Genetic contributions to variation in 1 human stature in prehistoric Europe 2 Samantha L. Cox^a, Christopher B. Ruff^b, Robert M. Maier^{c,d,e} & Iain Mathieson^{a,1} 3 4 5 ^a Department of Genetics, Perelman School of Medicine, University of Pennsylvania, Philadelphia PA. 6 ^b Center for Functional Anatomy and Evolution, Johns Hopkins University School of Medicine, Baltimore MD. 7 ^c Program in Medical and Population Genetics, Broad Institute of MIT and Harvard, Cambridge, MA 02142, USA 8 ^d Stanley Center for Psychiatric Research, Broad Institute of MIT and Harvard, Cambridge, MA 02142, USA 9 ^e Analytical and Translational Genetics Unit, Massachusetts General Hospital, Boston, MA 02114, USA 10 ¹ Correspondence to mathi@pennmedicine.upenn.edu

11 Abstract

12 The relative contributions of genetics and environment to temporal and geographic variation in 13 human height remain largely unknown. Ancient DNA has identified changes in genetic ancestry 14 over time, but it is not clear whether those changes in ancestry are associated with changes in 15 height. Here, we directly test whether changes over the past 38,000 years in European height 16 predicted using DNA from 1071 ancient individuals are consistent with changes observed in 17 1159 skeletal remains from comparable populations. We show that the observed decrease in 18 height between the Early Upper Paleolithic and the Mesolithic is qualitatively predicted by 19 genetics. Similarly, both skeletal and genetic height remained constant between the Mesolithic 20 and Neolithic and increased between the Neolithic and Bronze Age. Sitting height changes 21 much less than standing height-consistent with genetic predictions-although genetics predicts 22 a small Bronze Age increase that is not observed in skeletal remains. Geographic variation in 23 stature is also qualitatively consistent with genetic predictions, particularly with respect to 24 latitude. We find that the changes in genetic height between the Neolithic and Bronze Age may 25 be driven by polygenic adaptation. Finally, we hypothesize that an observed decrease in genetic 26 heel bone mineral density in the Neolithic reflects adaptation to the decreased mobility 27 indicated by decreased femoral bending strength. This study provides a model for interpreting 28 phenotypic changes predicted from ancient DNA and demonstrates how they can be combined 29 with phenotypic measurements to understand the relative contribution of genetic and 30 developmentally plastic responses to environmental change.

31 Introduction

32 Stature, or standing height, is one of the most heavily studied human phenotypes. It is easy to 33 measure in living individuals and relatively straightforward to estimate from skeletal remains. 34 As a consequence, geographic variation and temporal changes in stature are well documented 35 (1-3), particularly in western Europe, where there is a comprehensive record of prehistoric 36 changes (4). The earliest anatomically modern humans in Europe, present by 42-45,000 BP (5, 37 6), were relatively tall (mean adult male height in the Early Upper Paleolithic was ~174 cm). 38 Mean male stature then declined from the Paleolithic to the Mesolithic (~164 cm) before increasing to ~167 cm by the Bronze Age (4, 7). Subsequent changes, including the 20th century 39 40 secular trend increased height to \sim 170-180 cm (1, 4). It is broadly agreed that these changes are 41 likely to have been driven by a combination of environmental (e.g. climate or diet) and genetic 42 factors (4, 7-9), although the effects of these two variables cannot be separated based on 43 skeletal data alone. In this study, by combining the results of genome-wide association studies 44 (GWAS) with ancient DNA, we directly estimate the genetic component of stature and test 45 whether population-level skeletal changes between ~35,000 and 1,000 BP are consistent with 46 those predicted by genetics.

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48 Height is highly heritable (10-14), and therefore amenable to genetic analysis by genome-wide 49 association studies (GWAS). With sample sizes of hundreds of thousands of individuals, GWAS 50 have identified thousands of genomic variants that are significantly associated with the 51 phenotype (15-17). Though the individual effect of each of these variants is tiny (on the order of 52 +/- 1-2mm per variant (18)), their combination can be highly predictive. Polygenic risk scores 53 (PRS) constructed by summing together the effects of all height-associated variants carried by 54 an individual can now explain upwards of 30% of the phenotypic variance in populations of 55 European ancestry (16). In effect, the PRS can be thought of as an estimate of "genetic height" 56 that predicts phenotypic height, at least in populations closely related to those in which the 57 GWAS was performed. One major caveat is that the predictive power of PRS is much lower in 58 other populations (19). The extent to which differences in PRS between populations are 59 predictive of population-level differences in phenotype is currently unclear (20). Recent studies

have demonstrated that such differences may partly be artifacts of correlation between
environmental and genetic structure in the original GWAS (21, 22). These studies also
suggested best practices for PRS comparisons, including the use of GWAS summary statistics
from large homogenous studies (instead of meta-analyses), and replication of results using
summary statistics derived from within-family analyses that are robust to population
stratification.

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67 Bearing these caveats in mind, PRS can be applied to ancient populations thanks to recent 68 technological developments that have dramatically increased ancient DNA (aDNA) sample sizes. 69 These have provided remarkable insights into the demographic and evolutionary history of both 70 modern and archaic humans across the world (23-25), particularly in Europe, and allow us to 71 track the evolution of variants underlying phenotypes ranging from pigmentation to diet (26-72 29). In principle, PRS applied to ancient populations could similarly allow us to make inference 73 about the evolution of complex traits. A few studies have used PRS to make predictions about 74 the relative statures of ancient populations (29-31) but looked at only a few hundred samples in 75 total and did not compare their predictions with stature measured from skeletons. Here, we 76 compare measured skeletal data to genetic predictions and directly investigate the genetic 77 contribution to height independent of environmental effects acting during development.

78 Results

79 PRS and skeletal measurements

80 We collected published aDNA data from 1071 ancient individuals from Western Eurasia (west of 81 50° E), dated to between 38,000 and 1100 years before present (BP) (27, 29, 30, 32-57). Using 82 GWAS summary statistics for height from the UK Biobank (generated and made available by the 83 Neale lab: http://nealelab.is/), we computed height PRS for each individual, using a P-value cutoff of 10⁻⁶, clumping variants in 250kb windows, and replacing missing genotypes with the 84 85 mean across individuals (Methods). We refer to this as PRS(GWAS). Because of concerns about 86 GWAS effect sizes being inflated by residual population stratification, we also computed a PRS 87 where we used GWAS P-values to select SNPs, but computed the PRS using effect sizes

88 estimated from a within-family test for ~17,000 sibling pairs from UK Biobank (Methods) which 89 we refer to as PRS(GWAS/Sibs), and which should be unaffected by stratification. We also 90 obtained stature estimates from 1159 individuals dating to between 33,700 and 1100 BP taken 91 from a larger dataset of 2177 individuals with stature and body proportion estimates from 92 substantially complete skeletons (4, 58). There is limited overlap in these datasets (12 93 individuals), but they cover the same time periods and broadly the same geographic locations 94 (Supplementary Fig. 1), although the genetic data contain more individuals from further east 95 (30-50° E) compared to the skeletal data. We divided these individuals into five groups based 96 on date: Early Upper Paleolithic (>25,000 BP; EUP), Late Upper Paleolithic (25,000-11,000 BP; 97 LUP), Mesolithic (11,000-5500 BP), Neolithic (8500-3900 BP) and post-Neolithic (5000-1100 BP, 98 including the Copper and Bronze Ages, plus later periods). These groups broadly correspond to 99 transitions in both archaeological culture and genetic ancestry (33, 38, 59), and we resolved 100 individuals in the overlapping periods using either archaeological or genetic context (Methods). 101

102 Trends in PRS for height are largely consistent with trends in skeletal stature

103 We found a significant effect of group (time period) on mean PRS(GWAS) (ANOVA P= 1.9×10^{-9}). PRS(GWAS/Sibs) (P=0.045) and skeletal stature (P= 2.8×10^{-11}). There was no evidence of 104 105 difference between LUP, Mesolithic and Neolithic groups (Supplementary Fig. 2a-b), so we 106 merged these three groups (we refer to the merged group as LUP-Neolithic). We find that 107 PRS(GWAS) in the LUP-Neolithic period is 0.47 standard deviations (SD) lower than in the EUP (P=0.002), and 0.40 SD lower (P= 8.7×10^{-11}) than in the post-Neolithic period (Fig. 1a). 108 109 PRS(GWAS/Sib) shows a very similar pattern (Fig. 1b), demonstrating that this is not a result of 110 differential relatedness of the ancient individuals to the structured present-day GWAS 111 populations. Skeletal stature shows a qualitatively similar pattern to the genetic predictions, with a 1.5 SD (9.6cm; $P=2.9\times10^{-7}$) difference between EUP and LUP-Neolithic and a 0.27 SD 112 (1.8cm; P=3.6x10⁻⁵) difference between LUP-Neolithic and post-Neolithic. Broad patterns of 113 114 change in stature over time are therefore consistent with genetic predictions. 115

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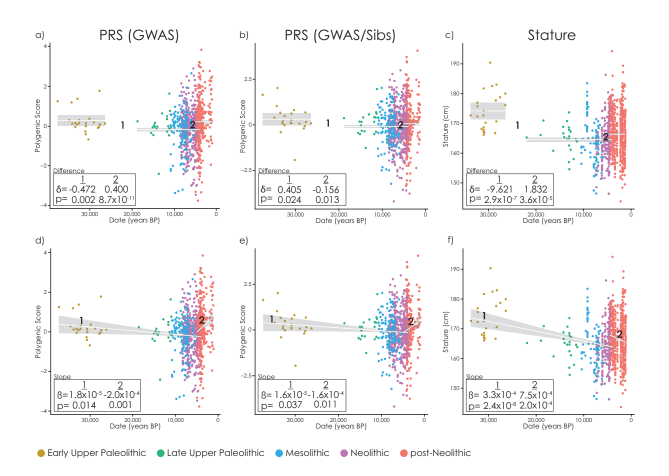


Figure 1: Changes in standing height PRS and stature though time. Each point is an ancient individual, white lines show fitted values, grey area is the 95% confidence interval, and boxes show parameter estimates and p-values for difference in means (δ) and slopes (β). **a-c**) PRS(GWAS) (a), PRS(GWAS/Sibs) (b) and skeletal stature (c) with constant values in the EUP, LUP-Neolithic and post-Neolithic. **d-e**) PRS(GWAS) (d), PRS(GWAS/Sibs) (e) and skeletal stature (f) showing a linear trend between EUP and Neolithic and a different trend in the post-Neolithic.

116 Additionally, we fit a piecewise linear model allowing PRS to decrease from the EUP to the 117 Neolithic and then increase and change slope in the post-Neolithic (Fig. 1d-f). In this model, PRS(GWAS) decreases by about 1.8×10^{-5} SD/year (P=0.014) from EUP to Neolithic, and increases 118 by 2.0×10⁻⁴ SD/year (P=0.001) post-Neolithic (Fig. 1d). PRS(GWAS/sib) decreases by about 119 1.6×10^{-5} SD/vear (P=0.037) from EUP to Neolithic, then increases by 1.6×10^{-4} SD/vear 120 throughout the period (P=0.011; Fig. 1e). Again, these changes are qualitatively consistent with 121 changes in stature (Fig. 1f), with a 4.7×10^{-5} SD/year (3.3×10^{-4} cm/year; P= 2.4×10^{-8}) decrease 122 from EUP to Mesolithic, and an increase of ~0.5 SD into the Neolithic. However, in this model 123 124 stature, unlike PRS, actually decreases during the post-Neolithic period $(7.5 \times 10^{-4} \text{ cm/year})$ $P=2.0\times10^{-4}$). 125

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127 To further explore these trends, we fitted a broader range of piecewise linear models to both 128 datasets (Methods; Supplementary Table 1; Supplementary Fig. 3-5). In the most general model 129 we allowed both the mean and the slope of PRS or stature with respect to time to vary between 130 groups. More constrained models fix some of these parameters to zero-eliminating change 131 over time-or merging two adjacent groups. We compared the fit of these nested models using 132 Akaike's Information Criterion (AIC, Supplementary Table 1). The linear model in Fig. 1d-f is one 133 of the best models in this analysis. In general, all the best-fitting models support the pattern-134 for both PRS and measured stature-of a decrease between the EUP and Mesolithic and an 135 increase between the Neolithic and post-Neolithic (Supplementary Fig. 3-5). Some models 136 suggest that the increase in stature-but not PRS-may have started during the Neolithic 137 (Supplementary Figure 5a-c). Finally, we confirmed that these results were robust to different 138 constructions of the PRS-using 100kb and 500kb clustering windows rather than 250kb 139 (Supplementary Fig. 6-7).

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141 Sitting height PRS is partially consistent with trends in body proportions

142 Standing height is made up of two components: leg length and sitting height (made up of the

143 length of the trunk, neck and head), with a partially overlapping genetic basis (60). Throughout

144 European prehistory, changes in leg length tended to be larger than changes in sitting height

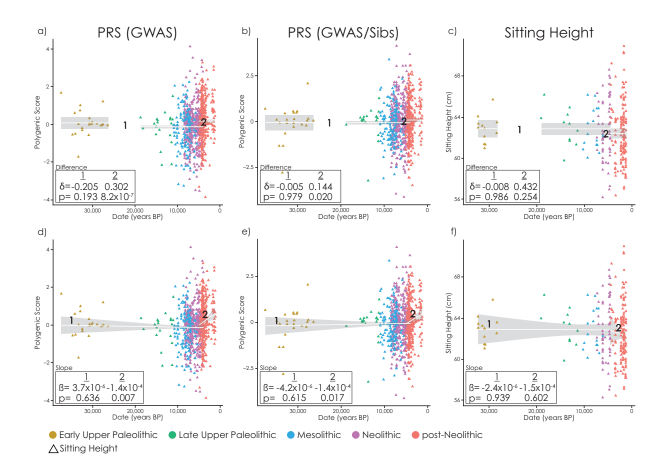


Figure 2: Changes in sitting height PRS and sitting height though time. Each point is an ancient individual, lines show fitted values, grey area is the 95% confidence interval, and boxes show parameter estimates and p-values for difference in means (δ) and slopes (β). **a-c**) PRS(GWAS) (a), PRS(GWAS/Sibs) (b) and skeletal sitting height, with constant values in the EUP, LUP-Neolithic and post-Neolithic. **d-e**) PRS(GWAS) (d), PRS(GWAS/Sibs) (e) and skeletal sitting height showing a linear trend between EUP and Neolithic and a different trend in the post-Neolithic.

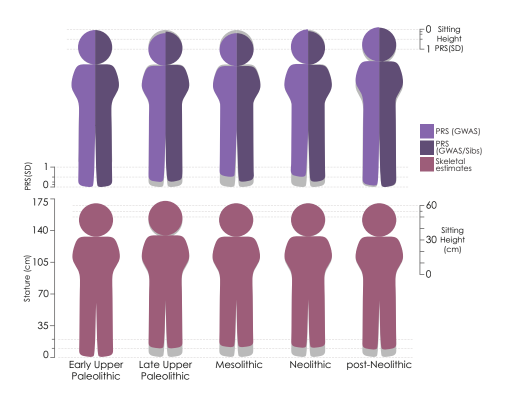


Figure 3: Changes in standing height and sitting height predicted using PRS (relative changes; upper row) and measured from skeletons (absolute values; lower row). Y-axes on the left represent changes in standing height and on the right represent changes in sitting height. Note that changes in PRS are exaggerated compared to the total height of the figures.

145 (4). We constructed PRS(GWAS) and PRS(GWAS/Sibs) for sitting height and analyzed them in 146 the same way as standing height (Fig. 2). In contrast to standing height, we find no evidence of 147 change between the EUP and Neolithic. Both PRS(GWAS) and PRS(GWAS/Sibs) do increase, 148 either between the Neolithic and post-Neolithic, or during the post-Neolithic period (Fig. 2a,b,d 149 & e). On the other hand, using only skeletons with complete torsos to estimate sitting height, 150 we find no evidence of change in any period. Thus, the skeletal data are consistent with the 151 genetic data for the EUP-Neolithic period, but inconsistent in the post-Neolithic period, where 152 PRS predicts an increase that is not reflected in the skeletons. This could be because of more 153 limited skeletal measurements (only 236 out of 1159 skeletons are sufficiently complete to 154 estimate sitting height directly), because the change in PRS is artefactual, it is being buffered by 155 non-genetic effects, or by opposing genetic effects which we do not capture. Overall, we find 156 mixed consistency between PRS and skeletal measurements (Fig. 3). The decrease in standing 157 but not sitting height between the EUP and Neolithic is consistent in both, as is the increase in 158 standing height between the Neolithic and post-Neolithic. However, PRS predicts a continued 159 increase in stature through the post-Neolithic period that is not seen in skeletal remains.

160

161 Geographic variation in standing height

162 As well as varying through time, human stature is stratified by geography, with trends related 163 to both longitude and latitude (61). North-South trends following Allen's (62) and Bergmann's 164 (63) rules are most often interpreted as environmental adaptations to the polar-equatorial 165 climate gradient. Today, Northern Europeans are generally taller than Southern Europeans (1), 166 a pattern which emerged between the Mesolithic and post-Neolithic (4, 7). Longitudinal 167 variation within Europe is present during the Mesolithic (64), though these trends are difficult 168 to interpret due to sampling bias across the time period (4). We therefore tested whether 169 geographic variation in PRS could explain these geographic trends, as it partially explains 170 temporal trends.

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We regressed the residuals from our fitted linear height model (the model shown in Fig. 1d-f) on longitude and latitude. Stature increases significantly with latitude ($P=1.2\times10^{-10}$) in the post-

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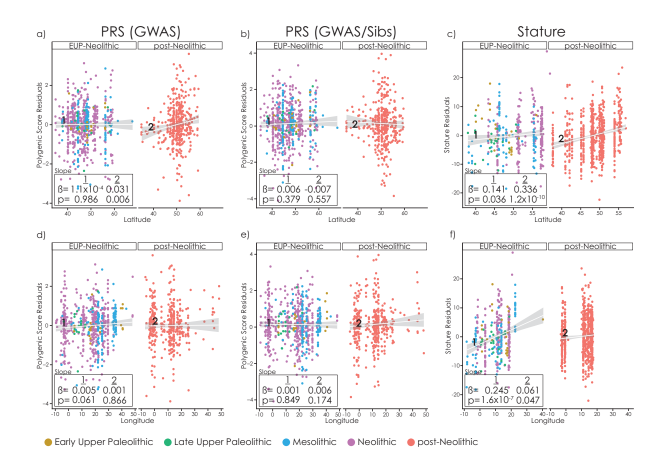


Figure 4: Geographic variation in PRS and skeletal standing height. Residuals for the linear height model (Fig. 1 d-f) against **a-c**) latitude and **d-f**) longitude. Each point is an ancient individual, lines show fitted values, grey area is the 95% confidence interval, and boxes show parameter estimates (β) and p-values for slopes.

Neolithic period. PRS(GWAS) increases in the post-Neolithic (P=0.006) although this is not
replicated by PRS(GWAS/Sibs) (P=0.557). PRS does not increase significantly with latitude in the
EUP-Neolithic period. There is some evidence of a modest trend in stature in the EUP-Neolithic
period (Fig. 4c). However, there is only evidence for this in the Neolithic, not in the EUPMesolithic (Supplementary Fig. 8a). Further, because time and geography are correlated in our
Neolithic sample, this can also be explained by a temporal increase during the Neolithic, in
which case there is no geographic trend (Supplementary Fig. 8b).

181

182 In contrast to latitude, there is a significant increasing trend of stature with longitude before but not during the Neolithic (0.36 cm/degree $P=1.6\times10^{-7}$; Fig. 4, Supplementary Fig. 8c). This 183 184 may be partly driven by a small number of samples from a single site, but still persists if these 185 samples are removed (0.20 standardized residuals per degree, P=0.004; Supplementary Fig. 8d). 186 There is little or no trend (0.06 cm/degree; P=0.047) in the post-Neolithic period (Figure 4f). We 187 find no evidence for longitudinal clines in PRS. In summary, we find that stature increases with 188 latitude in the post-Neolithic, possibly in the Neolithic, but not before. This cline may have a 189 genetic basis. Stature also increases with longitude, particularly in the Mesolithic, but this cline 190 is not predicted by genetics.

191

192 Correlated changes in bone density PRS and femoral bending strength

193 Beyond stature, we wanted to investigate the utility of using PRS to interpret other measurable 194 phenotypes in ancient individuals. Decreased mobility though time, associated with large-scale 195 lifestyle transitions between hunting-gathering, agriculture, and ultimately modern 196 industrialism, is well documented through declines in lower limb bone diaphyseal strength and 197 trabecular density (4, 65, 66). Today, heel bone mineral density (hBMD) is often used as an 198 indicator of general activity levels in younger people (67) and of osteoporosis in older 199 individuals (68, 69); UK Biobank has GWAS data for this trait, indirectly estimated by 200 ultrasound. However, evaluating differences in BMD in archaeological and paleontological 201 specimens can be problematic. In the short term soil leaches bone minerals, while later the 202 bone begins to fossilize, leading to unpredictable patterns of density in ancient remains (70)

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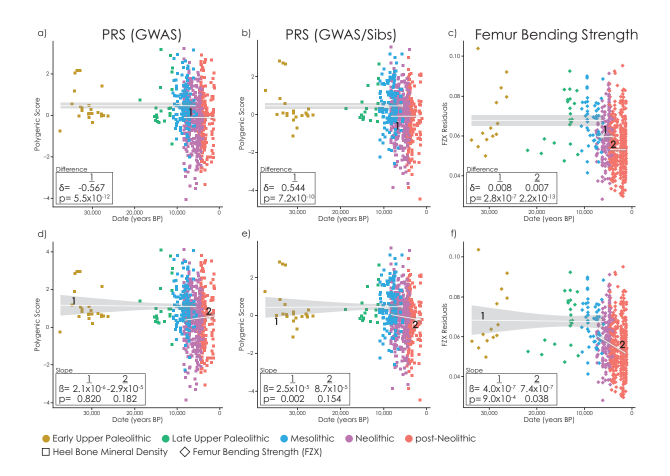


Figure 5: Changes in heel bone mineral density (hBMD) PRS and femur bending strength (FZx) though time. Each point is an ancient individual, lines show fitted values, grey area is the 95% confidence interval, and boxes show parameter estimates and p-values for difference in means (δ) and slopes (β). **a-b**) PRS(GWAS) (a) and PRS(GWAS/Sibs) (b) for hBMD, with constant values in the EUP-Mesolithic and Neolithic-post-Neolithic. **c**) FZx constant in the EUP-Mesolithic, Neolithic and post-Neolithic. **d-e**) PRS(GWAS) (d) and PRS(GWAS/Sibs) (e) for hBMD showing a linear trend between EUP and Mesolithic and a different trend in the Neolithic-post-Neolithic. **f**) FZx with a linear trend between EUP and Mesolithic and a different trend in the Neolithic-post-Neolithic.

203 and requiring special processing methods (65) that are difficult to apply to large samples. 204 However, femoral diaphyseal bending strength can be calculated from bone cross-sectional geometric measurements that are not as affected by bone preservation (71). Here we focus on 205 206 anteroposterior bending strength (section modulus) of the midshaft femur (FZx), which has 207 been linked specifically to mobility (72). Since both trabecular density and diaphyseal strength 208 should respond to mobility and activity levels, we reasoned that they would be likely to show 209 correlated patterns of temporal change. Following established protocols (71), we standardized 210 FZx first by sex, then the product of estimated body mass and femoral length (4). 211 Qualitatively, PRS(GWAS) and FZx show similar patterns, decreasing through time (Fig. 5, 212 Supplementary Figure 1g-i). There is a significant drop in FZx (Figure 5c) from the Mesolithic to Neolithic ($P=1.2\times10^{-8}$) and again from the Neolithic to post-Neolithic ($P=1.5\times10^{-13}$). PRS(GWAS) 213 for hBMD decreases significantly from the Mesolithic to Neolithic (Figure 5a; $P=5.5\times10^{-12}$). 214 which is replicated in PRS(GWAS/Sibs) (P=7.2x10⁻¹⁰; Figure 5b); neither PRS shows evidence of 215 216 decrease between the Neolithic and post-Neolithic. We hypothesize that both FZx and hBMD 217 responded to the reduction in mobility that accompanied the adoption of agriculture (72). In 218 particular, the lower genetic hBMD and skeletal FZx of Neolithic compared to Mesolithic 219 populations may represent adaptation to the same change in environment although we do not 220 know the extent to which the change in FZx was driven by genetic or plastic developmental 221 response to environmental change. On the other hand, FZx continues to decrease between the 222 Neolithic and post-Neolithc (Fig. 5c,f)–which is not reflected in the hBMD PRS (Fig. 5 a-b,d-e). 223 One possibility is that the two phenotypes responded differently to the post-Neolithic 224 intensification of agriculture. Another is that the non-genetic component of hBMD, which we 225 do not capture here, also continued to decrease.

226

227 Are changes in PRS driven by selection or genetic drift?

228 We tested whether there was evidence for selection on any of these traits, by computing the Q_x

statistic (73) for increasing numbers of SNPs from each PRS, with effect sizes taken from either

- 230 PRS(GWAS) (Fig. 6a-c) or PRS(GWAS/Sibs) (Fig. 6d-f). We computed the statistic between each
- pair of adjacent time periods, and over all time periods. We estimated empirical P-values by

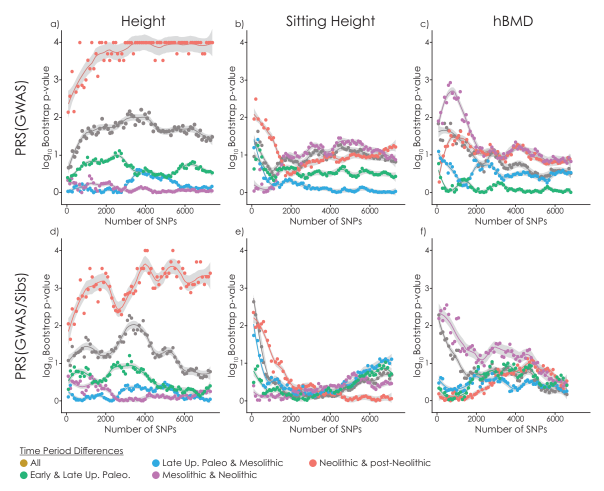


Figure 6: Signals of selection on standing height, sitting height and bone mineral density. We plot the log_{10} bootstrap P-values for the Q_x statistics (y-axis, capped at 4) for GWAS signals (top row) and GWAS/Sibs (bottom row). We tested each pair of adjacent populations, and the combination of all of them ("All"). We ordered PRS SNPs by increasing P-value and tested the significance of Q_x for increasing numbers of SNPs (x-axis).

232 sampling random frequency-matched SNPs from across the genome. Using GWAS effect sizes, we find selection between the Neolithic and Post-Neolithic for stature ($P<1 \times 10^{-4}$; Fig. 6a), 233 which replicates using effect sizes estimated within siblings $(10^{-4} < P < 10^{-2})$; Fig. 6d). The fact that 234 235 the signal is less strong for GWAS/Sibs than for GWAS could either indicate that some of the 236 signal is driven by stratification (21, 22), or that the power to detect selection for smaller effect 237 sizes is lower when using the nosier sibling effect sizes. We tested this by generating GWAS 238 results on a subsample of individuals, chosen so that the standard error of the effect sizes was 239 equal to those of the within-sibling effects. This produced similar results to the analysis using 240 the within-sibling effects (Supplementary Fig. 9), suggesting that the main reason for the 241 weaker signal is the reduction in sample size of the within-sibling analysis. 242

For sitting height, we find little evidence of selection in any time period ($P<10^{-2}$) We conclude 243 244 that there was most likely selection for increased standing but not sitting height in the Steppe 245 ancestors of Bronze Age European populations, as previously proposed (29). One potential 246 caveat is that, although we re-estimated effect sizes within siblings, we still used the GWAS 247 results to identify SNPs to include. This may introduce some subtle confounding, which remains 248 a question for future investigation. Finally, using GWAS effect sizes, we identify some evidence 249 of selection on heel BMD between when comparing Mesolithic and Neolithic populations (10⁻ 3 <P<10⁻²; Fig. 6c). However, this signal is relatively weak when using within-sibling effect sizes. 250 251 and disappears when we include more than about 2000 SNPs.

252 Discussion

We showed that the well-documented temporal and geographic trends in stature in Europe between the Early Upper Paleolithic and the post-Neolithic period are broadly consistent with those that would be predicted by polygenic risk scores (PRS) computed using present-day GWAS results combined with ancient DNA. However, because of the limited predictive power of current PRS, we cannot provide a quantitative estimate of how much of the variation in phenotype between populations might be explained by variation in PRS. Similarly, we cannot say whether the changes were continuous, reflecting evolution through time, or discrete,

reflecting changes associated with known episodes of replacement or admixture of populations
that have diverged genetically over time. Finally, we find cases where predicted genetic
changes are discordant with observed phenotypic changes–emphasizing the role of
developmental plasticity in response to environmental change and the difficulty in interpreting
differences in PRS in the absence of phenotypic data.

265

266 Our results indicate two major episodes of genetic change. First, there was a reduction in 267 stature PRS-but not sitting height PRS-between the Early Upper Paleolithic and Neolithic. 268 These genetic changes are consistent with the decrease in stature-driven by leg length-269 observed in skeletons during this time period (4, 64, 74, 75). This evolutionary change could 270 have been adaptive, driven by changes in resource availability (76) or to a colder climate (61). 271 Early Upper Paleolithic populations in Europe would have migrated relatively recently from 272 more southern latitudes and had body proportions that are typical of present-day tropical 273 populations (75). It is therefore plausible that they adapted to the colder climate of northern 274 latitudes throughout the Upper Paleolithic. Comparison between patterns of phenotypic and 275 genetic variation suggest that, on a broad scale, variation in body proportions among present-276 day people reflects adaptation to environment largely along latitudinal gradients (77, 78). On 277 the other hand, we do not find genetic evidence for selection on stature during this time 278 period-although with a small sample size we likely have very low power to detect it. Further, 279 the populations of Early Upper Paleolithic, Late Upper Paleolithic, Mesolithic and Neolithic 280 Europe are substantially discontinuous and deeply diverged genetically (33, 59). For example 281 the ancestors of Mesolithic and Neolithic Europeans are estimated to have diverged ~46,000 BP 282 (40). Therefore, if these genetic changes do reflect adaptation to climate, this adaptation must 283 have occurred at least partly independently in the ancestors of these populations.

284

The second episode of genetic change is either between the Neolithic and post-Neolithic, or during the post-Neolithic period. In genome-wide ancestry, this transition is characterized by the eastward movement of substantial amounts of "Steppe ancestry" into Central and Western Europe (27, 30, 38, 50). Our results are thus consistent with previous results that Bronze Age

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289 populations of the Eurasian steppe had been selected for increased height and that migration 290 and admixture of these populations with Neolithic European populations increased genetic 291 height in Europe (29, 30). There is no obvious climatic driver for this adaptation but one 292 possibility is that it represents adaptation to a change in social environment. Y chromosome 293 phylogenies suggests an increase in male reproductive variance at this time (29, 48, 50, 79, 80). 294 Culturally, the Bronze Age is characterized by increased social stratification (81) and the 295 introduction of patriarchal Indo-European culture (82). Perhaps these social changes implied 296 increased competition for resources and consequent selection for greater body size. The 297 geographic gradient of increasing skeletal stature is unclear in the Paleolithic, largely West-East 298 in the Mesolithic (7, 64) and largely South-North by the Bronze Age (4, 7, 9). Latitudinal, but not 299 longitudinal, patterns are qualitatively consistent with geographic patterns in PRS suggesting 300 that, like temporal variation, both genetics and environment contribute to geographic variation. 301

302 There is a major confounding factor in analysis of temporal and geographic variation in PRS, 303 particularly in the Bronze Age. Genetic population structure in present-day Europe is correlated 304 with geography (83) and largely driven by variation in proportions of Steppe ancestry, with 305 more Steppe ancestry in Northern Europe and less in Southern Europe (38). Suppose that 306 environmental variation in stature is also correlated with geography, and that Northern 307 Europeans are taller than Southern Europeans for entirely non-genetic reasons. Then, GWAS 308 that do not completely correct for stratification will find that genetic variants that are more 309 common in Steppe populations than Neolithic populations are associated with increased height. 310 When these GWAS results are then used to compute PRS for ancient populations, they will 311 predict that Steppe ancestry populations were genetically taller simply because they are more 312 closely related to present-day Northern Europeans (21, 22). In this study, we attempted to 313 avoid this confounding in two ways: first, by computing PRS using GWAS effect sizes from the 314 UK Biobank–a fairly homogenous dataset that should be well-controlled for population 315 stratification, and second, by replicating our results after re-estimating the effect sizes within 316 siblings, which should be robust to population stratification. The tradeoff between these two 317 methods is that the small sibling sample size means that effect size estimates are noisy, even

318 though they should be unbiased, and our results using sibling-estimated effects may miss subtle trends. However, we cannot exclude the possibility that some confounding remains, for 319 320 example because although we re-estimated effect sizes using the within-siblings design, we still 321 ascertained loci using the GWAS results. Residual confounding would also tend to create 322 spurious signals of polygenic adaption (21, 22). 323 324 As well as genetic contributions to phenotype, our results shed light on possible environmental 325 contributions. In some cases, we can make hypotheses about the relationship between 326 environmental or lifestyle changes, and genetic change. For example, if we interpret change in 327 femur bending strength as reflecting a decrease in mobility, the coincident Mesolithic/Neolithic 328 change in heel bone mineral density PRS can be seen as a genetic response to this change. 329 However, in the Neolithic/post-Neolithic periods, the two observations are decoupled. This

330 emphasizes the role of developmental plasticity in response to changes in environment, and of

331 joint interpretation of phenotypic and genetic variables. Even when looking at the same

332 phenotype, we find cases where genetic predictions and phenotypic data are discordant-for

333 example in post-Neolithic sitting height. We must therefore be cautious in the interpretation of

334 predicted genetic patterns where phenotypes cannot be directly measured, even if it is possible

to control stratification. Predicted genetic changes should be used as a baseline, against which

336 non-genetic effects can be measured and tested.

337 Methods

338 Ancient DNA and polygenic risk score construction

339 We collected published ancient DNA data from 1122 ancient individuals, taken from 29 340 publications. The majority of these individuals had been genotyped using an in-solution capture 341 reagent ("1240k") that targets 1.24 million single nucleotide polymorphisms (SNPs) across the 342 genome. Because of the low coverage of most of these samples, the genotype data are pseudo-343 haploid. That is, there is only a single allele present for each individual at each site, but alleles at 344 adjacent sites may come from either of the two chromosomes of the individual. For individuals 345 with shotgun sequence data, we selected a single read at each 1240k site. We obtained the 346 date of each individual from the original publication. Most of the samples have been directly 347 radiocarbon dated, or else are securely dated by context. 348

349 We obtained GWAS results from the Neale lab UK Biobank page (http://www.nealelab.is/uk-350 biobank/; Round 1, accessed February and April 2018). To compute PRS, we first took the 351 intersection of the 1240k sites and the association summary statistics. We then selected a list of SNPs to use in the PRS by selecting the SNP with the lowest P-value, removing all SNPs within 352 250kb, and repeating until there were no SNPs remaining with P-value less than 10⁻⁶. We then 353 354 computed PRS for each individual by taking the sum of genotype multiplied by effect size for all 355 included SNPs. Where an individual was missing data at a particular SNP, we replaced the SNP 356 with the average frequency of the SNP across the whole dataset. This has the effect of shrinking 357 the PRS towards the mean and should be conservative for the identification of differences in 358 PRS. We confirmed that there was no correlation between missingness and PRS, to make sure 359 that missing data did not bias the results (correlation between missingness and PRS ρ =0.02; 360 P=0.44, Supplementary Fig. 10). Finally, we normalized the PRS across individuals to have mean 361 0 and standard deviation 1.

362

We estimated within-family effect sizes from 17,358 sibling pairs in the UK Biobank to obtain
 effect estimates that are unaffected by stratification. Pairs of individuals were identified as
 siblings if estimates of IBSO were greater than 0.0018 and kinship coefficients were greater than

366 0.185. Of those pairs, we only retained those where both siblings were classified by UK Biobank 367 as "white British", and randomly picked two individuals from families with more than two 368 siblings. We used Hail (84) to estimate within-sibling pair effect sizes for 1,284,881 SNPs by 369 regressing pairwise phenotypic differences between siblings against the difference in genotype. 370 We included pairwise differences of sex (coded as 0/1) and age as covariates, and inverse-rank-371 normalized the phenotype before taking the differences between siblings. To combine the 372 GWAS and sibling results, we first restricted the GWAS results to sites where we had estimated 373 a sibling effect size and replaced the GWAS effect sizes by the sibling effects. We then restricted 374 to 1240k sites and constructed PRS in the same way as for the GWAS results.

375

To test whether the differences in the GWAS and GWAS/Sibs PRS results can be explained by differences in power, we created subsampled GWAS estimates which matched the sibling in the expected standard errors, by determining the equivalent sample size necessary and randomly sampling N_{sub} individuals. $N_{sub} = \frac{N_{sib}}{2 var(\delta_{sib})}$ where δ_{sib} is the difference in normalized phenotype between siblings after accounting for the covariates age and sex.

381

382 Stature data

383 We obtained stature data from Ruff (2018) (4) (data file and notes available at 384 http://www.hopkinsmedicine.org/fae/CBR.html), which also includes estimated body mass, 385 femoral midshaft anteroposterior strength (FZx), and other osteometric dimensions. Statures 386 and body masses were calculated from linear skeletal measurements using anatomical 387 reconstruction or sample-specific regression formulae (4, 58). We calculated sitting height as 388 basion-bregma (cranial) height (BBH) plus vertebral column length (VCL). We restricted analysis 389 to 1159 individuals dated earlier than 1165 BP (651 males and 508 females), of which 1130 had 390 estimates for stature, 1014 for FZx and 236 for sitting height. Sitting and standing height were 391 standardized for sex by adding the mean difference between male and female estimates to all 392 the female values. Sex differences in stature remain relatively constant over time (4), making it 393 reasonable to adjust all female heights by the same mean value. For FZx we first standardized

for sex as we did for stature then divided each by estimated body mass multiplied bybiomechanical femur length (4).

396

397 Grouping

398 We grouped individuals into broad categories based on date and, in some cases, archeological 399 and genetic context. All individuals were assigned to one time period group, based on median 400 age estimates of the sample obtained from the original publications. Date ranges for each time 401 period are based on a combination of historical, climatic, and archaeological factors. The Early 402 Upper Paleolithic comprises all samples older than 25,000 BP, which roughly coincides with the 403 end of the last glacial maximum (LGM). The Late Upper Paleolithic begins when the European 404 glaciers are beginning to recede (25,000 BP) and extends until 11,000 BP and a shift in lithic 405 technology that is traditionally used to delineate the beginning of the Mesolithic period. 406 Transitions between the Mesolithic, Neolithic, and Bronze Age are staggered throughout 407 Europe, so creating universally applicable date ranges is not possible. We instead defined 408 overlapping transition periods between the Mesolithic and Neolithic periods (8500-5500 BP) 409 and between the Neolithic and post-Neolithic (5000-3900 BP). For the genetic data, samples in 410 the overlapping periods were assigned based on genetic population affiliation, inferred using 411 supervised ADMIXTURE (48, 85, 86) which, in most of Western Europe, corresponds closely to 412 archaeological context (38, 48). In particular, the Mesolithic/Neolithic overlap was resolved 413 based on whether each individual had more (Neolithic) or less (Mesolithic) than 50% ancestry 414 related to northwest Anatolian Neolithic Farmers. The Neolithic/post-Neolithic overlap was 415 resolved based on whether individuals had more than 25% ancestry related to Bronze Age 416 Steppe populations ("Steppe ancestry"; See Ref. (86) for more details). For the skeletal data, 417 group assignment in the overlapping periods was determined by the archaeology of each site. 418 Broadly, sites belonging to the Neolithic have transitioned to agricultural subsistence. Similarly, 419 post-Neolithic populations are broadly defined by evidence of metal working (Copper, Bronze 420 and Iron Ages, and later periods). In particular, we included Late Eneolithic (Copper Age) sites 421 associated with Corded Ware and Bell Beaker material culture in the post-Neolithic category 422 but for consistency with the genetic classifications, we included 8 Early Eneolithic (before 4500

BP) individuals in the Neolithic category, since this precedes the appearance of Steppe ancestry
in Western Europe. We excluded samples more recent than 1165 BP.

425

426 Linear models

427 We fitted a series of linear models to changes in both PRS and stature data with time. In the 428 most general model, we allow both the intercept and slope to vary between groups. We then 429 either force some of the slopes to be zero, or some of the adjacent groups to have identical 430 parameters. We describe the models using underscores to indicate changes in parameters, 431 lowercase to indicate slopes (change with respect to time) fixed to zero, and upper case to 432 indicate free slopes (i.e. linear trends with time). For example, "E L M N B" is the most 433 general model, "elmnb" indicates that all groups have the same mean and there is no change 434 with time, and "ELMN B" indicates that the first four groups share the same parameters, and 435 the post-Neolithic has different parameters. The models shown in Figures 1 and 2 are "e Imn b" (panels a-b), "e Im nb" (panel c), "ELMN B" (panels d-e) and "ELM NB" (panel f). 436 437 To analyze geographic variation, we used the residuals of the "ELMN B" model for the PRS and 438 "ELM NB" for skeletal stature, and fitted regressions against latitude and longitude.

439

440 Polygenic selection test

We computed bootstap P-values for the Q_x statistic (73) by sampling random sets of SNPs in matched 5% frequency bins, and re-computing the statistic. Unlike for the PRS calculations, we ignored missing data, since the Q_x statistic uses only the population-level estimated allele frequencies and not individual-level data. We tested a series of nested sets of SNPs (x-axis in Fig. 6), adding SNPs in 100 SNP batches, ordered by increasing P-value, down to a P-value of 0.1.

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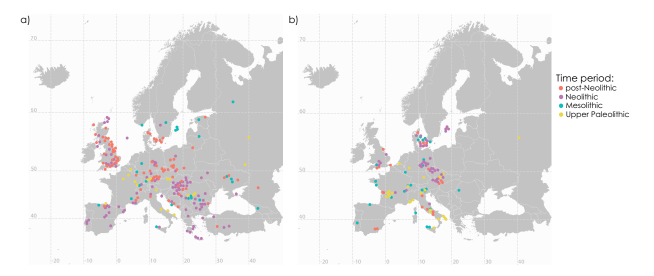
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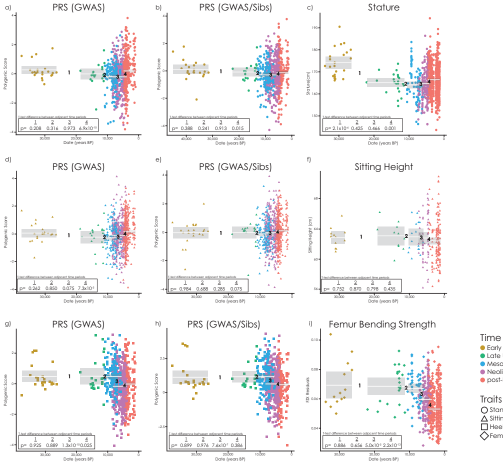
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Supplementary Figure 1: Locations of samples, colored by time period. a) ancient DNA samples;b) skeletal samples.

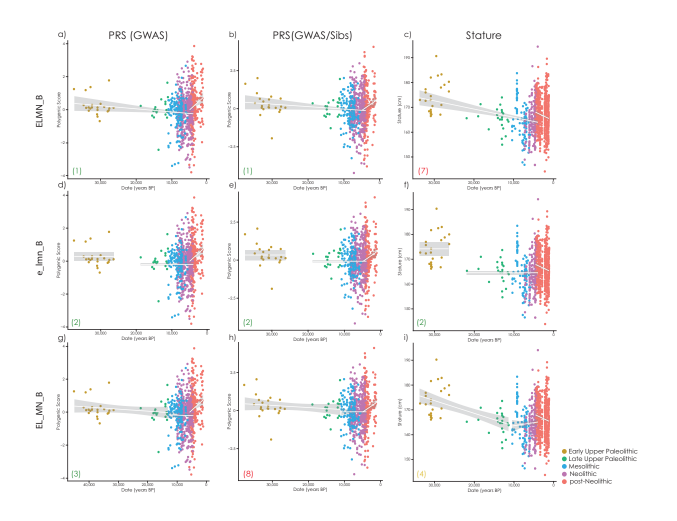


Time Periods • Early Upper Paleolithic • Late Upper Paleolithic • Mesolithic • Neolithic • post-Neolithic

O Standing Height △ Sitting Height Heel Bone Mineral Density ◇ Femur Bending Strength

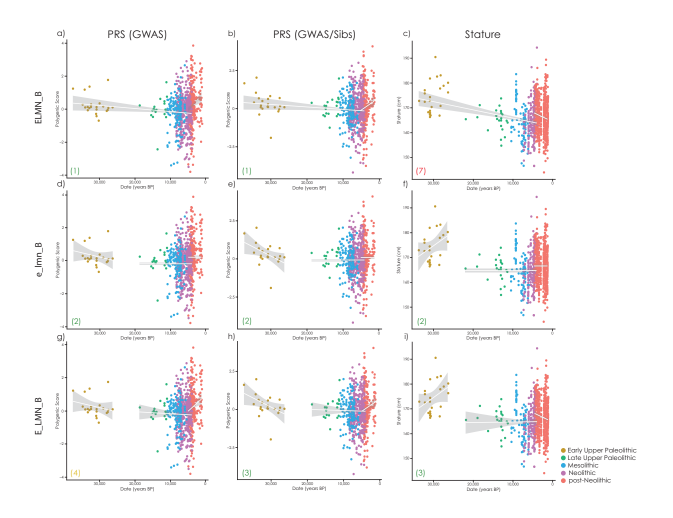
Supplementary Figure 2: Changes in PRS and skeletal phenotypes through time, with constant values in each time period. Each point is an ancient individual, lines show fitted values, grey area is the 95% confidence interval, and boxes show p-values for difference in means between adjacent groups. a) Standing height PRS(GWAS); b) Standing height PRS(GWAS/Sibs); c) Stature (skeletal); d) Sitting height PRS(GWAS); e) Sitting height PRS(GWAS/Sibs); f) Sitting height (skeletal); g) Heel bone mineral density PRS(GWAS); h) Heel bone mineral density PRS(GWAS); i) Femur bending strength (skeletal).

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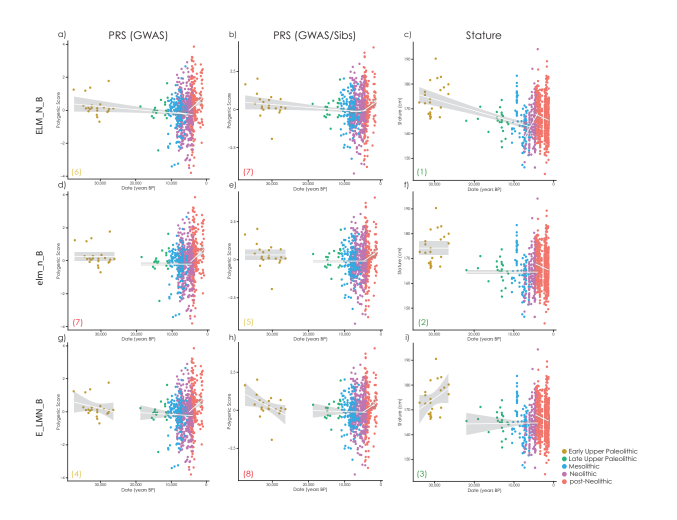
Supplementary Figure 3: Top three AIC models for PRS(GWAS), and the corresponding models for PRS(GWAS/Sibs) and skeletal stature. Row name indicates the model being tested, lowercase letters use fixed values for that time period, uppercase letters indicate the values were allowed to vary linearly with time. Number in the lower left corner of each plot indicates its place in the AIC ranking for PRS(GWAS), PRS(GWAS/Sibs) and Stature, green color indicates a good fit (rank 1-3/29 models), yellow a medium fit (rank 4-6/29 models), and red a poor fit (rank 7 or lower out of 29 models).

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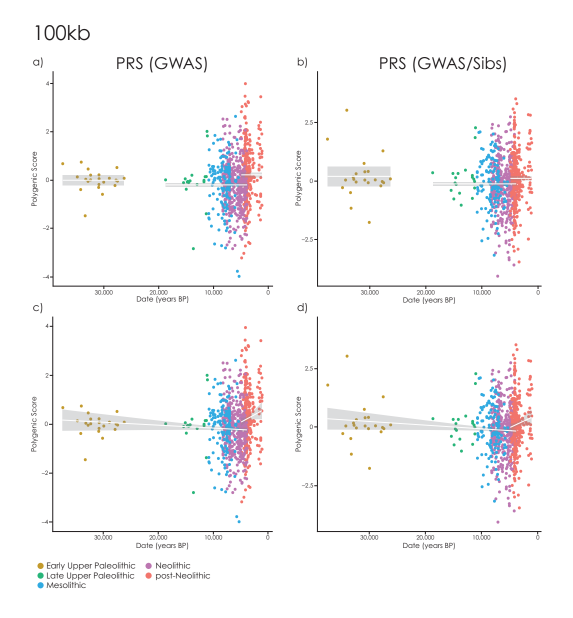


Supplementary Figure 4: Top three AIC models for PRS(GWAS/Sibs), and the corresponding models for PRS(GWAS) and skeletal stature. Row name indicates the model being tested, lowercase letters use fixed values for that time period, uppercase letters indicate the values were allowed to vary linearly with time. Number in the lower left corner of each plot indicates its place in the AIC ranking for PRS(GWAS), PRS(GWAS/Sibs) and Stature, green color indicates a good fit (rank 1-3/29 models), yellow a medium fit (rank 4-6/29 models), and red a poor fit (rank 7 or lower out of 29 models).

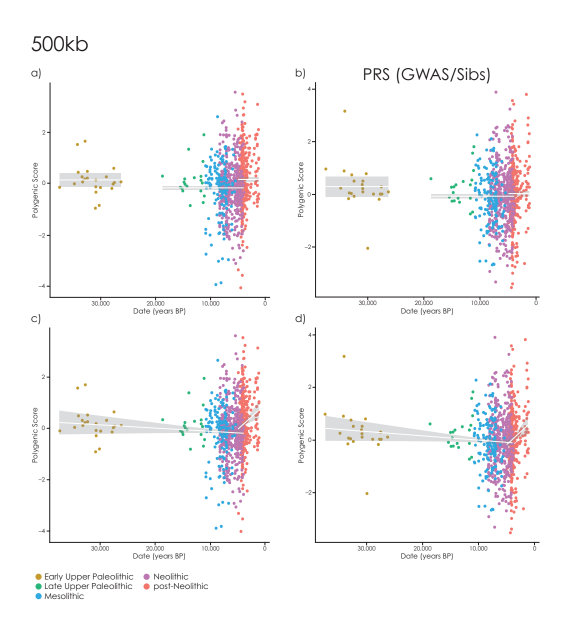
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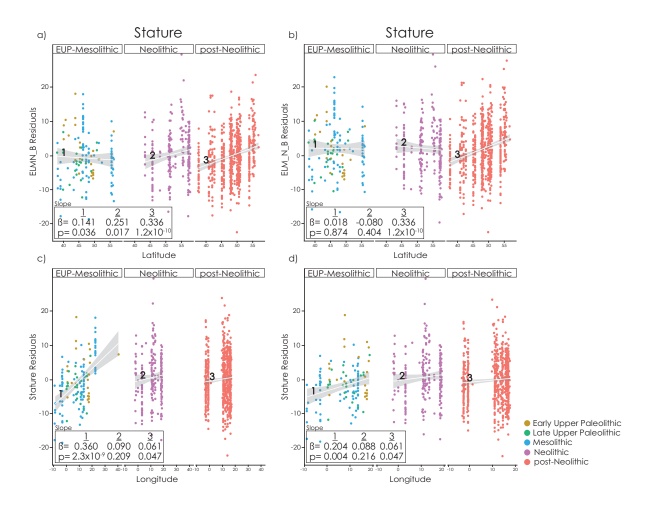
Supplementary Figure 5: Top three AIC models for stature, and the corresponding models for PRS(GWAS) and PRS(GWAS/Sibs). Row name indicates the model being tested, lowercase letters use fixed values for that time period, uppercase letters indicate the values were allowed to vary linearly with time. Number in the lower left corner of each plot indicates its place in the AIC ranking for PRS(GWAS), PRS(GWAS/Sibs) and Stature, green color indicates a good fit (rank 1-3/29 models), yellow a medium fit (rank 4-6/29 models), and red a poor fit (rank 7 or lower out of 29 models).



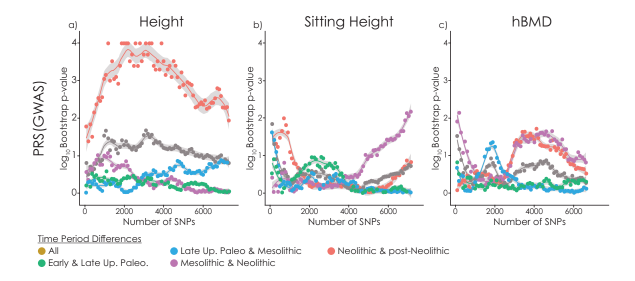
Supplementary Figure 6: Changes in standing height PRS though time with PRS constructed using 100kb clumping windows. Each point is an ancient individual, lines show fitted values, grey area is the 95% confidence interval. **a-b**) Constant values in the EUP, LUP-Neolithic and post-Neolithic; **c-d**) A linear trend with time between EUP-Neolithic and a different trend in the post-Neolithic.



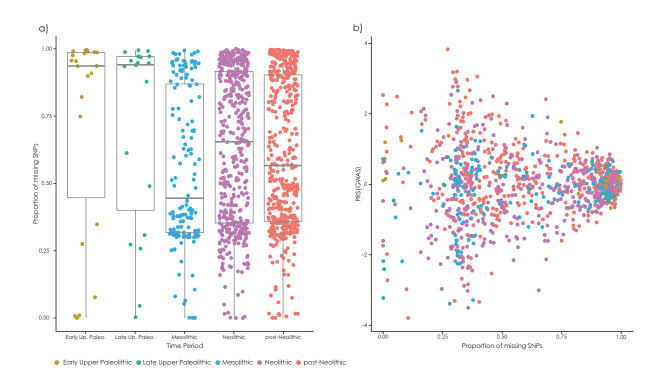
Supplementary Figure 7: Changes in standing height PRS though time with PRS constructed using 500kb clumping windows. Each point is an ancient individual, lines show fitted values, grey area is the 95% confidence interval. **a-b**) Constant values in the EUP, LUP-Neolithic and post-Neolithic; **c-d**) A linear trend with time between EUP-Neolithic and a different trend in the post-Neolithic.



Supplementary Figure 8: Geographic gradients in stature under different models. Here we show EUP-Mesolithic, Neolithic and post-Neolithic periods separately, instead of EUP-Neolithic and post-Neolithic as in the main text. **a**) Latitudinal gradient using residuals of the ELMN_B model (Fig. 1f). **b**) Latitudinal gradient using residuals of the ELM_N_B model (Supplementary Fig. 5c). Note that there is no longer a gradient in the Neolithic, so the apparent geographic gradient can equally be explained by temporal change interacting with sampling. **c**) Longitudinal gradient using residuals of the ELMN_B model (Fig. 1f); the gradient is steepest in the Mesolithic and earlier. **d**) Longitudinal gradient with relatively tall Eastern Mesolithic and Paleolithic samples removed.



Supplementary Figure 9: Selection test as in Figure 6, but using GWAS results generated on a subsample of individuals so that the standard error of the effect size estimates is the same as the standard error of the within-sibling pair estimates.



Supplementary Figure 10: Effect of missing data on PRS. **a**) proportion of missing data as a function of group. **b**) PRS(GWAS) as a function of missing data proportion.