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Melanesian mtDNA Complexity

Jonathan S. Friedlaender
Françoise R. Friedlaender
Jason A. Hodgson
Matthew Stoltz
George Koki

See next page for additional authors

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Melanesian mtDNA Complexity

Abstract
Melanesian populations are known for their diversity, but it has been hard to grasp the pattern of the variation or its underlying dynamic. Using 1,223 mitochondrial DNA (mtDNA) sequences from hypervariable regions 1 and 2 (HVR1 and HVR2) from 32 populations, we found the among-group variation is structured by island, island size, and also by language affiliation. The more isolated inland Papuan-speaking groups on the largest islands have the greatest distinctions, while shore dwelling populations are considerably less diverse (at the same time, within-group haplotype diversity is less in the most isolated groups). Persistent differences between shore and inland groups in effective population sizes and marital migration rates probably cause these differences. We also add 16 whole sequences to the Melanesian mtDNA phylogenies. We identify the likely origins of a number of the haplogroups and ancient branches in specific islands, point to some ancient mtDNA connections between Near Oceania and Australia, and show additional Holocene connections between Island Southeast Asia/Taiwan and Island Melanesia with branches of haplogroup E. Coalescence estimates based on synonymous transitions in the coding region suggest an initial settlement and expansion in the region at ~30–50,000 years before present (YBP), and a second important expansion from Island Southeast Asia/Taiwan during the interval ~3,500–8,000 YBP. However, there are some important variance components in molecular dating that have been overlooked, and the specific nature of ancestral (maternal) Austronesian influence in this region remains unresolved.

Keywords
archaeological dating, haplogroups, mitochondrial DNA, Oceania, Papua New Guinea, population genetics, Taiwan, United Kingdom

Disciplines
Anthropology | Biochemistry, Biophysics, and Structural Biology | Genetics and Genomics | Social and Behavioral Sciences

Author(s)

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1 Anthropology Department, Temple University, Philadelphia, Pennsylvania, United States of America, 2 Independent Researcher, Philadelphia, Pennsylvania, United States of America, 3 Anthropology Department, Binghamton University, Binghamton, New York, United States of America, 4 Institute for Medical Research, Goroka, Papua New Guinea, 5 Anthropology Department, University of Pennsylvania, Philadelphia, Pennsylvania, United States of America

Melanesian populations are known for their diversity, but it has been hard to grasp the pattern of the variation or its underlying dynamic. Using 1,223 mitochondrial DNA (mtDNA) sequences from hypervariable regions 1 and 2 (HVR1 and HVR2) from 32 populations, we found the among-group variation is structured by island, island size, and also by language affiliation. The more isolated inland Papuan-speaking groups on the largest islands have the greatest distinctions, while shore dwelling populations are considerably less diverse (at the same time, within-group haplotype diversity is less in the most isolated groups). Persistent differences between shore and inland groups in effective population sizes and marital migration rates probably cause these differences. We also add 16 whole sequences to the Melanesian mtDNA phylogenies. We identify the likely origins of a number of the haplogroups and ancient branches in specific islands, point to some ancient mtDNA connections between Near Oceania and Australia, and show additional Holocene connections between Island Southeast Asia/Taiwan and Island Melanesia with branches of haplogroup E. Coalescence estimates based on synonymous transitions in the coding region suggest an initial settlement and expansion in the region at ~30–50,000 years before present (YBP), and a second important expansion from Island Southeast Asia/Taiwan during the interval ~3,500–8,000 YBP. However, there are some important variance components in molecular dating that have been overlooked, and the specific nature of ancestral (maternal) Austronesian influence in this region remains unresolved.


INTRODUCTION

Northern Island Melanesia consists of the two archipelagos just to the east of New Guinea; the Bismarcks and Solomon Islands. Understanding the genetic diversity of its populations is important to prehistoric reconstructions across the Pacific because it was settled by some of the earliest human groups to enter the entire region, and was then the area from which the exploration and colonization of vast stretches of the Pacific commenced at a much later date.

Radiocarbon dates from archaeological sites indicate Northern Island Melanesia was first settled around 40,000 years ago, very soon after people reached the ancient continent of Sahul (present day New Guinea and Australia) [1,2]. During the next 35,000 years, it remained at a comparatively isolated edge of the human species range. The early populations in Northern Island Melanesia were very small groups of hunter-gatherers. For example, New Ireland, an island over 300 km long, is estimated to have had a pre-Neolithic carrying capacity of less than 1,200 people [3]. While they were dependent on marine resources at first, the people sometimes ventured into the large island interiors [2,4]. Isolation was not complete. Plant and animal introductions from New Guinea indicate continuing contacts at a very modest level [5]. Short voyages between islands have also been inferred [2,6], since people had made the windward crossing from New Ireland to Bougainville by 29,000 YBP, and there was a detectable and repeated trickle of New Britain obisidan to New Ireland between 20,000 YBP and ~7,000 YBP [5]. By extrapolation, movements between this region and as far west as Island Southeast Asia would also have been intermittent.

During the mid-to late-Holocene, at least one significant impulse of influence came from Island Southeast Asia that led to the development of the Lapita Cultural Complex in the Bismarck Archipelago, primarily on its small off-shore islands, at ~3,300 YBP [7,8]. A few hundred years after, people bearing the Lapita Cultural Complex had colonized the islands of the Pacific as far east as Tonga and in effect had become the Polynesians (a useful distinction is Remote Oceania, which refers to the Pacific islands beyond the central Solomons settled at ~3,200 YBP or later, versus Near Oceania, which includes New Guinea and Northern Island Melanesia [9]). In Northern Island Melanesia, variable contacts between the “Lapita People” and the native groups took place along the shorelines, and some later secondary population expansions have been detected in the region as well. As a result of this complex history, Northern Island Melanesian populations are linguistically extraordinarily diverse [10–12] as well as genetically heterogeneous.

Here we show through extensive hypervariable region and targeted complete sequencing of mtDNAs that the structure of (maternal) genetic variation in Northern Island Melanesia...
indicates a long history marked by very small population sizes and limited migration rates until relatively recently. The oldest haplogroups in the region have diversified in such a localized way that particular islands can be identified as their likely homelands. Within islands, the remote inland Papuan speaking groups have diverged the most, and the largest and most rugged islands contain the greatest distinctions among populations. People(s) who entered the region during the Holocene from Island Southeast Asia/Taiwan not only carried the mtDNA “Polynesian Motif” [13,14] or its precursor, but also branches of haplogroup E. We also identify problems with associating these young haplogroups to the appearance of the Lapita Cultural Complex, or the “Out of Taiwan” model for an Austronesian expansion.

RESULTS

This section presents the results of our extensive mtDNA survey throughout Near Oceania in the context of the published literature. This includes new whole genome sequences, an analysis of the geographic structure of the patterned variation, and the discussion of the revised estimation of expansion times.

A number of mtDNA haplogroups common in Near Oceania have not been found west of New Guinea [i.e., macrohaplogroups M27 and M29, and with some rare exceptions, P, Q, and M28 [15,16]]. On the other hand, many haplogroups present in Southeast Asia are missing east of the Wallace Line (most branches of M, as well as B4c, B5, C, D, G, and U). This pattern reflects the long isolation of the populations that entered Near Oceania. Two younger mtDNA lineages do occur in appreciable frequencies in both regions, namely B4a1a1 and branches of E.

Ancient Near Oceania haplogroups

Haplogroup P

Haplogroup P is the oldest branch of macrohaplogroup R in the region. Figure 1 shows the different known branches of P in Melanesia and their defining mutations (table S1 gives further details). Source references for the different branches, including the current study, are at the top of the figure.

Figure 1. Haplogroup P phylogeny for Near Oceania (branches shared with Australian Aborigines also shown). The branches of P found only in Australian Aborigines, and details of the P1 branches, are available in [17], supplementary materials. Control region mutations are in bold, those that recur in this phylogeny are underlined, those in blue are synonymous transitions, and transversions are noted with a base suffix. Asterisks denote substitutions that can be both synonymous and nonsynonymous because of gene overlap (nts 8563 and 8572). These were regarded as nonsynonymous. The dotted line in the tree denotes a missing control region sequence. The poly C regions in HVS1 and 2 as well as 16519 are excluded. Proveniences are listed at the top, abbreviated as follows: NG–New Guinea, TR–Trobriands, AUS–Australian Aborigine, NB–New Britain. Sample numbers, GenBank accession numbers, and sources are listed underneath. Source abbreviations are: SV–[18]; TK–[25]; MI–[19]; MP–[33].

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The branching of P1 is abbreviated since it has been detailed elsewhere [17]. The branching pattern at the base of P4 is ambiguous due to the apparent occurrence of back mutations at nucleotide sites (nts) 1719 and 5460. We have identified new branches of P2, P3 and P4. P3 and probably P4 retain old connections between Near Oceania and Australia, but branch P4a appears to be specific to Near Oceania, and branch P4b appears to be limited to Aboriginal Australia.

In addition to these two ancient connections between Australia and Near Oceania, there is a possible third one, inferred from shared HVSI transitions at 16104, 16223, 16256, and 16319. In Australia, this haplotype has been called haplogroup N(S) or AuA (Genbank AF346965) [18,19] and in New Guinea it was referred to as an unnamed branch under “cluster II” (Genbank AB119390, AB119397, AB119411, AB119420 and AB119397) [20]. It may also have been found in shorter Australian HVSI sequences, as in Genbank AF176175 [21] and “sample number 13.1”[22].

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The estimates for the Time to Most Recent Common Ancestor (TMRCA) for the P branches are in table 1, using the p technique [23] and two different mutation rates [24,25]. The rankings of the p values and TMRCA are the same for both techniques, i.e., P1>P1>P4a>P2, and the general conclusion is the same—that their founder ages in Near Oceania date to ~30–50,000 YBP.

Table S2 gives the distribution of the major haplogroups in our series. P has its highest frequency in New Guinea and P1, its most common branch, has its highest concentration and greatest diversity in the highlands. P2 and P4 are also more common in New Guinea than elsewhere.

**Macrohaplogroup M** Many deep branches of M have been found throughout Asia, especially India [26–32]. Pierson et al. [33] showed that all known branches of M diverged separately from the base, with the possible exception of Melanesian M29 and Q which may be somewhat more closely related.

Figure 2 shows the main branches of macrohaplogroup M that occur in Near Oceania, including new branches of M28 and M29 identified in this study. To date, there are no established links between Aboriginal Australia and Near Oceania within any M haplogroup. As with P, the Near Oceanic branches of M apparently developed around the time of initial settlement beginning before ~30,000 years ago [current study, 17,19,34–37]. The TMRCA is table 1 for these Near Oceanic M haplogroups and their branches suggest many are as old as those for P.

Haplogroup Q is the most common Near Oceanic subdivision of M (see tables S1 and S2). Q has a large number of defining mutations at its base and long internal branches (figure 3). The Q1 branch is especially common in West New Guinea, in the Markham Valley, throughout New Britain, and north Bougainville. Q2 is most common among certain inland Papuan groups of New Britain (Baining and Ata). Although we cannot be certain Q2 originated here, it clearly underwent an expansion among the inland Papuan groups of New Britain. Both Q1 and Q2 have been found as far to the east as Fiji [38]. We could identify only seven Q3 samples: two from the New Guinea highlands [also reported in 19,21] and five from West New Britain.

Other deep branches of macrohaplogroup M probably developed in Northern Island Melanesia (table S2) since they are most common and diverse there. Haplogroup M29 (possibly related to Q–see figure 2) is most common in East New Britain. M27 is centered in Bougainville, with the M27a branch common in north Bougainville, M27b most common in east New Britain, while M27c is more scattered. Haplogroup M28 is relatively

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**Table 1. Coding region age estimates for Haplogroups common in Near Oceania**

<table>
<thead>
<tr>
<th>Haplogroup</th>
<th>Coding Region nt577-nt16022a</th>
<th>Synonymous Transitions</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>p</td>
</tr>
<tr>
<td>P1</td>
<td>6</td>
<td>8.33</td>
</tr>
<tr>
<td>P2</td>
<td>6</td>
<td>3.17</td>
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<td>P3</td>
<td>5</td>
<td>11.60</td>
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<tr>
<td>P4a</td>
<td>4</td>
<td>6.25</td>
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<tr>
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</tr>
<tr>
<td>M28b</td>
<td>2</td>
<td>8.00</td>
</tr>
<tr>
<td>M29</td>
<td>4</td>
<td>5.00</td>
</tr>
<tr>
<td>Q</td>
<td>23</td>
<td>8.65</td>
</tr>
<tr>
<td>Q1&amp;Q2</td>
<td>16</td>
<td>7.69</td>
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<tr>
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<td>13</td>
<td>5.23</td>
</tr>
<tr>
<td>Q2</td>
<td>3</td>
<td>8.67</td>
</tr>
<tr>
<td>Q3</td>
<td>7</td>
<td>5.57</td>
</tr>
<tr>
<td>E1a</td>
<td>2</td>
<td>1.00</td>
</tr>
<tr>
<td>E1b</td>
<td>3</td>
<td>1.50</td>
</tr>
<tr>
<td>B “Pol.Motif”</td>
<td>13</td>
<td>1.54</td>
</tr>
</tbody>
</table>

*One substitution per 5,139 years (Mishmar et al. 2003).

*One synonymous transition per 6,764 years (Kivisild et al. 2006)

*Island Melanesia only

*Source: Pierson et al. (2006)

*doi:10.1371/journal.pone.0000248.t001
common, and a total of 8 M28 samples have been completely sequenced. More than 80% of our M28 samples were from New Britain, and its greatest diversity is there. The center of M28 diversity is among the Baining and Ata in East New Britain. While M28 is relatively rare in New Ireland, Bougainville, and the central Solomons, we found it at fairly high frequencies in some Remote Oceanic populations-Santa Cruz (29%), Vanuatu (30% of our series, and about the same in [38]), and also in New Caledonia, Fiji, and rarely in Polynesia [16,20]. This M28 distribution suggests a portion of Remote Oceanic and Polynesian mtDNA comes from a New Britain (and Papuan) origin.

Young haplogroups

As mentioned, a second set of mtDNA haplogroups is found in both Island Southeast Asia and Oceania and dates to the Holocene.

**Haplogroup B4a1a1**

Almost 40% of our samples are B4a1a1, which includes the so-called “Polynesian Motif” (table S2). This has been tied to an Austronesian expansion out of Taiwan that led to the development of the Lapita Cultural Complex in the Bismarcks, and finally to the settlement of Polynesia and Micronesia [13,33,39–41]. This association depended heavily on the haplogroup distribution. The “Motif” is very common in Polynesia, Micronesia, and many parts of Near Oceania, and is absent in the Papuan New Guinea highlands [13], as well as in some Papuan-speaking areas of Northern Island Melanesia (tables S2, S3, and figure 4a). The “Motif” has also been confirmed in central and eastern Indonesian populations in low frequencies [42], and it could have originated either there or in Near Oceania [see 33]. It was also carried to Madagascar [43]. Whole mtDNA sequencing has identified the immediate precursor to the “Motif” in Taiwan Aboriginal groups [40], apparently strengthening the “Out of Taiwan” hypothesis (N.B. the key difference between this precursor and the “Motif” is the transition at nts 14022; the transition at 16247 had been used to identify the “Motif” in many earlier studies, but it is hypermutable in our series and therefore is not reliable).

However, this association has its problems. As shown in figure 4a and 4b, haplogroup B4a1a1 is rare in Island Southeast Asia and is not particularly common in the New Britain vicinity, which is at the center of early Lapita sites (they are mostly on nearby small islands) [5]. The “Motif” becomes very common and almost reaches fixation in some New Ireland and Bougainville groups, but some of these groups speak Papuan-languages [44]. The “Motif” is also low in frequency in Vanuatu (an area of Remote Oceania settled first by Lapita people [5], as well as Fiji, before it reaches high frequencies again in Polynesia. Therefore, the distribution association is not so compelling as it once appeared. Also, the distribution of the “Motif” precursor is poorly understood. It could be considerably more widespread than Taiwan because its identification relies on sequencing nts 14022, which has not been generally done.

The TMRCA for both the precursor of the “Motif” in Taiwan (B4a1a1), and the “Motif” in Near Oceania are also troublesome.
for the genetic part of the “Out of Taiwan” hypothesis. The molecular estimates are older than the corresponding archaeological dates. The Neolithic period in Taiwan that could have led to a subsequent Austronesian expansion dates only to about 6,000 years ago [8], while the best coalescence date for the precursor B4a1a in Taiwan was estimated at 13,000±3,800 YBP [40]. In the Bismarck Archipelago, the Lapita cultural complex dates to no earlier than 3,300 YBP [5], but the TMRCA for the “Motif” in Papuans and Polynesians was 9,300±2,000 YBP (since then, the date for the “Motif” been estimated for 13 sequences [33], using two different methods, at 7,900±1700 YBP, and 6,200±1800 YBP). However, as covered in the Discussion, the variances of these coalescence estimates are greater than generally acknowledged, so that an accommodation with the archaeological dates remains a possibility.

Haplogroup E This relatively uncommon haplogroup, a subdivision of M9, was thought to be limited to Mainland and Island Southeast Asia [45]. Its E1a branch had been found in Thailand, in Sabah Aborigines, in Taiwan Aborigines, as well as across Indonesia. The E1b branch had been found in Indonesia and the Philippines (apparently absent in Taiwan). The rarer E2 had only been found in Taiwan Aborigines and Filipinos.

We have added 5 complete E sequences now identified in Northern Island Melanesia (figure 5). The Island Melanesian E1a branches share a key mutation with one from the Philippines (nts 373), and the Melanesian E1b branches share 4 mutations with another branch from the Philippines.

The distribution of E in our series was spotty (table S2). 75 samples were E1b, and most of these were from New Britain (the Papuan-speaking Ata and Sulka). The rest were scattered across a number of Oceanic speaking groups in the region. We have not been able to ascertain from the literature if E was dispersed to Remote Oceania, since its identification depends on sequencing nts 16390 in HVSI. Figures 6 and 7 give a sense of the heterogeneous distributions of E1a and E1b in Island Melanesia and the Southwest Pacific as currently understood (table S3 has the data references). We also found two E2s, which have been identified in Taiwan as well as Indonesia and the Philippines.

In sum, E is a second young haplogroup that was brought to Near Oceania, specifically to New Britain, from the west. Coalescence time estimates for E1a and E1b in our series suggest both branches are about the same age as the “Polynesian Motif,” or slightly younger (table 1). The distribution of E1b suggests a connection between Island Southeast Asia (excluding Taiwan) and Northern Island Melanesia.

The distributions in Near Oceania of the haplogroups B4b1, F, M7, and Y are too rare to be informative.

Analysis of molecular variance.

Figure 8 gives a sense of the haplogroup variation in the core region of Northern Island Melanesia (table S2 shows the actual haplogroup incidences). While there is an island-by-island distinction, New Britain is considerably more internally diverse.
than Bougainville, with both New Ireland and Malaita considerably less so.

An analysis of molecular variance (AMOVA) was performed on the HVS1 and HVS2 sequences to quantify the mtDNA population structure (table 2). Only 32 populations from the 4 largest islands in our Island Melanesian series were included, since these were from the most intensively sampled area and could be used to compare population variation within-and among-islands, as well as within-and among-language groups. The among-population variance represented a very large proportion of the total—almost 28%, reflecting the remarkable population structure in this small region of the Southwestern Pacific. Although comparing AMOVA results across studies is not straight-forward, this value for these 4 islands is unsurpassed in the mtDNA literature for among-population variation within entire continents. In a global survey, the general among-group, within-continent variance proportion for the mtDNA control region was reported as ~8% [46], and the most recent report on African among-group mtDNA variance components is ~20% [47]. Over the entirety of North and South America, where native populations have undergone extreme drift, the estimate of the total among-group mtDNA variance is 26% [48], close to the estimate for Northern Island Melanesian populations. This indication of very high mtDNA among-group variation is no aberration. A global survey

Figure 4. Spatial frequency distribution of haplogroup B4a* and B4a1a1 in Island Southeast Asia and the western Pacific, created using the Kriging algorithm of the Surfer package of haplogroups. Figure 4b presents the detailed distribution for Northern Island Melanesia. Data details are provided in table S3.

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of short tandem repeat polymorphism variation also suggested very high Melanesian diversity, even though only two Melanesian population samples were included [49].

Partitioning the variance showed that while the variation among the four island groups was significant (12%), the variation among-populations within-islands was even greater (17.8%). New Britain, the largest and most rugged island in our series, contributed disproportionately to the within-island variance component, and New Ireland, which is over 300 kilometers long but averages less than 10 kilometers in width along most of its length, contributed the least. The size and topographical complexity of the islands is related to the genetic diversity of their populations.

Within-population mtDNA diversity across the region is related to the same pattern affected by population size and isolation. As shown in table S4, the smallest population haplotype diversity values are for seven inland Papuan groups on different large islands (the Mali, Kaket, Ané, Ata, Aita, Rotokas, and Nagovisi), while 2/3 of the highest haplotype diversity values are for beach-dwelling Oceanic groups. By island, the average population haplotype diversities are lowest for New Britain and highest for New Ireland.

Partitioning the variance by the two major language families (Oceanic vs. Papuan) produced a non-significant between-group statistic (2.9%). This is not surprising since languages belonging to these families are spoken on different islands. However, the variation among Papuan-speaking groups (40.4%) was far greater than among Oceanic-speaking groups (14.9%)—an important distinction, since the Oceanic-speaking groups tend to be distributed along the coastlines, and they were introduced much more recently in the region.

**Multidimensional Scaling Plot (MDS)**

To visualize the population relationships, we performed a non-parametric multidimensional scaling (MDS) on the pairwise FST statistics (figure 9). This plot has a stress value of 0.123 with an r² of 0.95, and therefore is a good representation of the population distinctions. The general island-by-island clustering of the populations is apparent (with New Britain populations clustered to the left top quadrant, New Ireland to the upper right, and Bougainville populations generally in the bottom half). The extreme outliers are Papuan-speaking groups, consistent with the AMOVA results. The New Britain Ata, Mali, and Kaket form one extreme cluster that contrasts with the New Ireland Kuot and south Bougainville Nagovisi at the other extreme of the first dimension; the second dimension contrasts the same New Britain Papuan cluster with the north Bougainville Aita and Rotokas.
same populations have the lowest haplotype diversities (table S4). However the Anêm, who are Papuan-speaking, fall towards the middle of the distribution. Since they show signs of substantial linguistic borrowing from their Oceanic-speaking neighbors on the shore, the Kove [50], this is not surprising.

In order to identify important associations, we correlated the population scores on the two MDS dimensions with the population haplogroup frequencies. The first dimension was negatively correlated with the frequency of haplogroup B4a1a1 ($r = -0.95$) and positively correlated with frequencies of M28 ($r = 0.70$). This explains the contrast between the New Britain and Kuot/Nagovisi Papuan-speakers. Population scores on the second dimension are most strongly correlated with Q1 frequencies ($r = 0.83$), contrasting the Aita and Rotokas with the Baining and Ata cluster. The next strongest correlations for the second dimension scores are with M27 ($r = 0.50$) and M28 ($r = -0.49$).

**DISCUSSION**

This intensive sampling in the major islands of Northern Island Melanesia, combined with sequencing of HVSI, HVS2, and whole mtDNA genome sequencing for ambiguous haplogroups, indicates a remarkably structured pattern of population diversity. It is the difference among the remote inland Papuan-speaking

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**Figure 6.** Spatial frequency distribution of haplogroup E1a in Island Southeast Asia and the western Pacific (6a), and details for Northern Island Melanesia (6b) created using the Kriging algorithm of the Surfer package of haplogroups. Data details and references are provided in table S3. doi:10.1371/journal.pone.0000248.g006
clusters on separate islands that drives the pattern of overall mitochondrial variation across the region, while the coastal groups are more intermixed. This is apparent from the MDS plot of population distances and from the AMOVA results. The shoreline Oceanic speaking groups do have island-by-island distinctions, but much less than the Papuan isolates. This overall pattern has very likely been caused by differences between the shore and inland groups in effective population sizes and marital migration rates [35,51]. The larger and the more rugged the island, the more apparent is this “beach vs. bush” distinction. As mentioned, the islands of Remote Oceania are all considerably smaller and do not appear to have retained the same levels of among group diversity across whole archipelagos. Their within population haplotype diversities are also considerably lower [16], since the frequency of the full “Motif” in the central Pacific is very high.

In addition, the phylogeography of the ancient set of haplogroups suggests the ancient haplogroup variants originated in different Papuan-speaking areas and have tended to survive there. Taken together, while associating haplogroup trees and their TMRCAs to particular population histories is a complex endeavor (see [52] for a complex Pacific example), at least in this region it produces reasonable results that are also compatible with AMOVA and population distance analyses.

Figure 7. Spatial frequency distribution of haplogroup E1b in Island Southeast Asia and the western Pacific (7a), and details for Northern Island Melanesia (7b). Data details and references are provided in table S3.
doi:10.1371/journal.pone.0000248.g007
Molecular dating estimates remain approximations and should not be used alone to reject one prehistoric migration scenario in favor of another. The mtDNA methodology is being revised yet again [see, among others, 25,53–56]. At the moment, the most attractive technique focuses on the accumulation of synonymous transitions in the coding region [25], with an estimated average rate at $3.5 \times 10^{-8}$ per year (S.D. $0.1 \times 10^{-8}$), using a human-chimpanzee split time and TMRCA date of 6 and 6.5 million years ago, respectively, for calibration. These are the same calibration dates that have been used before (see [24], citing [57]).

However, these point estimates are less precise than acknowledged. The range of legitimate estimates for the human-chimpanzee split from the fossil record is from 4.98 to 7.2 million years, and for any of these estimates there is a 95% confidence interval of $-12\%$ to $+19\%$ [58]. The earliest fossil evidence for hominid erect bipedalism is *A. anamensis* at 4 million years, providing an unequivocal lower bound for the split. If the 4.98 million year split is used rather than the one at 6 million, all resulting age estimates will be 17% less, and there will be the additional variance component contributed by the 95% confi-

![Figure 8. Northern Island Melanesian Southwest mtDNA haplogroup frequency distribution taken from our series (table S2).](doi:10.1371/journal.pone.0000248.g008)

**Table 2. AMOVA based on mtDNA HVS1 and HVS2**

<table>
<thead>
<tr>
<th>Grouping</th>
<th>$n$</th>
<th>No. of Populations</th>
<th>No. of Groups</th>
<th>Variance Components (%)</th>
</tr>
</thead>
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<tr>
<td>No grouping (32 populations)</td>
<td>1223</td>
<td>32</td>
<td>1</td>
<td>Between Groups: 27.5</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Within Groups: 72.5</td>
</tr>
<tr>
<td>Geography (4 Islands)</td>
<td>1223</td>
<td>32</td>
<td>4</td>
<td>12.2</td>
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<td></td>
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*Groups for geography: New Ireland, New Britain, Bougainville and Malaita

Linguistic groups: Papuans, Oceanic

$p = 0.07$

doi:10.1371/journal.pone.0000248.t002
dence interval. For example, an estimated TMRCA of 6,000 YBP (as with the current estimate for the “Polynesian Motif” in table 1) could be more than 1,000 years too early or too late, with an uncertainty of ~10–15% added to the estimate and standard deviation. The same percentages would apply to the much older estimated TMRCA ages for the most ancient haplogroups as well.

Tying the B4a1a1 haplogroup, which predominates in Polynesia and Micronesia, with mid-Holocene population movements originating in Taiwan [59] or Wallacea [60] has gotten more complicated as coverage has improved [45,51]. What is clear is that precursors of the “Motif” originated to the west of Wallacea in the early Holocene; that the full “Motif” with the transition at 14022 developed in eastern Island Southeast Asia or Near Oceania; that its frequency varies a great deal across Island Southeast Asia, Near Oceania, and sections of Remote Oceania before becoming very common in central Polynesia; and that subtypes of haplogroup E, which also developed in Island Southeast Asia during the Holocene, also have a very spotty distribution and were carried to the Bismarcks but probably no further east. Also, the analyses of skeletal remains associated with Lapita or earliest Polynesian sites have still not yielded any B4a haplogroup identifications [61,62]. It must be remembered that mtDNA haplogroups (along with Y variants) are especially affected by genetic drift, since the effective sample size is only ¼ that of an autosomal marker in any population. Also, the small populations colonizing Near and Remote Oceania were probably subject to exaggerated drift distortions [53,56].

The main point is that mtDNA variation among populations in Northern Island Melanesia is extreme but understandable. It reflects the very ancient settlement of the region; the subsequent isolation and drift of many inland populations; some subsequent internal population expansions; the introduction of two haplogroups and populations in the mid Holocene, combined with some intermixture among many groups, especially those living along the shorelines. Because the mtDNA only reflects a very small (exclusively maternal) fraction of the heritage of an individual or population, it may yield a biased result, but this survey also shows its power in elucidating ancient population dynamics.

MATERIALS AND METHODS

The samples analyzed were selected from our Southwest Pacific collection. Its core consisted of blood samples collected in three recent field seasons in the Bismarck Archipelago. This primary set was augmented with plasmas and urines from older collections, described elsewhere [17]. Information on survey subjects included their language, a short genealogy, current residence, and familial birthplaces (used to assign location), although such details were not available from some of the other collections. The primary samples were collected, and all selected samples were analyzed, with informed consent protocols approved by the appropriate Human Subjects Ethical Committees of Papua New Guinea, the University of Michigan, Binghamton University, and Temple University.
One sample from each identified matriline was selected for the initial mtDNA control region analysis. The analysis of the samples occurred in three phases: (1) sequencing of hypervariable segments 1 and 2 (HVSI and HVSI); (2) for those samples not definitely assigned to a known haplogroup on this basis, RFLP screening for the two mutations defining macrohaplogroup M (DdeI 10394, Ahu 10397) and, depending on the presence or absence of these, additional RFLPs known to identify other haplogroups in the Southwest Pacific [37,63,64]; and (3) sequencing of the coding region on 16 representative samples from the remaining major haplogroups that could not be assigned to currently published sublineages.

DNA was extracted from 100 to 200 µl of buffy coat, plasma, or urine (depending on the source of the sample) by using either the guanine-silica based IsoQuick extraction kit (Ora Science, Bothell WA) or the column-based QiaGen extraction kit (Qiagen, Valencia CA). In preparation for sequencing, the mtDNAs were PCR amplified following standard protocols, and employing Platinum Taq Polymerase (Invitrogen, Carlsbad CA). The control region was amplified using primers spanning nps 15938 to 00429. The coding region was amplified using the PCR primers and conditions of Rieder et al. [65]. Successful amplification was verified by electrophoresis on 1% ethidium bromide stained agarose gels. Samples were prepared for sequencing by an ExonI digest followed by filtration through a Millipore 96-well filter plate (Millipore, Billerica MA) to remove single stranded DNA and unincorporated nucleotides. PCR products were sequenced using various versions of the BigDye Terminator Sequencing kits from ABI (Applied Biosystems Inc) on an ABI 377XL automated sequencer using conditions described previously [66]. Custom designed internal sequencing primers were used for all large PCR fragments to increase double-fold coverage.

Contig assemblage and sequence alignment was accomplished with Sequencer: Forensic Version 4.1.9 (GeneCodes, Ann Arbor MI). Sequencer was also used to determine the synonymous transitions. The phylogenetic tree was inferred from median-joining networks rooted to L3. The tree was hand-checked to resolve several homoplasies. A few ambiguities remained, and we tended to be conservative in interpreting those cases.

REFERENCES


