A dynastic elite in monumental Neolithic society

Author list
Lara M. Cassidy, Ros Ó Maoldúin, Thomas Kador, Ann Lynch, Carleton Jones, Peter C. Woodman, Eileen Murphy, Greer Ramsey, Marion Dowd, Alice Noonan, Ciarán Campbell, Eppie R. Jones, Valeria Mattiageli and Daniel G. Bradley

Affiliations:
1 Smurfit Institute of Genetics, Trinity College Dublin, Dublin 2, Ireland
2 The Irish Fieldschool of Prehistoric Archaeology, National University of Ireland, Galway, Ireland
3 Archaeological Management Solutions, Consultancy, Kilrush, Clare, Ireland
4 UCL Arts and Sciences, University College London, London, WC1E 6BT, UK
5 Former Chief Archaeologist, National Monuments Service, Department of Culture, Heritage and the Gaeltacht, Custom House, Dublin
6 School of Geography and Archaeology, National University of Ireland Galway, Galway, Ireland
7 Department of Archaeology, University College Cork, Cork, Ireland
8 Archaeology & Palaeoecology, School of Natural & Built Environment, Queen’s University Belfast, Belfast BT7 1NN, Northern Ireland
9 National Museums NI, Cultra, Holywood BT18 0EU, Northern Ireland
10 CERIS, School of Science, Institute of Technology Sligo, Sligo, Ireland
11 Department of Zoology, University of Cambridge, Downing Street, Cambridge CB2 3EJ, UK
† Deceased
The nature and distribution of political power in Neolithic Europe remains poorly understood\(^1\). During the period, many societies began to invest heavily in monument-building, suggesting an increase in social organisation. The scale and sophistication of megalithic architecture along the Atlantic seaboard is particularly impressive, culminating in the great passage tomb complexes\(^2\). While megalith builders have often been proposed as co-operative networks of independent communities, the human expenditure required for the largest monuments has led some to emphasize hierarchy\(^3\), the most extreme case being a small elite marshalling the labour of masses. Here we present evidence that such a social stratum was established during the Irish Neolithic period. In a sampling of 44 whole genomes, we identify the adult son of a first-degree incestuous union discovered within the most elaborate recess of the imposing Newgrange passage tomb. Socially sanctioned matings of this nature are highly rare and occur almost exclusively among politico-religious elites\(^4\), specifically within polygynous and patrilineal royal families headed by god-kings\(^5,6\). We identify relatives of this individual within two other major passage tomb complexes 150km to the west, as well as dietary differences and unprecedented fine-scale haplotypic structure between passage tomb samples and the larger population, implying hierarchy. This system emerged against a backdrop of rapid maritime colonisation that displaced a unique Mesolithic isolate, although Irish hunter-gatherer introgression is detected within the Neolithic population.\(^7\)
Ancient genomes have demonstrated common ancestry between the societies of the Atlantic Neolithic⁰⁻⁹, while recent modelling has defined repeat expansions of megalithic architecture from northwest France, at a pace implying more advanced maritime technology than previously assumed in these regions¹⁰. This includes the spread of passage tombs along the Atlantic façade during the 4th millennium BC, a period which also saw the arrival of agriculture to Ireland alongside other distinct megalithic traditions. These structures reached one of their highest known concentrations and diversities on the island. However the political systems underlying these societies remain obscure, as does the genetic input from indigenous Mesolithic hunter-gatherers.

To investigate, we shotgun sequenced individuals from the Irish Mesolithic (n=2) and Neolithic (n=42) to a median 1.1X coverage (Fig. 1a, Supplementary Tables 1, 2). We imputed 43 of these alongside relevant ancient genomes (Supplementary Table 3), including an additional 20 British and Irish individuals⁷,⁹,¹¹. These were merged with a published imputed ancient dataset¹² to allow for finescale haplotypic inference of population structure¹³ and estimation of inbreeding. Four key individuals were subsequently sequenced to higher (13-20X) coverage.

All major Irish Neolithic funerary traditions were sampled: court tombs, portal tombs, passage tombs, Linkardstown-type burials and natural sites (Fig. 1a, c; Supplementary Information section 1). Within this, the earliest Neolithic human remains from the island, interred at Poulnabrone portal tomb¹⁴, are of majority Early_Farmer ancestry and show no evidence of inbreeding (Extended Data Fig. 1; Fig. 1a), implying that from the very outset agriculture was accompanied by large-scale maritime colonisations. ADMIXTURE and ChromoPainter analyses do not distinguish between the Irish and British Neolithic populations and confirm⁷,⁸ the Spanish Early Neolithic as the best proxy source of their Early_Farmer ancestry (Fig. 1d, Extended Data Figs. 1, 2), emphasising the importance of Atlantic and Mediterranean waterways in their forebears’ expansions.

Overall, no increase in inbreeding is seen through time in Neolithic Ireland, indicating that communities maintained sufficient size and communication to avoid matings of 5th degree relatives or closer (Fig. 1a). However, we report a single extreme outlier interred within Newgrange passage tomb, a focal point of the UNESCO monumental landscape of Brú na Bóinne (Fig. 2a). Incorporating over 200,000 tonnes of earth and stone, this megalithic mound is one of the most spectacular of its kind in
While externally designed for public consumption, internally, the tomb consists of a single narrow passage with specialised ritual inventory, whose winter solstice solar alignment would have been viewed only by a select few. Unburnt, disarticulated human bone was found concentrated within the most elaborately decorated recess of the terminal cruciform chamber, including the cranial remains of an adult male (NG10; Fig. 2b; Supplementary Information 1.4.1). This exceptional location is matched by an unprecedented genomic heritage. He possessed multiple long runs of homozygosity (ROH), each comprising large fractions of individual chromosomes (Fig. 2c; Extended Data Fig. 3a) and totalling to a quarter of the genome (Inbreeding Coefficient = 0.25). This marks him as the offspring of a first order incestuous union, a near universal taboo for entwined biological and cultural reasons. However, given the auspicious nature of the interment, his parentage was very likely socially sanctioned.

While simulations cannot distinguish whether his parents were full siblings or parent and offspring (Extended Data Fig. 3), the only definitive acceptances of such matings occur among siblings, specifically within polygynous elites, as part of a rarely observed phenomenon known as “royal” or “dynastic” incest. In all documented cases (e.g. Hawaii, Inca Empire, Ancient Egypt), this behaviour co-occurs with the deification of political leaders and is typically limited to ruling families, who are exempt from social convention. Both full and half-sibling marriages are found most commonly during the earliest stages of state formation, and are generally viewed as a means to intensify hierarchy and legitimise power in the absence of more advanced bureaucratic systems, alongside tactics such as extravagant monumentalism and public ritual. We propose a comparable set of social dynamics was in operation in Ireland by the Middle Neolithic, and, given the construction of solstice-aligned passage tombs similar to Newgrange in Wales, Orkney and Brittany, may have occurred outside the island as well. Notably, levels of consanguinity are consistently low and decrease through time across our wider ancient dataset (Extended Data Fig. 4), with only one other incidence of close inbreeding detected, the son of 2nd-3rd degree relatives from a Swedish megalith.

The Brú na Bóinne passage tombs appear in Medieval mythology, which relates their construction to magical manipulations of the solar cycle by a tribe of gods, leading to unresolved speculation about the durability of oral tradition across millennia. Such longevity seems unlikely but,
surprisingly, our results strongly resonate with mythology first recorded in the 11th century AD, which has a builder-king copulate with his sister to restart the daily solar cycle. A Middle Irish placename for the Dowth passage tomb which neighbours Newgrange, Fertae Chula, is based on this lore and can be translated as “Hill of Sin” or “Hill of Incest”.

A second centre of the passage tomb tradition is found 150km west near the Atlantic coast. Here, the mega-cemeteries of Carrowmore and Carrowkeel have origins pre-dating the construction of Newgrange by several centuries, with depositions at Carrowkeel continuing until at least the end of the Neolithic. Using both SNP- and haplotype-sharing analysis, we uncover a web of relatedness connecting these sites to both Newgrange and the atypical Millin Bay megalith on the northeast coast, recognised as part of the passage tradition for its artwork and morphological features.

Firstly, using lcmlkin (Fig. 2c), we find the earliest passage tomb genome in the dataset (car004), interred within the focal monument at Carrowmore, has detectable distant kinship with NG10, as well as with other later individuals from Carrowkeel and Millin Bay (CAK533 and MB6). A similar kinship coefficient (≥6th degree) is also seen between NG10 and CAK532 (Extended Data Fig. 5a), demonstrating familial ties between several of the largest hubs of the tradition.

Secondly, in a fineSTRUCTURE analysis of Atlantic Neolithic genomes, samples from Newgrange, Carrowkeel and Millin Bay form a distinct cluster, which is split from a larger British and Irish grouping (Fig. 1d, e). The robustness of this cluster is confirmed using a larger ancient dataset (Extended Data Fig. 2). ChromoPainter also identifies excessive reciprocal haplotype donation specifically between NG10 and CAK532, confirming their kinship (Extended Data Fig. 5b). Evidence of more distant relatedness is seen between the inferred relatives of car004, who share elongated haplotypic chunks with one another; this signature of recent shared ancestry also links CAK530 to CAK533 and NG10 (Fig. 2d; labelled on Fig. 1d).

The earlier car004 genome is of low coverage (0.04X) and thus was excluded from ChromoPainter analysis. However, D-statistics demonstrate that this sample preferentially forms a clade with the passage cluster (Z > 3.4; Supplementary Table 10), despite being closer in time to the majority of samples from the larger British-Irish cluster. Moreover, this attraction is only partially driven by the aforementioned kin connections, which we further corroborate. Downsampling tests on the larger
dataset demonstrate D-statistic results for car004 to be highly significant (Supplementary Tables 10, 11).

Taken together, we favour that the haplotypic structure within our dataset is driven by excessive IBD-sharing between passage tomb samples, implying non-random mating across large territories of the island. A high degree of social complexity would be required to achieve this, as is predicted by the parentage of NG10. However, our non-passage tomb genomes are largely earlier in date and denser sampling of diverse sites from the Late Neolithic will be required to evaluate the contribution of temporal drift to the fineSTRUCTURE clustering. Stable isotope values also differentiate passage tomb interments (Fig. 1b). Their combination of high δ¹⁵N and depleted δ¹³C values is best explained by a more privileged diet of meat and animal products, although it remains to be seen how this relates to broader dietary change during the period.

Simpler court and portal tombs lack the artwork and prestigious grave-goods of the passage tradition, and are arguably a manifestation of smaller lineage-based societies. These architectures do not typically occur within passage tomb cemeteries, although exceptions exist, including a court tomb constructed beside Carrowmore, which showed a potential instance of inter-site kinship. We find evidence of both distant kinship (Supplementary Information section 6) and societal structure between another pair of distinct but neighbouring megaliths (10 km apart) - Poulnabrone portal tomb and Parknabinnia court tomb. Their majority male cohorts show a significant difference in the frequency of two Y chromosome haplogroups (P=0.035, Fisher exact test), as well as dietary difference (Fig. 1b, Extended Data Figs. 5, 6). Given the lack of close kin within either tomb, we exclude small family groups as their sole proprietors and interpret our results as the result of broader social differentiation with an emphasis on patrilineal descent. The double occurrence of a rare Y haplogroup (H2a) among the individualised male Linkardstown burials of the southeast provides further evidence of the importance of patrilineal ancestry in these societies, as does the predominance of a single Y haplogroup (I-M284) across the Irish and British Neolithic (Extended Data Fig. 7).

It is hypothesised that the spread of agriculture into Britain and Ireland was assisted by pre-existing Mesolithic maritime connections. However, our results suggest that prior to the Neolithic the Irish Sea posed a formidable barrier to gene flow. Irish hunter-gatherer (HG) genomes form a distinct
cluster within a wider grouping of Mesolithic HGs from northwest Europe\textsuperscript{11,28}, sharing excessive levels of drift with each other despite over half a millennium’s separation (Fig. 3a, Extended Data Fig. 2, Supplementary Information section 4). In contrast, British HGs show no differentiation from continental contemporaries\textsuperscript{11}. This accords with paleogeographic models positing a Doggerland bridge between Britain and the continent for most of the Mesolithic, but a pre-Holocene separation of Ireland\textsuperscript{29}. 

Irish HGs also exhibit the largest degree of short ROH (Fig. 3b) yet described in any ancient, or indeed modern genome, a signature of ancestral constriction that supports a prolonged island isolation. This implies that continental and British HGs lacked the technology or impetus required to maintain frequent contact with Ireland and reflects the relatively late Mesolithic colonisation of the island, followed by a sharp divergence in lithic assemblages\textsuperscript{30}. Nonetheless, with no signatures of recent inbreeding (Fig. 1a), it appears Irish HGs were capable of sustaining outbreeding networks within the island itself, despite an estimated carrying capacity of only 3,000-10,000 individuals\textsuperscript{30}. 

Ultimately, Irish HGs originate from sources related to Italian Upper Palaeolithic individuals\textsuperscript{28} (Fig. 3a), with no evidence of contribution from an earlier western lineage that persisted in Spain\textsuperscript{31}. However, we detect a significant excess of this ancestry in the Luxembourg Mesolithic relative to Irish and British HGs (Supplementary Table 9), demonstrating its survival outside Iberia. We also explore the genetic legacy of Irish HGs in the island’s Neolithic population and discover an incidence of direct ancestral contribution. Within a broader pattern of high haplotypic affinities among European farmers to local HG groups (Fig. 3c), we uncover an outlier from Parknabinnia (PB675) with a disproportionate and specifically Irish HG contribution. High variance in HG ancestry across the genome and an excess of elongated Irish HG haplotypes support a recent introgression (Extended Data Figs. 8, 9), estimated within four generations (Supplementary Information section 3).

This finding, taken together with evidence of local HG input into the Scottish Neolithic\textsuperscript{11}, implies recurring interactions between incoming farmers and the indigenous populations of the islands. Notably, a ~4th degree relative of PB675 was interred within the same tomb (Extended Data Figs. 5), implying this outlier was integrated within the community. An alternate instance of diversity in those selected for megalithic interment is seen in a male infant from Poulnabrone (PN07) with a dietary signature of breastfeeding (Fig. 1b, Extended Data Fig. 6). This individual has a clear trisomy of
overall, our results demonstrate the capacity of ancient genomes to shed light not only on population movements, but on political systems and social values where no written records exist. This is particularly true when imputation and haplotypic analyses are utilised, which we affirm outperform popular SNP-based methods in the resolution of ancient population structure (Extended Data Fig 10).

Together with estimations of inbreeding and kinship, these methods broaden the scope within which we can study the development of agricultural societies from chiefdoms to civilisations. Specifically, our findings support a re-evaluation of social stratification and political integration in the megalithic cultures of the Atlantic, with the passage tomb building societies of Ireland exhibiting several early attributes of state formation.
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Materials & Correspondence Requests: Please direct to Lara M. Cassidy or Daniel G. Bradley
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**Fig. 1. Fine-scale haplotypic and dietary structure in the Neolithic.** a, Timeline of analysed Irish genomes with inbreeding coefficients shown for those of sufficient coverage. All dates are direct excluding CAK534 (translucent). Sample site key follows panel e. The earliest widespread evidence of Neolithic activity (cereal/house horizon) is marked with a black line. The Irish Neolithic ends circa 2500 BC. b, Stable isotope values for Irish and British Neolithic samples (n=292). Irish sample key follows c, with those included in aDNA analysis outlined in black. British samples are shown as hollow shapes - Scotland: black, England/Wales: grey; circles: pre-3400 BC, squares: post-3400 BC. A high-trophic-level infant with Down syndrome is labelled. c, Site locations for Irish individuals sampled or included in this study coloured by burial type -court tomb (yellow), portal tomb (blue), Linkardstown-type (green), passage and related (magenta), natural sites (light pink) and the unclassified Ballynahatty megalith (light blue). Sites outlined in black were included in aDNA analysis. d, ChromoPainter\textsuperscript{13} PCA of Atlantic Neolithic genomes (n=57) generated using a matrix of haplotypic length-sharing. Passage tomb outliers in Fig. 2d are labelled. CAK532 is excluded due to kinship. e, fineSTRUCTURE dendrogram derived from the same matrix with five consistent clusters.

**Fig. 2. Genomic signals of dynasty among focal passage tomb interments.** a, Front elevation and interior of Newgrange passage tomb (photo credits: Fáilte Ireland; Photographic Unit, National Monuments Service) b, Plan of chamber after O’Kelly 1983. c, The coefficient of relatedness (\(\hat{P}_i\)) between another auspicious interment from the central monument at Carrowmore, car004, and 38 British and Irish Neolithic samples, with the top five hits labelled. d, Average length of donated haplotypic chunks between all reciprocal pairs of the ‘passage cluster’ (pink; n=42) and ‘British-Irish cluster’ (grey; n=1190) as defined by fineSTRUCTURE in Fig. 1e. Highest values for passage tomb pairs are marked along the x-axis, with an excess of longer chunks shared between the inferred kin of car004 (CAK533, MB6, NG10) in c. Darker lines link reciprocal donations. Combined symbols are used for inter-site pairs. e, A sliding window of heterozygosities is plotted for transversions along selected chromosomes of NG10, revealing extreme ROH.

**Fig. 3. Origins and Legacy of the Irish Mesolithic.** a, Estimates of shared drift between Irish, British
and continental HGs from the Mesolithic and Upper Paleolithic (triangles: Magdalenian culture). Top ten hits with sufficient coverage are cross-compared in a heatmap. b, Short and long ROH spectra in modern and ancient genomes. Hollow shapes indicate direct (rather than imputed) diploid calls. For four Irish samples both imputed and direct data are presented, showing close agreement. c, Normalised haplotypic length donations from HG populations to Neolithic individuals, arranged by geographic region (labelled). The top three donors are outlined for each individual. Donor population colour key follows that in panel b, with British and Northwestern HGs merged into a single population (blue).
Extended Data Fig. 1. Genomic Affinities of the Irish Neolithic. a, ADMIXTURE plots (K=14) for ancient Irish and British populations (first row), other ancient Eurasians (second and third rows) and global modern populations (fourth row). For components that reach their maximums in modern populations, the five individuals with highest values were selected for representation. If the majority of these come from a single population the block is labelled as such, otherwise it is labelled using the general geographic region from which these individuals originate. Three components reach their maximum in ancient populations and we label these European_HG (Red), Early_Farmer (Orange) and Steppe (teal). b, Box plot (Tukey method) showing the distribution of the European_HG component among British and Irish Neolithic individuals (n=50). c, Normalised haplotypic length contributions, estimated with ChromoPainter, from Early Neolithic populations to later Neolithic and Chalcolithic individuals. The top two donors are outlined in black for each individual. Given the unsupervised nature of the analysis, regional differences in overall haplotypic donation levels should be ignored, as larger populations have more opportunity for within group sharing.

Extended Data Fig. 2. Haplotypic structure among ancient populations. a, ChromoPainter PCA of diverse ancient genomes (n=149) generated using the output matrix of haplotypic lengths. Colour and shape key for Irish samples follows Fig. 1. b, fineSTRUCTURE dendrogram derived from the same matrix with the passage tomb cluster highlighted.

Extended Data Fig. 3. Inferring the relationship between NG10’s parents. a, Whole genome plot of heterozygosity in NG10, revealing extreme ROH. b, Nine matings (coloured lines) that can lead to an inbreeding coefficient of 25%. c, Number and average lengths of homozygous-by-descent (HBD) segments for each of these simulated scenarios (500 iterations) and the same values observed for NG10 genome. Box plots follow the Tukey method. Scenarios in i and ii best fit the HBD distribution of NG10, with ii being less parsimonious when anthropological and biological factors are taken into consideration.

Extended Data Fig. 4. Levels of inbreeding through time in ancient populations. Inbreeding coefficients for imputed ancient samples estimated by measuring the proportion of the genome that is homozygous-by-descent. Boxplots follow Tukey’s method. Individuals are binned according to archaeological period - UP-MS: Upper Palaeolithic to Mesolithic (n=24), EN: Early Neolithic (n=13), MN-CA: Middle Neolithic-Chalcolithic (n=69), BA: European Bronze Age (n=12), IA: IA-MA: Iron Age to Medieval (n=21), Steppe CA-BA (n=14). Outliers of...
note are labelled. The inferred degrees of relatedness between an individual’s parents are marked.

Extended Data Fig. 5. Detecting recent shared ancestry between pairs of British and Irish Neolithic samples. a, lcMLkin\textsuperscript{29} kinship coefficients between pairs of Irish and British Neolithic samples, jittered by a height of 0.00018 and width of 0.00036 for visualisation. Optimised duplicate tests are linked by dotted lines. Several standalone values are also shown (italics), where one duplicate did not meet the threshold of overlapping sites (>20,000). The MB6 and car004 pairing (19,850 sites) is shown as translucent point. An inset is shown for lower values of pi-HAT. Pairs over 5σ from the mean pi-HAT and K\textsubscript{0} for panel ii (marked with line) are highlighted using the same colour and shape key as Fig. 1. Combined symbols are used for inter-site pairs. b, Total haplotypic lengths donated between all pairs (n=2162) of British and Irish samples from the ChromoPainter analysis of diverse ancient samples (Extended Data Fig. 2). Outlying pairs (4σ above the mean) are labelled. c Outgroup f\textsubscript{ST} statistics measuring shared drift between pairs (n=2236) of Irish and British Neolithic samples (>25,000 informative sites). d, Total haplotypic lengths donated between all pairs of ‘passage cluster’ (pink; n=42) and ‘British-Irish cluster’ (grey; n=1190) samples from the ChromoPainter analysis of Atlantic Neolithic genomes (Fig. 1d, e).

Extended Data Fig. 6. Regional-scale diversity in the Irish Neolithic. a, Nitrogen stable isotope values, an indicator of trophic level, plotted across time for samples from the neighbouring sites of Poulnabrone (blue) and Parknabinnia (yellow). For male samples, the Y chromosome haplogroup is given. Distant kinship connections are marked with a dotted line, while a closer (3rd-4th degree) relationship is highlighted with a solid line. b, Boxplot (Tukey method) of normalised read coverage aligning to chromosome 21 for shotgun sequenced ancient samples (n=188), with a single trisomy outlier infant.

Extended Data Fig. 7. Subclade distributions of Y chromosome haplogroup I2a1 in Ireland, Britain and Europe from the Mesolithic to Bronze Age. a, Y haplogroups observed for Neolithic individuals in Britain and Ireland. Shape indicates the approximate time period within the Neolithic based on McLaughlin et al. 2016\textsuperscript{17}, while colour indicates haplogroup and follows the same keys as in b, c and d. Approximately 94% of the British/Irish Neolithic population belong to haplogroups I2a1b1 (45%), I2a1a1 (14%) and I2a1a2 (35%). Incidences of these haplogroups in European individuals from the Mesolithic to Bronze Age are shown in b (I2a1b1), c (I2a1a1) and d (I2a1a2). Haplogroup colour keys are shown with respect to phylogenetic placement, with those in bold observed within Britain and Ireland. European individuals who share an identical set of
haplotypic mutations (for sites covered) to an Irish Neolithic individual are highlighted with a black outline in e (I2a1a1) and d (I2a1a2).

Extended Data Fig. 8. Geographic and genomic distributions of northwestern European HG (NWHG) ancestry among British and Irish Neolithic individuals. a, Geographical distribution of NWHG introgression in Britain and Ireland across 103 Neolithic samples. Box plot (Tukey method) highlights four extreme outliers, three from the Early to Middle Neolithic of Argyll and one from Ireland - Parknabinnia675 (PB675). The next highest value is also from Parknabinnia, PB754. b, The same D-statistic run on separate chromosomes individuals of sufficient coverage (n=86). Outlying individuals are marked for each chromosome. Irish outliers follow the same shape and colour key as in Fig. 1 and outliers who are also outliers in the box plot in a are marked in bold. c, Box plot (Tukey method; n=86) of sample standard deviations across the chromosomes for the same D-statistic. Four outliers with high variance across the chromosomes are marked, including three samples from Parknabinnia, two of whom are also top hits in a.

Extended Data Fig. 9. Haplotypic affinities of the Irish and British Neolithic to northwestern hunter-gatherers. Colour and shape key follows Fig. 1. The outlying individual PB675 shows a preference for Irish HG haplotypes in all measures. Regression lines shown with 95% confidence interval shaded (sample size=47). a, Total haplotypic chunk counts donated from Irish HGs (n=2) and other northwestern HGs (n=3). PB675 shows a higher than expected number of Irish HG haplotypes. b, The sum total of Irish and northwestern HG haplotype lengths (y-axis) against the normalised ratio of these two totals (x-axis). PB675 has the highest overall HG haplotypic length contribution, with a ratio skewed towards Irish HG. c, PB675 also displays the longest average length of Irish HG haplotype chunks. This elongation is not seen for haplotypes from other northwestern HGs.

Extended Data Fig. 10. SNP-sharing analyses of autosomal structure in Atlantic Neolithic populations. a, PCA created using an identical sample (n=57) and SNP set (~488k sites; pseudoploidised) as that presented in Fig. 1d, e. b, Outgroup f2-statistics for all combinations of samples in a, using a reduced SNP set (~270k sites; pseudoploidised). Results are presented in heatmap and corresponding dendrogram.

Extended Data Fig. 11. Imputation accuracies for chromosome 22 of the high coverage NE1 genome
downsampled to 1X. The levels of accuracy seen across all SNPs (solid line, n=204,316) is compared to that seen for transversions only (dashed line, n=62,374). Accuracies at different genotype probability (GP) thresholds and minor allele frequency (MAF) filters are shown for the three different genotype categories. MAF filters are based on overall frequency in the 1000 Genomes phase 3 dataset.
Methods

Sampling and Sequencing

We sampled 54 petrous temporal bones and 12 teeth (Supplementary Table 1) sourced from 20 archaeological sites (Supplementary Information section 1). Two of these, PN10 and PN113, were later found to belong to the same individual. Processing was carried out in clean-room facilities dedicated to ancient DNA research at Trinity College Dublin. Photographs were taken prior to sample alteration and these are available upon request to the authors. The dense otic capsule region of petrous bones and the root cementum of teeth were targeted for sampling. Bone/tooth powder (130-150mg) was subject to a described silica-column method of DNA extraction with modifications. Three successive extractions were performed on samples (incubation times of 24 hours at 37°C). Five samples were subject to a modified protocol, with powder first washed twice with EDTA (0.5M) and then subject to a single extraction (incubation time of 48 hours at 37°C).

Select sample extracts, typically the 3rd, were purified at a volume of 55μl and NGS double-stranded libraries were created from 16.50μl aliquots, following previously described methods that are based on established protocol. Library amplification reactions were carried out using Accuprime Pfx Supermix (Life Technology), primer IS4 (0.2μM), a specific indexing primer (0.2μM) and 3μl of library as previously described, and DNA concentrations assessed on an Agilent 2100 Bioanalyzer. Amplified libraries were first screened for endogenous human content on an Illumina MiSeq platform (TrinSeq, Trinity College Dublin) using 65 or 70 bp single-end sequencing. Extracts with sufficient human endogenous content (>5%) and concentration (>0.5 ng/μl at 12 PCR cycles) were incubated with USER Enzyme (volume of 5μl to 16.50μl of extract) for 3 hours at 37°C, to repair post-mortem molecular damage. Following this, library creation and amplification was carried out as described above. USER-treated libraries from a total of 45 individuals were sent for higher coverage sequencing at Macrogen Inc., Seoul, Korea (100 bp single-end with the exception of JP14, for which 100bp paired-end data was also obtained). Detailed experimental and sequencing results are found in Supplementary Table 2.
Demultiplexed data returned in FASTQ format were subject to quality control using the FastQC suite\textsuperscript{17}. Residual adapter sequences were trimmed using cutadapt v1.2.1\textsuperscript{38}, with non-default parameters \texttt{-m 34} and \texttt{-O 1}. Quality trimming was performed on read ends where necessary. Paired-end reads from JP14 were merged and trimmed for adapters using the leeHom software \textsuperscript{39}. Trimmed reads were mapped to hg19/GRCh37 with the mitochondrial genome replaced with the revised Cambridge reference sequence (NC_012920.1). BWA version 0.7.5\textsuperscript{40} was used for alignment with non-default parameters \texttt{-l 16500, -n 0.02} and \texttt{-O 2}. Reads were sorted, filtered for a mapping quality (MQ) of 20 or above and PCR duplicates removed using Samtools v0.1.19\textsuperscript{41}. Quality trimming was performed on read ends where necessary. Paired-end reads from JP14 were merged and trimmed for adapters using the leeHom software \textsuperscript{39}. Trimmed reads were mapped to hg19/GRCh37 with the mitochondrial genome replaced with the revised Cambridge reference sequence (NC_012920.1). BWA version 0.7.5\textsuperscript{40} was used for alignment with non-default parameters \texttt{-l 16500, -n 0.02} and \texttt{-O 2}. Reads were sorted, filtered for a mapping quality (MQ) of 20 or above and PCR duplicates removed using Samtools v0.1.19\textsuperscript{41}. Read groups were added and BAM files merged to sample level using Picard Tools v1.101 (http://broadinstitute.github.io/picard/). GenomeAnalysisTK v2.4-7\textsuperscript{45} was used to locally realign reads. Two base pairs at both the 5' and 3' ends of reads had their qualities (BQs) reduced to a PHRED score of 2. Where necessary, published ancient data\textsuperscript{7,8,11,12,28,35,43–70} was realigned for use in downstream analyses from either unaligned FASTQ (when available) or aligned BAM following the same parameters described above and filtered in an identical manner.

**Radiocarbon Dating and isotope analysis.**

Direct radiocarbon dates were obtained for 27 samples from accelerator mass spectrometry facilities at Queen's University Belfast and the University of Oxford. All calibrated dates are taken from CALIB 7.1 after Reimer et al. 2013\textsuperscript{71} and reported at two standard deviations (95.4% confidence). The median probabilities (cal BC) have been used for plotting samples chronologically. Stable isotope ratios ($\delta^{13}$C and $\delta^{15}$N) are also reported for the 27 samples and compared with published stable isotope data from 85 Irish and 81 British samples\textsuperscript{8,9,11,14,24,68,72–89} (Fig. 1, Supplementary Table 4). The timeline in Fig. 1 is phased following McLaughlin et al. 2016\textsuperscript{90}.

**Molecular sexing and aneuploidy detection with read coverage**

Molecular sexing was done following two methods, one previously published\textsuperscript{91}, and one described as follows. The total number of X chromosome reads was divided by the length of the X chromosome. This value was then normalised using the median seen for the same calculation across the autosomal chromosomes. We call this Rx. A value above 0.9 was designated female and below 0.6 as male.
Chromosomal deletions or duplications of sufficient length can be detected by aberrations read coverage for shotgun data. We estimated the chromosomal coverages for 145 shotgun sequenced ancient individuals and 43 samples from the current study (>0.3X mean genome coverage) using Qualimap. To compare chromosomal coverages between samples, we normalised values by the mean autosomal coverage for each genome. An extreme outlier was observed for chromosome 21. To estimate the aberration in read coverage for this sample, we divided its normalised chromosome 21 coverage by the median for this value seen across all samples (Extended Data Fig. 6b).

**Mitochondrial analysis**

To determine mitochondrial coverages and haplogroups, reads aligned (no MQ filter) to the human reference genome and revised Cambridge mitochondrial reference sequence were realigned to the mitochondrial reference alone and re-processed as described above. Coverages were obtained using Qualimap v2.1.1. Consensus sequences were determined as previously described with Samtools mpileup (-B, -d6 and -Q 30) and vcftools.pl (vcf2fq)\(^4^1\). HaploFind\(^4^3\) was used to identify defining mutations and assign haplogroups (Supplementary Table 6). Mitochondrial contamination was estimated as previously described\(^7,^6^2\). Realigned data was not used for these, to avoid the confounding effects of misaligned NUMT sequences. Contamination estimates with and without the inclusion of potential damage sites are given (Supplementary Table 6).

**Genotype Calling**

As the majority of published ancient genomic data possess sequencing coverages too low for direct diploid genotype two alternative methods were employed, pseudo-diploid genotype calling and genotype imputation. In order to minimize the impact of reference bias previously observed in pseudo-diploidised data\(^1^2\), a relaxed MQ filter of 20 was applied during data processing. Randomised pseudo-diploid genotypes (BQ>30) were called following previously established methods\(^7\). Imputation was carried out using Beagle 4.0\(^4^4\) for 43 individuals from the current study (>0.4X) (Supplementary Data Table 2), and 51 published\(^1^8,^1^1,^2^8,^5^2,^5^4,^5^6,^5^8,^6^3,^6^4,^9^3\) ancient genomes (>0.66X) (Supplementary Data Table 3), following previously published method\(^4^2,^3^5,^4^9,^5^7\) with some modifications described below.
Genotype likelihoods for biallelic autosomal SNPs in the 1000 Genomes phase 3 dataset\textsuperscript{46} were called using the UnifiedGenotyper tool in GenomeAnalysisTK v2.4-7\textsuperscript{42}. These were filtered to add equal likelihoods for missing data and for genotypes which could be the result of post-mortem damage. Samples were merged by chromosome and imputed in 15,000 marker windows using the 1000G phase 3 haplotypic reference panel and genetic map files provided by the BEAGLE website (http://bochet.gcc.biostat.washington.edu/beagle/). To assess accuracy, imputed genotypes for chromosome 22 of the downsampled NE1\textsuperscript{35} genome (1X), were compared to direct diploid genotypes from the high coverage version (25X) (Extended Data Fig. 1). Optimal filters of >5% MAF, >99% GP and exclusion of transition sites were subsequently chosen for all downstream analysis. Six individuals, including three from the current study (ANN2, PB754 and PN16), were excluded from downstream analysis due to a high percentage of genotype missingness (>16%) after the imposition of the genotype probability filter. The remaining 88 individuals were combined with published imputed genotypes (filtered identically) from 67 ancient samples\textsuperscript{12}. Direct diploid genotype calling was also carried out for high coverage ancient genomes (>10X) at positions in the 1000G Phase 3 variant panel using the HaplotypeCaller tool in GenomeAnalysisTK v4.0\textsuperscript{42} with parameter \texttt{-mbq 30}. A minimum genotype quality of 30, a minimum depth of coverage of 10X, and a maximum depth of coverage twice that of the sample’s mean genomic coverage were required, with a more conservative minimum coverage filter of 15X used for assessment of imputation accuracy.

**Pigmentation profiles for high coverage genomes.**

We availed of the hIrisPlex-S system to predict hair, skin and eye colour in high coverage ancient samples\textsuperscript{97,98}. Diploid genotypes were called at the relevant variant sites and inputted into the hIrisPlex-S online tool (https://hirisplex.erasmusmc.nl). Imputed diploid genotypes (GP > 0.66) were also used for pigmentation prediction across the larger ancient dataset. Results are shown in Supplementary Table 12.
Population genetic analyses

Detailed descriptions for Y chromosome analysis, ADMIXTURE analysis\textsuperscript{99}, \(D\)- and \(f\)-statistics\textsuperscript{100,101} using the AdmixTools package\textsuperscript{102}, ChromoPainter and fineSTRUCTURE analysis\textsuperscript{11}, estimations of ROH, inbreeding coefficients and kinship determination with lcMLkin\textsuperscript{23} can be found in Supplementary Information sections 2-6. We used smartpca\textsuperscript{103,104} to construct the SNP-sharing PCA in Extended Data Fig. 12, using an identical sample and SNP set as that presented in Fig. 1d, e, with imputed genotypes converted randomly to homozygous to mimic pseudodiploid data.

Data availability

Raw FASTQ and aligned BAM files are available through the European Nucleotide Archive under accession number PRJEB36854.
