of two main biological components, and a major replacement of Paleoamericans by Amerindians during the Middle or Earlier Late Holocene, the “invading” population (i.e. Amerindians) could have had a genetic exchange with the local population (i.e. Paleoamericans). Currat and co-workers [51–53] showed that even the presence of low values of genetic exchange between local and “invading” populations can result in a major contribution of local neutral genes into the invader gene pool, and almost exclusively in this direction. This is important here because even if the Paleoamericans were replaced by a population of Amerindian morphology, mtDNA variation found in extant populations (of Amerindian morphology) could have been originated in Paleoamerican populations. These arguments contrast with the common suggestion that if a major replacement of Paleoamericans by Amerindians occurred during the Middle or Earlier Late Holocene [7–9], the variation among Later Late Holocene groups would not be relevant to discuss the early peopling.

Finally, we suggest that the lack of concordance between molecular evidence and morphological data could be explained if we take into consideration that craniofacial variation among human populations could mainly result from the action of non-random factors such as directional selection and phenotypic plasticity. These processes are suggested by the correspondence between craniofacial morphology and ecological variables (i.e. diet and climate) in South American samples. Specifically, Perez and Monterio [54] found a strong correspondence between brachycephalic crania and farmer groups, while the dolicocephalic ones associated with hunter-gatherers. If we take into account that all samples assigned as Paleoamericans belong to hunter-gatherers, and that changes in diet (i.e. production of domesticated resources) and food preparation technology (i.e. pottery and grinding use) took place between 8,000 and 2,000 years BP [55,56], the influence of ecological variables could be important to explain the morphological differences between Paleoamericans and Amerindians. A similar trend of change in craniofacial shape—mainly in the facial shape, but also in the cranial vault—has been associated with ecological factors in other world regions. Such factors include differences in the diet or masticatory activities related to the economies of hunter-gatherers vs. farmers [57–61]. The importance of environment-dependent phenotype expression during the ontogeny has been suggested to explain this morphological variation. Directional selection and/or phenotypic plasticity can generate fast morphological changes and account for the craniofacial variation found among American populations [7,54].

All this stresses the importance of elucidating the probable sources of variation of craniofacial morphology, including random and non-random factors, before being able to affirm that different traits reflect different ancestry.

Although we need more studies to discuss the alternative hypotheses about the discrepancy between mtDNA and cranio-metrical variation in South America, other molecular analyses suggest that such discrepancies could be mainly the result of non-random factors acting over the morphological divergence. Specifically, the Native American Y chromosome haplogroups...
are also originated in Central Asia and share similar coalescent dates, indicating that they have a single ancestral gene pool [12–14,37]. The American groups also share alleles at specific microsatellite loci that are not found in any Old World populations [14]. Therefore, the chromosomal and microsatellite loci molecular studies, along with the evidence that the Early Holocene samples with Paleoamerican cranial morphology carried the same mtDNA haplogroups as modern Amerindians, suggest that the Holocene East Central Argentina human populations did not have two different extra-American origins but a single one in Central or Northeast Asia.

In light of the results that were discussed here, the craniofacial variation found in samples from other South American regions also dated ca. 8,000 to 5,000 years BP [18], such as the Bogotá savannah in Colombia [9] and Lagoa Santa in Brazil [8], should be carefully interpreted. This is particularly true for the samples from the Bogotá savannah, with radiocarbon dates performed on human bones similar to Arroyo Seco 2 [18], and displaying morphologies attributable to Paleoamericans. The craniofacial comparison of these samples with samples displaying Amerindian like morphologies—as those assigned to Later Late Holocene from Perú—has been used up till now as evidence supporting the existence of two main biological components in the peopling of America [9]. However, craniofacial morphology in these regions could be also related to non-random factors, since the diachronic samples compared (e.g. Tequendama or Lagoa Santa vs. Peru sample; [8,9]) do not belong to the same area but come from regions not only geographically distant but also ecologically different (i.e. hunter-gatherers vs. farmer groups respectively).

The analysis of processes and events of population diversification involved in the American peopling has proven to be very difficult but could be addressed through the study of regional population histories, integrating diachronic skeletal samples with chronological control, mtDNA data of ancient and extant populations, archaeological records and ecological information, as well as quantitative descriptions of morphological variation. Following this approach we show here that even when the oldest samples display traits attributable to Paleoamerican crania, they present the same mtDNA haplogroups as later populations with Amerindian morphology. A possible explanation for these results could be that the craniofacial differentiation was a local phenomenon resulting from random (i.e. genetic drift) and non-

Figure 5. Relative Warp analysis of the cranial vault. A) Ordination of the individuals from East Central samples in the space of the first two relative warps, based on the partial warp and uniform component variables calculated for the vault. The circles (●) represent the individuals for Later Late Holocene sample (lH), stars (★) represent the individuals for each Earlier Late Holocene sample (eLH), and number sign (#) represents the individual for the Early Holocene sample from Arroyo Seco 2 (eH). The ellipses are the 95% confidence intervals of Earlier and Later Late Holocene mean samples. B and C) Vault shape changes implied by variation along the first relative warp axis is shown as deformation grids. Grids show shape changes for negative (B) and positive (C) deviations from the mean for RW1.

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random factors (e.g., selection and plasticity). Local processes of morphological differentiation in America are a probable scenario if we take into consideration the rapid peopling and the great ecological diversity of this continent.

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Author Contributions

Conceived and designed the experiments: SIP VB PNG. Performed the experiments: SIP. Analyzed the data: SIP. Contributed reagents/materials/analysis tools: GP. Wrote the paper: SIP VB PNG MS.

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