Population dynamics in the Japanese Archipelago since the Pleistocene revealed by the complete mitochondrial genome sequences

Fuzuki Mizuno1, Jun Gojobori2, Masahiko Kumagai3, Hisao Baba4, Yasuhiro Taniguchi5, Osamu Kondo6, Masami Matsushita7, Takayuki Matsushita7, Fumihiko Matsuda8, Koichiro Higasa9, Michiko Hayashi3, Li Wang10, Kunihiko Kurosaki1 & Shintaroh Ueda1,6

The Japanese Archipelago is widely covered with acidic soil made of volcanic ash, an environment which is detrimental to the preservation of ancient biomolecules. More than 10,000 Palaeolithic and Neolithic sites have been discovered nationwide, but few skeletal remains exist and preservation of DNA is poor. Despite these challenging circumstances, we succeeded in obtaining a complete mitogenome (mitochondrial genome) sequence from Palaeolithic human remains. We also obtained those of Neolithic (the hunting-gathering Jomon and the farming Yayoi cultures) remains, and over 2,000 present-day Japanese. The Palaeolithic mitogenome sequence was not found to be a direct ancestor of any of Jomon, Yayoi, and present-day Japanese people. However, it was an ancestral type of haplogroup M, a basal group of the haplogroup M. Therefore, our results indicate continuity in the maternal gene pool from the Palaeolithic to present-day Japanese. We also found that a vast increase of population size happened and has continued since the Yayoi period, characterized with paddy rice farming. It means that the cultural transition, i.e. rice agriculture, had significant impact on the demographic history of Japanese population.

The Japanese Archipelago located in the boundary of the Eurasian, the North American, the Pacific and the Philippine Sea Plates was created by separating from the Asian continent by 15 million years ago. Archaeological evidence indicates the first appearance of people in the Japanese Archipelago during the late Pleistocene between 40,000 and 30,000 years ago. The prehistoric era after the Palaeolithic era is typically split into two contrasting periods, Jomon and Yayoi. The Jomon period is characterized by hunter-gatherer subsistence that persisted for more than 10,000 years. The name of Jomon derived from the distinctive rope-pattern decorations on the pottery. The presence of pottery in Jomon period shows that making pottery was not just unique to agricultural peoples. On the other hand, the Yayoi period is characterized as paddy rice farming. And it is thought that this style of rice production was introduced into the Japanese Archipelago by migrants from mainland Asia and that the agricultural migrants came to the Japanese Archipelago in and after the Yayoi period. Since the middle nineteenth century several hypotheses have been proposed in regard to the population history of present-day Japanese, but it is now popularly accepted that the present-day Japanese consists of at least two ancestries, one from southeast Asia-originated Jomon people and the other from northeast Asia-originated Yayoi people, although scarce knowledge of Palaeolithic people.

1Department of Legal Medicine, Toho University School of Medicine, Tokyo, Japan. 2Department of Evolutionary Studies of Biosystems, SOKENDAI (The Graduate University for Advanced Studies), Hayama, Japan. 3Advanced Analysis Center, National Agriculture and Food Research Organization, Tsukuba, Japan. 4Department of Anthropology, National Museum of Nature and Science, Tokyo, Japan. 5Department of Anthropology, Faculty of Letters, Kokugakuin University, Tokyo, Japan. 6Department of Biological Sciences, Graduate School of Science, The University of Tokyo, Tokyo, Japan. 7The Organization of Anthropological Research, Tokyo, Japan. 8Graduate School of Medicine, Kyoto University, Kyoto, Japan. 9Department of Genome Analysis, Institute of Biomedical Science, Kansai Medical University, Hirakata, Japan. 10School of Medicine, Hangzhou Normal University, Hangzhou, China. *email: fuzuki.mizuno@med.toho-u.ac.jp; gojobori_jun@soken.ac.jp; liwang@hznu.edu.cn
The Japanese Archipelago is widely covered with acidic soil made of volcanic ash, making ancient DNA studies challenging. This is the first report of Palaeolithic mitogenome (mitochondrial genome) sequence in the Japanese Archipelago. Here we investigate the population dynamics in the Japanese Archipelago using complete mitogenome sequences from the Palaeolithic, Jomon, Yayoi and present-day Japanese, and find continuity in maternal gene pool among them from the point of view of haplogroup diversity. Through demographic analysis using over 2,000 present-day Japanese, we observe dramatic population explosion across cultural transition from hunter gathering to farming. This was a period in which the environment is known to have undergone short, rapid fluctuations in temperature. During the transition from the Last Glacial Period (LGP) to the present Holocene, there was also a temporary return to glacial conditions, known as the Younger Dryas. This rapid climate change has been shown to be strongly associated with the widespread extinction of megafauna.

Mitochondrial Haplogroup M is frequently observed in present-day Asian populations but is not found in present-day European populations, while analysis of mitochondrial DNA from archaeological human remains dating to the Late Pleistocene has identified individuals belonging to haplogroup M that populated Europe before the LGP. This indicates drastic change in the maternal gene pool. By obtaining complete mitogenome sequences, we here investigate genetic relationships among the Palaeolithic, the following hunter-gathering Jomon and agricultural Yayoi people, and furthermore disclose the past demographic history of present-day Japanese population. In case when we found a mitochondrial DNA whose haplogroup is not found in the present-day Asian population, there would be a possibility of drastic change of maternal gene pool as observed in Europe. But in case when we found a mitochondrial DNA whose sequence is closely related to the mitochondrial DNA found in present-day Asian population, there would be less possibility of drastic change of maternal gene pool, namely population continuity.

Results and discussion

Mitogenome of the Palaeolithic remains. We successfully determined a highly accurate complete mitogenome sequence of 20,000-year-old Minatogawa 1 (Minato1), a plausible direct descendant of the initial migration into the Japanese Archipelago (Fig. 1, Table 1 and Supplementary Table S1). Its sequence obtained with average depth of 52 was classified into haplogroup M, and carries no substitutions that are defining subgroups of haplogroup M. Haplogroup M is found at high frequency in present-day Asians, Australasians, and indigenous Americans. The sequence of this ancestral type of haplogroup M is not seen in any of 2,062 present-day Japanese samples newly determined in this study, 672 present-day Japanese, 21,668 Han Chinese, and present-day Japanese. But it located near the root of haplogroup M. This suggests that Minato1 belongs to the ancestral population of present-day Japanese but also to the ancestral population of present-day East Asians. A similar case was reported in Asian mainland, a 40,000-year-old Tianyuan individual excavated from northern China. It was reported to be the mitochondrial sequence of ancestral-type haplotype B having four singletons (private mutations), i.e. a common ancestor of present-day mitogenome belonging to haplogroup B. Due to fluctuating global warming, Palaeolithic period from the LGP is supposed to be a difficult time to survive in5,6, and changes in the gene pool are expected to occur various populations all over the world. However, the results of phylogenetic network and Neighbor-Joining phylogenetic tree including Minato1 and Tianyuan individuals (Fig. 4a,b) indicate that drastic changes in maternal gene pool during the LGP had not occurred in East Asia because it shows that both Minato1 and Tianyuan belong to the ancestral population of present-day East Asians.

Mitogenome of the hunting-gathering Jomon remains. We further successfully determined highly accurate complete mitogenome sequences with average depths of coverage from 11 to 1,577 using 8,300- to 3,700-year-old Jomon individuals (10 individuals from six archaeological sites, Fig. 1, Table 1 and Supplementary Table S1). All of the Jomon individuals fall into the same clusters with present-day Japanese in phylogenetic network, phylogenetic trees and MDS plot (Figs. 2, 3, 4). This shows population continuity from Jomon period to present-day in the Japanese Archipelago, meaning no dynamic change in the maternal gene pool of people living in the Japanese Archipelago from the Palaeolithic to Neolithic periods. The results also show that most of the Jomon period individuals in our collection are of haplogroups M or N, and many of them are of sub-haplogroups M7a or N9b in the narrow taxonomic group (Table 1 and Supplementary Table S1). This is consistent with the previous observations from PCR-based typing. Both haplogroups M7a and N9b characteristic of Jomon individuals have been inherited by the present-day Japanese individuals; 7.9% and 1.3% among the present-day Japanese population, respectively (Supplementary Table S2).

Mitogenome of the farming Yayoi remains. In addition, we successfully determined highly accurate complete mitogenome sequences of four individuals from Yayoi period with average depths of coverage from 13 to 5,891 (each two individuals from the Hanaura and Doigahama Yayoi sites, respectively. Figure 1, Table 1 and Supplementary Table S1). The Yayoi period started by the lifestyle of paddy rice farming that was brought into Japanese Archipelago by Yayoi migrants. The results show that three of the Yayoi individuals belong to haplogroup D4 (Table 1). D4 is the most common haplogroup in present-day Japanese (34.3%) (Supplementary Table S2), which is also common throughout East Asia. Like the Jomon individuals, all of the Yayoi individual fall into any of the clusters of the phylogenetic network, MDS, Bayesian and Neighbor-Joining phylogenetic trees that are constructed together with the present-day Japanese people, although the Jomon and Yayoi people have
some mitochondrial sub-haplogroups characteristic to each of them (Figs. 2, 3, 4). Combining with the results of Minato 1 and Jomon samples, it is shown that there is at least some population continuity from the late Pleistocene to present-day human populations in the Japanese Archipelago.

**Estimating past demographic trends.** Our results show that the gene pool of the present-day Japanese has been established, swallowing all of the genetic diversities of the people living in the Japanese Archipelago during more than 10,000 years, starting from the Palaeolithic period through the hunter-gatherer Jomon and farming Yayoi periods. Bayesian-Skyline Plot (BSP) analysis using 2,062 present-day Japanese found three large population increases, at 45,000–35,000 BP, 15,000–12,000 BP, and 3,000 BP (Fig. 5). These correspond with a rise in temperatures seen in the Late Pleistocene, the dawn of agriculture in East Asia, and the beginning of the Yayoi period, respectively. In a recent study relating to East Asia, demographic history was referred from com-
plete mitogenome sequences of 21,668 present-day Han Chinese individuals. The results showed that there was a population increase towards the end of the LGP around 45,000–35,000 BP, followed by another, more rapid increase during the Neolithic period 15,000–12,000 BP. The first two population increases are common among the present-day Japanese and Chinese populations, whereas the third one can be seen uniquely in the present-day Japanese. And the third phenomenon unique to the Japanese Archipelago, which took place after beginning of the Yayoi period, did great contribution to population size of the present-day Japanese. Combined with them, the first two population increases seen in the present-day Japanese populations should have mainly reflected ones occurred before the ancestors of Yayoi people migrated to the Japanese Archipelago, namely the increases taken place in the Asian mainland. It is easily expected that the Yayoi migrant bringing rice paddy farming should have a significant impact on the number of population and population structure in the Japanese Archipelago. It is suggested that Holocene climate shifts happened 2.8 ka or 4.2 ka BP events affected the population of Korean Peninsula and drove migration to Japanese Archipelago. Further population increase afterwards could be related to the introduction of ironware into the Japanese Archipelago, which allowed for more efficient rice paddy farming and a more stable food supply. Given all the findings obtained, our results show that genetic makeup of the present-day Japanese populations is built by migration events of Yayoi farmers and subsequent multiple migrations from the Asian mainland. But we cannot ignore contribution of the Jomon people to the present-day Japanese population structure.

We show (a) there were continuity in the maternal gene pool since the initial wave of migration into the Japanese Archipelago of Palaeolithic period from the point of view of haplogroup diversity, (b) ancestors of the present-day Japanese had experienced three large population increases, but the first two of them mainly occurred in the Asian mainland. The third increase was a sharp one occurred in relatively short period, (c) gene pool of the Jomon hunter-gatherers has survived, even after migration of the Yayoi farmers and its subsequent population explosion.

The following key questions remains open such as whether the migrating Yayoi farmers genetically mixed with the indigenous Jomon hunter-gatherers at each stage after the Yayoi migration (and, if so, what type of mix occurred). Our study includes only mitogenome and the number of samples is limited. Therefore, to reach more solid or concrete conclusion, we need to reveal more genome information of ancient people then we can obtain clearer evidence on the presence of friendly relationships between Jomon hunter-gatherers and Yayoi farmers. Further mitogenome investigation of other Palaeolithic remains and nuclear genome analysis of Minato 1 should give us more clues and details to disclose human history in Japanese Archipelago.

### Materials and methods

Approval for the present study was provided by the Ethics Committee of Toho University School of Medicine (A20110_A18099_A18056, A19095). Approval was also provided by the Ethics Committee of Department of Evolutionary Studies of Biosystems, SOKENDAI; The Graduate University for Advanced Studies (SKD2019GHG001). The Ethics Committee of Kyoto University Graduate School and Faculty of Medicine (G0751-4), the Ethical Review Board of the Nagahama Study, and the Nagahama Municipal Review Board of Personal Information

---

**Table 1.** Late Pleistocene and Holocene human remains of the Japanese Archipelago used in this study.

| Individual ID | Era       | Site               | Approximate date       | mtDNA haplogroup | References |
|---------------|-----------|--------------------|------------------------|------------------|------------|
| Minato1       | Palaeolithic | Minatogawa fissure  | 19,900 cal BP          | M                | This study |
| Iyai1         | Initial Jomon | Iyai rock-shelter  | 8,300–8,200 cal BP     | N9b              | Our study (Mizuno et al. 2020) |
| Iyai4         | Initial Jomon | Iyai rock-shelter  | 8,300–8,200 cal BP     | N9b             | Our study (Mizuno et al. 2020) |
| Iyai8         | Initial Jomon | Iyai rock-shelter  | 8,300–8,200 cal BP     | N9b             | Our study (Mizuno et al. 2020) |
| Higa002       | Initial Jomon | Higashimou shell midden | –                   | N9a2a           | This study |
| Higa006       | Initial Jomon | Higashimou shell midden | 7,934–7,792 cal BP    | M7a1a           | This study |
| Higa020       | Initial Jomon | Higashimou shell midden | –                   | M80’D           | This study |
| Todo5         | Early Jomon | Todoroki shell midden | 6,210–6,094 cal BP    | M7a1a           | This study |
| Kasa6         | Middle Jomon | Katori shell midden | –                   | M7a             | This study |
| Uba2          | Middle Jomon | Ubayama shell midden | –                   | N9b             | This study |
| MB-TB27       | Late Jomon | Mabuni hantabarai  | –                     | M7a1a           | This study |
| F5            | Late Jomon | Funadomari         | 3,800–3,500 cal BP     | N9b1            | Kanzawa-Kiryama et al. |
| F23           | Late Jomon | Funadomari         | 3,800–3,500 cal BP     | N9b1            | This study |
| Ik002         | Final Jomon | Ikawazu shell midden | 2,720 cal BP          | N9b1            | McColl et al. |
| DH-S01        | Middle Yayoi | Doigahama         | 2,306–2,238 cal BP     | D4b2b1          | This study |
| DH-A          | Middle Yayoi | Doigahama         | –                     | D4b2b1          | This study |
| HN-SJ001      | Middle Yayoi | Hanaura           | –                     | B4b1a1a         | This study |
| HN-SJ002      | Middle Yayoi | Hanaura           | –                     | D4b2a1          | This study |
**Figure 2.** Bayesian phylogenetic tree of complete mitogenome sequences of 18 ancient and 171 present-day individuals in the Japanese Archipelago. Coloured arrows show the positions of ancient individuals; red, yellow, and purple for Palaeolithic, Jomon, and Yayoi, respectively. The tree was rooted with mitogenome haplogroup L0a (accession number: EF184601.1). The estimated haplogroups for newly determined ancient samples are Minato1:M, Iyai1:N9b, Iyai3:N9b3, Iyai6:N9b, Higa002:N9a2a, Higa006:M7a1a, Todo5:M7a1a, Kaso6:M7a, Uba2:N9b, MB-TB27:M7a1a, DH-S01:D4b2b1, DH-A:D4b2b1, HN-SJ001:B4b1a1a and HN-SJ002:D4b2a1, respectively (Table 1 and Supplementary Table S3).
Protection approved all study procedures. All methods were performed in accordance with the relevant guidelines and regulations.

**Ancient samples.** Bones and teeth from human remain representing 15 individuals from nine archaeological sites were sampled for ancient genome analyses; Minatogawa fissure and Mabuni hantabaru (Okinawa Pref.), Iyai rock-shelter (Gunma Pref.), Higasimyou and Hanaura (Saga Pref.), Todoroki shell midden (Kumamoto Pref.), Kasori shell midden and Ubayama shell midden (Chiba Pref.), Doigahama (Yamaguchi Pref.) (Supplementary Table S1, Fig. 1). Dates of five samples are available, the others are referenced on pottery typological chronology. Minato1 indicates many archaic morphological features, such as a thick and inferiorly wide skull vault, a small frontal bone with extremely developed temporal fossae, and a rugged facial profile. It is reasonable to suppose that it was remnants of the earliest *Homo sapiens* in East Asia, who somehow migrated to Okinawa Islands around 30,000 or 40,000 years ago and survived at least until around 20,000 years ago, keeping these archaic features.

**DNA extraction, NGS library preparation, enrichment of the ancient mitochondrial genome and sequencing.** DNA was extracted from bones and teeth, and NGS library was constructed according to our previous methods with slight modifications as follows. After exhaustively brushing to eliminate dirt and exogenous contaminants, the outer bone surface was mechanically removed with a sanding machine (Dremel) to further remove surface contaminants. The clean bone was cut into pieces of approximately 1–2 cm³ using an electric drill cutter (Dremel). The bone fragments were cooled with liquid nitrogen, and fine powdered bone was obtained by grinding bone fragments in a mill (Multi-beads Shocker MB601U, YASUI KIKAI). During all steps of DNA extraction, NGS library preparation, and enrichment, we took all possible precautions to guard against contamination. Experiments were performed in a laboratory that is exclusively dedicated to ancient DNA work and is physically isolated from other molecular work laboratories. All manipulations were performed in a laminar flow cabinet routinely irradiated with UV light. Frequent surface cleaning was routinely performed before and after working. A facemask, head cap, and clean laboratory coat were always worn, and gloves were frequently replaced. All the procedures were conducted using new sterilized disposable tubes and filter pipette tips. All non-disposable glass and metallic materials were dry-heat sterilized at 160 °C for 2–4 h. The powdered samples (10 to 200 mg) were digested in 0.5 M EDTA (pH 8.0) for 2 h at 56 °C in a rotating hybridization oven, and the supernatant was removed by centrifugation. This decalcification step was repeated three times in total. DNA was extracted with phenol:chloroform:isoamyl alcohol (25:24:1), followed by extraction with an equal volume of...
Figure 4. (a) A median-joining network for whole coding mitogenome of 37 ancient individuals and 287 present-day Asians belonging to macrohaplogroups M and N. Number of mutations was shown as hatch marks on branches. (b) The phylogenetic tree inferred using the Neighbor-Joining method for world-wide 37 ancient individuals and 287 present-day Asians belonging to macrohaplogroups M and N. The samples used here belong to either of macrohaplogroup M (including C, D, G, M7, M8, M10, and Z), and macrohaplogroup N (including A, B, B5, F, N9, R9, R11, and Y). The evolutionary distances for whole coding region of mitochondrial genome, totally 14,918 bp, were computed using the Maximum Composite Likelihood method. The percentage (50% and greater) of replicate trees in which the associated taxa clustered together in the bootstrap test (1,000 replicates) are shown next to the branches. All positions containing gaps and missing data were eliminated (complete deletion option).
chloroform. After centrifugation, the aqueous solution was removed and subsequently concentrated by centrifugation dialysis using an Amicon Ultra-15 30 kDa centrifugal filter (Merck Millipore) to a final volume of 200 μL. The DNA solution was purified with silica-based MiniElute spin columns (Qiagen) according to manufacturer's protocol. The obtained DNA was quantified using Quant-iT dsDNA HS assay kit (Thermo Fisher Scientific).

Using the obtained DNA, we prepared single- or double-strand NGS libraries. For some samples, PreCR Repair Mix (New England BioLabs) treatment was done. Libraries were enriched for human mitochondrial genomes, using in-solution target enrichment (SureSelect, Agilent Technologies). The libraries were pooled sequenced on illumina MiSeq or HiSeq platform 1500 according to the manufacturer's instructions.

**NGS data processing and ancient DNA authentication.** Raw sequencing reads were trimmed by removing adapter sequences and low-complexity sequences with fastp ver. 0.20.0 (parameters: -l 30 -y –detectadapter_for_pe)\(^3\). The trimmed reads were mapped against the human mitogenome reference sequence rCRS\(^3\), using the Burrows-Wheeler Alignment tool (BWA)\(^3\) with optimal parameters for ancient DNA (parameters: -l 1024 -n 0.01 -o 2)\(^3\). Reads with a mapping quality lower than 30 were filtered out and high-quality mapped reads

---

**Figure 4.** (continued)
were retained using samtools ver. 1.9 \textsuperscript{36}. To minimize the effects of nucleotide mis-incorporations on building a consensus mitogenome sequence, the first two bases on each end of the read were clipped with BamUtil ver. 1.0.1437 according to our previous bioinformatics procedure\textsuperscript{38}. Deamination is considerably repaired by using PreCR Repair Mix, but even if it is repaired, it is difficult to completely repair it. Because the damage still remains, both ends are clipped. Then, reads with a significant hit (E-value \leq 1e^{-15}) to non-human genomes (e.g., fungal or bacterial genomes) were identified by BLASTN and then were also filtered out. In addition, reads mapped to human nuclear genome sequences of hg19 were also removed as numts (nuclear copies of mitogenome). Finally, consensus sequences were built using MitoSuite\textsuperscript{39}, and their mitogenome haplogroup assignments were called with HaploGrep\textsuperscript{240}. We checked variants using IGV software\textsuperscript{41} based on PhyloTree Build 17 (\url{http://www.phylo-tree.org/})\textsuperscript{42}. To assess whether the results could be affected by present-day human contamination, sequences with ancient DNA specific cytosine to thymine substitutions (C to T) at the 5’ or 3’ molecule end were selected and analysed separately (Supplementary Fig. S2). DNA contamination was estimated using MitoSuite ver.1.0.9\textsuperscript{39}.

Present-day Japanese samples. Present-day Japanese examined in this study are participants of the baseline survey in the Nagahama Study. The Nagahama Study is an ongoing community-based cohort study conducted by the Kyoto University Graduate School of Medicine and Nagahama City. The participants are members of the general population living in Nagahama City, a rural city of 125,000 inhabitants in Shiga Prefecture located in central Japan, aged 30 to 74 years old.

NGS data processing of present-day Japanese mitogenome DNA. Whole genome sequencing of present-day Japanese was conducted using the Illumina HiSeq X Ten sequencer (Illumina Inc., San Diego, CA, USA). After aligning the sequence reads onto the reference genome (GRCh37/hg19) using the Burrows-Wheeler Aligner with default parameters, according to the developer’s manuals\textsuperscript{34}. Then, duplicate reads were marked using Picard. A total of 2,062 subjects were included in the present study, of which complete mitogenome sequences are available with high depth of coverage. Finally, consensus sequences were built from the BAM files of chrM using MitoSuite\textsuperscript{39}, and their mitogenome haplogroup assignments were called with HaploGrep\textsuperscript{240} (Supplementary Table S2). We checked variants using IGV software\textsuperscript{41} based on PhyloTree Build 17\textsuperscript{42}.

Statistical phylogenetic analyses. To produce Bayesian phylogenetic tree, we used mrbayes v. 3.2\textsuperscript{43}. “GTR + I + Γ” model was used for substitution model. Model was selected by Maximum Likelihood fits function in MEGA X\textsuperscript{44}. We set the Markov Chain Monte Carlo (MCMC) chain length to 3.0 \times 10^6 with 7.5 \times 10^5 burn-in steps to collect sufficient samples for estimation. In this Bayesian tree, we selected 171 present-day Japanese mitochondrial genome sequences to cover major haplotypes belonging macro haplogroups M and N (see Supplementary Table S3 for the list of sequence used) for the ease of computation. Our newly determined 15 ancient mitochondrial genome sequences were aligned together and included in the tree estimation. The tree was estimated by using haplogroup L0a (accession number: EF184601.1) as an outgroup. For MDS, we used 2,062 complete mitogenome sequences from Japanese population, which were produced in this study. Sequences without “N” nucleotide were used. We used mafft v7.471 to align these 2,062 sequences with rCRS sequence\textsuperscript{45}. Mutations were scored relative to the rCRS and we extracted the coding regions. The 15 ancient mitochondrial genome sequences produced in this study were added to this multiple sequence alignment. MDS was performed on R software with cmdscale() function\textsuperscript{46}. We used p-distance based on the number of differences on nucleotide sequences for MDS. We used MEGAX for obtaining p-distances\textsuperscript{44}. A median-joining Network\textsuperscript{47} was constructed by using PopART v1.7\textsuperscript{48} with a parameter “epsilon = 0”. We selected 287 modern Asian mitochondrial genome sequences to cover major haplotypes belonging macro haplogroups M and N. Also, 22 ancient mitochondrial

![Figure 5. Bayesian Skyline Plot (BSP) of 2,062 present-day Japanese. Effective population size (y axis, log-scale) was plotted against time (x axis, years ago). Solid black line shows the mean effective population size from the posterior distribution. Two grey lines and grey coloured region within them show the 95% posterior density intervals.](image-url)
genomes available were obtained. Together with above dataset, our newly determined 15 ancient mitogenome sequences were aligned, and then, variant sites in coding region of mitogenome was used as an input for the network analysis (see Supplementary Tables S3 and S4 for the lists of sequence used). The resultant network was manually modified to improve visibility on PopART.

**Bayesian skyline plot (BSP).** To obtain BSP, we used BEAST v2.5.0 suite. We applied the general time reversible model mutation model using gamma-distributed rate. A clock model was used assuming a log-normal distribution. We set the Markov Chain Monte Carlo (MCMC) chain length to 1 x 10^8 with 1 x 10^8 burn-in steps to collect sufficient samples for parameter estimation. We performed the MCMC simulation twice independently to confirm that the simulation converged to the same state. The time was scaled by the number of mutations for BSP construction. To estimate time in year, we assumed a molecular clock of 2.77 x 10^{-8} and applied the correction method of Soares et al. To scale the effective population size (Ne), we assumed a generation time of 25 years. The simulation results were analysed using Tracer v1.7.1 (http://tree.bio.ed.ac.uk/software/tracer/).

**Data availability**
Sequencing data produced in this study have been deposited in the GenBank (accession no. LC521960-LC521962, LC597326-597337) and Human Genetic Variation Database (accession no. HG0000011, http://www.hgvd.genome.med.kyoto-u.ac.jp) for ancient samples and present-day Japanese, respectively.

Received: 12 January 2021; Accepted: 25 May 2021

Published online: 13 June 2021

**References**
1. Hanika, K. Dual structure model for the population history of the Japanese. Nichibunken Ipn. Rev. 2, 1–33 (1991).
2. Taniguchi, Y. The beginning of pottery technology in Japan: The dating and function of incipient Jomon pottery. Proc. Natl. Acad. Sci. U.S.A. 115, 7931–7936 (2018).
3. Miyamoto, K. The spread of rice agriculture during the Yayoi Period: From the Shandong peninsula to the Japanese Archipelago via the Korean Peninsula. Ipn. J. Archaeol. 6, 109–124 (2019).
4. Cooper, A. et al. Abrupt warming events drove late Pleistocene Holartic megafaunal turnover. Science 349, 602–606 (2015).
5. Broughton, J. M. & Weitzel, E. M. Population reconstructions for humans and megafauna suggest mixed causes for north American Pleistocene extinctions. Nat. Commun. 9, 5441 (2018).
6. Fu, Q. et al. The genetic history of ice age Europe. Nature 534, 200–205 (2016).
7. Pozhit, C. et al. Pleistocene mitochondrial genomics suggest a single major dispersal of non-Africans and a late glacial population turnover in Europe. Curr. Biol. 26, 827–833 (2016).
8. Bramanti, B. et al. Genetic discontinuity between local hunter-gatherers and central Europe’s first farmers. Science 326, 137–140 (2009).
9. Malmström, H. et al. Ancient DNA reveals lack of continuity between Neolithic hunter-gatherers and contemporary Scandinavians. Curr. Biol. 19, 1758–1762 (2009).
10. Haak, W. Massive migration from the steppe was a source for Indo-European languages in Europe. Nature 522, 207–211 (2015).
11. Brace, S. et al. Ancient genomes indicate population replacement in early Neolithic Britain. Nat. Ecol. Evol. 3, 765–771 (2019).
12. Kivisild, T. et al. The emerging limbs and twigs of the East Asian mtDNA tree. Mol. Biol. Evol. 19, 1737–1751 (2002).
13. Yao, Y. G. et al. Phyleogeographic differentiation of mitochondrial DNA in Han Chinese. Am. J. Hum. Genet. 70, 635–651 (2002).
14. Tanaka, M. et al. Mitochondrial genome variation in eastern Asia and the peopling of Japan. Genome Res. 14, 1832–1850 (2004).
15. Malhi, R. S. et al. Mitochondrial haplogroup M discovered in prehistoric North Americans. J. Archaeol. Sci. 34, 642–648 (2007).
16. Li, Y. et al. River valleys shaped the maternal genetic landscape of Han Chinese. Mol. Biol. Evol. 36, 1643–1652 (2019).
17. Fu, Q. et al. DNA analysis of an early modern human from Tianyuan cave, China. Proc. Natl. Acad. Sci. U.S.A. 110, 2223–2227 (2013).
18. Yang, M. A. et al. 40,000-year-old individual from Asia provides insight into early population structure in Eurasia. Curr. Biol. 27, 3202–3208.e9 (2017).
19. Shinozuka, K. & Kanai, S. Intracemetery genetic analysis at the Nakazuma Jomon site in Japan by mitochondrial DNA sequencing. Anthropol. Sci. 107, 129–140 (1999).
20. Adachi, N. et al. Mitochondrial DNA analysis of Hokkaido Jomon skeletal: Remnants of archaic maternal lineages at the southwestern edge of former Beringia. Am. J. Phys. Anthropol. 146, 346–360 (2011).
21. Park, J. et al. A abrupt Holocene climate shifts in coastal East Asia, including the 8.2 ka, 4.2 ka, and 2.8 ka BP events, and societal responses on the Korean peninsula. Sci. Rep. 9, 10806 (2019).
22. Gutaker, R. M. et al. Genomic history and ecology of the geographic spread of rice. Nat. Plants 6, 492–502 (2020).
23. Fuji, S. Rethinking the iron historical view in the Yayoi culture. Bull. Natl. Museum Ipn. Hist. 185, 155–182 (2014).
24. Lipson, M. et al. Parallel Palaeogenomic transects reveal complex genetic history of early European farmers. Nature 551, 368–372 (2017).
25. Mathieson, I. et al. The genomic history of Southeastern Europe. Nature 555, 197–203 (2018).
26. Øversti, S. et al. Human mitochondrial DNA lineages in iron-age Fennoscandia suggest incipient admixture and eastern introduction of farming-related maternal ancestry. Sci. Rep. 9, 16883 (2019).
27. Baba, H & Endo, B. Postcranial skeleton of the Minatogawa Man. In: Suzuki, H. & Hanihara, K. (eds.), The Minatogawa Man - the Upper Pleistocene Man from the island of Okinawa. University. Mus., Univ. Tokyo Bull. 19, 61–195 (1982).
28. Kaffu, Y. & Fujita, M. Fossil record of early modern humans in east Asia. Quatern. Int. 248, 2–11 (2012).
29. Kihana, M. et al. Emulsion PCR-coupled target enrichment: An effective fishing method for high-throughput sequencing of poorly preserved ancient DNA. Gene 528, 347–351 (2015).
30. Mizuno, F. et al. A study of 6,300-year-old Jomon human remains in Japan using complete mitogenome sequences obtained by next-generation sequencing. Am. Hum. Biol. 47, 555–559 (2020).
31. Chen, S. et al. Fastp: An ultra-fast all-in-one FASTQ Preprocessor. Bioinformatics 34, 1884–1890 (2018).
32. Andrews, R. M. et al. Reanalysis and revision of the Cambridge reference sequence for human mitochondrial DNA. Nat. Genet. 23, 147 (1999).
33. Li, H & Durbin, R. Fast and accurate short read alignment with Burrows-Wheeler transform. Bioinformatics 25, 1754–1760 (2009).
34. Schubert, M. et al. Improving ancient DNA read mapping against modern reference genomes. BMC Genom. 13, 178 (2012).
36. Li, H. et al. The sequence alignment/map format and SAMtools. *Bioinformatics* **25**, 2078–2079 (2009).
37. Jun, G. et al. An efficient and scalable analysis framework for variant extraction and refinement from population-scale DNA sequence data. *Genome Res.* **25**, 918–925 (2015).
38. Ishiya, K. et al. MitoIMP: A computational framework for imputation of missing data in low-coverage human mitochondrial genome. *Bioinform. Biol. Insights* **13**, 1177932219783884 (2019).
39. Ishiya, K. & Ueda, S. MitoSuite: A graphical tool for human mitochondrial genome profiling in massive parallel sequencing. *PeerJ* **5**, e3406 (2017).
40. Weissensteiner, H. et al. HaploGrep 2: Mitochondrial haplogroup classification in the era of high-throughput sequencing. *Nucleic Acids Res.* **44**, W58–W63 (2016).
41. Thorvaldsdóttir, H. et al. Integrative genomics viewer (IGV): High-performance genomics data visualization and exploration. *Brief. Bioinform.* **14**, 178–192 (2013).
42. Van Oven, M. PhyloTree build 17: Growing the human mitochondrial DNA tree. *Forens. Sci. Int.* **5**, e392–e394 (2015).
43. Ronquist, F. et al. MrBayes 3.2: Efficient Bayesian phylogenetic inference and model choice across a large model space. *Syst. Biol.* **61**, 539–542 (2012).
44. Kumar, S. et al. MEGA X: Molecular evolutionary genetics analysis across computing platforms. *Mol. Biol. Evol.* **35**, 1547–1549 (2018).
45. Katoh, K. et al. MAFFT: A novel method for rapid multiple sequence alignment based on fast Fourier transform. *Nucleic Acids Res.* **30**, 3059–3066 (2002).
46. R Core Team. R: A language and environment for statistical computing. *R* Foundation for Statistical Computing, Vienna, Austria. URL https://www.R-project.org/ (2020).
47. Bandelt, H. et al. Median-joining networks for inferring intraspecific phylogenies. *Mol. Biol. Evol.* **16**, 37–48 (1999).
48. Leigh, J. W. & Bryant, D. POPART: Full-feature software for haplotype network construction. *Methods Ecol. Evol.* **6**, 1110–1116 (2015).
49. Bouckaert, R. et al. BEAST 2.5: An advanced software platform for Bayesian evolutionary analysis. *PLoS Comput. Biol.* **15**, e1006650 (2019).
50. Soares, P. et al. Correcting for purifying selection: An improved human mitochondrial molecular clock. *Am. J. Hum. Genet.* **84**, 740–759 (2009).
51. Kanzawa-Kiriyama, H. et al. Late Jomon male and female genome sequences from the Funadomari site in Hokkaido, Japan. *Anthropol. Sci.* **127**, 83–108 (2019).
52. McColl, H. et al. The prehistoric peopling of southeast Asia. *Science* **361**, 88–92 (2018).
53. Reuter, H. I. A. et al. An evaluation of void filling interpolation methods for SRTM data. *Int. J. Geogr. Inf. Sci.* **21**, 983–1008 (2007).

Acknowledgements
We would like to thank Koji Ishiya for processing and curating of NGS data, Masaki Fujita for valuable discussions, Katherine Hampson for English expressions, Koichi Matsuo for bone anatomy, Oki Nakamura and Kou-saku Matsumoto for drawing Fig. 1. We also thank Aiko Saso and Gen Suwa for access to the samples, Hideaki Kanzawa-Kiriyama for data of Funadomari, Hiroki Ota and Takashi Gakuhari for data of Ikawazu. This work was supported by Grant-in-Aid for JSPS KAKENHI Grant Number 17K07588, 19H05737, 20H01376, 20K15884 to F. Mi., 18H05511 (to J.G.), 17K07255 (to K.H.), and 17H00939 (to S.U.). This work was supported by The SOKENDAI Advanced Sciences Synergy Program (SASSP) (to J.G.).

Author contributions
F.Mi., J.G., M.K., L.W., K.K., and S.U. designed this study and performed research. H.B., Y.T., O.K., M.M., and T.M. provided ancient samples and assembled archaeological and anthropological information. F.Mi. M.H., L.W., K.K., and S.U. performed ancient laboratory work. F.Ma. and K.H. provided present-day Japanese data and analyses genetic data. F.Mi., J.G., M.K., H.B., K.H., and S.U. wrote the manuscript. All authors reviewed the manuscript.

Competing interests
The authors declare no competing interests.

Additional information

Supplementary Information The online version contains supplementary material available at https://doi.org/10.1038/s41598-021-91357-2.

Correspondence and requests for materials should be addressed to F.Mi., J.G. or L.W.

Reprints and permissions information is available at www.nature.com/reprints.

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

© The Author(s) 2021