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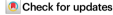
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Human genetic structure in Northwest France provides new insights into West European historical demography

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A list of authors and their affiliations appears at the end of the paper

The demographical history of France remains largely understudied despite its central role toward understanding modern population structure across Western Europe. Here, by exploring publicly available Europe-wide genotype datasets together with the genomes of 3234 present-day and six newly sequenced medieval individuals from Northern France, we found extensive fine-scale population structure across Brittany and the downstream Loire basin and increased population differentiation between the northern and southern sides of the river Loire, associated with higher proportions of steppe vs. Neolithic-related ancestry. We also found increased allele sharing between individuals from Western Brittany and those associated with the Bell Beaker complex. Our results emphasise the need for investigating local populations to better understand the distribution of rare (putatively deleterious) variants across space and the importance of common genetic legacy in understanding the sharing of disease-related alleles between Brittany and people from western Britain and Ireland.

Understanding how genetic diversity is distributed within and between human populations (i.e., population structure) can shed light on the history of our species, and inform studies that search for genetic associations with traits and diseases¹. The increasing interest in elucidating the role of rare variation in complex traits, the consequent demand for surveying multiple local populations and the need for a detailed understanding of their evolutionary history has motivated multiple population genomic studies at a country-wide scale²⁻⁵.

Located on one of the edges of the European landmass, Northwestern France includes the peninsula of Brittany, whose westernmost tip is named *Finistère* (end of land), and the surrounding region of *Pays-de-la-Loire*. The region is bordered by the English Channel to the north and the Bay of Biscay to the south, and is crossed by the Loire River - the largest river in France. From a linguistic point of view, this region has been the stage of long-term contacts between Celtic and Gallo-Romance spoken forms. The first evidence of modern human occupation of Northwestern France is associated with the Early Aurignacian culture, dating between about 43,000 and 37,000 years ago⁶. However, population occupation in the region may have been scarce up to the Middle/Late Neolithic. It is during this period that megalithic

construction appears in the archaeological record⁶. The peninsula of Brittany hosts the oldest⁷ and some of the biggest megalithic monuments (stone rows, long barrow and passage tombs) and one of the highest concentrations thereof. The Neolithic culture in Northwestern France largely derives from the Danubian Wave, also known as the Linear Pottery culture^{8,9}, although the presence of pottery with shell impressions in regions south of the river Loire raises the possibility of Mediterranean influence in the neolithization of the region⁹. Genetic evidence suggests that the introduction of the France Neolithic lifestyle occurred through a complex interaction between the two Neolithic waves - the Danubian and the Mediterranean wave - and local, likely genetically structured, populations of hunter-gatherers¹⁰⁻¹². Both archaeological and genetic evidence suggest strong connectivity among the populations along the Atlantic façade from 4500 BC onwards, from Northern Iberia to Ireland and Western Britain¹³. Although Northwestern France was likely part of the Atlantic façade with respect to its genetic landscape, there is no ancient DNA data currently available from the Neolithic period.

The connectivity along the western fringes of the European continent continued during the Early Bronze Age as suggested by the

e-mail: richard.redon@univ-nantes.fr; christian.dina@univ-nantes.fr

presence of a common Bell Beaker pottery style, usually known as the "Maritime" style^{14,15}, and further related artefacts¹⁶. In addition, the Bell Beaker complex in Northwestern Europe is closely associated with the appearance of copper metallurgy¹⁷. In Britain, the arrival of the Bell Beaker complex has been recently associated with a major genetic ancestry shift disrupting the preceding genetic homogeneity of the Atlantic façade and introducing genetic ancestry related to the Yamnaya migration from the Eurasian Steppes¹⁸. These findings contrast with what has been found in Iberia, where Beaker-complex-associated individuals show low genetic affinity with those from central Europe suggesting considerable heterogeneity underlying the mode of transmission of the Bell Beaker complex. Within France, Yamnaya-related ancestry has been primarily found among Bell-Beaker associated individuals from the Northeastern and Southern parts of the country, although ancestry proportions widely vary across samples 10,11. However, the lack of samples from Northern and Northwestern France before the Iron Age hinders the full understanding of the introduction of Steppe ancestry along the westernmost part of the North Sea and the English Channel. In Northern France, the Iron Age is genetically characterised by high levels of Steppe-ancestry and a homogenisation of it, suggesting continuous cultural change instead of a massive migration underlying the arrival of iron technology¹⁹. In the Iron Age, Brittany was called Aremorica and was part of the Gaul. It was home to multiple Celtic-speaking tribes, like the Veneti and Osismii in the west of the peninsula²⁰.

During the Roman Empire, Roman influence reached Brittany, as attested by the presence of Roman-style villas and sanctuaries. However, such influence was far smaller in Northwestern France than in other parts of the Gaul²¹. With the decline of the Roman Empire (around the 3rd century CE), Northern France became progressively ruled by the early mediaeval kingdoms²². However, the history of Brittany remains largely unknown from the late Antiquity (4th-5th century CE) up to the 9th century CE, when it was conquered by Frankish Carolingian Emperors and put under native rulership²³. Interestingly, it was during this period that the peninsula acquired the name of "Britannia". Linguistically, places' names and the still-surviving language closely related to Cornish and Welsh imply that extensive connections took place with Western Britain, Nevertheless, whether such a process involved small or large settlements from the British Isles, as often suggested, or is the basis of the Breton language is still under debate²⁴⁻²⁷. Archaeological evidence displays changes in architecture and funerary rituals during this time. However, they reflect a mixture of influences, rather than a single major contribution, from the British Isles and French regions along the British Channel as well as from Germany²⁸.

Northwestern France has been the end point of multiple continental prehistoric migrations and occupied a central place in trading routes along the Atlantic façade²⁹. Understanding the genetic makeup of Northwestern France can inform on the complex interaction between the European-wide demographic events that have culminated in the present-day genetic landscape. However, a systematic assessment of patterns of population structure in Northwestern France and an identification of the past demographic events that have shaped them is still lacking. In this study, we analysed 3234 genome-wide genotyped samples together with 620 high-coverage whole-genome sequences (WGS) from Brittany and Pays-de-la-Loire. Also, to put the genetics of Northwestern France into context we used an additional set of 236 WGS from other regions in France and merged the WGS with available datasets encompassing a geographically diverse set of present-day and ancient European samples. Last but not least, we analysed, for the first time, a set of western French mediaeval samples dating from the 4th to the 11th century in order to fill in the gap with respect to the lack of publicly available genome-wide data from the last two millennia.

Results

Haplotype-based population structure in Northwestern France

To explore patterns of genetic structure among the population in Northwestern France, we first applied the haplotype-based approach implemented in fineSTRUCTURE (vs4.1)³⁰ on 210,171 SNPs genotypes from 3234 individuals from Brittany and *Pays-de-la-Loire* (see *Methods*). Genome-wide data revealed extensive population structure in Northwestern France with 154 clusters inferred, of which 78 contain more than 10 individuals (Fig. S1). We investigated whether recent ancestry due to the sampling scheme carried out in this study impacts the clustering patterns and we found no evidence of such an effect (see Methods and Fig. S2–S4). Pairs of individuals within clusters are not more closely related than pairs of individuals across the whole dataset.

At the coarser level *k* = 3 (Fig. 1a) of the fineSTRUCTURE tree (FS tree), we found that the distribution of one of the clusters, hereafter referred to as the "Western Brittany" (WBR) cluster, broadly overlaps with the linguistic distribution of the Breton language. The cluster's border falls between the easternmost (Loth line 1) and the westernmost (Loth line 2) estimated historical bounds of the Breton language (Fig. 1b). A second cluster, hereafter referred as "Eastern Brittany and Pays-de-la-Loire" (EBP), encompasses individuals from eastern Brittany (*département* Ille-et-Vilaine) and the area of *Pays-de-la-Loire* situated north of the river Loire. A third cluster, hereafter referred to as "South Loire" (SLO), covers the part of the *Pays-de-la-Loire* situated south of the river Loire.

The relative distributions of these clusters mimic those of surnames across the entire region, as inferred through a Neighbour Joining Tree (Fig. 1c). The three surname-based clusters are highly supported by the bootstrap analysis (>75%). Interestingly, the surnames from the northern part of *Pays-de-la-Loire* are more closely related to those from south Loire than to those from Brittany. Furthermore, while the highest level of differentiation is observed between WBR and SLO (F_{ST} = 0.00107), F_{ST} values indicate larger population differentiation between EBP and SLO than between EBP and WBR (respectively 0.00042 vs. 0.00028, Fig. S5). Together we found evidence that the river Loire influences population differentiation, likely acting as a barrier to gene flow.

For higher levels of resolution (k > 3), because the FS tree depends on the sample sizes30 and might not reflect haplotype sharing differences between pairs of clusters³¹, we computed the clustering tree based on the total variation distances (TVD tree^{5,31}). Indeed, when assessing the performance of the two tree-building algorithms (see Methods for details) by computing cluster assignment confidence⁵, we found that for the same k the FS-tree shows lower cluster assignment confidence than the TVD-based tree (Fig. S6). To ease visualisation we chose k = 39 and merged small-sized clusters (1–5 individuals) with the closest large cluster (>= 31 individuals) resulting in 18 clusters (>31 individuals each). As a result, the north Loire region splits into nine clusters, four of which in the westernmost part of Brittany, while individuals from south Loire split also into nine clusters (Fig. 1d). Overall, we retrieved a TVD-tree largely consistent with the fineStructure tree at k=3, with the exception of samples located at the borders between the three main clusters (e.g., individuals assigned to the "Malo-Rennais" cluster of the TVD-tree). In the resulting 18 clusters, individuals from the westernmost part of Brittany are grouped into four clusters: "Cornouaille", "Lèon", "Vannes" and "Bretagne-Centre", while individuals mostly from south of the river Loire cluster into nine clusters (Fig. 1d). Samples geographically located between these regions are grouped into five clusters: "Malo-Rennais", "Nantes", "Guérande", "Maine-Anjou" and "Ancenis". The coancestry matrix for the 18 clusters (Fig. S7) reveals a global pattern of increased coancestry within clusters, as expected after a history of shared drift between individuals in the same group, with the exceptions of the clusters "Sèvre" and "Vendée-Est".

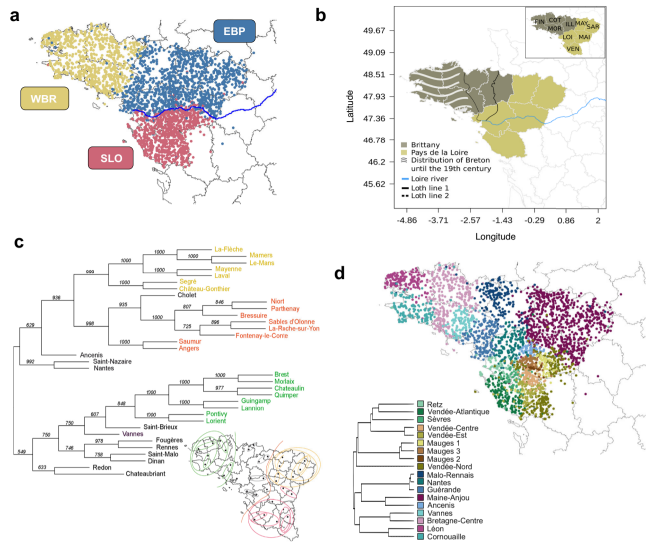


Fig. 1 | **Population structure, linguistics, and patronym distribution in Northwestern France.** a fineSTRUCTURE clustering of 3234 samples from Brittany and the downstream Loire basin at k = 3. WBR stands for Western Brittany, EBP for Eastern Brittany and *Pays-de-la-Loire* and SLO for South Loire. **b** Distribution of the Breton language over time⁹³. Loth line 1, attested by the border between Celtic place names in -ac (on the west side) and Romance place names in $-\acute{e}$ (on the east side), represents the westernmost boundary between Armorican Gaulish and Gallo-Romance spoken varieties (25). Loth line 2 represents the Celto-Romance linguistic border in the 16th century CE. Grey lines encircle political territorial divisions called *départements*. Only *départements* belonging to the regions of Brittany and *Pays-de-la-Loire* are coloured. In the inset, the *départements* are abbreviated as FIN (Finistère), COT (Côtes-d'Armor), MOR (Morbihan), ILL (Ille-et-Vilaine), MAY (Mayenne), SAR (Sarthe), MAI (Maine-et-Loire), LOI (Loire-Atlantique), VEN (Vendée).

c Neighbour Joining tree from pairwise Arccos distances (68) computed on the distribution of surnames between pairs of *arrondissements* from Northwestern France. The support of the tree nodes was obtained by bootstrapping 1000 times the distance matrices. **d** 18 genetic clusters identified using total variation distances (TVD) from the fineSTRUCTURE results (see Material and Methods for details). The TVD-based tree was cut at k = 39 and to ease visualisation we merged clusters with <5 individuals into the closest cluster (>= 31 individuals) resulting in a tree with 18 clusters. All the clusters have at least 31 individuals. The TVD-based tree is sample size independent and is more informative on cluster genetic affinities³¹. The tree was built using the *clustree* and *ggtree* packages of R-statistical software⁸⁹ and the map with the following R libraries: sp as in (**a**, **b**). In (**a**) and (**c**), individuals are coloured according to their cluster assignment. Individuals' coordinates correspond to the most common grandparents' birthplace.

The cluster "Maine-Anjou" encompasses the largest number of individuals (n = 798) and covers by far the largest geographical area, revealing broader genetic homogeneity relative to the other regions. Most of the other clusters appear to have contributed to the coancestry of this large cluster, indicating that haplotype homogeneity within "Maine-Anjou" may result from repeated admixture into the region (Fig. S7).

The second largest cluster (n = 344), "Bretagne-Centre", stretches from the northern to the southern coast of Brittany indicating the existence of a corridor of relative genetic homogeneity connecting the two sides of the peninsula (see Fig. S12). While the neighbouring clusters "Léon", "Cornouaille" and "Vannes" display large within-cluster

but low between-cluster coancestry (Fig. S7), despite their close geographic proximity, each of them appears to have contributed to the coancestry of the "Bretagne-Centre" cluster. When considering the area located south to the river Loire, most clusters— and particularly "Mauges 1" and "Mauges 2" - appear to have largely contributed to the coancestry of the "Sèvres" cluster in eastern Vendée. In contrast, the "Mauges [1–3]" and "North-Vendée" clusters show low coancestry from most of the other clusters and large within-cluster coancestry, likely indicating stronger genetic drift within these groups. Finally, we observed a low contribution of the clusters south of the river Loire to the coancestry of those from the westernmost part of Brittany, indicating that gene flow between these two areas has been relatively

limited, in agreement with the F_{ST} values obtained between the 18 clusters (Fig. S8).

We then explored identity-by-descent (IBD) sharing between pairs of individuals belonging to the 18 clusters (Fig. S9), and measured average length of runs of homozygosity (ROH) across individuals within administrative circumscriptions in Northwestern France (Fig. S10). Higher levels of identity-IBD sharing for chromosomal segments of any size as well as increased length of ROH were observed between the clusters "Mauges [1–3]" and "Vendée-Nord", indicating that low effective population sizes may explain the particular population structure found in this restricted area. Similar patterns were observed within Brittany. However, considering that increased IBD sharing within the westernmost clusters of Brittany is only observed for chromosome segments under 7 cM (Fig. S9), lower effective population sizes within this cluster are likely more ancient than for "Mauges [1–3]" and "Vendée-Nord". Overall, these results fit with those based on the coancestry matrices and F_{ST} values (Figs. S7, and S8).

Fine-scale genetic structure, linguistics and geography

In an attempt to shed light on the origin of the observed genetic structure, we used partial Mantel tests to correlate pairwise F_{ST} values and surname-based distances (Fig. S11). We found such correlation to be high even after correcting for geographical distances (Partial Spearman correlation = 0.68). To illustrate this finding, the "Léon" and "Malo-Rennais" cluster locations match those of the surname clusters Brest/Morlaix and Saint-Malo/Dinan, respectively (Fig. 1c). Concordant with an association between genetics and language, we observed that the cluster "Bretagne-Centre" overlaps with the dialectal area featured by the usage of two initial consonants - the aspirated [h] instead of an unaspirated, and the alveolar fricative [z] instead of [s] (Fig. S12 and Supplementary Discussion for more details). Similarly, the cluster "Cornouaille" apparently overlaps with the area with palatalisation of -h- [h] into -y- [j] (Fig. S13; see Supplementary Discussion for more details). Although these observations are not supported by statistical tests, they are consistent with a spatial correspondence between genetic clusters and dialectal areas.

Given that geographical features (e.g., rivers) could also explain fine-scale population structure³, we investigated whether rivers, other than the Loire, overlap with cluster borders. We observed overlaps, although not systematically, between river courses and cluster borders (Fig. S14). For instance, the clusters "Lèon" and "Cornouaille" are respectively delimited to the east by the Morlaix-Jarlot River and Laïta-Ellé rivers, respectively, and separated from each other by the river Aulne. "Vannes" is enclosed to the west by the Blavet river and to the north by the Oust. The "Nantes" cluster is bordered by the rivers Semnon and Vilaine to the north and the west, respectively, while the "Malo-Rennais" cluster's borders co-localise with the Gouessant and the Yvel - Hivet to the west. In South Loire, the cluster "Vendée-Atlantique" is bordered to the southwest by the river Lay (Fig. S14). Overlaps between regions of low migration and rivers also appear, although not systematically, in the effective migration surfaces retrieved with the EEMS³² (Fig. S15). Specifically, we noted a significant decrease in migration across the Loire River (dark blue) and areas of low effective migration coincide with relatively large rivers such as Aulne (separating "Cornouaille" and "Léon") and Laïta (between "Cornouaille" and "Bretagne-Centre"), primarily in the westernmost part of Brittany. On the other hand, while the Vilaine River coincides with a region of low effective migration, its downstream part is in the centre of one cluster ("Guérande", close to the sea). However, it matches the North-West border of cluster "Nantes" where migration rates were inferred to be lower than expected. Recurring matches between watercourses and genetic cluster borders, while not formally and quantitatively tested, suggest an impact of rivers on local demographics and consequently on genetic structure and the dialectal distribution in Northwestern France.

Evolution over time of effective population sizes

We estimated the historical dynamics of effective population size (Ne) using IBDNe³³, and observed 12-, 5- and 5.2-fold increases in Ne, respectively for WBR, EBP and SLO, in the last 1450 years (assuming a generation time of 29 years; Fig. S16a). Such expansions have led to present-day population sizes of around 10^{5.5} to 10^{5.7} individuals for the three clusters, which is broadly consistent with the explosive population growth estimated for other European populations³⁴⁻³⁶. However, the Ne trajectories differ between clusters (Fig. S16a). Indeed, while WBR shows the smallest Ne in the Early Mediaeval Period (~5th century CE), its effective population size surpassed that of SLO by the High Mediaeval Period (~1000 CE). This suggests that population expansion started earlier in WBR than in SLO, which kept a stable Ne until the 10 last generations (i.e., 17th century CE). Reduced Ne at recent times explain the stronger IBD sharing observed between the clusters "Mauges [1-3]" and "Vendée-Nord" for chromosome segments >7 cM. Finally, the Ne trajectories in EBP and SLO show a slight and short population decline, starting around 1230-1350 CE and lasting for almost ~300 years (Fig. S16a). Similar patterns have been previously described in French and other European populations and putatively associated with the Black Death^{34,37}. However, we caution that the Ne profiles suggesting a bottleneck are not consistently observed across different thresholds for the minimum centimorgan (cM) length (Fig. S15b-d). In addition, our computer simulations suggest that alternative demographic scenarios involving recent migration or population structure within the EBP and SLO clusters generate similar Ne profiles (see Supplementary Discussion for details, Fig. S31). Given that a large part of the "Maine-Anjou" cluster has received genetic ancestry from most of the surrounding areas (Fig. S7, coancestry matrix) and the region south of the river Loire encompasses the largest number of subclusters, such alternative scenarios cannot be excluded. Historical evidence that the Black Death had a lesser demographic impact in Western Brittany, as the Ne trajectory suggests, is lacking, although recent studies refining its impact through pollen record support a lighter effect of the epidemics in Brittany³⁸.

Fine-scale population structure based on rare variants

Population stratification based on rare variation is typically stronger than with common polymorphism^{1,39} and rare variants are particularly informative to infer recent fine-scale population structure. Here, we computed allele sharing between pairs of individuals originating from Brittany and Pays-de-la-Loire using genotypes from 620 high-coverage whole-genome sequences (see Methods, Fig. S17). We computed independently two matrices, reporting genotypes with minor allele counts equal to two (MAC 2) and ranging from 3 to 10 (MAC 3-10), respectively. We applied hierarchical clustering to both matrices to cluster individuals according to patterns of allele sharing. When assuming three population groups (k=3), most individuals from the three westernmost départements of Brittany are clustered into a single group, both for MAC 2 and MAC 3-10 (Fig. 2). The proportion of individuals assigned to this cluster decreases as one moves away from Western Brittany, regardless of the allele count category, and the two alternative clusters become more prevalent. Similarly, to the fineS-TRUCTURE findings, these results provide evidence for relative differentiation between traditionally Breton-speaking populations in Western Brittany and their Gallo-speaking neighbours.

Interestingly, we identified a genetic component restricted to the *départements* located south to the river Loire for k=4 (cluster_2) for MAC 3–10 (Fig. 2, Figs. S18 and S19). For MAC 2 alleles we found this cluster only at k=9 (cluster_2). Assuming that alleles with MAC 2 (minor allele frequency ≈ 0.0016) tend to be more recent, these results suggest that population structure does not result from reduced gene flow between the northern and southern shores of the river in the very near past (average doubleton age -500 years⁴⁰). Clustering patterns within Brittany are, on the other hand, consistent across the full range

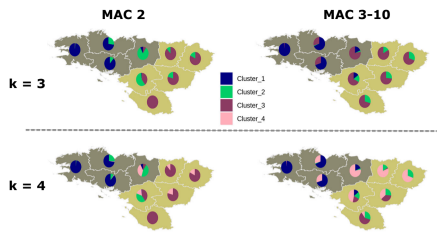


Fig. 2 | **Allele sharing within and between populations.** Proportions of individuals assigned to different clusters for each *département* in Northwestern France (see Fig. 1b), using hierarchical clustering on levels of rare allele sharing between pairs of individuals. Allele-sharing matrices were computed for alleles present in

two chromosomes (MAC 2) and three to 10 chromosome copies (MAC 3–10) across 620 individuals from Northwestern France and from one million randomly selected sites. Maps were generated with the R statistical package together with the following libraries: *ComplexHeatmap, rgdal, sp, broom, ggplot2 and scatterpie.*

of allele counts, indicating that population differentiation associated with traditionally Breton-speaking groups has persisted to modern times.

In general with MAC 2 alleles, increasing k from 3 to 10 assign individuals from neighbouring *départements* into seven additional geographically restricted clusters (Figs. S18 and S19), suggesting similar patterns of population structure as found with fineSTRUCTURE (Fig. 1d). Although an exhaustive comparison with fineSTRUCTURE results is beyond the scope of this study, this general concordance in clustering patterns emphasises the power of rare variants to infer fine-scale population structure. With MAC 3–10 alleles, increasing k tends to generate smaller clusters with relatively large geographical distribution, likely reflecting a relative lack of resolution to detect very recent population structure (Figs. S18 and S19).

Brittany in the context of France

To investigate population structure on a larger geographical scale, we enriched our dataset with additional WGS-based genotypes from 223 individuals originating from neighbouring French areas (see Methods, Fig. S17). We first performed principal component analysis (PCA) based on common and low-frequency variants (Fig. S20). While population differentiation based on common variants separates mostly Brittany from the remaining regions, low-frequency variants disclose a more subtle population structure, as previously reported disclose a more subtle population structure, as previously reported France (Fig. S20). Pairwise $F_{\rm ST}$ values also support stronger differentiation between WBR and the remaining populations (Fig. S5). In subsequent analyses, we will consider individuals from Brittany and *Pays-de-la-Loire* according to the three clusters shown in Fig. 1a (WBR, EBP and SLO), unless stated otherwise.

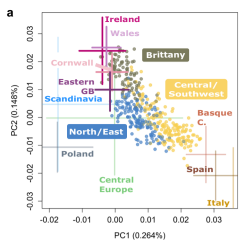
Brittany in the European context

To further investigate the genetic history of people from Northwestern France, we first performed a PCA on our entire WGS dataset merged with genotype data from 20 diverse Northern and Western European populations^{42,43}. In agreement with the previously reported isolation-by-distance pattern in Europe⁴⁴, we found that French samples are continuously distributed along the axis connecting Southwestern (i.e., Spain) and Northwestern (i.e., Ireland/UK) Europe (Fig. 3a, Fig. S21). Individuals from Central/Southwestern France appear closer to samples from Spain whereas individuals from Brittany appear at the other extreme and closer to samples from Ireland/UK. While individuals

from Brittany fall onto the axis and overlap with the Irish, Welsh and Cornish samples, samples from Eastern Great Britain show a slight shift towards Central Europe. These results support the idea of increased genetic proximity between Brittany and Ireland, as previously suggested⁴⁵. In contrast with Brittany and similarly to what we observe for samples from Eastern Great Britain, individuals from Northern and Eastern France show a slight shift towards Central Europe (represented here mainly by Germany).

To further investigate the contribution of other European populations to the genetic makeup of Northwestern France we used the regression-based statistical approach implemented in GLOBE-TROTTER software⁴⁶. We found that all of the seven French populations showed evidence of admixture (P < 0.01, computed from the proportion of inferred dates outside of the limits, i.e., <=1 or >=400). In agreement with the PCA results, three main sources of ancestry -North/Central, Northwestern, and Southwestern Europe - were consistently found across these seven populations. The North/Central component derives mainly from Denmark and/or Belgium, the Northwestern component from Ireland and the Southwestern one from Spain and/or Italy (Fig. S22). The population of Brittany (BRI) was found to trace back ~23.5% of its DNA to Ireland vs. 14% or less found in any other French population. Brittany is also the region that received the largest ancestry contribution from Cornwall (-9% vs. 3.1-3.4% across the remaining populations). While Hauts-de-France (HAU) and Grand Est (GRA) ancestries show evidence of single admixture events (Table S1), with Italy (Southern) and Kent (Northwestern Europe) as possible sources of admixture, the remaining populations (Bretagne, BRE; Normandie, NOR; Pays-de-la-Loire, PAY; Nouvelle-Aquitaine, NOU) show evidence of multiple waves of admixture (the best-guess model could not be inferred for Centre - Val de Loire, CEN) mostly from the same source populations. Estimates of admixture times varied between 1035 and 1540 years ago assuming a generation time of 29 years/generation.

To confirm the proportions of ancestry in relation with these external groups, we performed supervised clustering considering Ireland, Germany and Spain as putative sources of ancestry for the French samples. We found that the Irish component accounts for >75% of the ancestry in -50% of the individuals from Brittany (Fig. 3b), while it contributes to a much lesser extent to the genetic composition of the other regions (8–33%). On the other side of the spectrum, a Spanish-like component accounts for the largest proportion (-70%) in NOU, while German-like ancestry is predominant among individuals



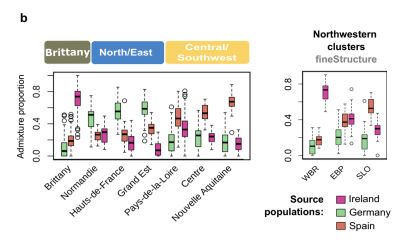


Fig. 3 | **PCA** and supervised admixture analysis of present-day western Europeans. a The two first principal components of genetic variation in the "modern merged dataset", which includes 843 French WGSs for which the four grand-parent origin was concordant, and genome-wide data from 20 central and western European populations ^{12,43} (population acronyms below, see Methods for further details). To avoid bias due to sample size differences among regions, all locations are represented by a maximum of 100 individuals resulting in a total of 2070 samples and 201,999 independent SNPs (MAF > 4%). For simpler visualisation samples from Norway, Sweden, and Denmark were labelled as "Scandinavia", samples from Belgium and Germany were labelled as "Central Europe". France-GenRef samples are represented by dots and labelled with boxes while non-French

samples are represented by 2*sd of the two PCs. UK samples other than those from the POBI dataset, whose origin of the grandparents is known, are not shown. Similarly, French samples from public datasets are also not shown. To better visualise the distribution of our samples the plot is a zoom of the full PCA (Fig. S20). **b** Ancestry proportions retrieved from a Supervised Admixture analysis considering Ireland, Spain, and Germany as proxy source populations on the basis of their polarised positions relative to the distribution of the 843 French samples on (a). In the boxplot the central mark and edges indicate the 50th (median), 25th and 75th percentiles. Whiskers encompass the largest or smallest point comprised within 1.5× of the interquartile range from both edges.

from Northern and Eastern France (NOR, HAU, GRA), with its proportion increasing with the geographical proximity to the German border.

When considering the three clusters inferred by fineSTRUCTURE in Northwestern France, we observe the same ancestry pattern for WBR as for Brittany, with a major Irish component. Consistently with this result, the smallest F_{ST} values between the WBR group and other non-French populations were retrieved with the Irish and Northern Irish populations (0.00057 and 0.00062, respectively, Fig. S23). The only French population showing lower pairwise F_{ST} with WBR is the neighbouring EBP (F_{ST} = 0.00028, Fig. S23). Finally, among the three clusters, SLO carries the largest Spanish-related ancestry while EBP carries similar proportions of the Irish- and Spanish-related components, reflecting its intermediary position between WBR and SLO. In summary, our results indicate that people from Brittany show strong genetic affinities with populations from Western Britain and Ireland, although separated by the Celtic Sea.

To measure the relationship between the genetic clusters found in Northwestern France and other European populations, we used the outgroup f₃-statistics to assess the genetic drift shared by pairs of populations relative to the outgroup population (Mbuti)^{47,48}. Given that this statistic reflects the length of the branch from the internal node to the outgroup (connecting the pair of populations being tested), it is not affected by lineage-specific genetic drift, contrarily to F_{ST}. We found that most French clusters located north of the river Loire share the largest drift with Southwestern Welsh populations (i.e., from Dyfed), whereas those located south to the river Loire share the largest drift with the Basques (Fig. 4a, Supplementary Data 1). The f_4 -statistics of the form f_4 (Mbuti, French subgroup; Dyfed, X), which should produce significantly positive values when the tested population shares more alleles with X than Dyfed, show that the French subgroups from areas south to the river Loire consistently share more alleles with the Basques than with the Southwestern Welsh. Conversely, those located north to the river Loire share the largest amount of alleles with Southwestern Welsh and other populations from Great Britain and Scandinavia (Fig. 4b, Fig. S24).

Genetic continuity since mediaeval times

To disentangle the sources of ancestry contributing to the genetic makeup of present-day French, we merged our WGS data with a large publicly available collection of ancient (>3000 samples⁴⁹ including ~400 ancient Vikings⁵⁰) and >5000 present-day samples⁴⁹ (see studyspecific references in SOM). In addition, to increase our resolution in detecting changes in ancestry during the Mediaeval Period, we sequenced DNA libraries of four ancient samples with radiocarbon dates ranging from the 375-1024 cal. AD, from Pays-de-la-Loire (Fig. 5a) and two samples from archaeological sites dated to the 6-7th century. Pseudo-haploid calls were obtained for the set of autosomal SNPs within the Human Origins Array (HOA - 593,124 SNPs). Average coverage across HOA sites ranged from 0.12 to 4.15x and the number of covered SNPs varied from 64,438 to 574,582 (Table S2). PCA resulting from projecting the ancient individuals onto the principal component space of modern variation shows that most of the samples fall well within the distribution of present-day French (Fig. S25). Out of the six individuals, one (fra009, Table S2) likely represents a migrant with genetic affinities to present-day North Africans. This individual, whose radiocarbon dates are 414-548 cal. AD, was found in an archaeological site located in an ancient town likely built during the Roman period (see SOM, Supplementary Archaeological Details). The exact reason for the presence of migrants of North African ancestry so far north is difficult to pinpoint. Nevertheless, it agrees with the recently reported high levels of mobility in historical Europe⁵¹. Although two of our samples (fra008 and fra017) appear to have levels of contamination >10%, this is not the case for fra009 (contamination ~4%, Table S2), which led us to exclude contamination as a source of its outlier position. To test whether French Mediaeval samples from the 4 to 6th century and samples from the 6 to 7th century significantly differ in their genetic affinities to other ancient European populations we computed the f_4 -statistics of the form f_4 (Mbuti, ancient European sample; sLoire France 4-6th, sLoire France 6-7th). We found no significant differences in allele sharing between individuals from early (fra001 and fra004) and later Mediaeval Period (fra016 and fra017, Supplementary Data 2). Therefore, we considered individuals from

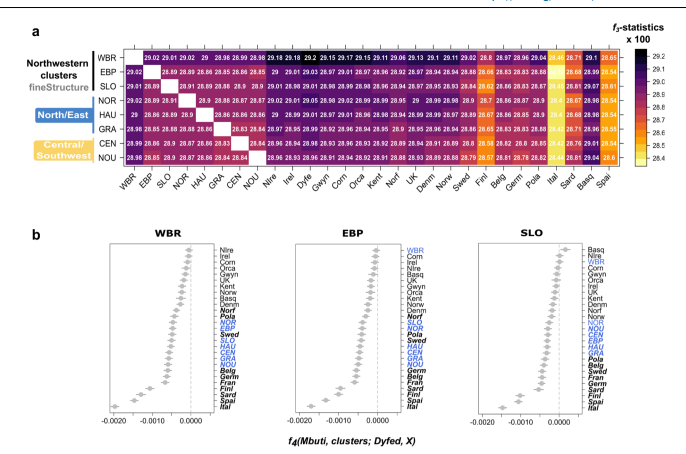


Fig. 4 | **Genetic affinities between Brittany and Ireland/West British Isles from** *f-statistics*. **a** Heatmap reporting outgroup *f3-statistics* in the form f3(Mbuti; French population/cluster, X), where French population/cluster is represented on the y-axis and X on the x-axis. **b** f_4 -statistics of the form f_4 (Mbuti, French cluster; Dyfed, X), where X are the populations on the right side of the plot. Due to disparities in samples sizes, both the outgroup f_3 - and f_4 -statistics were computed on a n=25 samples/population, excepting for the Mbuti (n=10), Basque (n=20), Normandy (n=19) and the Orcadians (n=13). In bold are represented f4-statistics values with a corresponding |Z| > 3 and ranges correspond the standard deviation (sd) from *weighted block jackknife*. Cluster acronyms: western Brittany (WBR),

eastern Brittany and *Pays-de-la-Loire* (EBP), south Loire (SLO). Region/country acronyms: Brittany (BRI); *Pays-de-la-Loire* (PAY); *Normandie* (NOR); *Hauts-de-France* (HAU); *Grand Est* (GRA); *Centre-Val de Loire* (CEN); *Nouvelle-Aquitaine* (NOU), Basque Country (Basq), Belgium (Belg), Cornwall (Corn), Denmark (Denm), Dyfed (Dyfe), Finland (Finl), Germany (Germ), Gwynedd (Gwyn), Ireland (Irel), Italy (Ital), Kent (Kent), Norfolk (Norf), Northern Ireland (NIrel), Norway (Norw), Orkney Islands (Orca), Poland (Pola), Sardinia (Sard), Spain (Spai), Sweden (Swed), United Kingdom (UK). Plots were generated with the R statistical package⁸⁹ and the heatmap with the library *lattice*.

both periods to represent the same population and refer to them as "Mediaeval French".

To check for ancestry changes since the Mediaeval Period, we tested whether Mediaeval French individuals are a good proxy source for the ancestry of present-day Northwestern French. To do so we used the modelling approach implemented in qpAdm⁵² and we tested oneway models (i.e., genetic continuity) using the five Mediaeval French individuals (all but the fra009) as surrogate source population for the present-day French. Our results support genetic continuity between the eight present-day French populations (WBR, EBP, SLO, NOR, HAU, GRA, CEN and NOU) and Mediaeval French (P > 0.05, Supplementary Data 3). qpAdm modelling results are consistent with the PCA showing that Mediaeval French from Pays-de-la-Loire fall within the distribution of modern individuals from the same region. Due to constraints in sample overlap between our SNP-array and WGS datasets, the randomly selected individuals within WBR do not cover the full geographic distribution of the initial cluster (départments: Morbihan, Finistère and Côtes-d'Armor) with the *départment* of Finistère being mostly underrepresented. Hence, we tested a one-way model using only individuals from the westernmost département (Finistère) of Brittany and we found evidence (P = 0.0169) for discontinuity between Mediaeval samples and present-day individuals from Finistère. Due to the fact that model fit p values can be affected by factors such as sample size and coverage, p values comparison should be avoided. Nevertheless, given that we kept the same sample sizes (n = 25 with the exception of NOR with n = 19) and differences in coverage should be minimal due to the quality of our sequencing, it is tempting to argue that the lack of continuity when using only samples from Finistère might reflect ancestry variation already present during mediaeval times. An alternative scenario could, for instance, include later migrations from a non-French source likely through the sea. We also found that one-way models using ancient Northwestern Europeans, such as those dating to the Roman period in Great Britain, fit the ancestry of French populations north of the Loire (0.074 < P < 0.936; WBR, EBP, NOR, HAU, and GRA; Table S3) but do not fit the ancestry of those south of the river (P < 0.05, CEN, SLO and NOU). Furthermore, Italians dating from the early modern period with a central European ancestry fit modern French from south of the river Loire (P > 0.05). Interestingly, ancient Spanish reported as Celtiberians or from the Mediaeval period do not generally fit contemporary French (with the exception of NOU) and only Spanish individuals associated with the Germanic invasions, especially individuals associated with Visigoth or Carolingian archaeological remains, are suitable proxies for contemporary French ancestry (Table S3).

We also tested two-way models combining Mediaeval French with other ancient or present-day (non-French) populations

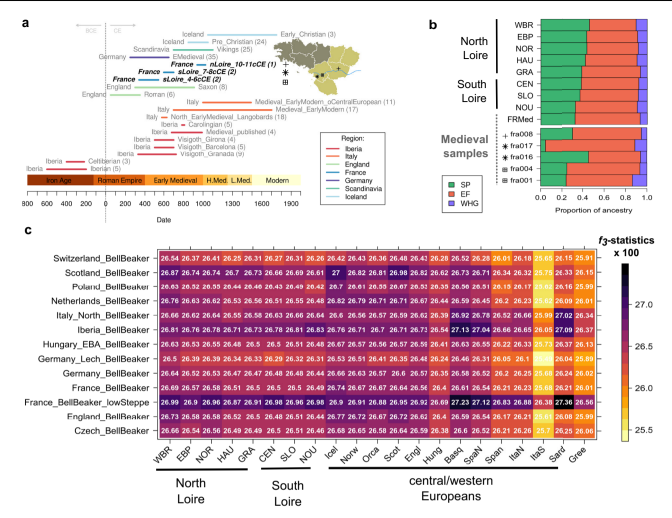


Fig. 5 | **Genetic continuity, admixture and relationship between modern and ancient French populations.** a Timeline of the ancient individuals used to test one- and two-way models of admixture for the ancestry of present-day French using *qpAdm*. The numbers in brackets represent the sample sizes. **b** Mixture proportions from the three major populations contributing to the European ancestry using three-way admixture model, where each population on the left (y-axis) is modelled as a mixture of Mesolithic Western Hunter-Gatherer- (WHG), Neolithic early

farmers- (EF) and Early Bronze Age Steppe- (SP) derived ancestry. $\bf c$ Shared drift between Bell-Beaker-associated individuals and present-day Europeans by means of the outgroup f_3 -statistics of the form f_3 (Mbuti; Bell Beaker Pop, X), where X is the population on the x-axis. These analyses were performed on the "ancient merged dataset" (49 , see Material and Methods for details). Plots were generated with the R statistical package 89 and the heatmap as explained in Fig. 5a.

(Supplementary Data 3). We found that models fitting the WBR ancestry involve populations from the British Isles, Norway or Iceland (0.27 < P < 0.96). Indeed, the proportion of ancestry derived from present-day English or English Romans as proxy sources exceeds 0.90 (standard error of 0.26 and 0.23, respectively) versus less than 0.10 from Mediaeval French. Moreover, we observed that two-way models involving Northwestern Europeans/Scandinavians and Mediaeval French fit the ancestry of the other present-day French (EBP, NOR and HAU) on the English Channel coast (P > 0.13, Supplementary Data 3). However, the Scandinavian or Northwestern European contribution in these groups is smaller than that found in WBR. For EBP, NOR, HAU, GRA, CEN and SLO, we noted that two-way models including a central European proxy source - such as early Mediaeval individuals from Germany or modern Hungarians - also fit the data with estimated contributions as large as 0.70 and 0.63 to NOR and EBP, respectively. However, they do not represent a good fit for the ancestry of WBR or NOU, in agreement with the low proportions of Germany-related ancestry previously observed with the supervised clustering analysis (Fig. 3b). In sum, *qpAdm* modelling shows relative population continuity between the Mediaeval period and present-day times in Northwestern France while emphasising a north/south Loire split with respect to the genetic contribution from Northwestern or Mediterranean Europe, respectively. Although genetic ancestry from countries in Northwestern European is found across the multiple populations dwelling north to the river Loire, WBR is the population in which such ancestry is found at its maximum.

Large genetic affinities between Brittany and ancient Bell Beakers

Using a three-way admixture model, we then estimated the ancestry contributions from the three major ancient populations that spread across Europe - western hunter-gatherers (WHG), early farmers (EF) and steppe pastoralists (SP) – to present-day French and confirmed the aforementioned results (Fig. 5b and Table S4). SP ancestry proportion for regions bordering the Channel (HAU, NOR, EBP and WBR) varies from 42% to 46% versus less than 40% among CEN, SLO, NOU and GRA, and reaches its maximum in WBR where it is similar, if not larger than EF ancestry (46% \pm 2.1% and 43.6% \pm 2%, respectively, Table S4). Consistently, significant negative values (Z-score < -4, Table S5) for f_4 statistics of the form f_4 (Mbuti, Russia EBA Yamnaya Samara; WBR, other

modern French) revealed increased allele sharing between Early Bronze Age Yamnava pastoralists from the Eurasian steppe and WBR relative to other present-day French samples. In agreement with these elevated levels of steppe-like ancestry found in WBR, the results of the outgroup f_3 -statistics of the form f_3 (Mbuti; Bell Beaker, present-day Europeans) and the f_4 -statistics of the form f_4 (Mbuti, Bell Beaker; WBR, other modern French) show that Bell Beaker-associated individuals from Northwestern Europe - who are reported to carry large amounts of steppe-related ancestry - share large levels of drift and significantly increased allele sharing (Z-score < -3) with WBR relatively to other present-day French samples (Fig. 5c and Fig. S26). f₄ statistics of the form f_4 (Mbuti, Western Hunter-gatherer; WBR, other modern French) were mostly not significant (except when the other present-day French was either NOU or GRA, see Table S5), which allow us to exclude the possibility that the genetic affinities between WBR and ancient Bell-Beaker-associated individuals could be caused by a significantly increased WHG ancestry uniquely in WBR. Altogether, these results indicate that the largest contribution of steppe-related ancestry in present-day France is found across the regions located north to the river Loire, and that WBR shares levels of steppe ancestry only comparable to other populations living along the northwestern shores of the European continent (e.g., Ireland, Scotland, Orkney Islands, Iceland and Norway).

Discussion

By exploring whole genomes of present-day individuals originating from Northwestern France in comparison to those from neighbouring French and European populations, we provide novel insights into the demographic events shaping the genetic makeup of the Northwestern edges of the European landmass.

First, we found that population structure in the northern part of France is accompanied with variation in genetic affinities with Northwestern versus Southwestern Europeans. In particular, observations based on supervised admixture analysis, f_3 -outgroup statistics and pairwise F_{ST} values consistently attest connections between Western Brittany, Western Great Britain (e.g., Wales and Cornwall) and Ireland (Fig. 4). These findings confirm previous reports of strong genetic affinities between Brittany and Ireland relative to the general population of Great Britain⁴⁵. They were further supported by haplotypebased methods, which are more powerful than allele frequency-based methods to capture differential contributions from closely related populations^{30,53}. Indeed, by inferring ancestry profiles from a set of surrogate sources with GLOBETROTTER, we highlighted the ancestry contribution from Irish populations into present-day Bretons (~24%) and found relatively smaller contributions from Wales or Cornwall (~9and ~3-fold, respectively). Importantly, we detected some Irish ancestry across all the French regions we surveyed, leading us to hypothesise a long history of shared ancestry likely on the basis of the genetic makeup of the Celtic (Iron-Age) Gaul. The ancestry contributions of Wales and Cornwall exclusively found in Brittany, albeit very limited, are likely associated with more recent migrations. The importance of ancient migrations between the British Isles and Northwestern France has been previously proposed based on ubiquitous haplotype sharing between samples from France and the British Isles, especially samples from Western Great Britain and Northern Ireland⁵. Such results fit with the reported higher frequencies of mutations associated with cystic fibrosis, hemochromatosis and lactase persistence shared between Brittany and Ireland^{45,54,55}. In parallel, our observations reveal larger genetic affinities between populations from Northern France and present-day individuals from Germany (Figs. 3 and 4). As for presentday French populations located south to the river Loire, genetic proximity is observed with contemporary Spanish, and signals of shared ancestry with the Basques. Basque people have recently been described as a typical Iberian Iron Age population whose ancestry was not influenced by later admixture into Iberia⁵⁶. These results are compatible with a scenario where Atlantic France south of the Loire shares the Iron Age legacy of the Basques, while it diverges from the Basques likely due to higher levels of gene flow associated with later incoming migrations (e.g., Germanic invasions) or simply through isolation by distance from other regions to the north and to the east.

Considering the relative ancestry contributions from ancient populations prior to 2000 BCE that spread across Europe, we provided evidence for differential distribution of steppe vs. Early Neolithic ancestry between the two sides of the river Loire, with Neolithic ancestry being more prevalent south of the river. These findings likely explain the shared ancestry between present-day Atlantic French located south of the river Loire (such as Nouvelle-Aquitaine) and the Neolithic-enriched Basque population. Similarly, the relatively high proportions of steppe ancestry found north to the river Loire and along the coast of the English Channel (Fig. 5b and Table S4) - and in particular in Brittany - might explain the genetic affinities observed with Ireland and the Western Great Britain (Figs. 3 and 4). Present-day Irish exhibit a strong signal of continuity with the geographically close Early Bronze Age individuals carrying high levels of steppe ancestry $(-39 \pm 8\%)$ and among whom was found the earliest presence of the hemochromatosis mutation⁵⁷. In Ireland and in the rest of the British Isles, the introduction of steppe ancestry, which led to a considerable turnover of the contemporaneous genetic makeup, has been recently linked to the migration of individuals associated with the Bell Beaker complex from Northern/Central Europe^{18,57}. The lack of human ancient DNA from northern France - dating from the period between Copper Age and Early Iron Age - has hampered our understanding of such genetic turnover across the northern coast of France. Given (i) the elevated levels of steppe ancestry found among present-day French located along the English Channel coast and (ii) the high degree of allele sharing observed between present-day individuals from Brittany and Bell-Beakers-associated individuals carrying high levels of steppe ancestry (Fig. 5), we hypothesise that a similar degree of turnover has reached northern France and Brittany after the arrival of steppe ancestry. Previously reported Late Iron Age individuals from the region of Normandy were found to carry considerable levels of mitochondrial steppe ancestry and strong affinities with Bronze Age samples from the British Isles¹⁹, suggesting extensive gene flow across the English Channel during the Bronze Age. This hypothesis is supported by metalwork-based archaeological evidence dating from the Middle Bronze Age, which points to the existence of a British Channel metalworking core area⁵⁸. Indeed, metal-based relationships between Ireland and Brittany may have started earlier. In Ireland, Bell Beaker pottery is closely associated with the introduction of copper metallurgy ~2400 BC. However, while Irish Bell Beaker culture shows strong influence from Britain, the introduction of metalwork appears to have occurred through Atlantic Europe¹⁷. Metalwork identified in Brittany from the Chalcolithic and Early Bronze Age suggests exchanges of metals such as copper and tin between the regions of Brittany, Ireland⁵⁹ and Cornwall¹⁷. Archaeological evidence exists for extensive connections between Brittany and Ireland/Western Britain dating as early as 4500 BCE and related to the phenomenon of Megalithic monument construction. However, genetic data presented in this study do not support a scenario where the close relationship between Brittany and Ireland/Western Britain can be simply explained by a larger sharing of alleles pre-dating the arrival of the steppe ancestry. Samples associated with the Megalithic period from the British Isles exhibit similar levels of allele sharing with all the present-day samples from France (Table S6), in contrast with increased allele sharing with Corded-Ware- and steppe-enriched Bell Beakers-associated samples uniquely found in Brittany (Fig. S26 and Table S7). Furthermore, we found no differences in genetic affinities between present-day French and the ancient samples related to the two Neolithic waves - the Mediterranean and the Danube waves - that could alternatively explain

both the differences between north and south of the river Loire and the close relationship between Brittany and Ireland (Table S8).

In contrast with the pattern found for the distribution of Neolithic and steppe ancestries, we found that fractions of western hunter gatherer (WHG) ancestry do not differ across the two margins of the river but instead tend to decrease inland (Fig. 5b. Table S4). Larger WHG proportions in present-day coastal French populations might reflect a longer coexistence between late Mesolithic populations and newly incoming farmers due to the significant exploitation of marine resources by the former communities⁹, as it seems to have been the case in Brittany. However, a denser sampling of present-day populations from central and eastern France is required to test this hypothesis. Together our results point to a scenario where the migration of people associated with the Bell Beaker complex culture from North/ Central Europe considerably influenced the genetic makeup of Northwestern France during the Bronze Age by introducing stepperelated ancestry all along the French coast of the English Channel. However, whether this occurred through direct influence of the Bell Beaker incomers from the east or through extensive contacts with the British Isles is an important question that can only be addressed through the study of human ancient DNA from Northern France from the Copper to the Early Iron Age. Importantly, Brittany is the place where the genetic legacy associated with the introduction of steppe ancestry is currently the strongest. This likely indicates relative isolation from later continental migrations, which seem to have increased Neolithic ancestry eastwards. Our admixture modelling approach lends support to this view by showing a lack of genetic affinities between Central European and Brittany, as one- or two-way models involving such populations (e.g., Hungarians, Germans) do not fit the data for Western Brittany (Supplementary Data 3). This contrasts with the results for all the remaining present-day French populations. The introduction of this Central European component, probably less rich in steppe ancestry, might reflect the influence of Germanic peoples in present-day France during the first millennium CE similarly to what has recently been reported in England⁶⁰. Nevertheless, we do not exclude the possibility that the shift towards a Central European component, which we observed across multiple analyses with the "merged-modern dataset" (Figs. 3 and 4), could be explained by earlier migrations.

We found Mediaeval samples from Western France to carry generally less steppe ancestry than their geographical close present-day populations (Fig. 5b). This is consistent with a model in which the introduction of steppe ancestry in Northern French ~2000 BCE remained restricted to the coastal or near coastal regions for centuries. Nevertheless, we recall that Early Mediaeval samples display substantial genetic heterogeneity as two samples carry contrasting proportions of steppe- vs. Neolithic-ancestry (fra016 and fra017, with the latter showing high levels of contamination ~10%) and one sample (fra009) did not fit the genetic diversity of present-day France. Instead, this sample seems to originate from North Africa and provides evidence for long-distance migration between the northern part of France and northern Africa, as early as the Early Mediaeval period (~5-6 century CE). Finally, we found a lack of genetic continuity between Mediaeval French and Iberian populations dating from the first millennium BCE (Supplementary Data 3, P < 0.05). Signals of genetic continuity were only found with Iberian individuals archaeologically associated with Germanic invasions, suggesting that until late Antiquity and Early Mediaeval Period (3th-10th century CE) French and Iberians might have kept low levels of gene flow.

Focusing on the genetic structure of the present-day populations from Northwestern France, global clustering patterns based on fineS-TRUCTURE overlap with the distribution of surnames as well as the linguistic boundaries between varieties of Breton, which are spoken in the westernmost part of the Peninsula of Brittany, and Gallo-Romance varieties, spoken across the eastern part of the peninsula and the whole neighbouring region of *Pays-de-la-Loire* (Fig. 1). The role of

linguistic relationships in the worldwide patterns of structure of human populations has long been recognised⁶¹. Within-country studies, such as the one we present here, have not only validated previous claims but also showed that such a relationship holds at local geographical scales, as it has recently been shown in Northern Spain and in Great Britain^{3,5}. While we found that Western Brittany forms a distinct cluster based on haplotype similarity and rare allele sharing, it is worth noting that genetic differentiation increases from the westernmost tip of the peninsula of Brittany southwards, and especially south of the river Loire. The differentiation between Western Brittany and regions south of the river Loire is larger than that found along the west-east axis (between the *département* of *Finistère* and that of *Sarthe*) north of the river Loire, despite similar geographic distances (~350 km). The role of the river Loire as a geographical barrier to the movement of people has been recently proposed due to the overlap between genetic cluster boundaries, inferred across the whole country, and the natural occurrence of the river³⁷. In the Netherlands, rivers have also been shown to locally restrict gene flow⁶², but the general role of water bodies in promoting or impeding human movements remains unclear. A good understanding of the role of the river Loire on the local patterns of migration requires associating the river's features with genetic differentiation measures along its full extension, which, in our case, is not possible due to the absence of samples along its upstream drainage area. Altogether, the structure patterns observed within Northwestern France contrast with the simple vision that genetic differentiation increases with the geographical distance⁴⁴.

At finer scales, present-day data revealed extensive population structure across Brittany and Pays-de-la-Loire, despite the overall low levels of differentiation (F_{ST} = 0.00043). Within Brittany, at intermediate resolution levels, cluster borders broadly overlap with watercourses (Fig. S14 and Supplementary Discussion). Furthermore, low population differentiation and increased IBD sharing between the cluster "Vannes" and other clusters eastwards on one side, and between the westernmost clusters "Léon" and "Cornouaille" on the other side, suggest a dichotomy between northwest and southeast. Such division is in agreement with the accruing geo-linguistic evidence for a bipartition of Breton varieties into two groups - the Breton varieties spoken in the northwest and those spoken in the southeast of the peninsula of Brittany, whose distributions coincide with the ranges occupied by the Gaulish (Celtic) Ossimii and Veneti, respectively⁶³. Noteworthy, the cluster "Bretagne-centre" roughly overlaps with the border of the Ossimii and Veneti territories, suggesting relative genetic homogeneity likely due to extensive gene flow at this border. Such homogeneity is also recognisable at the level of linguistic features (Figs. S12, 13 and S26, see Supplementary Discussion for details). Finally, we also found genetic clusters with no apparent geographical barrier at their border, such as "Vannes" and "Guérande". Overall, the genetic structure observed within Brittany appears consistent with a complex scenario of interaction between geographical, cultural and likely economic factors influencing population connectivity. Located south to the river Loire, the region of Mauges - which exhibits the highest density of clusters - is situated at the southeastern tip of the Armorican Massif and encompasses hedged farmlands crossed by steep valleys. We hypothesise that this landscape contributed to relative isolation and increased genetic drift and lasted until about 300 years ago, as estimated in the dataset (assuming a generation time of 29 years/generation, Fig. S16). Such a scenario could explain the increase in haplotype sharing for long, and likely recent, chromosomal segments (Fig. S9). Other population-specific features, such as recent ancestry caused by inbreeding, could equally explain such strong clustering patterns. However, levels of relatedness were not found significantly larger than in other clusters (Figs. S2-4). This ultra-finescale structure with geographically restricted clusters resembles that observed in the region of Galicia, Spain³ and emphasises the need for whole-genome sequencing studies on local populations in order to

better understand the distribution of rare, and likely more deleterious, variants across space^{1,64}.

In conclusion, the patterns of genetic structure observed across Brittany and the downstream Loire basin mainly reflect the likely combined effect of linguistics and geographic features. This lends further support to the idea that local population differentiation exists. as it has been shown within other countries^{3,5}, and can be detected at geographical distances as small as a few tens of kilometres even in the absence of major geographical barriers. Within this structured genetic landscape, Brittany reveals a history of isolation from the rest of France together with a strong legacy of the important genetic changes (i.e., the introduction of steppe ancestry) that followed the arrival, in Northwestern Europe, of people associated with the Bell Beaker complex from north-central Europe ~2500 BCE. A similar scenario has been proposed for the present-day Celtic populations in the western part of the British Isles - the Welsh and Irish - who display elevated haplotype sharing with Bronze Age samples⁵⁷ and strong genetic affinities with pre-Anglo-Saxon samples from Britain⁶⁵. Our results imply that: (1) a scenario in which Mediaeval population movements along the English Channel and the Atlantic facade, such as those related to incoming Western Britons in Brittany during the 4-6th century CE or the Viking incursions ~8-9th century CE, are unlikely to be the main explanation for the close genetic between the northwestern edges of the European continent; and (2) a long-history of shared ancestry between Brittany and Ireland/Western Britain followed by their relative isolation explains the sharing of disease-related alleles such as those associated with hemochromatosis, cystic fibrosis and lactase persistence.

Methods

France administrative division overview

This study relies on a dense sampling of the geographical region of western France, which crosses multiple administrative subdivisions of the territory. Among these subdivisions are the (1) regions - the largest administrative units - which are subdivided into (2) *départements*. A *département* is subdivided into (3) *arrondissements*, which at the smallest scale are divided into (4) town centres. Given the specificity of the administrative system we kept the French name for the *département* and *arrondissement* along the manuscript.

Cohort description

The PREGO study ("Population de référence du Grand Ouest", https:// umr1087.univ-nantes.fr/prego-biobank) collected DNA from 5707 healthy persons originating from western France (Regions Pays de la Loire and Bretagne). Individuals were recruited during 295 blood drives organised by the French Blood Service (EFS in French) carried out between February 2014 and March 2017, with a mean of 19 donors per blood drive. Blood drives were spatially and temporally sampled in order to obtain a coverage as homogeneous as possible of the nine départements included in the study. Priority was given to blood drives taking place in rural areas. Participants should be native of western France. Individuals' birthplace was assessed by the place of birth of the four grandparents. Only individuals whose four grandparents were born in western France and preferably within a radius of 30 km were included in this study. From the 3234 individuals included in the present study, 25%, 50% and 75% have their grandparents born 3.25, 6.38 and 12.33 kms from each other, respectively. Venous blood samples (6 ml) were collected from recruited individuals by venipuncture into Vacutainer tubes. Participants filled out a questionnaire providing grandparents', parents' and their own birthplaces, residence, age, sex and information about previous participation in the study (of the individual itself or another member of the family). Neither phenotypic nor clinical data was collected in the present study. The PREGO study received approvals from the local ethical committee of Nantes (Comité de Protection des Personnes), the Advisory Committee on Information Processing for Health Research (CCTIRS) and the National Commission on Informatics and Liberty (CNIL). Participants signed a written informed consent for participation in the study, inclusion in bioresource and personal data processing.

The FranceGenRef study aims to describe patterns of population diversity across metropolitan France at the beginning of the 20th century. Thus, individuals were sampled based on the birthplace of their grandparents, whose distance should not exceed 30 kms. FranceGenRef includes 862 individuals satisfying the aforementioned criteria: 354 blood donors from the PREGO cohort (described above) with origin in the départements of Côtes d'Armor (COT), Ille-et-Vilaine (ILL), Morbihan (MOR), Loire-Atlantique (LOI), Maine-et-Loire (MAI), Mayenne (MAY), Sarthe (SAR) and Vendée (VEN); 50 blood donors from Finistère (FIN); 458 individuals from the GAZEL cohort (www.gazel.inserm.fr/ en)66,67 originating from five other regions of France: Normandie, Hauts-de-France, Grand East, Centre-Val de Loire and Nouvelle-Aguitaine. The GAZEL study received approvals from the National Commission for Data Processing and Freedoms (CNIL), the National Medical Council and the National Consultative Committee of Ethics. All individuals signed informed consent for genetic studies at the time of enrolment. DNA samples from GAZEL samples were extracted at the CEPH Biobank on an automated system either Autopure (Qiagen) or Chemagic Prime (PerkinElmer) using respectively the salting out method or magnetic beads and were quantified using fluorimetry (Quant-iT DNA Assay kit, Broad Range, Thermo Fisher Scientific).

Genotyping, whole-genome sequencing and quality control (QC)

SNP array genotype processing. Under the scope of PREGO, genomic DNA was extracted from peripheral blood lymphocytes by standard protocols. Out of 5707 collected samples, 3385 were genotyped on the Axiom TM Precision Medicine Research Array (~920,000 markers, ThermoFisher). Standard QC was performed using SNPolisher software (http://tools.thermofisher.com) and SNPs not passing the QC report were removed according to the manufacturer's instructions. SNPs with a missing rate >5%, minor allele frequency <10% and not in Hardy-Weinberg Equilibrium ($p < 10^{-6}$) were excluded from the dataset resulting in a total of 210.171 SNPs. Relatedness was assessed via the PI HAT statistics, which provides an estimation of the proportion of the genome shared by two pairs of individuals (i.e., identity-by-descent, IBD). PI_HAT statistics was computed using PLINK (vs.1.9). After removing related individuals with a PI_HAT >= 0.08, there were 3234 samples left for analysis. Genotyping was conducted in three batches of 971, 1266, and 997 individuals, respectively. To investigate potential batch effects, we employed linear regression using a batch indicator variable on the first five principal components. We found no significant association at the significance level of 0.05.

Whole-genome sequencing and variant calling. Whole-genome sequencing of 856 individuals was performed at CNRGH (Evry, France) using their standard workflow (www.cnrgh.fr). All the samples were sequenced at high coverage (average coverage 30x). Read processing was performed with GATK 3.8 following the "best practices" NGS pipeline recommendations of the Broad Institute (https:// software.broadinstitute.org/gatk/best-practices). Reads were mapped on the GRCh37 reference genome using bwa-mem, duplicates were removed and reads were then realigned and recalibrated according to GATK best practices. Variant calling was performed with GenotypeGVCF. GVCF files were then merged using GATK CombineGVCF function, recalibrated and annotated with SnpEff⁶⁸ and the gnomAD database (https://gnomad.broadinstitute.org/). The resulting GVCF files were filtered out in order to only keep SNPs with high mapping quality (MQ > 30). Furthermore, only SNPs in Hardy-Weinberg equilibrium (HWE, $p = 10^{-5}$) and less than 10% of missing data were kept for analysis. Related samples were identified using PI HAT statistics

computed with PLINK (vs.1.9) and excluded when values were >0.10. Individuals whose grand-parents were not born in the same *département* were also excluded leaving us with 843 samples. The sizes of the samples used in each analysis are specified below and in every corresponding figure.

aDNA Mediaeval samples

We generated whole-genome data for six ancient individuals from three different archaeological sites in western France, specifically from the region of Pays-de-la-Loire. Samples fra001, fra004, fra008 and fra009 were dated using radiocarbon methods and estimates vary from 375 to 1024 cal. AD (Table S2, Figs. S27-S29). This interval corresponds to the early and High Mediaeval periods. The dates of the other two samples - fra016 and fra017 - are based on the archaeological context⁶⁹. Out of the six ancient individuals, four (fra001, fra004, fra016, fra017) were sampled south of the river Loire while two were sampled north of the Loire (fra008, fra009, Fig. 5). The archaeological excavations in Saint-Lupien, Rezé (south shore of the Loire) took place between 2005 and 2016 and were led by the team of Mikaël Rouzic under the request of the city of Rezé. The site located in Chaussé Saint-Pierre, Angers (north shore of the Loire), was excavated by the team of Martin Pithon, from the French Institute for Preventive Archaeological Research (INRAP) and the excavations occurred between July and August 2009. Based on the archaeological remains and radiocarbon dates, the site shows evidence of occupation from the beginning of the Roman Empire to modern times. The project was initialised under the request of the city of Angers given the plan for public construction affecting the archaeological site. The archaeological study of the site was authorised by the Regional Division for Cultural Affairs (Délégation Régional des affaires culturelles) and the INRAP. Finally, the excavations in the archaeological site in Chéméré (south shore of the Loire) started in the 60 s but the two individuals sequenced in this study belong to a group of 181 individuals found in the last excavations in 2007. These excavations were led by an INRAP archaeological team under a project of preventive archaeology before construction of a pavilion. For more details on the description of the sites see Supplementary Material Online (Supplementary Archaeological Details).

aDNA library preparation and bioinformatic processing

One DNA extraction (following a modified version of the extraction method from⁷⁰⁻⁷²) and one to two single indexed blunt end libraries⁷³ were prepared for each of the six ancient samples (fra001, fra004, fra008, fra009, fra016 and fra017). All DNA libraries were doublestranded. DNA extractions and library preparations were performed in the dedicated clean aDNA-laboratory at the Uppsala University, Sweden. The DNA libraries were sequenced in two batches and over multiple lanes, first as a pilot run at the SciLife Sequencing Centre in Uppsala, Sweden, using Illumina HiSeq 2500 with paired-end 125 bp chemistry, and later in more depth at the CNRGH (Evry, France) using Illumina HiSeq X and with a paired-end 150 bp chemistry. Raw reads were trimmed with CutAdapt version 2.3⁷⁴ using the parameters: --quality-base 33, --quality-cutoff 15, -e 0.2, --trim-n and --minimumlength 15. Overlapping read pairs were then merged using FLASH version 1.2.11⁷⁵ and with parameters: --min-overlap 11, --max-overlap 150 and --allow-outies. Merged fastq files were mapped to the human reference genome hs37d5 as single end reads using bwa-aln version 0.7.1776 and parameters -l 16500, -n 0.01 and -o 2, as suggested for ancient DNA77,78.

BAM files were merged to a per sample library level using the merge command in Samtools version 1.5⁷⁹ followed by PCR duplicates removal (reads with identical start and end positions were identified and collapsed) using a modified version of FilterUniqSAMPCons_cc.py, which ensures random assignment of bases in a 50/50 case, as described in ref. 80. All reads longer than 35 base pairs and with less than 10% mismatches to the reference were kept, and a final merging

step was performed for those samples with two sequenced libraries by merging the processed sample library BAMs to a final sample BAM.

Processed sample BAM files were then used to call pseudo-haploid genotypes using Samtools and the option *mpileup -R -B -q30 -Q30*. In order to merge with available datasets, the pseudo-haploid calling was carried out on the 593,124 autosomal sites on the Human Origins Array (Affymetrix). Contamination was estimated based on the X-chromosome and mitochondrial DNA using ANGSD⁸¹ and schmutzi⁸², respectively. Sample quality, inferred sex and contamination estimates are shown in Table S2.

Merging with publicly available datasets

Publicly available datasets of Western Europeans. To investigate the relationship of modern individuals from the north part of France and other European populations we merged the WGS dataset with three available genome-wide datasets encompassing a large number of Western European samples: (1) the EGAD0000000120 from the The International Multiple Sclerosis Genetics Consortium and the Wellcome Trust Case Control Consortium 2 (referred to hereafter as the MS dataset)⁴³, (2) the EGAD00010000124 from the Genetic Analysis of Psoriasis Consortium & the Wellcome Trust Case Control Consortium (referred to hereafter as PS dataset)⁴², and (3) the EGAD00010000632 from the Peopling of the British Isles (referred to hereafter as POBI⁵). In the MS and PS datasets samples were genotyped on the Human670-QuadCustom SNPchip encompassing 580,030 autosomal sites and the datasets include 11,376 and 2622 individuals, respectively. MS dataset includes samples from: Australia, Belgium, Denmark, Germany, Finland, France, Italy, New Zealand, Northern Ireland, Norway, Poland, Spain, Sweden, US and UK, whereas the PS dataset includes samples from UK and Ireland. In the POBI dataset, 2912 individuals from the UK were genotyped on the Human1-2M-DuoCustom SNPchip encompassing 1,115,428 autosomal sites. For the three datasets, original genotype likelihood files (.gen) were converted into plink format files with gtool (vs. 0.7.5). Only individuals and sites passing quality criteria thresholds (provided with the datasets) in the original studies were kept. Genotypes were called using a probability cut-off of 0.90. Sites containing alleles in the negative strand were flipped according to the corresponding strand file https://www.well. ox.ac.uk/~wrayner/strand/ using PLINK vs.1.9. First, we checked whether the alleles were in the illumina TOP configuration, as required to flip based on the strand file. Sites not found in TOP configuration were removed from the datasets. We used the liftOver tool (https://genome. ucsc.edu/cgi-bin/hgLiftOver) to convert the physical coordinates to hg19 as they were originally in hg18 in the MS, PS and POBI datasets. We merged the samples from Belgium, Denmark, Germany, Finland, France, Italy, Northern Ireland, Norway, Poland, Spain, Sweden and UK with the Irish samples from the PS dataset. In a second step, we merged this dataset with a subset (to keep the sample size computationally tractable) of the POBI dataset comprising two samples from Wales (Dyfed and Gwynedd), the sample from Cornwall and two samples from Norfolk and Kent (eastern UK). Finally, we merged the dataset above with seventy samples from the available Human Genome Diversity Project dataset (https://www.hagsc.org/hgdp/files.html) genotyped on the Illumina 650Y SNPchip array and originated from Sardinia, the Basque Country, Orkney Islands and the Mbuti (to use as an outgroup). Chromosomal coordinates for the HGDP dataset were lifted over as described above. From this merged dataset we filtered out sites not in Hardy-Weinberg Equilibrium (HWE, $p = 10^{-5}$), removed those falling into the HLA region (chr6:29,691,116-33,054,976) and removed all the multiallelic sites. The final dataset, hereafter referred to as "merged modern dataset", encompassed 9704 samples and 433,940 SNPs.

Human origins array and Viking datasets. We merged the French WGS dataset with the Human Origins Array (HOA) dataset

V42.4 (https://reich.hms.harvard.edu/allen-ancient-dna-resource-aadrdownloadable-genotypes-present-day-and-ancient-dna-data) encompassing 3589 ancient and 6472 present-day individuals genotyped for 593.124 autosomal SNPs. From the original dataset we extracted European samples (Austria, Belgium, Czech Republic, Denmark, France, Germany, Great Britain, Greece, Hungary, Iceland, Ireland, Italy, Luxembourg, Netherlands, Norway, Poland, Portugal, Russia, Spain, Sweden, Switzerland, Turkey) with PASS tag under the "Assessment" field in the annotation file [see references in the Supplementary Discussion]. From this subset we removed samples whose individuals' ID contain *Ignore* and with less than 50,000 genotypes called to avoid potential bias. This dataset was then merged with 405 ancient DNA samples belonging to Vikings from the study: "Population Genomics of the Viking world"50 and previously called for HOA set of SNPs. This dataset is referred to hereafter as "merged ancient dataset". Mbuti samples (HGDP-CEPH, Human Genome Diversity Project-Centre d'Etude du polymorphisme Humain - https://cephb.fr/en/hgdp panel. php) from the HOA were also extracted to compute f-statistics (see below).

ChromoPainter and fineSTRUCTURE

PREGO's SNP-array dataset (post-QC) was phased with SHAPEIT v2.r79083 using the genetic map build 37 provided with the software and no reference panel. Phased genotype files were converted into CHROMOPAINTER⁴⁶ format using the impute2 chromopainter2.pl script. Switch (global Ne) and emission rates (µ) were estimated with CHROMOPAINTER version 2 using data on chromosomes 1, 4, 10, and 15 from 330 individuals (around 10% of the total sample). The estimated parameters were then used to run CHROMOPAINTER on the full data. A Principal Component Analysis (PCA) on the coancestry matrix (chunkcounts output) was performed in R. Coancestry matrix estimates the proportion of the genome of each individual that is most closely related to every other individual in the matrix. In particular, chunkcounts matrix is based on the number of copied haplotype chunks. We assigned individuals to clusters using the model-based approach implemented in fineSTRUCTURE version 2.1.3. We ran fineSTRUCTURE version 2.1.3 on the coancestry matrix with 10.000.000 burn-in iterations and 1.000.000 MCMC iterations, from which every 10,000th iteration was recorded. We kept the default values for the other options. The tree was built using 100,000 tree comparisons and 10,000,000 additional optimisation steps. MCMC convergence was assessed by comparing the individual assignment to clusters in a second independent chain. Confidence measures of cluster assignment and visualisation of coancestry matrix were performed with the help of the R library FinestructureLibrary provided with the software. The tree provided by fineSTRUCTURE software is based on posterior probabilities of the population configuration (ie. individual partition across clusters), which is highly dependent on the sample sizes and does not reflect differences between pairs of clusters^{31,84}. Therefore, as an alternative we computed a tree based on the total variation distance (TVD) as firstly described by Kerminen and colleagues³¹. TVD measures the distance between the clusters and it is not affected by differences in sample sizes. TVD values are computed from the copying profiles of all individuals, estimated with CHRO-MOPAINTER, and the cluster assignment inferred with fineS-TRUCTURE as described in ref. 31. It results in a final symmetrical matrix with as many rows and columns as the number of clusters inferred (K), where each entry represents the average closest ancestry contribution in cluster k coming from other clusters. We computed the TVD matrix on the final fineSTRUCTURE partition (k = <154, finest level of FS-tree) and the TVD-tree was obtained using complete linkage hierarchical clustering.

We assessed the performance of the two tree building approaches on the PREGO dataset. The performance was measured by comparing cluster assignment confidence as computed in

Leslie et al.⁵. As expected, cluster assignment confidence decreases with increasing number of clusters (Fig. S6). Moreover, we found that for the same k the FS-tree shows lower cluster assignment confidence than the TVD-based tree, confirming the better performance of the latter. To visualise the relationship between the clusters (Fig. 1d), we arbitrarily chose k = 39 because it provides a large enough number of clusters to access fine-scale structure while keeping cluster assignment confidence >90% (Fig. S6). At this level of the tree, clusters containing 1-5 individuals were merged into the closest cluster with >=31 individuals or removed if the closest cluster included itself <4 individuals. This approach resulted in a TVD-based tree of 18 clusters performing better than FS tree, which achieves similar values of cluster assignment confidence for only 12 clusters. Finally, we tested whether inferred clusters capture significant differences in ancestry by following the approach of ref. 5. To do so, we randomly reassigned the individuals to clusters, maintaining the cluster sizes, and computed the p-value of the population configurations produced by the k = 18. The TVD-based tree was less likely than a random distribution of individuals across pairs of clusters. We performed 1000 permutations to obtain p values, which were computed as the number of permutations where random assignment resulted in a higher value of TVD between any pair of clusters and the total number of permutations. For the level of k=18, p values were <0.001 for all pairs of clusters.

Assessing sampling effect on fineSTRUCTURE clustering. We investigated whether the sampling scheme carried out in this study, which densely selects individuals representing every location over four generations, increases the probability of having recently related individuals. This recent ancestry may affect clustering patterns. To check the effect of recent ancestry on the fineSTRUCTURE clusters we computed a relatedness measure based on identity by state (IBS) within clusters, for k = 3, 18 and 78 (only for those clusters with n > 10 at the finest scale level k = 154), and compared it with the overall distribution. With the exception of k = 154, where six clusters have a median IBS-statistics larger than the 75% of overall IBD distribution, across the other values of k, most of the clusters do not show increased relatedness (Figs. S2–S4) and therefore, we conclude that recent ancestry is not driving the results.

Surname analysis

The list of surnames associated with birth registration data and occurring in two periods (period one: 1891-1915 and period two: 1816–1940) were retrieved from the French Institute for Statistics and Economic Studies, the INSEE. Surname lists are available at the town level. To analyse the distribution of surnames we proceed as follows. (1) For each town, we chose the surnames present in the two periods and registered at least four times, ie. given to four newborns. Such an approach removes very rare names often associated with spelling errors in the registration and rare emigrants from other regions of France or elsewhere. (2) summed the number of surname occurrences over the two periods at the arrondissement level. There are 35 arrondissements within the ten selected départments. (3) We computed the Arccos distance (68) between the 35 arrondissements and (4) constructed a consensus tree based on 1000 bootstrapped distance matrices obtained by the Neighbour Joining method. Finally, we built a map linking the arrondissements in a nested way according to the bootstrap values of the NJ tree. The threshold for grouping arrondissements was 90% bootstrap robustness while the threshold for higher order grouping is 85%.

Correlation between surname based Arccos distance and Fst was performed using Mantel distance matrix correlation test using Spearman rank correlation. The physical distance between arrondissements was done using partial correlation. Surname distribution diversity index were entropy and the Barrai index. Let N be the population size

in a geographical area and S the number of different surnames and p_i the probability of Surname *i*..entropy = $\left[-\sum_{i=1}^{S} pi * Ln(pi)\right]$

Effective migration surface (EEMS) analysis

We estimated effective migration surfaces using the software EEMS³². The matrix of average pairwise genetic dissimilarities was generated for 82,362 SNPs (after prunning: --indep-pairwise 50 5 0.2) and 1414 individuals using the bed2diffs software included in the EEMS package. Samples were assigned to the nearest of 300 demes. We run ten independent MCMC chains, each with a random seed, for 10,000,000 iterations, including 9,900,000 burn-in iterations, thinning every 200 iterations. From the chain with the highest final log-likelihood, and we started a second round of ten EEMS chains using this chain as a starting point for 1,000,000 additional sampling iterations, thinning every 9999 iterations. Log- posterior trace of ten replicate MCMC chains from the last round shows mixing and convergence of the independent EEMS runs. Plots were generated in R statistical software using the *rEEMSplots* package.

PCA analysis and Fst computation

PCAs was carried out using the smartpca software from the EIGEN-SOFT package version $6.1.4^{86}$. F_{ST} values were obtained by setting up the option *fsthiprecision: YES*. Both analyses were performed after pruning for linkage disequilibrium using a sliding window of 50 SNPs in steps of 5 SNPs and keeping those SNPs with $r^2 < 0.50$ (Figs. 3a and S20). The regions previously reported as exhibiting long-range linkage-disequilibrium⁸⁷ were excluded before performing the PCAs.

Runs of homozygosity (ROH), Identity by descent (IBD) and IBDNe estimations

Individuals' runs of homozygosity (ROH) and identity by descent (IBD) segments were calculated using RefinedIBD (version from 23rd December 2017, with default settings) based on the PREGO dataset. Individuals' ROH were computed by summing the total length in ROH segments per individual and averaged across individuals within *départments* and *arrondissements*. We computed IBD sharing by counting the number of IBD segments shared between pairs of individuals assigned to the 18 clusters inferred by fineSTRUCTURE. This was done independently for IBD segments of length: 1–2 cM, 2–7 cM and longer than 7 cM.

We estimated the trajectories of effective population size with IBDNe³³ (version released on 7th May 2018). IBD segments were identified with IBDseq (version r1206, with default settings). As suggested by Browning and Browning 2018⁸⁸, we removed regions with excess of IBD to avoid potential bias, such as the Major Histocompatibility Complex (MHC) on chromosome 6 (chr6:26291527-33464061). We thus split chromosome 6 into two continuous parts. To evaluate the robustness of the effective population size trajectories across different IBD segment sizes, we varied the *mincm* parameter, which sets up the minimum length of IBD segments used by IBDNe. Value of *mincm* should be chosen considering SNP density on the SNP array.

Rare variation analysis

We investigated allele sharing patterns using doubletons (alleles that are present in only two chromosomes or minor allele count MAC = 2) and variants with MACs between 3 and 10 computed from the WGS dataset encompassing 620 individuals from Northwestern France. We first randomly selected 1 million sites from each dataset and allele sharing matrices were computed by summing all the variants shared between pairs of chromosomes of different individuals from Brittany and *Pays-de-la-Loire*. We plotted the results as a heatmap and performed hierarchical clustering (*hclust* function in R with method = "complete") to identify clusters of highly correlated allele sharing. The resulting tree was cut at multiple levels (k = 2-10) and the proportion

of individuals, within each *département*, assigned to the alternative clusters was plotted onto the map of Northwestern France. These analyses were performed using the R statistical package⁸⁹ together with the following libraries: *ComplexHeatmap*, *rgdal*, *sp*, *broom*, *ggplot2* and scatterpie.

Supervised admixture analysis

The supervised clustering analysis was performed using ADMIXTURE vs1.3 software⁹⁰ on the WGS dataset including the 843 French samples used in the PCA. We assumed that modern French originate from three modern source populations: Spain, Germany and Ireland, which lie in the extremes of the distribution of modern French individuals in the PCA (Fig. 3a). Given sample size differences (see section *Publicly available datasets of western Europeans*), we down-sampled the source populations from Germany and Ireland to 350 individuals. SNPs in linkage disequilibrium were removed by following the recommended practises in the software manual, ie. PLINK option --indep-pairwise 50 10 0.1. Only sites with minimum genotype rates <10% were analysed.

Ancestry profiles with GLOBETROTTER

To further investigate the contribution of external populations to the genetic makeup of France we estimated ancestry contributions from their European neighbours using the statistical approach implemented in GLOBETROTTER software⁴⁶. This method identifies and describes recent (<4.5 ky old) admixture events, by providing admixture times and proportions, explaining the haplotypic ancestry of a target population. Given that GLOBETROTTER infers admixture events from patterns of haplotype sharing it requires two input files: the "copy vectors" and "painting samples" generated by the CHROMOPAINTER. To obtain these files we ran CHROMOPAINTER following the author's recommendations (section 6.2 "GLOBETROTTER") on the phased "merged modern dataset". The "copy vectors" files (*.chunklengths.out) were combined using the scripts provided with the software package. To detect signals of admixture we first ran GLOBETROTTER with the option *null.ind*: 1 and then computed the number of bootstraps providing a date of admixture >= 1 and <= 400 generations. This step provides a p value that can be interpreted as the evidence of "detectable admixture". P values were computed using 100 bootstraps. Confidence intervals for the inferred admixture times were estimated by running 100 additional bootstraps and with the option: null.ind: 0. We phased the "merged modern dataset" using SHAPEIT and no reference panel, in a similar manner to what was done with the PREGO dataset. To keep the analyses computationally tractable and due to imbalanced sample sizes, we randomly sampled 350 individuals from countries with large sample sizes such as Italy, Germany, Norway and Belgium. To represent the haplotypic diversity of the UK and the Scandinavian Peninsula we only kept the samples from PoBI and Norway, respectively. A total of 3,598 samples were used to estimate ancestry contributions. Proportions of admixture were plotted with the R statistical package⁸⁹.

Admixture analysis: f_3 -, f_4 -statistics and qpAdm estimates

We computed the outgroup f_3 -statistics of the form f_3 (Outgroup; Pop1, Pop2), as implemented in Admixtools⁴⁷, using the Mbuti population as an outgroup. Pop1 and Pop2 were either a pairwise combination of a modern French and any other European population or a Bell Beaker sample and a present-day western European. The outgroup f_3 -statistics quantifies the amount of shared history between pairs of populations in relation to an outgroup under a three-population tree. Provided that the outgroup is equally distant from Pop1 and Pop2, the outgroup f_3 -statistics represents the length of the branch from the outgroup to the internal node if no gene flow occurred between Pop1 and Pop2. To statistically assess the presence of admixture we computed f_4 -statistics using the same software package. Admixture was also computed in order to test whether modern and ancient Mediaeval French show

signals of admixture f_3 -statistics by setting these populations as the target. Admixture f_3 -statistics was computed using the option inbreed: YES in the case of the ancient Mediaeval as recommended by the authors of the programme.

gpAdm, implemented in the Admixtools, is a computer programme that assesses the goodness of fit of admixture models, involving a user-defined set of commonly known as "left" and "right" populations, and estimates the proportion of admixture within a regression framework. The set of "left" populations encompasses the target population and the possible sources of admixture, whereas the "right" populations include a set of populations distantly related to the "left". Importantly, the model assumes that the "right" populations are differentially related to the "left" and exchange of genes between "left" and "right" must not have occurred after the event of admixture, i.e., genetic drift shared between "left" and "right" derives from deep shared evolutionary history. The method implemented in *qpAdm* is built on a matrix of f_4 -statistics computed across all possible combinations of "left" and "right" populations (f_4 (target, source; R1, R2), where R1 and R2 is any possible pair of "right" populations). P > 0.05together with admixture proportions varying between 0 and 1 are indicative of a good fitting model.

We performed the *qpAdm* analyses to estimate the contribution of the three major European migrations^{52,91} using western huntergatherers (WHG), Neolithic early-farmers (EF) and Bronze Age steppe pastoralists (SP) as "left" populations together with the Mediaeval and modern French populations. We grouped ancient individuals into WHG, EF and SP based on the ancestry proportions inferred in Mathieson et al. 92. Only individuals with ancestry proportions >0.90 were included into the three groups giving origin to sample sizes of 73, 42 and 17 for the EF, WHG and SP respectively. Admixture proportion estimates were computed using 25 randomly selected modern individuals for the three genetic clusters inferred by fineSTRUCTURE (WBR, EBP, and SLO) and the remaining five regions of France (NOR, HAU, GRA, CEN and NOU). With respect to the French Mediaeval samples we inferred admixture proportions at the population (five Medieval French, the outlier was removed from this computation) and individual level. The set of *right* populations used in these analyses was the following: Mbuti, Han, AfontovaGora3, ElMiron, Karitiana, Kontenki14, MA1, Mota, Papuan, Ust Ishim, Vestonice16, Villabruna, GoyetQ116-1, Morocco_Iberomaurusian.

In order to test for continuity of the genetic ancestry of Mediaeval and present-day French populations we tested one-way models. In the case of Mediaeval French, the single source is a neighbouring population (no post-Neolithic sample from France is available in the merged dataset) of the same period or of the preceding one. In the case of present-day French, the single source included the Mediaeval French and all the possible sources tested for the Mediaeval French. Furthermore, we also tested two-way models where the genetic ancestry of modern French is assumed to result from a mixture between the Mediaeval French and any other ancient population from the neighbouring regions from Mediaeval to present-day times. Given the larger availability of Iberian samples and the close geographical proximity with France we included in the aforementioned analyses all the Iberian samples from the 5th century BCE to the Mediaeval period (up to the 10th century CE). Only targets and source populations with a sample size larger than two were used in these analyses.

The early Neolithic period is characterised by the arrival of the first farmers in Western Europe and the Early Bronze Age is characterised by the arrival of populations related to or carrying a large proportion of their ancestry from pastoralist populations from the Eurasian steppes^{52,91}. Therefore, we added Anatolia_N and Russia_E-BA_Yamnaya_Samara to the set of *right* populations: Mbuti, Han, AfontovaGora3, ElMiron, Karitiana, Kontenki14, MAI, Mota, Papuan, Ust_Ishim, Vestonice16, Villabruna, GoyetQ116-1, Morocco_Iberomaurusian, Anatolia N and Russia EBA Yamnaya Samara.

Reporting summary

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

Data availability

Data from the International Multiple Sclerosis Genetics Consortium & the Wellcome Trust Case Control Consortium 2 (EGAD00000000120). the Genetic Analysis of Psoriasis Consortium & the Wellcome Trust Case Control Consortium 2 (EGAD00010000124), and the Peopling of the British Isles (EGAD00010000632) are available in the EGA repository. The HGDP data set is available at https://www.hagsc.org/hgdp/files.html. The HOA SNP genotypes for the Viking samples are now available in version 54.1 of the Allen Ancient DNA Resource (https://reich.hms. harvard.edu/allen-ancient-dna-resource-aadr-downloadable-genotypespresent-day-and-ancient-dna-data). The WGS data on ancient DNA produced in this study (bam files) are available from the European Nucleotide Archive (ENA) under accession number PRJEB71835. Genetic data on contemporary human individuals are subject to restrictions in accordance with French legal requirements regarding the protection of identifiable personal data. Access to the data will be provided under the terms of a signed agreement as described at https://umr1087.univnantes.fr/prego-biobank. In short, the PREGO array genotyping data are available from the European Genome-Phenome Archive (EGA) under accession number EGAD00010002661. Access to WGS data from the FranceGenRef study - subject to compliance with French legislation on identifiable personal data and its compatibility with the legislation of the requesting country - should be requested from richard.redon@univnantes.fr.

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Author contributions

I.A., J. G., M.G.B.B., C.Férec., J.J.S., H.L.M., M.Jakobsson., E.G., J.F.D., R.R., and C.D. contributed to the conception and study design. C.Bernhardsson., S.C., A.S.P., A.F.H., R.O., M.M., V.G., E.Cabot., E.S., D.B.D., E.B., C.Besse., H.B., O.B., A.B., S.B., E.Charpentier., P.D., C.D.R., C.Fruchet., S.L., E.L.F., T.E.L., G.M., V.M., F.R., K.R., F.S., J.J.S., L.V.S., J.V., E.W., M.G., M.Z., H.L.M., M.Jakobsson, E.G., J.F.D., R.R. and C.D. contributed to data collection. I.A., J.G., C.Bernhardsson., M.K., E.S., C.Berthellier., E.Charpentier., E.Q., P.D., C.D. contributed to data analysis. I.A., J.G., M.G.B.B., A.F.H., M.M., V.G., E.Cabot., F.R., Y.C., M.Jézéquel., D.L.B., C.N., Y.P., P.D., R.R., C.D. contributed to data interpretation. I.A., J.G., R.R., C.D. drafted the manuscript. I.A., J.G., M.G.B.B., C.Bernhardsson., S.C., M.K., A.S.P., A.F.H., R.O., O.B., A.B., E.Charpentier., F.R., J.J.S., L.V.S., M.Jézéquel., D.L.B., C.N., Y.P., M.Jakobsson., P.D., E.G., J.F.D., R.R., C.D. contributed to and revised the manuscript.

Competing interests

The authors declare no competing interests.

Additional information

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Correspondence and requests for materials should be addressed to Richard Redon or Christian Dina.

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Isabel Alves ® ^{1,22,23}, Joanna Giemza ® ^{1,23}, Michael G. B. Blum², Carolina Bernhardsson ® ³, Stéphanie Chatel¹, Matilde Karakachoff¹,⁴, Aude Saint Pierre⁵, Anthony F. Herzig ® ⁵, Robert Olaso ® ⁶,ⁿ, Martial Monteil³, Véronique Gallien 9,¹0, Elodie Cabot 9,¹1, Emma Svensson³, Delphine Bacq ® ⁶,ⁿ, Estelle Baron ® ¹, Charlotte Berthelier ® ¹, Céline Besse ⁶,ⁿ, Hélène Blanché ® ¹², Ozvan Bocher⁵, Anne Boland ® ⁶,ⁿ, Stéphanie Bonnaud ® ¹, Eric Charpentier¹, Claire Dandine-Roulland ⁶,ⁿ, Claude Férec ⁵,¹¹³, Christine Fruchet¹, Simon Lecointe¹, Edith Le Floch ⁶,ⁿ, Thomas E. Ludwig ® ⁵,¹¹³, Gaëlle Marenne ® ⁵, Vincent Meyer ⁶, Elisabeth Quellery¹, Fernando Racimo ® ¹⁴, Karen Rouault ⁵,¹³, Florian Sandron ⁶,ⁿ, Jean-Jacques Schott¹, Lourdes Velo-Suarez ⁵, Jade Violleau¹, Eske Willerslev ® ¹⁵,¹6, Yves Coativy ® ¹ⁿ, Mael Jézéquel¹ⁿ, Daniel Le Bris¹¹, Clément Nicolas¹³, Yvan Pailler¹9, Marcel Goldberg ® ²⁰, Marie Zins ® ²⁰, Hervé Le Marec ® ¹, Mattias Jakobsson ® ³, Pierre Darlu²¹, Emmanuelle Génin ® ⁵,¹³, Jean-François Deleuze ® ⁶,ⁿ,¹², Richard Redon ® ¹,²⁴ ⋈ & Christian Dina ® ¹,²⁴ ⋈

¹Nantes Université, CHU Nantes, CNRS, INSERM, l'institut du thorax, Nantes, France. ²TIMC-IMAG, UMR 5525 CNRS, Univ. Grenoble Alpes, Grenoble, France. ³Department of Organismal Biology, Human Evolution, Uppsala University, Uppsala, Sweden. ⁴Nantes Université, CHU Nantes, Pôle Hospitalo-Universitaire 11: Santé Publique, Clinique des données, INSERMCIC 1413, Nantes, France. ⁵Univ Brest, Inserm, EFS, UMR 1078, GGB, Brest, France. ⁶Université Paris-Saclay, CEA, Centre National de Recherche en Génomique Humaine (CNRGH), Evry, France. ⁷Labex GenMed, Evry, France. ⁸Nantes Université, CNRS, Ministère de la Culture, CReAAH, LARA, Nantes, France. ⁹INRAP - Institut national de recherches archéologiques préventives, Paris, France. ¹⁰CEPAM UMR7264 - Culture et Environnements, Préhistoire, Antiquité, Moyen-Age, Nice, France. ¹¹Anthropologie Bio-Culturelle, Droit, Ethique et Santé, Faculté de Médecine Site Nord, Marseille, France. ¹²Fondation Jean Dausset, CEPH, Paris, France. ¹³CHRU Brest, Brest, France. ¹⁴Section for Molecular Ecology and Evolution, Globe Institute, University of Copenhagen, Copenhagen, Denmark. ¹⁶Department of Zoology, University of Cambridge, Cambridge, UK. ¹⁷Centre de Recherche Bretonne et Celtique, UR 4451, Université de Bretagne Occidentale, Brest, France. ¹⁸CNRS UMR 8215 Trajectoires, Université Paris 1 Panthéon-Sorbonne, Centre Malher, 9 rue Malher, Paris, France. ¹⁹CPJ ArMeRIE UBO, UMR 6554 LETG, CNRS, Université de Brest, Université de Nantes, Université de Rennes 2, Institut Universitaire Européen de la Mer, Plouzané, France. ²⁰Université Paris Cité, "Population-based Cohorts Unit", INSERM, Paris Saclay University, UVSQ, Paris, France. ²¹UMR 7206 Eco-anthropologie, Musée de l'Homme, MNHN, CNRS, Université de Paris Cité, Paris, France. ²²Present address: Université de Strasbourg, CNRS, GMGM, Strasbourg, France. ²³These authors contributed equally: Isabel Alves, Joanna Giemza. ²⁴These authors jointly supervised this work: Richard Redon, Christian Dina.