nature communications



Article

https://doi.org/10.1038/s41467-024-54052-0

Genetic legacy of ancient hunter-gatherer Jomon in Japanese populations

Received: 6 December 2023

Accepted: 30 October 2024

Published online: 12 November 2024

Check for updates

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The tripartite ancestral structure is a recently proposed model for the genetic origin of modern Japanese, comprising indigenous Jomon hunter-gatherers and two additional continental ancestors from Northeast Asia and East Asia. To investigate the impact of the tripartite structure on genetic and phenotypic variation today, we conducted biobank-scale analyses by merging Biobank Japan (BBJ; n = 171,287) with ancient Japanese and Eurasian genomes (n = 22). We demonstrate the applicability of the tripartite model to Japanese populations throughout the archipelago, with an extremely strong correlation between Jomon ancestry and genomic variation among individuals. We also find that the genetic legacy of Jomon ancestry underlies an elevated body mass index (BMI). Genome-wide association analysis with rigorous adjustments for geographical and ancestral substructures identifies 132 variants that are informative for predicting individual Jomon ancestry. This prediction model is validated using independent Japanese cohorts (Nagahama cohort, n = 2993; the second cohort of BBJ, n = 72,695). We further confirm the phenotypic association between Jomon ancestry and BMI using East Asian individuals from UK Biobank (n = 2286). Our extensive analysis of ancient and modern genomes, involving over 250,000 participants, provides valuable insights into the genetic legacy of ancient hunter-gatherers in contemporary populations.

Anatomically modern humans, who originated in Africa, began a global dispersal 50–60 thousand years ago (kya) through a series of migrations, settlements, and admixture^{1,2}. The arrival in East Asia can be traced back to at least 40–50 kya, as they gradually spread across the region³. A crucial event in human history was the encounter between indigenous hunter-gatherers and immigrant farmers, subsequently leading to significant shifts in lifestyle⁴. While this transition to farming occurred on a global scale, its timing and process varied from one

region to another; the agricultural revolution in continental East Eurasia dates back to around 10 kya^s.

Archeological evidence suggests that humans occupied the Japanese archipelago, an insular region of East Eurasia, as early as 38 kya, during the Paleolithic period⁶. Still, our understanding of their ancestral connection to modern populations is limited due to the scarcity of ancient DNA data⁷. Among the well-studied ancestral groups in Japan are the Jomon, a cultural group of

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hunter-gatherer-fishers who inhabited the archipelago as far back as 16.5 kya^{8,9}. The Jomon are notable for their pioneering use of pottery, which is among the earliest instances in the world¹⁰. The Jomon period lasted until -3 kya, when immigrants from the continent introduced rice cultivation during the Yayoi period that spanned from 3 to 1.7 kya. This agricultural revolution prompted sociopolitical developments, leading to the establishment of the Japanese state in the Kofun period, which began around 1.7 kya and endured for 200–300 years¹¹.

A long-standing model of the origin of Japanese populations is a dual-ancestral structure^{12,13}. This model states that modern Japanese people are a mixture of indigenous Iomon hunter-gatherers from Southeast Asia and immigrant farmers from Northeast Asia. However, a recent ancient DNA study provides compelling evidence that the genetic origin of Japanese populations consists of three distinct ancestors (i.e., tripartite ancestral structure): (1) ancient huntergatherer Jomon, (2) Northeast Asian ancestry introduced during the agrarian phase, the Yayoi period, and (3) East Asian ancestry brought in the state formation phase, the Kofun period¹⁴. This tripartite structure, which was established during the Kofun period, persists in contemporary populations¹⁴⁻¹⁶. Japanese populations are well characterized with their north-to-south gradient of genomic variation¹⁷⁻¹⁹. However, the applicability of the tripartite model and variability of the three distinct ancestral components throughout the archipelago remains unknown due to limitations in samples and geographic representation of modern individuals used in previous studies^{14,15}. It is thus crucial to model this ancestral structure using comprehensive, population-level genomic data.

Recent advancements in paleogenomics have made it possible not only to identify diverse genetic ancestors^{20–23}, both locally and globally, but also to uncover their impact on health and disease in contemporary populations, such as the exacerbation and resistance of coronavirus disease 2019 through Neanderthal-introgressed genes^{24,25} or the genetic predisposition to multiple sclerosis brought by Steppe ancestry²⁶. However, the extent to which the human past has shaped phenotypic variation today remains poorly understood, especially in non-European contexts²⁷. Here, we focus on the Japanese archipelago, where the hunting-gathering Jomon period lasted for more than 10,000 years in the insular environment, and where the genetic remnants of this ancient hunter-gatherer ancestry persist in present-day populations.

In this study, we present an integrated analysis of ancient human genomes (n = 22) and modern Japanese genomes, utilizing the first cohort of Biobank Japan (BBJ, n = 171,287), one of the largest population-based cohorts in a non-European population, encompassing participants from all regions of the archipelago. Our approaches involve five key steps: First, we evaluate the applicability of the tripartite ancestral model to different subpopulations defined by their geography or genetic clusters. Second, we quantify the impact of Jomon ancestry on phenotypic variation among Japanese individuals. Third, we identify Jomon-related genetic variants by employing a genome-wide association study (GWAS) method, with robust adjustment for geographic and genetic substructures, and control for genomic inflation of test statistics. Fourth, we demonstrate the predictive power of Jomon-related variants in estimating an individual's Jomon ancestry using independent Japanese cohorts (the Nagahama cohort n = 2993 and the second cohort of BBJ n = 72,695). Finally, we apply the Jomon predictive model to East Asian (EAS) individuals within UK Biobank and replicate the phenotypic impact of Jomon ancestry (UKB EAS, n = 2286).

Our study provides a comprehensive understanding of the genetic legacy of ancient hunter-gatherers in contemporary descendants throughout the Japanese archipelago and highlights its impact on phenotypic variation today.

Results

Inference for the tripartite ancestral structure in Biobank Japan dataset

To assess the fit of the tripartite model in contemporary populations. we used BBI GWAS data, comprising participants from hospitals across seven geographic regions in Japan^{28,29} (Fig. 1a and Supplementary Data 1). The total number of participants is 171,287, with regional distribution from northeast to southwest throughout the archipelago as follows: Hokkaido = 7955, Tohoku = 11,013, Kanto-Koshinetsu = 94,981, Chubu-Hokuriku = 9489, Kinki = 25,200, Kvushu = 15.962. Okinawa = 5804. Our PCA defines distinct clusters (Fig. 1b), with a separation of the Ryukyu cluster, primarily including individuals from Okinawa, from the rest of the populations as reported in previous studies^{17,18}. There are also regional clusters observable from Tohoku, Kanto-Koshinetsu, Kinki, and Kyushu respectively. Based on the PCA results, we define five distinct genetic clusters in the Japanese populations: EastAsian admix (EA admix; n = 1019), Mainland (n = 159,642), Ryukyu admix (n = 640), Ryukyu (n = 9847), and Hokkaido sub (n=139) (Fig. 1c). Population stratification is further evident even within the Ryukyu Islands, reflecting their geographic affinities in this local insular context (Fig. 1d, e, Yakushima; n = 431, Amami; n = 1531, Kikai; n = 561, Okinoerabu; n = 845, Tokunoshima; n = 476, Yoron; n = 167, Okinawa; n = 4795, Miyako; n = 827).

We then merged this diverse set of Japanese individuals with ancient genomic data from Japan and the Eurasian continent¹⁴. To find the sites that are present in both the array-typing genome data from BBJ and the pseudo-diploid data from ancient humans, we converted the highly accurate BBJ imputed dosage data ($Rsq \ge 0.7$) into genotype data (details in "Methods"). This conversion resulted in 2,038,260 shared variants (n = 171,287 of BBJ and n = 22 of ancient genomes). We subsequently applied qpAdm in AdmixTools^{30,31}, based on a set of source populations defined in a previous study¹⁴ (see Supplemental Data 2), to evaluate the fit of admixture models and to estimate admixture proportions at both population and individual levels in the biobank. To comprehensively capture the geographic and genetic diversity of Japanese populations, we fitted the tripartite model to geographically-defined populations throughout the Japanese archipelago and the Ryukyu Islands (Fig. 1b, e), five genetically-defined populations (Fig. 1c), as well as the entire BBJ dataset. This analysis demonstrates that the tripartite structure provides a better fit for all populations, both at broad and local scales, when compared to all possible dual-ancestral structure models (Supplementary Data 3). The only exception is EA_admix, where the population can be sufficiently explained by a two-way admixture between Northeast Asian and East Asian ancestry.

In the whole BBJ dataset, the proportions of the three distinct ancestral components closely align with those reported in the previous study (Jomon: 12.4%, Northeast Asia: 21.2%, and East Asia: 66.4%)¹⁴. However, Jomon ancestry exhibits regional variation, ranging from 9.8% in Kinki to 26.1% in Okinawa (Fig. 2a). Within the Ryukyu Islands, there is an elevated level of Jomon ancestry, with the highest proportion observed on Yoron Island (Fig. 2b). Jomon ancestry is even higher in one of the genetically-defined populations, Hokkaido sub (31.6%, Fig. 2c and Supplementary Data 3), which primarily includes a subset of individuals from Hokkaido. In contrast, EA_admix, possibly representing continental individuals from East Asia, has very little Jomon ancestry; this may explain why the admixture model without Jomon ancestry is preferred. The Mainland cluster mirrors the proportion of the entire BBJ as it includes the majority of the samples in the data (159,642 out of 171,287). Even when we separate individuals from this cluster based on their geographic origins (i.e., where the sample collection took place), this proportion remains relatively consistent across different regions (Supplementary Fig. 1). The Ryukyu cluster represents the ancestral composition from Okinawa, while the

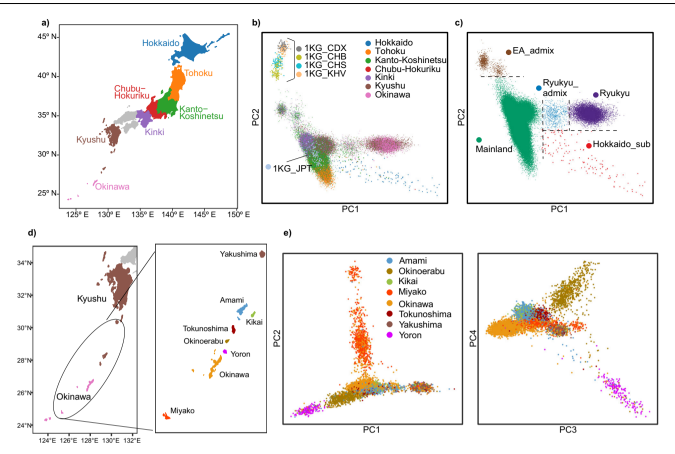


Fig. 1 | **Population structure of Biobank Japan. a** Seven geographic regions, each represented by a different color, indicate the locations where participants were registered at local hospitals. **b** A scatter plot of PCA for BBJ participants with 1KG EAS samples. The colors of the BBJ participants correspond to those used in (a). **c** Clustering results of BBJ participants based on PCA. **d** Geographic locations of the Ryukyu Islands. **e** Scatter plots of PCA for BBJ participants from the Ryukyu Islands.

The individuals are colored differently according to the locations of their registered hospitals. In (**a**, **d**) the map of Japan is drawn using the R package "jpndistrict" (https://github.com/uribo/jpndistrict). PC Principal component, 1KG 1000 Genomes Project, JPT Japanese in Tokyo, CDX Chinese Dai in Xishuangbanna, CHB Han Chinese in Beijing, CHS Han Chinese South, KHV Kinh in Ho Chi Minh City, NEA Northeast Asian, EA East Asian.

proportions in the Ryukyu_admix cluster fall between those of the Mainland and Ryukyu clusters (Supplementary Data 3).

Next, we asked whether Jomon ancestry is uniquely present in the Japanese populations or whether this ancestry is also observable in continental populations using f_4 -statistics with the form of f_4 (Mbuti, Jomon; Han, X). Our target populations include those in the Simon Genome Diversity Project (SGDP) panel, the 1000 Genomes Project (1KG), and the subpopulations within BBJ. As shown in previous studies^{8,9,14}, there is no extra genetic affinity between the Jomon and any of the populations tested, except for Ulchi in the SGDP or the East Asians (EAS) within the 1KG (Supplementary Fig. 2). Among the 1KG EAS population, only Japanese in Tokyo (JPT) show a significant affinity to the Jomon. Within the BBJ participants, the Hokkaido_sub and Ryukyu subpopulations exhibit an extremely strong affinity with the Jomon as consistent with the higher Jomon ancestry in our admixture modeling (Fig. 1e), as well as previous studies^{8,9}.

Overall, our analysis provides detailed pictures of regional variation in the tripartite ancestral structures throughout the archipelago.

Variation in the tripartite structure among individuals

Our admixture modeling reveals that two subpopulations, Ryukyu and Hokkaido_sub, have higher proportions of Jomon ancestry compared to the other subpopulations (Fig. 2). To visualize the genetic distance between ancient and present-day populations, we projected the ancient or modern individuals, who represent the three distinct ancestors underlying the tripartite structure of modern Japanese populations, along with additional ancient Japanese individuals (i.e.,

Yayoi and Kofun) onto the PCA plots. While the Kofun are included within variation in the present-day populations, the Jomon are clustered in a position extending from the Ryukyu and Hokkaido_sub subpopulations. The two individuals from the Yayoi period, who are morphologically considered as the Jomon³² but genetically admixed between the Jomon and continental ancestry¹⁴, are positioned between the Jomon and the present-day populations (Supplementary Fig. 3a).

Given that the population-based admixture modeling only represents an averaged pattern in a group of individuals, we then apply the tripartite model to each participant in BBJ individually. This model fits 154,339 out of the 171,287 individuals (90.1%), with varying proportions of three ancestral components. The proportions of these three ancestors are negatively correlated with each other (Supplementary Fig. 3b, c), supporting a previously proposed scenario that two continental ancestors, Northeast Asian and East Asian, are likely to have arrived in the archipelago independently¹⁴. Approximately 5% of the individuals did not conform to the tripartite model (8932 individuals), showing a better fit for a two-way admixture involving either Jomon and East Asian ancestry or Northeast Asian and East Asian ancestry. However, 28 individuals were excluded from subsequent analysis due to insufficient support for the dual ancestry models over the tripartite model, as indicated by nested p < 0.05. There was also a minor exception, where a model of East Asian ancestry alone, rather than any dual structure models, was preferred in ten individuals (Supplementary Fig. 4). It's important to note that none of the models could adequately fit the remaining individuals, comprising nearly 5% (7962 individuals). This is likely due to the use of a tail probability

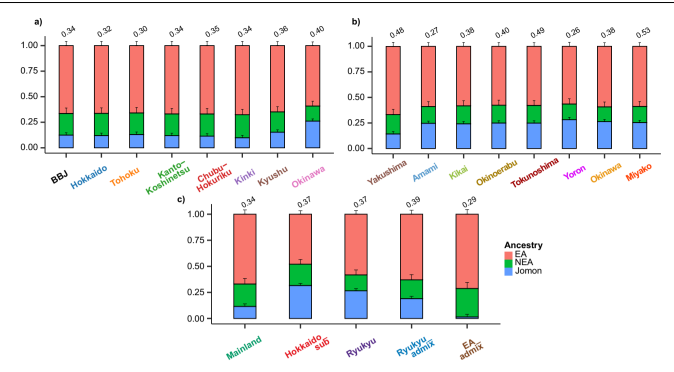


Fig. 2 | **Variation in tripartite ancestry structures across groups of Biobank Japan participants.** Bar plots show the proportions of three distinct ancestors. **a** Populations are defined either as a group of the entire BBJ samples or by the regions where participants were registered (Hokkaido = 7955, Tohoku = 11,013, Kanto-Koshinetsu = 94,981, Chubu-Hokuriku = 9489, Kinki = 25,200, Kyushu = 15,962, and Okinawa = 5804). **b** Individuals recruited from the Ryukyu Islands are grouped by their specific islands (Yakushima = 431, Amami = 1531, Kikai = 561, Okinoerabu = 845, Tokunoshima = 476, Yoron = 167, Okinawa = 4795,

and Miyako = 827). c The BBJ samples are split into five different insular populations based on their PCA clusters shown in Fig. 1 (Mainland = 159,642, Hokkaido_sub = 139, Ryukyu = 9847, Ryukyu_admix = 640, and EA_admix = 1019). Proportions in the bar plots represent the mean values of three ancestral components estimated using qpAdm, with error bars indicating standard errors. Values at the top of each bar indicate the tail probabilities of the tripartite model for each group. NEA Northeast Asian, EA East Asian.

cut-off of 5%, which accounts for the inherent variability in the data, as a defined proportion of all tests would be expected to deviate from the models tested.

By incorporating the proportion of Jomon ancestry onto the PCA plots, it becomes evident that there is a noticeable gradient in the Jomon proportion along the first and second principal components (Fig. 3a). Indeed, the Jomon proportion has a remarkably strong correlation with the first two principal components (Fig. 3b; |R| = 0.61 and 0.09 for PC1 and PC2, respectively; see Supplementary Data 4). Similar patterns of correlation are also observable between Northeast Asian or East Asian ancestry and the PCs; however, the strength of correlation is not as pronounced as that observed for Jomon (|R| = 0.14 for Northeast Asian and PC1; |R| = 0.26 for East Asian and PC1). The correlation between Jomon ancestry and PCs is still evident even upon focusing solely on individuals from the Mainland or Ryukyu cluster (Supplementary Fig. 5). These findings strongly indicate that Jomon ancestry has a critical role in shaping genomic variation among Japanese individuals at the level of PCA. Furthermore, these individual-based estimates not only mirror the population-based patterns in terms of their means but also highlight significant variation in Jomon ancestry across the Japanese archipelago (Fig. 3c, d).

Impacts of Jomon ancestry on phenotypic variation in Japanese populations

Next, we explored whether or not Jomon ancestry has any phenotypic impact on present-day populations. Among 163,243 individuals from BBJ, who were successfully modeled for their genetic ancestry, the average proportion of the Jomon component is $12.5 \pm 6.3\%$ (mean \pm SD). There is no substantial difference in the Jomon proportion between ages or between genders (Supplementary Fig. 6).

We then tested associations of Iomon ancestry with 80 different complex traits by making robust adjustments for genetic and geographic subpopulations (see "Methods" and Supplementary Data 5). To account for type I error, we set a threshold of statistical significance at $P < 0.05/80 = 6.3 \times 10^{-4}$, based on the Bonferroni correction. We confirm that this threshold was calibrated by simulating 10 dummy heritable phenotypes as negative controls ("Methods"). Our analysis with all BBJ participants identifies significant associations with an increase in body mass index (BMI) (Fig. 4a and Supplementary Data 6; Beta = 0.012, Standard error [SE] = 0.003, $P = 3.0 \times 10^{-5}$). However, it is important to note that regional disparity, such as the higher BMI in Okinawa compared to the other regions³³, may potentially confound this association, even when adjusting for geographic factors as covariates. To address this concern, we confined our analysis to individuals included in the Mainland cluster (n = 152,148; see Fig. 1c). The association with BMI remains statistically significant (Fig. 4b and Supplementary Data 7; Beta = 0.012, SE = 0.003, $P = 7.9 \times 10^{-5}$). We further confirmed the significance of the association with BMI regardless of sex or age (see Supplementary Fig. 7).

While it may be argued that our approach is conservative, we prioritize addressing any potential inflation of association signals stemming from population stratification³⁴. Therefore, it is crucial to account for its effect by incorporating PCs as covariates, despite the stratification originating from variation in Jomon ancestry across individuals to some degree. To mitigate this issue, we also employed an alternative methodology where phenotypes underwent correction by regressing their measures with all covariates, including PCs, before being tested for associations with Jomon ancestry. We observed a significant association in BMI, while none of the other traits reached statistical significance (Supplementary Fig. 8, and Supplementary

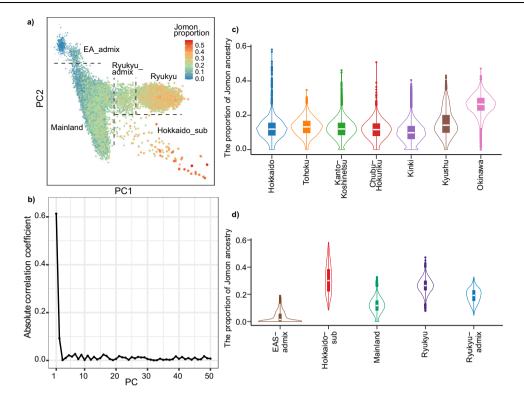


Fig. 3 | **Genetic legacy of Jomon ancestry throughout Japanese populations.** Proportions of Jomon ancestry are estimated through admixture modeling under models optimal for each individual (n = 163,243). a Projection of the Jomon proportions onto the PCA plots. b Absolute values of the correlation coefficients between the Jomon proportions and PCs. The correlation coefficient is calculated using Pearson's method. c Boxplots representing variation in the Jomon proportion

across BBJ participants within each of their registered regions. **d** Boxplots representing variation in the Jomon proportion across the participants within each of different genetic clusters defined by PCA. In (**c**, **d**), boxes denote the interquartile range (IQR) and the median is shown as white horizontal bars; whiskers extend to 1.5 times the IQR; outliers are shown as individual points. PC Principal component, EA East Asian.

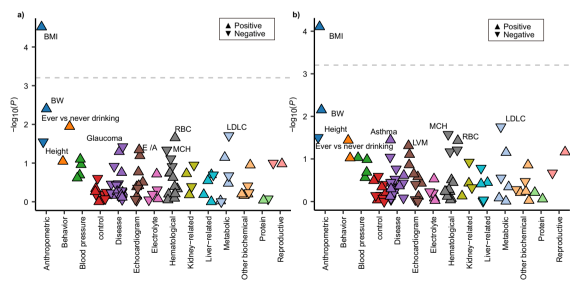


Fig. 4 | **Associations of Jomon ancestry with 80 complex traits.** Associations are tested using a generalized linear model in (**a**) the entire BBJ participants (n = 163,243) and (**b**) the participants only in the Mainland cluster (n = 152,148). The quantitative traits are modeled by linear regression, while the binary traits are analyzed with logistic regression. The direction of the triangle corresponds to the sign of the beta coefficient for each trait. The gray dashed line represents the statistical significance based on the Bonferroni correction ($P < 0.05/80 = 6.3 \times 10^{-4}$).

Control indicates 10 dummy phenotypes. All traits labeled on the plots have nominal significance with P < 0.05. Details of statistical test results are presented in Supplementary Data 6 and 7. P-values are computed by linear regression or logistic regression. All statistical tests are two-sided and unadjusted for multiple comparisons. BMI Body mass index, BW Body weight, LVM Left ventricular mass, E/A E/A ratio, RBC Red blood cell count, MCH Mean corpuscular volume, LDLC low-density lipoprotein cholesterol.

Data 8 and 9). Taking all results together, our analyses consistently demonstrate the robustness of the BMI signals, regardless of the methods used for correcting population stratification (see Fig. 4 and Supplementary Fig. 8).

To assess the impact of Jomon ancestry on BMI, we incorporated the proportion of Jomon ancestry as a covariate in our GWAS for BMI (Supplementary Fig. 9a and Supplementary Data 11). While the majority of lead SNPs remain consistent, we observed a reduction in the effect of Jomon ancestry on polygenic scores (PGS) of BMI (Supplementary Fig. 9b). It is worth noting that this effect persists to some extent, indicating a functional correlation between Jomon ancestry and BMI-PGS. Still, the PGS tends to be inflated when Jomon ancestry is not accounted for in GWAS (Supplementary Fig. 9c), with the degree of inflation weakly correlating with the proportion of Jomon ancestry (Supplementary Fig. 9d). These findings suggest that the effect sizes estimated from GWAS without including Jomon ancestry as a covariate may be biased due to residual confounding.

Our analysis also includes height, which has been shown to exhibit a north-to-south gradient in Japanese populations 35 . The association of Jomon ancestry with the decrease of height is observable only if PCs are not accounted for in the test (Supplementary Fig. 10 and Supplementary Data 10), indicating that this association can be confounded by population stratification. Overall, these results suggest that the genetic legacy of the ancient hunter-gatherer Jomon significantly influences BMI across populations today, regardless of geographic differences, which may consequently contribute to an increased risk of obesity. Additionally, we tested the effect of the use of the Jomon proportion on the predictive power of the polygenic score (PGS) for BMI (see "Method"). The incremental predictive performance is limited with a magnitude of -2.8×10^{-3} .

Identification of genetic variants underlying variation in Jomon ancestry

We performed a genome-wide investigation of variants associated with variation in Iomon ancestry among individuals. In this analysis, an individual proportion of Iomon ancestry is considered as a proxy phenotype. However, a conventional null hypothesis, as used in standard genotypephenotype association studies, is not directly applicable to this phenotype since it is inferred from the genomic data. To ensure robust correction for genomic and geographic subpopulations and to control for genomic inflation of the test statistics, we adopted a double genomic control correction method and a mixed linear model (MLM) approach that includes PCs as covariates³⁶. Furthermore, we conducted a metaanalysis of the association tests for individuals within the Mainland and Ryukyu clusters separately, as the Jomon proportions significantly differ between these groups (Fig. 3d). To gain biological insights into the genetic signals enriched for Jomon ancestry, we conducted stratified linkage disequilibrium score regression (S-LDSC) on the genome-wide meta-analyzed result. Our S-LDSC identified a significant enrichment of the heritability in skeletal muscle cells across major cell groups (Fig. 5a).

Based on our association analysis, we classify a variant as Jomon-related if it has a positive *Z*-score and reaches a genome-wide significance ($P < 5.0 \times 10^{-8}$) after rigorous control measures. This results in 132 independent variants identified from LD clumping (details in "Methods"; Table 1 and Supplementary Data 12). To explore evolutionary contexts of these Jomon-related variants, we examined haplotype structures of the genomic regions containing Jomon-related variants. Contrasting these haplotype structures with those of allele frequency-matched non-Jomon-related variants (i.e., variants with P > 0.05), we find that the Jomon-related variants exhibit significantly longer haplotypes than non-Jomon-related variants ($P = 3.1 \times 10^{-32}$) (Fig. 5b, c and Supplementary Data 12). This strongly supports a Jomon origin for these extended haplotypes, supported by strong linkage disequilibrium throughout the genome, possibly due to the high genetic homogeneity and small effective population size (-1000)

within the Jomon population^{14,37}. Furthermore, the persistence of these long Jomon-derived haplotypes in the contemporary populations can be attributed to the relatively recent admixture with the continental ancestors¹⁴, coupled with insufficient time for recombination to break apart the haplotypes. Leveraging selection scan results based on singleton density score (SDS)¹⁷, which detects selection over approximately the past 100 generations³⁸, we further observe an enrichment of selection signals in the Jomon-derived haplotypes (P = 0.008). These results support the idea that the Jomon-related variants can serve as markers for quantifying Jomon ancestry, tag Jomon-derived haplotypes, and potentially have been subject to recent selective pressures.

These Jomon-related variants are significantly more frequent in JPT compared to the other East Asian populations within the 1KG dataset (Table 1, Supplementary Fig. 11, and Supplementary Data 12). Even within the BBJ population, the Ryukyu group exhibits higher frequencies of these variants than the mainland group, supporting their strong linkage with the Jomon-derived segments. We measured the fixation index (F_{ST}) for the 132 Jomon-related variants in order to assess their specificity in the Japanese population. While there is no major difference in F_{ST} across the East Asian populations, rs536618 at 4p12 and rs2871660 at 1q31 show relatively high F_{ST} values in the Japanese population (Supplementary Figs. 12 and 13; Supplementary Data 13). Notably, the top variant, rs13017060, is located in the intron region of the NBAS gene, exhibiting pleiotropic effects on the increase of BMI and body weight in modern Japanese populations²⁹. It also serves as an expression quantitative trait locus (eQTL) for NBAS in heart muscle, as evidenced by GTEx data³⁹. Moreover, the same is associated with increased leg fat mass in the UK Biobank^{40,41}. These findings provide further evidence supporting the significant association between Jomon ancestry and BMI.

Validation of the tripartite model and Jomon-related variants using independent Japanese cohorts

To validate our admixture modeling of the tripartite structure, we utilized two independent Japanese cohorts, the Nagahama cohort and the second cohort of BBJ (BBJ-2nd). The Nagahama cohort includes participants from Nagahama city, Shiga Prefecture, within the Kinki area of Japan (comprising Nagahama A with n=1549 and Nagahama B with n=1444)⁴². BBJ-2nd is an additional independent cohort, which is distinct from the first BBJ cohort, representing various regions across the Japanese archipelago (n=72,695).

Our validation process consists of two different approaches: (i) estimating the proportion of Jomon ancestry at the individual level and (ii) deriving the Jomon ancestry prediction score as a form of PGS. The PCA of BBJ-2nd confirms the significant correlations of Jomon ancestry with the first two PCs (Fig. 6a; Supplementary Data 4), as observed in the first BBJ cohort (Fig. 3a). We then derived the prediction scores of Jomon ancestry using 132 Jomon-related variants identified from the first BBJ cohort. Under the robust adjustment for population stratification, the variance of Jomon ancestry explained by the prediction scores are as follows: 014, 0.12, and 0.13 in Nagahama A, Nagahama B, and BBJ-2nd respectively ($P_{Nagahama}$ $_A = 2.5 \times 10^{-50}$, $P_{Nagahama}$ $_B = 1.3 \times 10^{-40}$, and $P_{BBJ-2nd} < 1.0 \times 10^{-300}$). Notably, the observed Jomon proportions gradually increase in line with the prediction scores across the different cohorts when the scores are grouped into deciles (Fig. 6b and Supplementary Fig. 14).

The predictive power of Jomon-related variants shows an increase when the p-value cut-off is relaxed from 5.0×10^{-8} to 1.0×10^{-3} , resulting in the variance of 0.27 (Nagahama A), 0.26 (Nagahama B), and 0.24 (BBJ-2nd) respectively for Jomon ancestry explained by genetic variants (Supplementary Fig. 15). However, it is crucial to note that this clumping and thresholding method has been demonstrated to potentially overfit the data, particularly when relaxing the p-value cut-offs⁴³. Therefore, we adhere to a p-value cut-off of 5×10^{-8} for predicting Jomon ancestry in subsequent analysis.

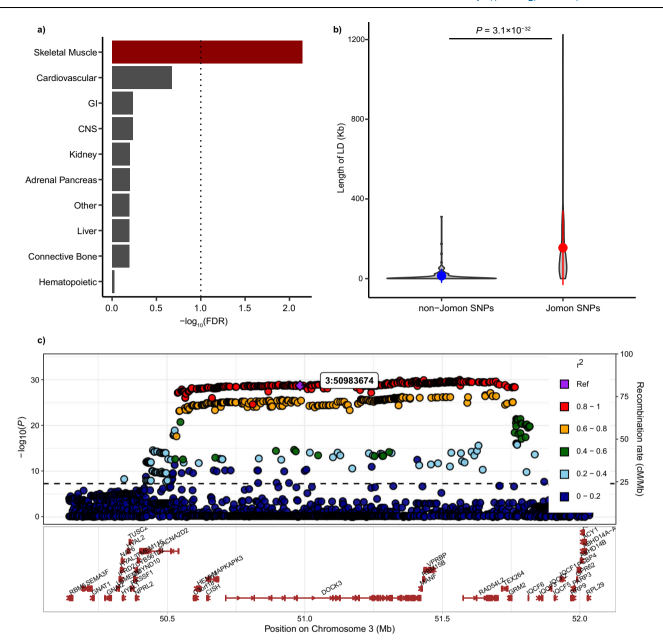


Fig. 5 | **Functional and genomic characterization of Jomon-related marker SNPs. a** Stratified LD score regression analysis is conducted with major 10 cell type groups. False discovery rate (FDR) is calculated using the Benjamini-Hochberg method. A dotted vertical line represents the FDR of 0.1. **b** Violin plots compare the lengths of regions in strong LD with each of 132 Jomon-related SNPs ($r^2 > 0.8$) to those of 132 frequency-matched non-Jomon-related SNPs. Dots represent the mean lengths, with bars showing the standard deviations. A p-value (3.1×10^{-32}) is calculated by the Wilcoxon rank sum test, reflecting a two-sided test.

c A locus plot highlights a Jomon-related SNP, which is shown as a purple diamond in the upper plot, that is linked with the longest haplotype among all 132 variants. A dashed line defines a statistical significance as a p-value of 5.0×10^{-8} . Lower plots represent the protein-coding genes, based on GENCODE, that are present in the highlighted region. The p-value is computed by inverse variance weighted meta-analysis using METAL. All statistical tests are two-sided and unadjusted for multiple comparisons. GI Gastrointestinal, CNS Central nerve system, LD Linkage disequilibrium, SNP Single nucleotide polymorphism.

Replication of the association between Jomon ancestry and BMI using East Asian individuals from UK Biobank

We sought to replicate our finding on the link between Jomon ancestry and the increase of BMI using a completely independent cohort. Our focus is on East Asian individuals within UK Biobank (UKB EAS). We first selected the individuals of UKB EAS based on the PCA plots (n = 2286; Supplementary Fig. 16). The f_4 -test, in the form of f_4 (Mbuti, Jomon; Han, UKB EAS), shows that the Jomon are symmetrically related to Han and UKB EAS (Z = 0.93). We then split UKB EAS into different groups based on their self-reported ethnic backgrounds and performed f_4 -test for each group individually (Supplementary Fig. 17). This analysis

identifies Ethnic Group (EG) 6 (i.e., the self-reported Other ethnic population, n = 541) that shows a higher genetic affinity to Jomon than Han with a *Z*-score of 5.4. None of the other EGs support a significant affinity with Jomon, including EG5, consisting of self-reported Chinese individuals (n = 1375). These results suggest that participants in EG6 carry Jomon ancestry and are a suitable subpopulation for our replication analysis.

By fitting the admixture models to the UKB EAS cohort (n = 2286), we were able to quantify Jomon ancestry for 566 individuals. The predictive power of the scores based on 132 Jomon-related variants account for -2% of the total variance in the observed Jomon proportion

able 1 | List of the top 10 Jomon-related variants ranked by statistical significance in BBJ

								Frequency	Frequency of alternative allele	ive allel	d)					F _{ST} be	tween JF	F _{ST} between JPT and EAS	AS
								BBJ		1KG pc	IKG populations	SL			Jomon				
rsID	ਨੁੱ	Chr Pos (hg19)	Ref/Alt Loci	Loci	Nearest gene	٩	PheWAS	Mainland	Ryukyu	EAS	AFR	AMR	EUR	SAS		CDX	CHB	CHS	KHV
rs28729170	_	188,505,272	C/A	1q31	PLA2G4A	8.1×10 ⁻³³		0.78	0.83	0.72	0.74	99.0	0.79	0.85	0.86	0.01	0.03	0.03	0.00
rs13017060 2	2	15,641,500	A/C	2p24	NBAS	1.7 × 10-32	BMI, Body weight	0.29	0.38	0.16	0.20	0.52	0.63	0.41	0.86	0.10	0.03	0.13	0.04
rs2645158	19	31,885,791	A/G	19q12	TSHZ3	5.4×10 ⁻³⁰	T2D	0.38	0.36	0.42	0.82	0.58	99.0	0.62		0.00	0.00	0.00	0.01
rs6446239	ო	50,983,674	2/9	3p21	роскз	2.0×10 ⁻²⁹	Height, Body weight	0.68	0.73	0.65	0.62	0.71	0.97	0.95	0.83	0.01	0.02	0.00	0.02
rs1026980	9	47,624,988	C/T	6p12	ADGRF2	3.8×10^{-26}		0.34	0.39	0.31	0.54	0.47	0.68	0.72		0.00	0.00	0.00	0.00
rs2057165	9	64,966,049	T/A	6q12	EYS	3.5×10^{25}	1	0.50	0.53	0.50	0.71	99.0	0.71	0.73	0.56	0.02	0.09	0.03	0.04
rs4302225r	2	84,050,402	C/T	2p11	SUCLG1	2.8 × 10 ⁻²³		0.50	0.54	0.50	0.39	0.58	0.72	0.62		0.00	0.00	0.00	0.00
rs4981864	14	32,086,838	G/T	14q12	NUBPL	2.1×10^{-24}		0.55	0.56	0.61	0.19	0.54	0.58	0.53	0.83	0.05	0.00	0.00	0.00
rs629577	10	38,335,178	A/G	10p11	ZNF33A	3.4×10^{-22}		0.49	0.55	0.42	0.46	0.48	0.51	0.74		0.00	0.01	0.00	0.02
rs6097031	20	51,188,901	O/0	20	ZFP64	5.2×10^{-21}		0.35	0.46	0.32	0.88	0.35	0.30	0.41	0.91	0.00	0.00	0.01	0.01
All 132 Jomon-r	elated va	riants are listed in	Supplements	ary Data 11,	with their Fsr values	shown in Sup	132 Jomon-related variants are listed in Supplementary Data 11, with their Fex values shown in Supplementary Data 12. The P-value is computed by inverse variance weighted meta-analysis using METAL. All statistical tests are two-sided and unadjusted for	alue is compute	d by inverse	variance v	veighted	neta-analy	sis usina	METAL. A	ll statistical	tests are t	wo-sided	and unadi	usted for

Ref/Alt Reference/Alternative allele (effect allele is alternative allele). Alt freq. the alternative allele frequency, BBJ Biobank Japan, 1KG 1000 Genomes Project, 72D type 2 diabetes

 $(R^2 = 0.02 \text{ and } P = 2.7 \times 10^{-4}; \text{ Fig. 7a})$. This power substantially increases when only focusing on EG 6 within UKB EAS $(R^2 = 0.06 \text{ and } P = 3.8 \times 10^{-4}; \text{ Fig. 7b})$. In contrast, this prediction becomes less effective in EG5 $(R^2 = 0.03, \text{ and } P = 0.002 \text{ Fig. 7c})$, aligning with the apparent scarcity of Jomon ancestry in this group (Supplementary Fig. 17).

Finally, we tested the effect of Jomon ancestry on BMI in UKB EAS. There is no significant association either in the entire UKB EAS or EG5 of UKB EAS (Fig. 7d; Beta = 0.11, SE = 0.58, P = 0.86 in the entire UKB EA; Beta = -0.83, SE = 1.30, P = 0.52 in EG5). However, when focusing exclusively on EG6, which exhibits a higher Jomon genetic influence, we identify a significant association between Jomon ancestry and an increase in BMI (Beta = 2.2, SE = 0.99, P = 0.03).

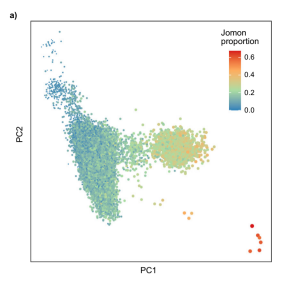
Overall, these results highlight the potential influence of Jomon ancestry on the risk of obesity in contemporary populations regardless of differences in their living environments, as observed between UK and Japan.

Discussion

Our study provides comprehensive insights into the genetic legacy of ancient hunter-gatherers in the Japanese archipelago by combining ancient human genomes from Japan and continental East Asia with modern genomic data of more than 250,000 Japanese individuals. Leveraging this extensive biobank-scale dataset, we demonstrate that the recently proposed model for the genetic origin of modern Japanese populations, known as the tripartite ancestry structure¹⁴, widely and consistently fits better than the dual structure model across the Japanese archipelago. The composition of the three distinct ancestors varies among different geographic and genetic subpopulations within Japan. Among these ancestral components, the ancient huntergatherer lomon stands out as the most influential in increasing BMI in contemporary individuals, as well as in shaping genomic variation, both at the individual and population levels. Our novel GWAS approach, coupled with rigorous controls for genomic inflation factors, effectively identifies genetic variants associated with the Jomon ancestry that modern Japanese possess. We have then pinpointed a set of 132 independent variants as genetic markers for predicting an individual's Jomon ancestry. The predictive power of these markers is validated with independent cohorts of Japanese populations. Furthermore, we successfully replicate the phenotypic impact of Jomon ancestry by studying a cohort of individuals who have Jomon components as their genetic ancestor, residing in the United Kingdom.

Our analysis presents the first in-depth characterization of the tripartite structure across the entire archipelago. It unveils substantial variation in the proportion of Jomon ancestry, mirroring the genetic ancestry continuum observed in present-day populations. This proportion is extremely high in individuals from the Ryukyu and Hokkaido_sub clusters, as indicated in previous studies^{8,9}. The continental ancestor (i.e., Northeast Asian and East Asian) is also observable in individuals from Okinawa. However, it is important to note that these continental components were not directly introduced from the continent but by immigrants from the main islands of Japan, who already possessed the tripartite ancestor¹⁵. This migration has been estimated to have occurred around the eleventh century AD, marking the end of the prehistoric period in the area. Until this transition, it is widely accepted that people with Jomon-like genetic characteristics continued to inhabit the region for at least several thousand years⁴⁴. Therefore, the elevated levels of Jomon ancestry in Okinawa can be attributed to this historical event.

In contrast to the Ryukyu or Hokkaido_sub clusters, the Kinki region exhibits a relatively low level of Jomon ancestry. Historical records indicate the enduring presence of governmental center in this area, implying more frequent interactions with people from the Asian continent than in other regions⁴⁵. Our biobank-scale analysis provides a refined picture of the genetic makeup of people throughout the lapanese archipelago both at broad and local scales.



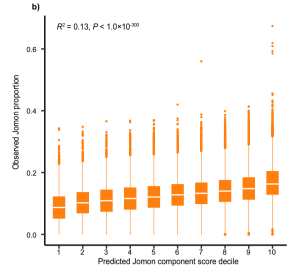


Fig. 6 | Validation of the genetic legacy of Jomon ancestry and the predictive power of 132 Jomon-related variants using an independent cohort of BBJ-2nd. a The PCA plots include all individuals from BBJ-2nd (n = 68,632), with their respective Jomon proportions overlaid. b The Jomon component prediction scores measured from all individuals within the cohort (n = 72,695) are split into the deciles. The boxplot shows the distribution of observed Jomon proportions in each

decile of the predicted scores. R^2 indicates the squared value of the Pearson's correlation coefficient between the prediction score and the residuals of the Jomon proportion regressed out with 10 PCs. P-value reflects a two-sided test. In (\mathbf{b}), boxes denote the interquartile range (IQR) and the median is shown as white horizontal bars; whiskers extend to 1.5 times the IQR; outliers are shown as individual points. PC Principal component.

The finding on the significant association between increased Jomon ancestry and the increase in BMI aligns with the observations that people from Okinawa have not only elevated levels of BMI but also a higher propensity for obesity compared to those from the main island³³. The obesogenic environment in present-day populations is certainly a substantial contributor to the ongoing obesity epidemics. Nonetheless, we identify this phenotypic impact of Jomon ancestry in the populations from Japan, as well as the UK, where individuals are exposed to varying degrees of obesogenic environments. Therefore, the ancient hunter-gatherer ancestry plays a key role in increasing BMI in Japan, which could also be linked to the disparities in obesity prevalence among Asian populations residing in Western countries⁴⁶. By emphasizing the importance of incorporating Jomon ancestry as a confounding factor in GWAS for BMI, this analysis provides a proof-ofconcept for research that bridges our human past with current health challenges.

Given that the set of genetic variants associated with the proportion of Jomon ancestry likely tags genomic segments descended from the Jomon, they could serve as ancestral informative markers (AIMs), offering a convenient way to predict an individual's Jomon ancestry. We demonstrate the predictive power of these markers by leveraging three additional validation cohorts. Furthermore, the Jomon-related variants are significantly enriched with active functions in skeletal muscle cells, some of which are also associated with increased BMI, body weight, and height. These findings suggest potential adaptations related to the high physical activity required for hunting and gathering lifestyles⁴⁷, as supported by selection scans in the Jomon showing that the top variants are associated with increased bone mineral density³⁷. However, the genetic legacy of this selection may now pose a risk factor for elevated levels of BMI through the interaction with modern environments. Our study provides evidence of how the past action of natural selection has shaped present-day disease risks, primarily due to rapid changes in diet and societal lifestyle. It is important to note that further analysis with post-Jomon genomes (e.g., Yayoi or Kofun genomes) is crucial to uncover how Jomon segments have been inherited by descendants as lifestyles transformed from hunting and gathering to farming.

Nonetheless, our research has several caveats. First, we employed modern genomic data converted from imputed genome-wide array data. Consequently, our modern data remains ascertained, and the use of large-scale whole-genome sequence data could offer an unbiased set of variants for analysis. Regarding ancient genomic data, all Jomon data used in this study are shotgun-sequenced, rather than capture-sequenced. Sp.14,48. Still, most of the data are of low-coverage and were analyzed as pseudo-haploid data alongside the modern dataset. This may potentially restrict our ability to establish associations between modern and ancient genomes at a finer resolution⁴⁹. However, the emerging genotype imputation for ancient genomes may provide an innovative solution to enhance the depth of genotypic profiles derived from such low-coverage data^{37,50,51}.

In summary, our integrated analysis of modern and ancient genomes unveils the genetic legacy of ancient hunter-gatherers in populations today and its impact on phenotypic variation, shedding light on the significance of understanding a person's genetic ancestry not only for tracing their genetic origins but also for controlling confounding effects in GWAS. The field of ancient genomics is rapidly evolving, and future research that encompasses a diverse range of ancient humans across various time periods and geographic locations will be essential in providing a more comprehensive understanding of the extent to which the human past has shaped genomic and phenotypic variation in contemporary populations.

Methods

Biobank Japan (BBJ) dataset

We used 171,287 individuals from the first cohort of BBJ, which enrolled participants between 2003 and 2007. BBJ is a hospital-based genome cohort that included participants affected by at least one of 47 diseases from 12 medical institutes located in seven regions in Japan²⁸. All participants provided written informed consent, which was approved by the ethical committees of the Institute of Medical Science, the University of Tokyo. This study was approved by the ethical committees of Osaka University Graduate School of Medicine and Graduate School of Medicine, the University of Tokyo.

We excluded (i) individuals with lower call rates (<99%), (ii) closely related individuals with genetic relatedness ≥ 0.178 calculated from a

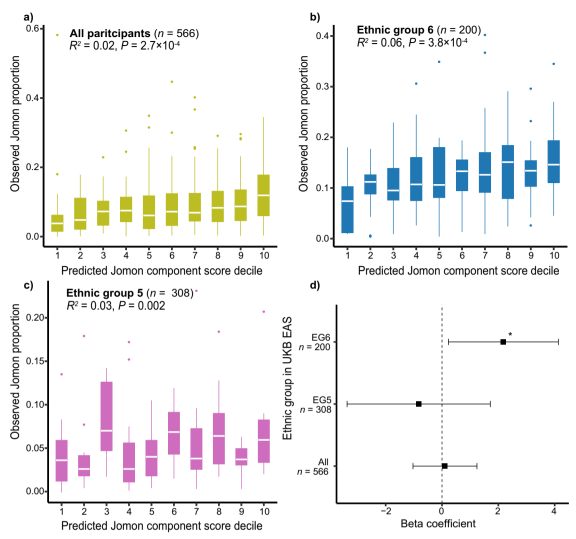


Fig. 7 | **Prediction of Jomon ancestry and replication of phenotypic association using UKB EAS.** Jomon component prediction scores are split into deciles. The boxplot shows the distribution of observed Jomon proportions in a given decile of the predicted scores based on (a) all UKB EAS participants (n = 566), (b) EG6 (self-reported Other ethnic population; n = 200), and (c) EG5 (self-reported Chinese population; n = 308). The EG number is referred from Data-coding of UKB. **d** The forest plots represent the effect size of Jomon ancestry on BMI in UKB EAS, EG5, and EG6. Squares indicate the point estimates, while error bars indicate 95% confidence

intervals. Asterisks represent P < 0.05 (P = 0.03 in EG6). In (**a**, **b**, **c**), boxes denote the interquartile range (IQR) and the median is shown as white horizontal bars; whiskers extend to 1.5 times the IQR; outliers are shown as individual points. R^2 indicates the squared value of the Pearson's correlation coefficient between the prediction score and the residuals of the Jomon proportion regressed out with 10 PCs. P-values reflect two-sided tests. In (**d**), P-values are computed by linear regression. All statistical tests are two-sided and unadjusted for multiple comparisons.

genetic related matrix (GRM) by GCTA (version 1.93.3beta2), or (iii) individuals were excluded if they were positioned far from the Japanese cluster defined within the 1000 Genomes Project (1KG) dataset in PCA plots using PLINK2 (v2.00a2.3). Based on where participants were registered, we defined seven geographically-defined populations for the Japanese archipelago (i.e., Hokkaido, Tohoku, Kanto-Koshinetsu, Chubu-Hokuriku, Kinki, Kyushu, and Okinawa, which are ordered from northeast to southwest Japan) and eight populations for the Ryukyu Islands (i.e., Yakushima, Amami, Kikai, Okinoerabu, Tokunoshima, Yoron, Okinawa, and Miyako). The five subpopulations were identified from visual inspection of the PCA plots (i.e., Mainland, Ryukyu, Ryukyu_admix, EA_admix, and Hokkaido_sub).

BBJ GWAS data were genotyped using the Illumina HumanOmniExpressExome BeadChip or a combination of the Illumina HumanOmniExpress and HumanExome BeadChips. The quality control of the genotypes has been described elsewhere 29,52 . In brief, we excluded variants with the following criteria: (i) call rate <99%, (ii) P value for Hardy-Weinberg equilibrium (HWE) < 1.0×10^{-6} , (iii) number

of heterozygotes <5, and (iv) a concordance rate <99.5% or a non-reference concordance rate between GWAS array and whole genome sequencing. The genotype data were phased by Eagle v2 and imputed with WGS merged the 1000 Genomes Project Phase3v5 (n = 2504) and BBJIK WGS (n = 1037) using Mimimac3 software (2.0.1).

Data merging between modern and ancient genomic data

We used ancient genomes from the Japanese archipelago and East Eurasian continent, which had been previously compiled and integrated with the Simons Genome Diversity Project (SGDP) panel (SGDP_Ancient)¹⁴. This SGDP_Ancinet dataset includes 14 ancient Japanese that were all shotgun sequenced. Under the stringent quality control (e.g., ancient DNA damage or low-coverage data), the ancient genomes (n = 22) were pseudo-diploid called for total 3,867,366 sites that were transversion only and a minor allele frequency of $1\%^{14}$. To further merge ancient genome data with the BBJ genotype data, we converted the accurately imputed dosage data of BBJ (Rsq \geq 0.7) to genotype data using PLINK2 with the following options:

fill-missing-from-dosage and hard-call-threshold 0.499. Then, we extracted sites that are present in both SGDP_Ancient and BBJ genotype data using PLINK (v1.90b4.4), resulting in the final merged dataset containing a total of 2,038,260 sites.

Admixture modeling and f_4 tests

We applied qpAdm in AdmixTools (version 7.0.2)³⁰ to the final merged data. In line with a previous study¹⁴, our analysis only used transversion sites with global minor allele frequencies of 1%, coupled with the option of "allsnps: NO." In qpAdm, we set the left and right populations as the source of admixture and reference populations, respectively. We then selected nine Eurasians as a right population; Sardinian (n = 3), Kusunda (n = 2), Papuan (n = 14), Dai (n = 4), Ami (n = 2), Naxi (n = 3)⁵³, Tianyuan (n = 1)^{3,54}, Chokhopani (n = 1)^{3,54}, and Mal'ta (n = 1)⁵⁵. As the left population, we set three ancient populations: Jomon (n = 12)^{8,9,14,48}, Northeast Asian (n = 2; WLR_BA_o and HMMH_MN, Bronze Age and Middle Neolithic individuals from the West Liao River basin)⁵⁶, and Han (n = 3; the SGDP panel, details in Supplementary Data 2)¹⁴.

We evaluated whether the tripartite structure provides a better fit to the data than any other possible scenarios at population levels based on p-values for nested models with a cut-off of 0.05. These alternative models include dual structure models involving any combination of Jomon, Northeast Asian, and East Asian ancestry, as well as single ancestry models. When conducting admixture modeling at individual levels, we considered the tripartite model to be a good fit if (i) a tail probability was >0.05 and/or (ii) estimates of admixture fractions were feasible (i.e., >0.0 and <1.0). The correlation of admixture proportions of the three distinct ancestors was calculated using Pearson's method. For individuals who did not support the tripartite model due to the tail probability being below 5%, we sought to identify an alternative dual ancestry model with the highest tail probability, subsequently confirmed through nested p > 0.05 by comparing the tripartite and dual structure models. If a single ancestry model was plausible, we further assessed whether a specific ancestor alone can adequately explain the individual using the nested p-values.

The f_4 -statistics were measured in the form of f_4 (Mbuti, Jomon; Han, X) using qpDstat with the f_4 mode in AdmixTools. We used the BBJ and the populations in the 1KG (n = 2504) or the SGDP as target populations (X in the form).

Curation of phenotypes in Biobank Japan

BBJ collected clinical conditions, laboratory data, and behavioral habit information for all participants through interviews and reviews of medical records using a standardized questionnaire. We selected 80 traits (3 anthropometrics, 55 biomarkers, 2 behavioral habits, 2 reproductive traits, and 18 diseases). We utilized the data for individuals over the age of 18 years, but only included the drinking and smoking traits for those over 20 years. For quantitative biomarkers, we applied the same processing and quality control methods as previously reported (Supplementary Data 4)^{19,29}. In brief, we used the laboratory values measured during the participants' first visit to the recruitment center and adjusted the values based on the type of medication used. We then applied a rank-based inverse normal transformation to normalize biomarker traits. Behavioral traits, including drinking and smoking history (ever versus never drinking and ever versus never smoking), were analyzed as binary phenotypes^{57,58}. Reproductive traits were coded as age at menarche and age at menopause⁵⁹. We excluded individuals if their age at menarche was below 10 or above 20 years, or if their age at menopause was below 40 or above 60 years. Patients with myocardial infarction, stable angina, and unstable angina were reclassified as having coronary artery disease (CAD)⁶⁰. 18 diseases were selected from a group of target diseases in BBJ, with sufficient numbers of cases and controls (Arrhythmia, Asthma, Cataract, CAD, Dyslipidemia, Ischemic stroke, Congestive heart failure, Osteoporosis, Glaucoma, Chronic hepatitis C, Colorectal cancer, Gastric cancer, Pollinosis, Urolithiasis, Rheumatoid arthritis, Prostate cancer, Breast cancer, and Type 2 diabetes). Individuals who were not affected by a particular disease under study were treated as controls.

In addition, we set a dummy phenotype as a negative control. We simulated 10 phenotypes with predefined heritability ($h^2 = 0.5$) from 10,000 causal variants randomly sampled from BBJ GWAS data using the GCTA GWAS simulation method⁶¹. The values were normalized by applying rank-based inverse normal transformation.

Phenotypic impact of Jomon ancestry in Japanese populations

Associations between Jomon ancestry and traits were tested using a glm() function implemented in R software (version 4.1.0). We applied a linear regression model to quantitative traits and a logistic regression model to diseases and behavioral habits, with adjustment for covariates. The proportion of Jomon ancestry was normalized using the rank-based inverse normal transformation method. As covariates, we included sex, age, age squared, top 20 PCs, 45 disease statuses, geographic regions, PCA clusters, and trait specific covariates listed in Supplementary Data 5. For BMI, we further tested the association with Jomon ancestry, stratified by sex and age groups, with the age threshold set at 65 years old, which reflects a mean age among the BBJ participants.

To address potential multicollinearity issues between Jomon ancestry and PCs, we employed an alternative approach. Initially, we conducted regression analysis of the quantitative trait measurements on all covariates, including PCs. Subsequently, we tested the associations between Jomon ancestry and the residuals derived from this regression using a linear regression model.

Assessment of genome-wide estimation and polygenic prediction with and without Jomon ancestry

We investigated the influence of the Jomon proportions on GWAS for BMI and the accuracy of BMI PGS. We performed BMI GWAS on all BBJ participants using a generalized MLM approach of GCTA-fastGWA⁶². We employed a sparse GRM constructed with variants subject to minimal LD pruning. Covariates in the original BMI GWAS included sex, age, age squared, the top 20 PCs, and 45 disease statuses. To evaluate the impact of Jomon ancestry, we introduced Jomon proportions as an additional covariate and compared the results with those obtained from the original GWAS.

To compute PGS, we adopted a five-fold leave-one-group-out GWAS method due to the absence of independent external reference for GWASs or genotype data with Japanese ancestry^{19,63}. In brief, we initially split the BBJ individuals randomly into five subsets. We then performed BMI GWAS on samples, excluding the subset under investigation (i.e., target subset), using GCTA-fastGWA. To estimate the posterior effects of SNPs from GWAS summary data, we utilized PRS-CS-auto (version Jun 4, 2021) with the HapMap3 LD reference panel of 1KG EAS^{64,65}. These posteriors of effect sizes were estimated both from the original GWAS and from the GWAS with the covariate of Jomon ancestry. Finally, we applied the PLINK2 score function to compute PGS for individuals within the target subset, comparing those scores with and without the Jomon ancestry covariate.

The incremental prediction performance of the Jomon ancestry was quantified with the difference in R^2 as follows:

$$\left(R_{PGS+fomon}^2/R_{PGS}^2\right) - 1\tag{1}$$

where R^2_{PGS} is the R^2 of PGS modeled by trait -PGS+covariates (sex, age, age squared, the top 20 PCs, 45 disease status, geographic regions, and PCA clusters), while $R^2_{PGS+Jomon}$ is the R^2 of the same PGS model but with the additional inclusion of the Jomon proportion.

Identification of Jomon-related variants

To detect genetic variants associated with Jomon ancestry, we conducted a genome-wide association analysis between the Jomon proportions and 6,861,976 biallelic variants (MAF \geq 1% and Rsq \geq 0.7). To strictly control the genomic inflation factor of test statistics, we adopted the following correction methods: (i) MLM-based approach using GCTA-fastGWA with the adjustment of covariates: age, age squared, sex, the top 20 PCs, 45 disease status, geographic regions, and PCA clusters; (ii) fixed-effect meta-analysis of Mainland summary data including individuals from the Mainland and EA_admix clusters (n = 151,075) and of Ryukyu summary data including individuals from the Ryukyu, Ryukyu admix, and Hokkaido_sub clusters (n = 10,080) using METAL (version 2020-05-05); and (iii) double genomic control correction method using METAL³⁶. We then computed a Z score for each variant by considering the sign of the beta coefficient and the associated p-value.

To identify independent variants, we performed LD clumping on those variants with positive Z scores using BBJ1K and 1KG EAS as reference populations, with the following PLINK parameters: p1=1, p2 = 1, r2 = 0.01, kb = 2000. Jomon-related variants (total 132 variants) were defined as those that are independent, not located in the HLA loci, and satisfy a genome-wide significance ($P < 5.0 \times 10^{-8}$). We calculated the F_{ST} value for the Jomon-related variants among the EAS populations in 1KG using Hudson's F_{ST} implemented in PLINK2⁶⁶. Variants with p-values greater than 0.05 were classified as non-Jomon-related. We then identified 132 non-Jomon-related variants matching allele frequencies of the Jomon-related variants using a nearest neighbor matching method, contrasting the Jomon-related and non-Jomonrelated variants by examining their frequency differences between the Japanese and continental East Asian populations, as well as their haplotypic lengths with LD greater than 0.8. The enrichment of selection signals in the Jomon-related variants was tested based on Z-scores of SDSs estimated in a previous study¹⁷. The sum of squared values of rank-based normalized Z-scores were compared to a chi-squared distribution with the degree of freedom equal to the number of available variants.

Stratified LD score regression was applied to the meta-analyzed summary data of the Jomon proportion, which estimated cell group enrichment with the recommended baseline LD model⁶⁷. For LD score regression, we adopted the HapMap3 SNPs, excluding those within the *HLA* region, and used pre-computed LD scores among the 1KG EAS populations, which were obtained from the LDSC software website⁶⁸. We investigated the pleiotropic effects of Jomon-related variants using BioBank Japan PheWeb for the Japanese population (https://pheweb.jp/) and Open Targets Genetics for Europeans (https://genetics.opentargets.org/). We also evaluated eQTL effects of the variants using GTEx (https://gtexportal.org/home/).

Validation with independent Japanese cohorts

As independent replication cohorts, we used the Nagahama cohort study and the second BBJ cohort (BBJ-2nd). The Nagahama cohort study is a community-based and recruited participants from Nagahama City, Shiga Prefecture, Japan⁴². The cohort was genotyped using six different genotyping arrays. We then selected two platforms (Nagahama A; Illumina Human610-Quad Beadchip, and Nagahama B; Illumina HumanOmni2.5-4v1 Beadchip) with a large number of samples. We excluded individuals with low call rates, high heterozygosity rates, closely related individuals, and PCA outliers from the EAS populations (n = 1591 in Nagahama A and n = 1444 in Nagahama B)⁶⁹. We also excluded variants with (i) call rate < 0.98, (ii) MAF < 1%, and (iii) HWE $P < 1.0 \times 10^{-6}$. Genotype data were phased by Eagle v2 and imputed with the reference panel from the 1000 Genomes Project Phase3v5 and BBJ1K using Mimimac3. As described above, we converted the high-quality imputed dosage data to genotype data and them with ancient genome data. We obtained 1,982,989 shared variants in Nagahama A and 2,109,225 shared variants in Nagahama B.

The BBI-2nd is an additional cohort of independent participants from the first cohort of BBI. The BBI-2nd consisted of ~80,000 individuals collected between 2013 and 2018. The subjects were genotyped using an Illumina Asian Screening Array chip. As in the first cohort, we applied stringent QC filters to both participants and SNPs as described elsewhere $\bar{^{70,71}}$. Briefly, we excluded individuals with a low call rate (<0.98), closely related individuals (King's kinship index \geq 0.0884), and outliers from the EAS cluster in the PCA with the samples of the HapMap3 project (n = 72,695). We also excluded variants with call rate < 0.99, minor allele count < 5, HWE P < 1.0 × 10⁻¹⁰ and >0.05 of allele frequency difference when compared with the Japanese WGS reference panels. The genotype data was phased by SHAPEIT (version 4.2.1) and imputed with the reference panel from the 1000 Genomes Project Phase3v5 and BBJ1K using Mimimac4 (version 1.0.1). Converting the high-quality imputed dosage data $(Rsq \ge 0.7)$ to genotype data, we merged the ancient genome data to obtain 1,940,657 shared variants.

We estimated proportions of Jomon ancestry for individuals in each cohort by using qpAdm in the same way as the first cohort of BBJ. We derived a score for predicting Jomon ancestry based on 132 independent Jomon-related variants by using the PLINK2 score option, which is defined as Jomon component prediction score. After regressing the Jomon proportion out with 10 PCs as covariates, we estimated the variance explained by the scaled Jomon component prediction score using Pearson's correlation. In each cohort, principal components were computed using the projection method on the PCs of the first cohort of BBJ. To evaluate the power of the Jomon component prediction score, we employed a clumping and thresholding approach. We built the Jomon component score for BBJ-2nd individuals using genetic variants meeting the following p-value thresholds: 5×10^{-8} , 5×10^{-7} , 1×10^{-6} , 1×10^{-5} , 1×10^{-4} , 1×10^{-3} , 0.01, 0.05, 0.1. To assess the proportion of variance explained by the Jomon component score, we calculated the adjusted R^2 from a full model that included the score and all covariates, compared to a null model without the score.

Replication analysis using East Asians individuals in UK Biobank

As an independent source of an East Asian (EAS) population, we focused on EAS individuals from UK Biobank (UKB), which is a population-based cohort that recruited 500,000 individuals between 40 and 69 years old from across the United Kingdom⁷². EAS individuals were extracted by visual inspection on the PCA plots including 1KG populations as the reference for EAS ancestry (n = 2286; Supplementary Fig. 11). UKB was genotyped with either the Applied Biosystems UK BilLEVE Axiom Array or the Applied Biosystems UKB Axiom Array. The genotypes were imputed using the Haplotype Reference Consortium, UK10K, and the 1000 Genomes Phase 3 reference panel using IMPUTE4⁷². The detailed characteristics of the cohort have been described elsewhere⁷². For UKB EAS individuals, we converted the variants with INFO score ≥ 0.8 to genotype data and obtained 3,610,183 sites shared with ancient genome data.

The genetic affinity between the Jomon and UKB EAS or between the Jomon and each of different ethnic background groups within UKB EAS was tested using f_4 -statistics with the form of f_4 (Mbuti, Jomon; Han, X). The breakdown of self-reported ethnic backgrounds at the participants' initial visit (Data-Field 21000) is as follows: Data-coding 1 for White (n=1), Data-coding 1001 for British (n=1), Data-coding 2003 for White and Asian (n=2), Data-coding 2004 for Any other mixed background (n=3), Data-coding 3 for Asian or Asian British (n=2), Data-coding 3004 for Any other Asian background (n=300), Data-coding 5 for Chinese (n=1375), and Data-coding 6 for Other EG (n=541).

We estimated the Jomon proportion for individuals in each cohort by using qpAdm. In cases where individuals did not support the tripartite ancestry structure, we applied the same procedure used for BBJ to identify a plausible model with a two-way admixture, where Jomon served as one of the source ancestors. Jomon component prediction scores based on 132 independent Jomon-related variants were calculated using PLINK2.

To assess the association between Jomon ancestry and BMI in UKB EAS, we computed the mean BMI for participants with measurements taken two or three times. The association tests were adjusted for sex, age, age squared, ascertainment center information, batch information, and ethnic backgrounds. This approach was also applied to EGS (i.e., self-reported Chinese, n = 1375) and 6 (i.e., self-reported Other ethnic group, n = 541) with the same covariates, except for ethnic backgrounds.

Reporting summary

Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

Data availability

The summary statistics from the genome-wide association analysis for Jomon ancestry have been deposited in the National Bioscience Database Center (NBDC) Human Database (https://humandbs.dbcls.jp/en/) under accession code hum0197 (https://humandbs.dbcls.jp/en/hum0197-latest). The genotype data of BBJ are available from the NBDC Human Database (research ID: hum0014 and hum0311). The genotype data of the Nagahama cohort study are available from the NBDC Human Database (research ID: hum0012). UKB analysis was conducted using application number 47821 (https://www.ukbiobank.ac.uk/). All ancient genomic data used in this study were previously compiled¹⁴.

Code availability

We used publicly available software for analysis. The software used is described in the "Methods" section.

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Acknowledgements

We thank all participants and investigators of the Biobank Japan Project, Nagahama cohort study, and UK Biobank. We would like to express our deepest gratitude to our co-author T.T., who passed away in July of 2024, for his invaluable contribution to BioBank Japan Project. K.Y. was supported by AMED (JP223fa627002) and Center for Advanced Modality and DDS (CAMaD), Osaka University. K. Suzuki., T.Y. and T.K. were supported AMED (JP20km0405202, JP23tm0424218). T.G was supported by SAKIGAKE Project in Kanazawa University, MEXT KAKENHI Grant Number (20H05822), JSPS KAKENHI Grant Number (21H04358). S.Nakagome. was supported by JSPS KANENHI (22H02711) and Wellcome Trust ISSF Award. Y.O. was supported by JSPS KAKENHI (22H00476), and AMED (JP23km0405211, JP23km0405217, JP23ek0109594, JP23ek0410113, JP23kk0305022, JP223fa627002, JP223fa627010, JP233fa627011, JP23zf0127008, JP23tm0524002), JST Moonshot R&D (JPMJMS2021, JPMJMS2024), Takeda Science Foundation, Bioinformatics Initiative of Osaka University Graduate School of Medicine, Institute for Open and Transdisciplinary Research Initiatives, Center for Infectious Disease Education and Research (CiDER), and Center for Advanced Modality and DDS (CAMaD), Osaka University.

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K.Y., S.Nakagome., and Y.O. designed this study and wrote the manuscript. K.Y., S.Namba., K.Sonehara., S.S., and N.P.C. conducted data curation and bioinformatics analysis. K. Suzuki., S.H., S.K., H.A., K. Matsuura., Y.M., Y.F., T.T., K. Matsuda., T.G., T.Y., T.K., and S.Nakagome. collected the samples. S.Nakagome. and Y.O. supervised this study.

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/s41467-024-54052-0.

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Peer review information *Nature Communications* thanks William Barrie and the other, anonymous, reviewers for their contribution to the peer review of this work. A peer review file is available.

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