Low Prevalence of Lactase Persistence in Bronze Age Europe Indicates Ongoing Strong Selection over the Last 3,000 Years

Highlights

- Genomic data from Tollense, the oldest large-scale conflict site north of the Alps
- Novel method indicates that Bronze Age warriors represent an unstructured population
- Lactase persistence frequency in Tollense (7.1%) is significantly lower than today
- Selection coefficient estimate of 6% over the last 3,000 years

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In Brief

Burger et al. report the first genomic data from the oldest known battlefield north of the Alps. With additional data from 55 individuals from sites in Southern and Eastern Europe dating to the Bronze Age, they find evidence for a strong and ongoing selection on lactase persistence in various parts of Europe over the last 3,000 years.



Report

Low Prevalence of Lactase Persistence in Bronze Age Europe Indicates Ongoing Strong Selection over the Last 3,000 Years

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SUMMARY

Lactase persistence (LP), the continued expression of lactase into adulthood, is the most strongly selected single gene trait over the last 10,000 years in multiple human populations. It has been posited that the primary allele causing LP among Eurasians, rs4988235-A [1], only rose to appreciable frequencies during the Bronze and Iron Ages [2, 3], long after humans started consuming milk from domesticated animals. This rapid rise has been attributed to an influx of people from the Pontic-Caspian steppe that began around 5,000 years ago [4, 5]. We investigate the spatiotemporal spread of LP through an analysis of 14 warriors from the Tollense Bronze Age battlefield in northern Germany (\sim 3,200 before present, BP), the oldest large-scale conflict site north of the Alps. Genetic data indicate that these individuals represent a single unstructured Central/Northern European population. We complemented these data with genotypes of 18 individuals from the Bronze Age site Mokrin in Serbia (\sim 4,100 to \sim 3,700 BP) and 37 individuals from Eastern Europe and the Pontic-Caspian Steppe region, predating both Bronze Age sites (\sim 5,980 to \sim 3,980 BP). We infer low LP in all three regions, i.e., in northern Germany and South-eastern and Eastern Europe, suggesting that the surge of rs4988235 in Central and Northern Europe was unlikely caused by Steppe expansions. We estimate a selection coefficient of 0.06 and conclude that the selection was ongoing in various parts of Europe over the last 3,000 years.

RESULTS AND DISCUSSION

The Majority of the Tollense Samples Are Male and Unrelated

We enriched DNA from 21 samples from the Tollense battlefield for 5 MB of putatively neutral regions and 487 phenotypically informative loci associated with metabolic syndrome, adult lactase persistence (LP), non-infectious and inflammatory diseases, and eye, skin, and hair pigmentation [6]. Of the 21 samples, two showed evidence of high contamination (>9% based on mtDNA) and one was not involved in the battle but rather dated to the Neolithic period. We analyzed the data of 14 of

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	Sample	Reference	Sample Age (calBP)	n	13,910*T Allele Frequency	90% CI	Genotyping
East European (Steppe)	Eneolithic	this study	5,200–5,978	2	0	0.000-0.602	PCR
	Usatovo	this study	4,950–5,450	1	0	0.000-0.842	PCR
	Yamnaya	this study	4,480–5,378	13	0	0.000-0.132	PCR
	Early catacomb culture	this study	4,341–4,674	8	0	0.000-0.206	PCF
	Developed catacomb culture	this study	4,250–4,564	2	0	0.000-0.602	PCR
	Yamnaya-Poltavkinskaya	this study	4,450-4,884	6	0	0.000-0.265	PCF
	Late catacomb culture	this study	3,975–4,706	5	0	0.000-0.3085	PCF
	East European Steppe: 3,600-2,300 BCE	[2–4, 7–12]	4,277–5,471	37	0	0.000-0.040	NGS (PH)
	Area of Corded Ware Culture: 2,900-2,300 BCE	[2–4, 9–11, 13–18]	4,250–4,833	55	0.018	0.000-0.027	NGS (PH
lest of the	Mokrin (Bronze Age Serbia)	[19]	3,700–4,100	18 ^a	0.046	0.001–0.145 ^a	NGS
ontinent	Tollense	this study	3,200–3,300	14	0.071	0.009–0.235	NGS
	Prague (Jinonice, Zahradnictví and Kobylisy, Ke Stírce Street)	[3]	3,700–4,200	14	0.101	0.031–0.227	NGS
	Bedfordshire, Biddenham Loop	[3]	3,130–3,206	6	0.170	0.028–0.358	NGS
	Lichtenstein Cave, Late Bronze Age in Germany	[20, 21]	2,700–3,000	34 ^a	0.294	0.2–0.42 ^a	PCF
	Kivutkalns, Baltic Bronze Age in Latvia	[22]	2,560–2,730	8	0.575	0.364–0.77	NGS
	Early Medieval Bavarians	[6]	~1,500	21	0.524	0.4–0.647	NGS
	Szolad (northern ancestry), Early Medieval in Hungary	[23]	1,500	13	0.727	0.567–0.856	NGS

Frequencies including 90% credible intervals (CIs) for next-generation sequencing (NGS) data were estimated based on genotype likelihoods using ATLAS. Frequencies based on pseudo-haploid (PH) calls and PCR data were estimated from allele counts taken from genotypes, and a 90% CI was calculated based on the beta distribution. Sample location as well as replication status of the LP genotype are given in Table S3. n, number of individuals.

^aFamily graves with high degrees of relatedness (see also Table S1)

the remaining individuals for which we successfully enriched DNA at targeted regions to a mean depth $>4\times$ (Tables 1 and S1). We detected no close relatives among these 14 individuals (Figure S1). Surprisingly, two of these individuals were women, consistent with a male-dominated, but not exclusively male, battle.

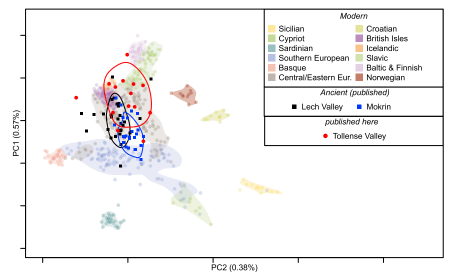
Tollense Sample Shows No Structure

Projected onto a principle-component analysis (PCA) plot trained on modern samples, all Tollense individuals fall within the range of Central and Northern European variation, suggestive of little or no genetic substructure (Figure 1). Multiple lines of evidence corroborate that impression: first, the spread of the Tollense samples on the PCA matches that of other ancient population samples of similar age, namely the Lech Valley in Bavaria (~4,700 to ~3,250 BP) [13] and Mokrin in Serbia (~4,100 to ~3,700 BP) [19]. Second, only seven (0.4%) out of the 1,638 D-statistics [24] of the form D (Tollense1 and Tollense2; Test_Population and Yoruba, [YRI]) were statistically significant with an absolute Z score cutoff of 3, and none were significant after correcting for multiple testing (Table

S2). Among the ten highest D-statistic values, six indicated that WEZ35 was more distant from various European population samples than the other battlefield samples. We additionally tested for differences between visually suggestive upper and lower clusters in the PCA, but no D-statistic of the form D (YRI, Test_Population, Tollense_lower, and Tollense_upper) was significant. Third, FST between random partitions of the Tollense sample were generally lower than those between random partitions of a modern 1000 Genomes Central European (CEU) sample of the same size (P[F_{ST} Tollense < F_{ST} CEU] = 65.57%, 65.42%, 65.65%, and 65.35% for partitions of 4/10, 5/9, 6/8, or 7/7 individuals). Finally, we failed to reject Hardy-Weinberg equilibrium (HWE) using a novel method that accounts for genotyping uncertainty in ancient samples (see STAR Methods). Specifically, we tested for a deficit of heterozygous genotypes as quantified by the inbreeding coefficient F, which is expected to be positive among samples from a structured population due to the Wahlund effect but for which we estimated F = 0.0 (maximum a posteriori [MAP], 90% credible interval [CI] 0.0-1.6.10⁻⁴; Figure 2A). Contrasting this model with strict HWE, the latter received 99.94% posterior support.



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The Tollense individuals thus conform to a sample from a single unstructured population.

Tollense Population Is from Central/Northern Europe

The ancestry of the 14 Tollense individuals is Central to Northern European, as indicated by several analyses. First, the 70% density contour in the PCA shows a slightly more northern center for the Tollense sample than for those of the Bavarian Lech Valley and Serbian Mokrin individuals (Figure 1). Second, the Tollense sample appears closest to 5th century Bavarians [6] and modern Central and Northern Europeans when quantified with F_{ST} , although all European populations have overlapping confidence intervals (Figure 2B). Finally, the same clustering was obtained with a TreeMix analysis (Figures 2C and S2) at the maximum-likelihood estimate of five migration events, of which an event between the Finnish and an East Asian population was the only one involving Europeans.

Based on archeological finds at the battlefield, it has been hypothesized that some of the warriors came from southern Central Europe, i.e., South-eastern Germany and Bohemia [25–27]. Carbon and strontium isotope ratios further suggest a mixture of locals and non-locals [28]. Although it is possible that our data are insufficient to resolve genetic affinities at such a small geographic scale, it appears likely that the Tollense individuals were sampled from a relatively homogeneous population with a high degree of continuity to those living in the same broad region today. But we caution that a strong correspondence between the material culture and genetic ancestry is only expected under conditions of strict and enduring population separation.

Major Allele Frequency Increase before the Medieval Period

To compare LP frequencies temporally and spatially, we determined rs4988235-A allele frequencies (also known as $-13,910^{*}T$) using a Bayesian estimator in various ancient, post-Neolithic population samples with more than five individuals per site. We estimated low frequencies for both Tollense (7.1%; n = 14) and Mokrin (4.6%; n = 18) samples (P[f_{Mokrin} < f_{Tollense}] = 0.889). However, we caution that various individuals at

Figure 1. PCA of Modern and Ancient Europeans

Black squares, ancient samples from the Lech Valley in Bavaria; blue squares, ancient samples from Mokrin in Serbia; red circles, Tollense samples from this study. Ancient samples are projected onto modern reference sample space. Colored circles correspond to the 70% density contour lines of the ancient groups.

the Mokrin site are close relatives. These estimates are consistent with those of other Bronze Age populations, albeit with a lack of high-quality data for comparison (Table 1). For instance, data from somewhat older sites in the UK (n= 6) [3] and Czechia (n = 14) [3] indicate allele frequencies of 17% and 10%, respectively, which are not significantly

different from those of Tollense (P[$f_{UK} < f_{Tollense}$] = 0.225; P [$f_{Czechia} < f_{Tollense}$] = 0.33). However, these samples do not originate from a single time point or location and are based on extremely low allelic depth, few individuals, or both. The highest Bronze Age frequency estimate (29%; n = 34) [20, 21] was observed for the Lichtenstein Cave in Germany, a family grave site a few centuries younger than Tollense.

Markedly higher frequencies are observed from more recent samples, suggesting ongoing strong selection on the rs4988235-A allele during the intermediate period. A sample from Latvia, dating to about 2,730–2,560 BP (P[f_{Latvia} < f_{Tollense}] < 10^{-3} ; n = 8) [22], and a Medieval sample from Southern Germany (~1,500 BP; P[f_{Germany} < f_{Tollense}] < 10^{-3} ; n = 21) [6] both have a frequency of roughly 57%, and a sample with Northern European ancestry from an Early Medieval cemetery in Hungary has a frequency of 73% (P[f_{Latvia} < f_{Tollense}] < 10^{-3} ; n = 13) [23].

These frequency estimates indicate that, although the rs4988235-A allele had reached frequencies that make it detectable in small-sample-size ancient DNA studies by the beginning of the Bronze Age in various parts of Europe, it had not reached the frequencies observed in population samples dating from the Iron Age or later. Such a pattern is consistent with selection starting as early as the Neolithic but also indicates continuing strong selection during and particularly after the Bronze Age.

Selection Is Ongoing and Strong after the Neolithic Period

To quantify the selection strength on the rs4988235-A allele since the Bronze Age, we treated the Tollense individuals as sampled from an unstructured population with broad continuity to modern Central/Northern Europeans and examined allele frequency changes in the intervening 130 generations for all phenotypically relevant loci genotyped. We used the unrelated 1000 Genomes CEU samples as a modern reference population sample and employed two complementary methods: (1) a Bayesian approach (ApproxWF) [29] under an additive model (dominance coefficient of h = 0.5) and a constant population, the size of which we estimated based on allele frequencies at neutral loci, and (2) forward simulations under a model of exponential population growth [30].



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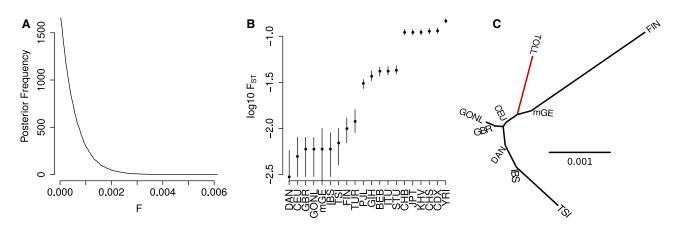


Figure 2. Tollense Samples Show No Population Structure (A) Posterior density of inbreeding coefficient F (see also Figure S3).

(B) F_{ST} between Tollense samples at 5K neutralome against 5th century Bavarians (mGE), Dutch (GONL), Danish (DAN), Turkish (TUR), and Eurasian+YRI 1000 Genomes populations, excluding transition SNPs,

(C) TreeMix analysis with zoom on European populations allowing five migration events with an unrooted tree. CEU, Utah residents with northern and western European ancestry (see also Figure S2).

Both methods identify two alleles influenced by strong positive selection: the LP causing allele rs4988235-A and rs7570971-C, which occurs almost exclusively on the same haplotype as rs4988235-A and additionally has been hypothesised to play a role in total serum cholesterol reduction [31]. Persuasive evidence for a functional role in LP strongly indicates that selection is acting on the rs4988235-A allele rather than on the linked rs7570971-C allele [32, 33]. Within 3,000 years, sample counts for these alleles increased from 2/26 to 144/192 and 2/28 to 134/190, respectively. ApproxWF estimates a selection coefficient of around 5% for rs7570971-C and 6% for rs4988235-A, also when assuming dominance (h = 1; Figures 3B and 3C). Consistent with the low frequency we report here for the Bronze Age, these estimates are a bit higher than recent estimates from modern (e.g., 1.6%) [34] and ancient data (e.g., 1.8%) [35].

The forward simulation approach additionally rejected genetic drift as the only explanation for the inferred frequency change of the derived allele at rs5743810 (p < 0.001). This locus is located in the toll-like receptor gene complex (TLR6) and is associated with pathogen pattern recognition and innate immune response. Notably, none of the other 440 tested loci showed a significant signal of selection between the period of the Tollense site (3,200 BP) and today.

Eastern Europeans Steppes Are Not the Source for Lactase Persistence

The spread of the rs4988235-A allele has previously been attributed to a Steppe-associated expansion during the early Bronze Age [36]. Based on imputed data, Allentoft et al. [4] reported a low allele frequency (5%) of the rs4988235-A allele in Bronze Age Europeans, similar to that reported here, but a much higher frequency (~25%) among Bronze Age samples from the Pontic-Caspian Steppe, indicating a possible Steppe origin of lactase persistence. Because imputing allele presence in ancient samples using modern reference individuals may be problematic in regions of strong recent selection, we investigated this hypothesis by genotyping the rs4988235 locus using PCR in Eneolithic and Early Bronze Age samples from Eastern Europe and the Steppe region. The majority of the sampled individuals were buried in barrow graves dating from the end of the 4th millenium BCE to the end of the 2nd millenium BCE and are representative for the Early Bronze Age in Eastern Europe (Figure 3A; Table 1). We could not detect the rs4988235-A allele among any of these samples (n = 37), suggesting that the frequency of this allele was very low, possibly close to zero, and almost certainly lower than the 5.4% previously reported for a geographically, culturally, and temporally diverse sample with "Steppe ancestry" [2]. Additionally, we re-analyzed published data from the Eastern European steppe area (5,600-4,300 BP) and that of the Corded Ware Culture in Central and North-eastern Europe (4,900-4,300 BP)-based on pseudo-haploid random allele picking-obtaining frequencies of 0% and 1.8%, respectively; this corresponds to a single LP-associated allele in 92 individuals (Table 1). Although these estimates are not directly informative about the origin of the rs4988235-A allele, they appear inconsistent with a major contribution of the Steppe-associated expansion to the high frequencies observed after the Bronze Age in Europe.

The Time Course of Selection

Although the LP-causing rs4988235-A allele has been under very strong natural selection at the broad geographic scale of Western Eurasia, considerable uncertainty and debate remains concerning the underlying drivers and spatiotemporal distribution of that selection and the role demographic processes played in shaping allele frequencies [37]. When considering the underlying drivers of selection, it is important to recognize that the date of origin of the rs4988235-A allele, the timing of selection on LP (whether constant from some point in the past or episodic), and the first observation of that allele in ancient DNA data are distinct and probably all separated by thousands of years. Linkage disequilibrium studies [38–40] indicate an allele origin in the last 20,000 years, perhaps in the Holocene. Evidence of milk consumption from analysis of fatty acids deposited on pot



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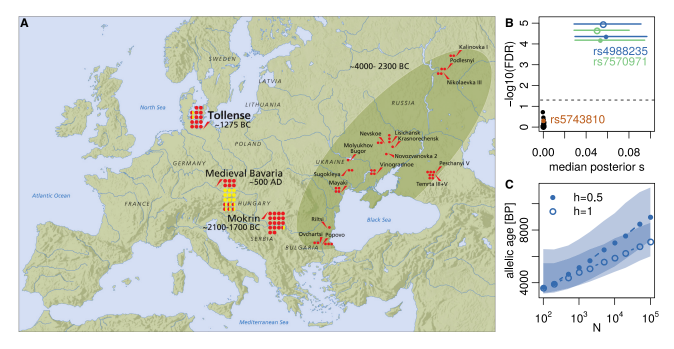


Figure 3. Lactase Persistence Distribution and Selection in Europe

(A) Map of lactase persistence in the Bronze Ages sites of Tollense and Mokrin, in Eneolithic and Early Bronze Age samples from Eastern Europe and the Pontic-Caspian Steppe area, and an Early Medieval sample from Bavaria. Red corresponds to the ancestral and yellow to the derived allele associated with LP (see also Table S3). The shaded shape encompasses the samples from the east European Steppe.

(B) For each phenotypic locus, we show the posterior selection coefficient and the false discovery rate (FDR) as inferred with ApproxWF. Loci identified as under selection by forward simulations approach are colored. Closed circles show estimates obtained under an additive model (h=0.5), and open circles show estimates obtained under a dominant model (h=1). The dashed line indicates the 5% FDR cutoff used.

(C) Posterior densities estimated by ApproxWF under the additive (h = 0.5, solid) and dominant (h = 1, dashed) model for the loci identified as under selection by both methods.

sherds [41–43] and from archaeological herd kill-off profiles [44] dates back to the Early Neolithic in Anatolia, the Levant, and South-eastern Europe. However, even if strong natural selection favoring LP has been operating since the Early Neolithic, we would expect sigmoidal allele frequency trajectories and so low and barely detectable frequencies for up to several thousand years. Thus, low-frequency estimates until the later Bronze Age [2–4, 13, 34, 45, 46] (this study) are not necessarily inconsistent with selection on LP starting when or shortly after milk became a significant dietary component.

To illustrate this, we inferred the age of the rs4988235-A allele from the Tollense data using a novel Bayesian method and assuming a hard sweep [36] with selection coefficient of 6% since the mutation occurred (Figure 3C). The inferred allele age is highly dependent on the unknown effective population size between the time of mutation and the Bronze age, as well as the dominance coefficient h. Assuming dominance (h = 1), the MAP estimate varied between 3,550 (90% CI 3,200-5,100) and 7,140 (90% CI 5,525-7,650) years BP for population sizes 2N = 10^2 and $2N = 10^5$, respectively, and was about 20% older when assuming additivity (h = 0.5). In line with the low frequency for the Tollense sample, these estimates are at the lower end of the age range [36] (3,280-23,100 BP) or considerably younger [34] (17,500 BP) than recent estimates from modern data. However, we note that all such estimates make strong assumptions about the demographic history, the strength of selection, as well as the origin of the selected allele.

The Bronze Age therefore represents an important waypoint at which the rs4988235-A allele became sufficiently common to be detectable in reasonably sized ancient DNA samples. Data from the Bronze Age can then serve as a starting point to quantify selection reliably in subsequent times. The Tollense sample, consisting of individuals that most likely died in a battle within a short period of time, possibly on the same day, is well suited for this. The inferred selection coefficient of 6% between Tollense (\sim 3,200 BP) and today is high, especially considering that the advancement of agricultural skills and the increase in dietary breadth since the Neolithic should have produced alternatives to milk consumption.

Drivers of Selection

Beyond milk being a nutrient-dense and relatively nutrientbalanced food, various explanations have been offered for the strong selection inferred for LP. These include improved calcium absorption by supplementing vitamin-D-poor diets at high latitudes [47], the supply of a relatively pathogen-free fluid [48– 50], the suppression of malaria symptoms through a reduction of p-aminobenzoic acid consumption [51, 52], improvements in gut health through galactose and galacto-oligosaccharides reshaping the colonic microbiome [37, 46, 53–56], avoidance of diarrhea under famine conditions [57], and increased economic efficiency of calorie production for dairy farming [58]. It is unlikely that any single one of these factors has acted alone over the whole period from the Neolithic to modern times. However, the



inference of an ongoing and possibly increased selection coefficient from the Bronze Age to at least Medieval times could be interpreted as favoring factors related to increases in population and settlement density, such as those concerning pathogen loads [37, 57]. In this context, it is interesting to note that, among the other >400 functional loci we examined, the only other allele reaching significance in terms of a signature of selection is the derived allele at rs5743810 in the toll-like receptor gene complex (TLR6), which is associated with pathogen pattern recognition and innate immune response. Although difficult to test explicitly, there remains a strong possibility that selection on the rs4988235-A allele is modulated by other genetic factors through a pleiotropic network, possibly in relation to epidemic disease resistance in the context of milk drinking.

Demography and Selection

Aside from strong natural selection, demographic processes, such as migration and population range expansion—leading to allele surfing [59, 60]—will have shaped the distribution of the rs4988235-A allele [61]. It is not clear whether the northwestern distribution of this allele in Europe is primarily the result of these processes or spatially structured selection strengths shaped by, for example, different traditions of milk use, different climates and ecologies, or different levels of incident ultraviolet radiation. However, it is noteworthy that there has been little genetic turnover between the Tollense population sampled in this study and populations in the same region today, yet we still infer a high selection coefficient. The same applies to the Mokrin site in Serbia, where a major population shift is equally unlikely—leaving natural selection as the main explanation for the observed allele frequency change over the last few millennia.

Final Remark

The somewhat similar inferred rs4988235-A allele frequencies between the early Middle Ages and today at different locations does not exclude the possibility of ongoing selection during that period [62], because this likely represents the transitional phase of the sigmoid frequency trajectory expected for a dominant allele. For LP, it now seems likely that the phase of most rapid frequency rise was between 4,000 BP and 1,500 BP. We contend that research should be focused on this phase to better understand the evolutionary history of the most strongly selected single gene trait in Holocene Western Eurasia and many other parts of the world.

STAR*METHODS

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SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j. cub.2020.08.033.

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AUTHOR CONTRIBUTIONS

J. Burger, D.W., T.T., and R.B. initiated this research. A. Schulz, A.Ž., L.W., and R.B. produced data. C.S., J. Blöcher, K.R.V., V.L., J. Burger, Z.H., J.O., A. Scheu, T.B., V.B., Z.P., Y.D., and C.L. analyzed data. J. Burger, V.L., D.W., K.R.V., and M.G.T. wrote the article with the help of all co-authors.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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STAR***METHODS**

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Deposited Data		
Tollense capture raw data	This study	PRJEB38406
PCR genotype estimates of 37 individuals from the following cultures: Eneolithic, Usatovo, Yamnaya, Early Catacomb Culture, Developed Catacomb Culture, Yamnaya- Poltavkinskaja, Late Catacomb Culture	This study	Table S1
Human reference genome NCBI build 37	[63]	GRCh37
Unrelated CEU from 1000 Genomes	[64]	NA06984, NA06985, NA06986, NA06989, NA06994, NA07000, NA07037, NA07048, NA07051, NA07056, NA07346, NA07347, NA07357, NA10847, NA10851, NA11829, NA11830, NA11831, NA11832, NA11840, NA11843, NA11881, NA11892, NA11893, NA11894, NA11918, NA11919, NA11920, NA11930, NA11931, NA11932, NA11933, NA11992, NA11993, NA11994, NA11995, NA12003, NA12004, NA12005, NA12006, NA12043, NA12044, NA12045, NA12046, NA12058, NA12144, NA12154, NA12155, NA12156, NA12234, NA12249, NA12273, NA12275, NA12282, NA12283, NA12286, NA12287, NA12340, NA12341, NA12342, NA12347, NA12348, NA12383, NA12399, NA12400, NA12413, NA12414, NA12489, NA12546, NA12716, NA12717, NA12718, NA12748, NA12775, NA12750, NA12751, NA12760, NA12763, NA12775, NA12778, NA12813, NA12814, NA12815, NA12843, NA12827, NA12829, NA12830, NA12842, NA12843, NA12872, NA12873, NA12874, NA12889, NA12890, NA12891, NA12892
1000 Genomes Eurasian Yoruba	[64]	NA18486, NA18488, NA18489, NA18498, NA18499, NA18501, NA18502, NA18504, NA18505, NA18507, NA18508, NA18510, NA18511, NA18516, NA18517, NA18519, NA18520, NA18522, NA18523, NA18853, NA18856, NA18858, NA18861, NA18864, NA18865, NA18867, NA18868, NA18870, NA18871, NA18873, NA18874, NA18876, NA18877, NA18878, NA18879, NA1881, NA18907, NA18908, NA18909, NA18910, NA18912, NA18915, NA18916, NA18917, NA18923, NA18924, NA18933, NA18934, NA19092, NA19093, NA19095, NA19096, NA19098, NA19099, NA19102, NA19107, NA19108, NA19113, NA19114, NA19116, NA19117, NA19118, NA19119, NA19121, NA19129, NA19130, NA19131, NA19137, NA19138, NA19141, NA19143, NA19144, NA19146, NA19147, NA19149, NA19152, NA19175, NA19159, NA19160, NA19171, NA19172, NA19177, NA19188, NA19200, NA19201, NA19109, NA19130, NA19206, NA19207, NA19209, NA19201, NA19204, NA19206, NA19227, NA19209, NA19210, NA19235, NA19236, NA19226, NA19223, NA19225, NA19235, NA19236, NA19226, NA19239, NA19247, NA19248, NA19256, NA19257



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REAGENT or RESOURCE	SOURCE	IDENTIFIER
Turkish genomes	[65]	06A010111, 08P210611, 24D220611, 25A220611, 31P140611, 32A140611, 33M140611, 34S291210, 35C240511, 38l220611, 42S291210, 48S210611, 50G301210, 52C130611, 57M220611, 65A220611
5th century Bavarian individuals	[6]	AED_106, AED_1119, AED_1135, AED_204, AED_249, AED_432, AED_92, ALH_10, ALH_1, ALH_2, ALH_3, NW_255, STR_241, STR_248, STR_266, STR_300, STR_316, STR_393, STR_480, STR_486, STR_502
Prague (Jinonice, Zahradnictví and Kobylisy, Ke Stírce Street)	[3]	17195, 17196, 17197, 17198, 17199, 17200, 17201, 17202, 17203, 14886, 14888, 14889, 14890, 14891, 14945
Bedfordshire, Biddenham Loop	[3]	17575, 17576, 17577, 17578, 17580, 17626, 17628
Kivutkalns, Baltic Bronze Age in Latvia	[22]	Kivutkalns153, Kivutkalns164, Kivutkalns194, Kivutkalns19, Kivutkalns207, Kivutkalns215, Kivutkalns222, Kivutkalns25, Kivutkalns42
Szolad (northern ancestry), Early Medieval in Hungary	[23]	SZ2, SZ3, SZ4, SZ5, SZ9, SZ11, SZ13, SZ15, SZ16, SZ18, SZ30, SZ38, SZ45
Genome DK (Danish)	[66]	only allele frequencies available
GONL (Dutch)	[67]	gonl-100c, gonl-101c, gonl-102c, gonl-103c, gonl-104c, gonl-105c, gonl-106c, gonl-107c, gonl-108c, gonl-109c, gonl-110c, gonl-110c, gonl-111c, gonl-112c, gonl-113c, gonl-114c, gonl-115c, gonl-112c, gonl-117c, gonl-118c, gonl-126c, gonl-123c, gonl-124c, gonl-125c, gonl-126c, gonl-127c, gonl-128c, gonl-129c, gonl-126c, gonl-130c, gonl-131c, gonl-132c, gonl-133c, gonl-134c, gonl-135c, gonl-132c, gonl-137c, gonl-134c, gonl-135c, gonl-136c, gonl-137c, gonl-134c, gonl-135c, gonl-136c, gonl-140c, gonl-145c, gonl-142c, gonl-134c, gonl-140c, gonl-145c, gonl-142c, gonl-134c, gonl-146c, gonl-145c, gonl-146c, gonl-147c, gonl-148c, gonl-152c, gonl-153c, gonl-154c, gonl-155c, gonl-152c, gonl-160c, gonl-154c, gonl-155c, gonl-152c, gonl-166c, gonl-161c, gonl-165c, gonl-167c, gonl-167c, gonl-168c, gonl-162c, gonl-167c, gonl-167c, gonl-168c, gonl-166c, gonl-167c, gonl-177c, gonl-168c, gonl-166c, gonl-177c, gonl-177c, gonl-178c, gonl-181c, gonl-179c, gonl-177c, gonl-180c, gonl-181c, gonl-182c, gonl-183c, gonl-184c, gonl-185c, gonl-182c, gonl-183c, gonl-184c, gonl-185c, gonl-180c, gonl-1910c, gonl-191c, gonl-182c, gonl-180c, gonl-191c, gonl-195c, gonl-196c, gonl-197c, gonl-200c, gonl-201c, gonl-202c, gonl-203c, gonl-204c, gonl-205c, gonl-202c, gonl-205c, gonl-204c, gonl-205c, gonl-206c, gonl-207c, gonl-208c, gonl-207c, gonl-214c, gonl-215c, gonl-216c, gonl-217c, gonl-214c, gonl-215c, gonl-216c, gonl-217c, gonl-214c, gonl-215c, gonl-226c, gonl-227c, gonl-226c, gonl-226c, gonl-227c, gonl-228c, gonl-226c, gonl-227c, gonl-228c, gonl-228c, gonl-226c, gonl-227c,



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REAGENT or RESOURCE	SOURCE	IDENTIFIER
		gonl-250c, gonl-25c, gonl-26c, gonl-27c, gonl-28c, gonl-29c, gonl-2c, gonl-30c, gonl-31c, gonl-32c, gonl-33c, gonl-34c, gonl-35c, gonl-36c, gonl-37c, gonl-38c, gonl-39c, gonl-3c, gonl-40c, gonl-41c, gonl-42c, gonl-43c, gonl-44c, gonl-45c, gonl-65c, gonl-77c, gonl-48c, gonl-49c, gonl-4c, gonl-50c, gonl-51c, gonl-52c, gonl-53c, gonl-54c, gonl-55c, gonl-66c, gonl-57c, gonl-58c, gonl-59c, gonl-64c, gonl-65c, gonl-61c, gonl-62c, gonl-63c, gonl-69c, gonl-65c, gonl-70c, gonl-71c, gonl-72c, gonl-73c, gonl-74c, gonl-75c, gonl-76c, gonl-77c, gonl-78c, gonl-79c, gonl-76c, gonl-81c, gonl-82c, gonl-83c, gonl-84c, gonl-85c, gonl-86c, gonl-87c, gonl-88c, gonl-89c, gonl-86c, gonl-97c, gonl-97c, gonl-95c, gonl-90c, gonl-91c, gonl-92c, gonl-93c, gonl-94c, gonl-95c, gonl-96c, gonl-97c, gonl-98c, gonl-99c, gonl-96c
Lech Valley Bavarians	[13]	AITI_119, AITI_2, AITI_40, AITI_43, AITI_50, AITI_72, AITI_78, AITI_98, AITI_65adult, AITI_66, AITI_92, AITI_95, OBKR_80, OBKR_117, OBKR_2, OBKR_50, OBKR_67, OBKR_9A, POST_28, POST_44, POST_50, POST_6, POST_35, POST_99, UNTA58_153, UNTA85_1412, WEHR_1192SkB, WEHR_1415adult, WEHR_1586
Mokrin	[19]	MOK10B, MOK12, MOK13, MOK14, MOK15, MOK16A, MOK17A, MOK18A, MOK19A, MOK20, MOK21A, MOK22, MOK23, MOK24A, MOK25A, MOK26A, MOK27, MOK28A, MOK29A, MOK30, MOK31, MOK32, MOK33, MOK9B
Ancient Greek individuals used in the D-statistics analysis	[68]	10070 10071 10073 10074 19005 19006 19010 19033 19041 19123 19127 19128 19129 19130 19131
Ancient Europeanindividuals used in the D-statistics analysis	[2]	10047 10059 10060 10099 10112 10113 10115 10116 10117 10164 10171 10803 10804 10805 10806 11271 11276 11280 11282 11284 11300 11303 11314 11502 11504 11532 11534 11536 11538 11540 11542 11544 11546 11549 RISE109 RISE150 RISE154 RISE175 RISE179 RISE210 RISE247 RISE254 RISE276 RISE349 RISE371 RISE373 RISE374 RISE431 RISE47 RISE486 RISE71 RISE97 RISE98
Ancient Baltic Sea Region individuals used in the D-statistics analysis	[22]	Kivutkalns153 Kivutkalns164 Kivutkalns19 Kivutkalns194 Kivutkalns207 Kivutkalns209 Kivutkalns215 Kivutkalns222 Kivutkalns25 Kivutkalns42 Turlojiske1 Turlojiske1932 Turlojiske3 Turlojiske5
Ancient European individuals used in the D-statistics analysis	[3]	E09537_d E09538 E09568_d E09569 E09613_d E09614_d 10258 10260 10261 10262 10263 10453 10459 10460 10461 10823 10825 10839 10840 11381 11388 11390 11391 11553 11765_d 11767 11770 11775 11970 11979 12364 12365 12416 12418 12421 12445 12446 12447 12450 12452 12453 12454 12455 12457 12458 12459 12461 12462 12463 12464 12477 12478 12565 12567 12568 12569 12573 12574 12575 12597 12598 12601 12602 12604 12609 12610 12612 12618 12639 12653 12654 12655 12656_d 12741 12786 12787 12859 12860



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REAGENT or RESOURCE	SOURCE	IDENTIFIER
		I2861 I2981 I3082 I3130 I3132 I3255 I3529 I3587 I3588 I3589 I3590 I3592 I3594 I3596 I3600 I3601 I3602 I3604 I3607 I3874 I3875 I4067 I4069 I4070 I4071 I4073 I4075 I4076 I4178 I4229 I4245 I4247 I4249 I4250 I4251 I4252 I4253 I4884 I4886 I4888 I4889 I4890 I4891 I4892 I4895 I4896 I4930 I4933 I4936 I4945 I4946 I4947 I4950 I5014 I5015 I5017 I5019 I5020 I5022 I5023 I5024 I5025 I5035 I5037 I5042 I5043 I5044 I5364 I5367 I5373 I5376 I5377 I5379 I5382 I5383 I5385 I5441 I5469 I5470 I5471 I5473 I5512 I5513 I5514 I5515 I5516 I5519 I5521 I5523 I5524 I5525 I5527 I5529 I5530 I5531 I5655 I5656 I5658 I5659 I5661 I5663 I5665 I5666 I5748 I5750 I5755 I5757 I5759 I5833 I5834 I5836 I6466 I6467 I6468 I6471 I6472 I6475 I6476 I6480 I6481 I6482 I6531 I6534 I6537 I6538 I6539 I6542 I6543 I6579 I6590 I6521 I6523 I6584 I6585 I6587 I6589 I6590 I6591 I6596 I6604 I6605 I6608 I6609 I6610 I6612 I6613 I6617 I6622 I6623 I6624 I6628 I6629 I6679 I6680 I6774 I6775 I6778 I7040 I7042 I7043 I7044 I7045 I7195 I7196 I7197 I7198 I7199 I7200 I7201 I7202 I7203 I7205 I7210 I7211 I7213 I7249 I7250 I7251 I7269 I7270 I7272 I7275 I7276 I7278 I7279 I7280 I7281 I7282 I7286 I7287 I7288 I7289 I7290 I7568 I559 I7570 I7571 I7572 I7573 I574 I5757 I5757 I5757 I5757 I7578 I7580 I7580 I758 I7639 I7580 I7580 I7580 I7562 I7270 I7272 I7275 I7276 I7278 I7279 I7280 I7280 I7280 I7280 I7270 I7272 I7275 I7276 I7278 I7279 I7280 I7280 I7280 I7280 I7281 I7282 I7286 I7287 I7288 I7289 I7290 I7568 I7569 I7570 I7571 I7572 I7573 I7574 I7575 I576 I7577 I7578 I7580 I7626 I7627 I7628 I7630 I7635 I7638 I7639 I7640
East European Steppe: 3,600-2,300 BCE	[2-4, 7-12]	 RISE547.SG, RISE552.SG, I0374, I1917, I2105, I4110, I5118, I5119, ILK001, ILK002, ILK003, I5882, I8745, I11501, I11531, I11732, I11735, I11736, I11734, RISE1166.SG, AY2001, AY2003, RK1001, RK1003, RK4001, RK4002, SA6001, SA6002, SA6003, SA6004, SA6010, SA6013, SIJ002, SIJ003, ZO2002, I2791_published, I5884_publishedRISE547.SG, RISE552.SG, I0374, I1917, I2105, I4110, I5118, I5119, ILK001, ILK002, ILK003, I5882, I8745, I11501, I11531, I11732, I11735, I11736, I11734, RISE1166.SG, AY2001, AY2003, RK1001, RK1003, RK4001, RK4002, SA6001, SA6002, SA6003, SA6004, SA6010, SA6013, SIJ002, SIJ003, ZO2002, I2791_published, I5884_published
Area of Corded Ware Culture: 2,900-2,300 BCE	[2–4, 9–11, 13–18]	 I7207, I7208, I7209, I7278, I7289, I7212, I6695, I6696, I7040, I7044, I7045, I7195, I7196, I7200, I7201, I7202, I7203, I7213, I7286, I0805, Ajvide52.SG, Ajvide58.SG, RISE1.SG, RISE94.SG, I3529, I2365, I4178, I2741, I4629, I5015, I5043, I5520, I5521, I5523, I5525, I5529, I6534, I6581, I2786, N44.SG, N45.SG, N47.SG, N49.SG, RISE1159.SG, RISE1162.SG, RISE1163.SG, RISE1164.SG, RISE1167.SG, RISE1170.SG, RISE1171.SG, RISE1172.SG, RISE1173.SG, RISE1248.SG, RISE1249.SG, RISE1250.SG, ans016.SG
Ancient individuals used in D-Statistics	[2, 3, 9, 22, 68]	Kivutkalns209, I5769, Turlojiske5, Kivutkalns164, Kivutkalns19, Kivutkalns207, Kivutkalns215, I5364, Kivutkalns153, Kivutkalns194, Kivutkalns25, I3313, Kivutkalns222, RISE276, Kivutkalns42, I2859, Turlojiske1, I2860, I2861, I3130, I2656_d, Turlojiske3, I7580, I7575, I5383, I7577, I0099, I7626, RISE175, I1504, I9010, I9041, I7576, I7628, I7627, I7578, I2574, I7574, Turlojiske1932, I7571, I9006, I2655, I9033, I7640, I2458, I2653, I2654, I7568, I2573, I7572, RISE210, RISE47, I9123, I3082, I2639, I5470, I7569, I2981,



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REAGENT or RESOURCE	SOURCE	IDENTIFIER
		 Ibelrin Itali Ibelrin Itali Ibelrin Itali Ibelrin Italia Ibelrin Italia
Oligonucleotides		10825, 11970, 16466, 16467, 16596, 16617, 16628, 16629
MYBait kit	Arbor biosciences; https://	N/A
	arborbiosci.com/genomics/ targeted-sequencing/ mybaits/mybaits-custom/	
P5 and P7	[69] IDT, Leuven, Belgium	N/A
IS4, IS5, IS6 and IS7	[69] IDT, Leuven, Belgium	N/A
Software and Algorithms		
ATLAS	[70]	N/A
TreeMix	[71]	N/A

Continued		
REAGENT or RESOURCE	SOURCE	IDENTIFIER
plink2	[72]	N/A
approxWF	[29]	N/A
laser2	[73]	N/A
ADMIXTOOLS	[74]	N/A
yHaplo*	[75]	N/A
scipy.stats python library	https://docs.scipy.org/doc	N/A
numpy python library	https://numpy.org/	N/A
bwa aln		N/A
GATK	[76]	N/A
Samtools	[77] [78]	N/A
VCFtools	[79]	N/A
picard-tools	https://broadinstitute. github.io/picard/	N/A
contamMix	[80]	N/A
Other	1.1.1	
Agilent 2100 Expert Bioanalyzer System	Agilent Technologies	N/A
and High Sensitivity DNA Analysis Kit		
Amicon Ultra-15 Centrifugal Filter Units	Merck Millipore, Darmstadt, Germany	N/A
AmpliTaq Gold ® Buffer II (10x)	Life Technologies TM	N/A
AmpliTaq Gold ® DNA Polymerase	Life Technologies TM	N/A
ATP Solution (100 mM)	Life Technologies [™]	N/A
Bovine Serum Albumin (BSA) (20 mg/ml)	Roche Diagnostics	N/A
<i>Bst</i> Polymerase, Large Fragment (8 U/μl)	New England Biolabs GmbH	N/A
dNTPs (each 10 mM)	QIAGEN, Hilden, Germany	N/A
dNTPs (each 25 mM)	Agilent Technologies	N/A
EDTA (0.5 M), pH 8.0	Ambion/Applied Biosystems, Life Technologies [™] , Darmstadt, Germany	N/A
Herculase II Fusion	Agilent Technologies	N/A
Herculase II Reaction Buffer	Agilent Technologies	N/A
MgCl ₂ (25 mM)	Life Technologies [™]	N/A
MinElute	QIAGEN, Hilden, Germany	N/A
MSB ® Spin PCRapace	Invitek, Stratec Molecular, Berlin, Germany	N/A
Sodium N-lauryl sarcosinate	Merck Millipore, Darmstadt, Germany	N/A
Nuclease-free H ₂ O	Life Technologies [™]	N/A
PEG-4000	Thermo Scientific [™]	N/A
Proteinase K	Roche Diagnostics, Mannheim, Germany	N/A
Phenol/chloroform/isoamylalcohol (25:24:1)	Roth, Karlsruhe, Germany	N/A
Qubit ® Fluorometric quantitation and dsDNA HS Assay Kit	Invitrogen TM	N/A
T4 DNA Ligase (5 U/μl)	Thermo Scientific [™]	N/A
T4 DNA Ligase Buffer (10X)	Thermo Scientific [™]	N/A
T4 DNA Polymerase (5 U/μl)	Thermo Scientific [™]	N/A
T4 Polynucleotide Kinase	Invitrogen TM	N/A
Tango Buffer (10x)	Life Technologies TM	N/A



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REAGENT or RESOURCE	SOURCE	IDENTIFIER
ThermoPol Buffer (10X)	New England Biolabs GmbH	N/A
Trichlormethan/Chloroform	Roth, Karlsruhe, Germany	N/A

RESOURCE AVAILABILITY

Lead Contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contacts, Joachim Burger (jburger@uni-mainz.de) and Daniel Wegmann (daniel.wegmann@unifr.ch).

Materials Availability

Genomic data are available at the European Nucleotide Archive under the accession number PRJEB38406 in BAM and FASTQ format.

Data and Code Availability

All genomic data used in this study is publicly available at the European Nucleotide Archive under the accession number PRJEB38406 or at the sources listed in the Key Resources Table. The code used to plot Figure S1 is available on Bitbucket at https://bitbucket.org/wegmannlab/atlas/downloads/Relatedness.R, the code used to estimate allelic age is available at https://bitbucket.org/wegmannlab/atlas/downloads/Allele_Age_MC_ABC_Sampling_V2.R.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Archaeological and morphological information

Sometime after 1,300 BC, the Tollense Valley saw an armed conflict of a scale hitherto unimagined for the European Bronze Age. Likely more than 1,000, perhaps even more than 2,000 combatants fought each other in the river valley. Long-distance as well as close-range weapons (swords, spears/lances, bow and arrow, wooden clubs and hammers) were used, and the number of casualties, attested to by a large number of perimortem lesions, was high [81]. The bones show perimortem as well as healed lesions, documenting the actual fighting in the valley, but in some cases also violent encounters in previous life. Cases include arrowheads still embedded in bone, small cuts and funnel-shaped lesions indicating arrow shots and stab wounds, as well as large impression fractures, caused by blunt objects [82–84]. These lesions prove hand to hand combat as well as long distance arrow shots. According to morphological criteria the skeletons are very heterogeneous and consist mainly of young adult males. According to the number of individuals and supported by Sr-isotope analyses the skeletal samples include local and non-local warriors [28].

The Tollense Valley (Mecklenburg-Western Pomerania) is characterized by a low moorland environment. Neighboring ground moraine areas have been favored settlement areas since the Neolithic. In 2007, systematic archaeological research in the valley was initiated, including excavations, underwater surveys, and metal detector surveys, as well as scientific analyses on bones and other finds. Human skeletal remains are discovered from a stretch of river of 2.5 km length. Based on a series of more than 100 radio-carbon dates on skeletal and wooden remains from the Tollense Valley, the finds horizon of the Bronze Age battle can now be dated to the first half of the 13th century BC. With one exception (WEZ16 was directly C¹⁴-dated to the Neolithic period) all samples are associated with the Bronze Age (c. 1,300- 1,200 calBC). The event is interpreted as one major conflict, not a series of chronologically divergent skirmishes, due to dating results and the general appearance of the finds layer with comparable disarticulation as well as lesion patterns of the human skeletal remains at all analyzed sites [85]. From 2009 to 2015, extensive excavations funded by the German Research Foundation (DFG) were conducted at the main site Weltzin 20 (exposing an area of c. 462 m²), less extensive excavations were conducted at Weltzin 32; in both cases supported by comprehensive underwater surveys. Archaeological research on the Bronze Age site in the Tollense Valley has uncovered remains of a minimum number of more than 140 individuals thus far, predominantly of young adult men. These remains represent a special Bronze Age skeletal sample due to the large number of individuals and their mortality profile.

The sampled burials from the Eneolithic and Bronze Age period in the Steppe region date from the second half of the 4th millennium and the 3rd millennium B.C. They are associated with the cultures of the Eneolithic, Yamnaya, and Catacomb Grave. Yamnaya Culture graves from the 1st half of 3rd mill. BCE were sampled in a region between the Urals and the Balkan (Figure 3; Table S3). Skeletons associated with the Catacomb Culture were sampled from the area between the rivers Wolga and Prut. While the early phase of the Catacomb Grave Culture is still synchronous to late Yamnaya, the developed and late phases of the Catacomb Culture date to 2,500-2,000 BCE. There is still an ongoing debate on the cultural interpretation of burials in the Middle Wolga region which are often described as Poltavkinskaya Culture to emphasize characteristic differences regarding funerary customs [86].

All sampled burials of the East European Steppe date to the 3rd mill. BCE except three earlier graves attributed to the Late Eneolithic, including the Usatovo Culture. Only one burial (sample No MOB 1) was a flat grave, all others were excavated in burial mounds (Table S3).



METHOD DETAILS

Sample preparation

Sample preparation and paleogenetic analyses were conducted at the Institute of Organismic and Molecular Evolution (iomE) at Johannes Gutenberg University, Mainz. Strategies to avoid and detect contaminations were applied as described elsewhere [87, 88] and included sample analysis in a dedicated ancient DNA facility separated from post-PCR laboratories, decontamination of samples, workspace and labware, independent DNA extraction as well as co-analysis of blank controls.

Tooth material or pieces of right femora were used for ancient DNA analyses to avoid double typing. For the isolation of DNA molecules 0.2-0.5 g of bone powder was incubated for 48h in 7 mL lysis buffer containing 0.48 M EDTA (pH 8.0), 0.2% sodium N-lauryl sarcosinate, and 0.1 mg/ml Proteinase K at 37°C followed by phenol-chloroform extraction. Samples were washed with nucleasefree water and concentrated on 50-kDA Amicon Ultra-15 Centrifugal Filter Units (Millipore) to a final volume of 150-350 µl.

Library construction and amplification

Libraries were constructed as detailed in Meyer and Kircher [69] with minor changes. After blunt end repair, samples were purified with the MinElute PCR purification kit (QIAGEN) following the manufacturer's instructions. *Bst* polymerase was heat inactivated for 20 minutes at 80°C [15]. Indexing PCR was performed with 1x Ampli Taq Gold ® Buffer II, 2.5 mM MgCl₂, 0.4 mg/ml BSA, 200 μ M dNTPs, 0.05 U/ μ l AmpliTaq Gold® Polymerase, 0.2 μ M index primer and either 0.2 μ M IS4 or IS7, and 8 μ l of DNA library in a final volume of 50 μ l. Libraries were amplified in five parallels and 10 to 15 PCR cycles followed by a second amplification in a final volume of 100 μ l with 1x Herculase II Reaction Buffer (Agilent Technologies), 250 μ M dNTPs (Agilent Technologies), 0.3 μ M Primer, 1 μ l of Herculase II Fusion® DNA Polymerase (Agilent Technologies) and 10 μ l of DNA library. Thermal conditions were 2 minutes at 95°C followed by eight cycles at 95°C, 60°C, and 72°C for 30 s each, and a final elongation at 72°C for 5 minutes. The DNA libraries were quantified using Qubit ® Fluorometric quantitation with the dsDNA HS Assay kit and their quality was assessed on the Agilent 2100 Expert Bioanalyzer using the High Sensitivity DNA Analysis kit according to the manufacturer's instructions.

Illumina sequencing

Prior to capture enrichment, DNA libraries were screened for endogenous DNA content on an Illumina MiSeqTM platform at StarSEQ GmbH (Mainz, Germany). Libraries were pooled equimolar and sequenced in a 50 bp single-end run. Endogenous DNA content varied between 0.4 and 48%. Libraries enriched for nuclear loci were pooled to 200 ng and sequenced on an Illumina HiSeqTM platform at the sequencing facilities of the Johannes Gutenberg University (Institute of Molecular Genetics, Mainz, Germany).

5 MB capture assay

The capture followed the protocol described in Veeramah et al. [6]. It was designed to cover 4,905,256 nucleotides (0.16% of the whole genome), including 4,687 independent neutral 1-kb regions in addition to 429 neutral regions of 500 bp length, as well as 486 phenotypic informative markers correlating with pigmentation, metabolic syndrome, diabetes risk, lactase persistence, and markers associated with non-infectious and inflammatory diseases.

Read mapping, alignment cleaning and merging

Residual adaptor sequences were trimmed from the raw reads with a custom made python script. Only reads with a minimum of 95% of all bases with quality score > 15 were kept. Reads originating from paired-end sequencing were joined using fastq-join from eautils [89] with default parameters. All reads were aligned with bwa aln [76] to hg19. During SAM/BAM conversion, alignments were filtered and only reads with a minimum mapping quality of 25 were kept. PCR duplicates were removed from the files using Mark-Duplicates from picard-tools (https://broadinstitute.github.io/picard/), prior to local re-alignment with GATK [77].

QUANTIFICATION AND STATISTICAL ANALYSIS

Sex Determination

We determined the molecular sex of each sample following Skoglund et al. [90] by calculating the number of reads aligned to the Y chromosome as a fraction of the total number of reads aligned to both sex chromosomes.

Ancient DNA Authenticity

To estimate the authenticity of the ancient sample DNA, we used contamMix, as described in Fu et al. [80]. Therefore, all reads mapping to the mitochondrial genome were extracted from the alignments and realigned against the rCRS (NC_012920.1).

Haplogroups of uniparental Markers

mt-DNA

In order to determine haplogroups for uniparental markers, majority allele calls were created with ATLAS ([70], commit 401af70, task = majorityBase). The resulting VCF files for the mitochondrial genomes were merged and uploaded to the Haplogrep 2.0 web page (https://haplogrep.uibk.ac.at/). See Table S1 for details.



Y chromosome haplogroups

Y chromosome haplogroups were determined based on majority allele calls, created by ATLAS ([70], commit 401af70, task = majorityBase). The resulting VCF files were converted to the text based plink format [91], which is the base for the input files necessary for the yhaplo script [75]. Prior to the conversion, the VCF files were filtered in two steps, with a minimum depth of two as well as five alleles per position. Haplogroup determination was therefore done three times, with all available SNPs and for each of the two filter steps for each sample. See Table S1 for details.

Genotype likelihoods in Tollense samples

Post mortem damage patterns

We used ATLAS ([70], commit d5b0aee, task = PMD) to infer empirical PMD patterns as described in Kousathanas et al. [92] from the tabulated mismatches between the raw reads and the reference genome. These patterns were inferred independently for all individuals and read groups.

Base quality score recalibration

We used ATLAS ([70], commit d5b0aee, task = recal) to recalibrate base quality scores with the method described in Kousathanas et al. [92]. This approach exploits known homozygous sites. As known homozygous sites, we used the 10 million sites with highest GERP conservation scores as calculated across the multiple sequence alignments of 88 mammals and provided by Ensembl (http:// ftp.ensembl.org/pub/release-96/bed/ensembl-compara/88_mammals.gerp_constrained_element/). For recalibration, we pooled all samples to increase power and estimated common recalibration parameters for all samples generated with the same sequencing run (with option poolReadGroups), but used individual damage patterns.

Genotype likelihoods

We used ATLAS ([70], commit 52a2276, task = GLF) to obtain genotype likelihoods for all individuals at all sites. We included the damage patterns and the recalibration parameters. We then created a set of GLF files for the phenotypic SNP's and a set of GLF files for the captured regions, restricting the analysis to all Bronze Age Tollense samples with low contamination levels and depth > 4x (denoted as Tollense > 4x in the following). For the male samples we further specified chromosomes X,Y and MT to be haploid and for the female samples chromosomes Y and MT.

Pseudo-haploid calls

Post mortem damage patterns

We used ATLAS ([70], commit 401af70, task = PMD) to infer empirical PMD patterns as described in Kousathanas et al. [92] from the tabulated mismatches between the raw reads and the reference genome. These patterns were inferred independently for all individuals and all read groups.

Base quality score recalibration

We used ATLAS ([70], commit 401af70, task = recal) to recalibrate base quality scores with the method described in Kousathanas et al. [92]. This approach exploits known homozygous sites and we used the UCNEbase regions [93]. For recalibration, we pooled all samples to increase power and estimated common recalibration parameters for all samples generated with the same sequencing run (with option poolReadGroups), but used individual damage patterns.

Genotype calling

We generated two sets of calls for all Tollense > 4x to be used in downstream analyses, both based on genotype likelihoods calculated with ATLAS ([70], commit 401af70, task = call, method = MLE): 1) Diploid Maximum Likelihood calls and 2) pseudo-haploid calls by identifying the allele with the highest support as given by the genotype likelihoods.

Kinship analysis

We used ATLAS ([70], commit d5b0aee, task = geneticDist) to estimate the squared genetic distances between all pairs of Tollense > 4x samples, using the GLF files created in step "Estimate genotype likelihoods." We then used a custom R script to apply the method of Waples et al. [94] to transform these distances into estimates of genetic relatedness. This script can be found on the ATLAS wiki (https://bitbucket.org/wegmannlab/atlas/downloads/Relatedness.R). For more details see Figure S1.

Allele count estimation

Under the assumption of population continuity, we identified loci under selection by comparing allele frequencies in the Tollense Valley population to those of a modern reference population using two complementary methods (see below). In absence of direct data from northern or Central Germany, we used the CEU population from the 1000Genomes project [64] as a modern proxy.

We downloaded the BAM files for 96 unrelated CEU genomes from EBI (http://ftp.1000genomes.ebi.ac.uk/vol1/ftp/ data_collections/1000_genomes_project/). These BAM files were based on the GRCh38 human reference while the Tollense samples were based on GRCh37. Thus, we used the UCSC Genome Browser (https://genome.ucsc.edu/) liftover tool to translate the coordinates of our capture loci from GRCh37 to GRCh38. We used GRCh38 coordinates to extract the reads that mapped to our regions of interest from the CEU BAM files, and mapped these reads to the GRCh37 reference using BWA mem [95]. Since the downloaded BAM files already had recalibrated quality scores, we did not recalibrate the BAM files with the ATLAS pipeline but directly created GLF files, as specified above for the Tollense samples.

In a first step, we used ATLAS ([70], commit 52a2276, task=majorMinor) to identify the major and minor allele and to create a VCF containing genotype likelihoods of all Tollense > 4x and CEU individuals. Then, we used ATLAS ([70], commit 52a2276, task=allele-Counts) to estimate allele counts in the Tollense > 4x and CEU populations separately.



To ensure positive selection coefficients, we estimated selection coefficients for the allele with a higher frequency in CEU than Tollense > 4x.

Selection inference with ApproxWF

Population size estimation

We used the estimated allele counts at the putatively neutral regions to estimate the population size and sequencing error rate with ApproxWF ([29], commit a19e439) under a constant size model and assuming no selection. We assumed that 130 generations have passed between the Tollense battle and the modern CEU reference population, implying a generation time of roughly 25 years. We ran the MCMC for 100,000 iterations and specified that the error rate should be estimated. This resulted in posterior medians for the population size of 2,503 and the error rate of 0.02.

Estimating selection coefficients

We used ApproxWF to estimate selection coefficients for all loci while fixing the population size and sequencing error rate to the values inferred from neutral loci (see above). We ran the estimation assuming a dominance coefficient h = 0.5 (additive) for all loci. We then reran the estimation also with h = 1 (dominant) for the loci associated with lactase persistence.

Selection inference with forward simulations

To test if allele frequency changes over time are the result of drift, or if selection has to be invoked, a forward simulation approach assuming exponential population growth was used, as described in Wilde et al. [30]. For each simulation run, the initial allele frequencies were drawn at random from a beta ($n_p + 1$, $n_q + 1$) distribution, where n_p and n_q denote the estimated allele counts in the Tollense > 4x population. Allele frequencies in each following generation were simulated by binomial sampling from the previous generation, while the estimations for the final generation were saved for each simulation run.

Since the size of the population from which the Tollense > 4x sample originates is unknown, we ran the analysis for 50 initial population sizes 10^2 and 10^5 , evenly spaced on the log scale. Population sizes were then assumed to grow exponentially over 130 discrete generations. The final population size was set to $8 \cdot 10^6$, which corresponds to approximately one tenth of the Census population of Germany (~83 million, 31.12.2018, https://de.wikipedia.org/wiki/Deutschland).

Under this demographic model, we generated 10^4 allele frequency trajectories for each combination of an initial population size and 51 evenly spaced selection coefficients between -10 and 10. Finally, we calculated a Z-score to determine if the frequency in the modern reference population was outside the distribution of allele frequencies obtained with these simulations. The Z-score was then translated into a two-tailed p value.

Principal component analysis

We generated PCA's using LASER (version 2.04 [73];). In a first step, a reference space was generated, with a principal components analysis on the genotype data of modern individuals, with imputation of missing entries by averaging over all individual genotypes at that position (data published as part of reference [96]; Southern European [Italian North/South, Spanish /North, Canary Islander, Maltese, Greek], Sicilian, Basque, Sardinian, Cypriot, Central- and Eastern European [Albanian, Bulgarian, Romanian, Hungarian, Czech, German, French], Croatian, Slavic [Russian, Ukrainian, Belarusian, Polish, Sorb, Mordovian], Baltic and Finnish [Estonian, Lithuanian, Finnish], British Isles [English, Orcadian, Scottish, Irish /Ulster, Shetlander], Icelandic, Norwegian). In the following step, samples with depth > 4x were projected onto the reference space based on ten replicates, using pileup files generated with Samtools (version 1.9 [78],) with the suggested filter criteria of minimum mapping quality 30 and minimum base quality 20.

F_{ST} and TreeMix

Reference samples

We extracted all SNP's from whole genome sequences generated from European (CEU, FIN, GBR, IBS and TSI, together named EUR), SouthEast Asian (GIH, PJL, BEB, STU, and ITU, together named SAS), East Asian (CHB, JPT, CHS, CDX and KHV, together named EAS) and Yoruban (YRI) populations that are part of the 1000 Genomes Project. 1000 Genomes SNPs were obtained using VCFtools [79]. We complemented this dataset with the Genome DK (Danish), GONL (Dutch) and Turkish modern samples [65], as well as 5th century Bavarians [6]. Datasets were merged using PLINK [72] and custom Python scripts.

We restricted our analysis to transversion SNPs within the 5K neutralome and to those whose frequencies are likely to vary due to drift (rather than novel mutations) by ascertaining variable sites in the Yoruba population (n = 31,494 SNPs). We used pseudo-haploid calls (see above) for the Tollense > 4x sample, but diploid calls for the reference populations.

F_{ST} analysis

We estimated genome-wide SNP based pairwise F_{ST} between the Tollense > 4x and reference samples as described in Bhatia et al. [97]. Standard errors and 95% confidence intervals were estimated using the delete-m jackknife method [98], with all m-SNPs within each of the 1kb locus successively removed (see also Figure S2).

Treemix analysis

We inferred a phylogenetic tree relating all populations using TreeMix 1.12 [71]. Though we recognize that there is unlikely to be a strict bifurcating structure to the population history of this data, this analysis can help extract useful relationships among these populations. To ensure there were no biases due to sample size differences among populations, all allele frequencies were resampled

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down to 14 haploid alleles to match the Tollense > 4x dataset. We assumed no migration and grouped SNPs into blocks of 100 to control for linkage disequilibrium (option -k 100) (see also Figure S2).

Current Biology

Report

D-statistics with ancient individuals

In order to examine differentiation among Tollense > 4x individuals in their relationship to other populations, we performed D-statistic analyses with ADMIXTOOLS 5.1 [74] in the form of D(Tollense1, Tollense2; Test_Population, YRI) for each Tollense pair and a reference population. The reference populations were individuals from our ancient reference set used for F_{ST} and TreeMix with additional ancient reference samples [2, 3, 9, 22, 68], with ages within 1,000 years of the Tollense samples grouped based on modern country borders (see also Table S2). To be consistent with how the ancient reference samples were called, pseudo-haploid genotypes were called for each Tollense individual by randomly sampling a read at each of the 1.2 million target SNPs. We note that this analysis relies on off-target reads in each Tollense sample, and as a consequence the SNP numbers involved in each test are low (mean 11,078.2 SNPs). We used a custom python script that allowed multithreading to perform all tests. In addition, the same reference populations were used to test the partition of our samples suggested by the PCA into upper (WEZ24, WEZ40, WEZ51, WEZ56, WEZ58, WEZ71, WEZ83) and lower (WEZ35, WEZ39, WEZ54, WEZ57, WEZ59, WEZ61) clusters.

F_{ST} permutation test

We created a multisample vcf. file from the MLE-calls for the Samples > 4x to calculate F_{ST} 's for every possible combination of samples in group sizes from four to seven, toward the remaining samples using Hudson's estimator [97]. Therefore a set of SNPs was selected for which a minimal depth of two reads, a minimal genotype quality of 15, a minor allele frequency of ten or higher, as well as a call rate of 100% was found (1947 SNPs). For comparison the same was done for 14 samples drawn at random from CEU from the 1000 Genomes project (NA06986, NA07357, NA10847, NA11840, NA11881, NA11994, NA11995, NA12286, NA12287, NA12347, NA12414, NA12760, NA12777, NA12873). Results for the ancient samples were compared to the results for CEU by randomly sampling a F_{ST} from each distribution of results and subtracting the F_{ST} found in CEU from the F_{ST} found in the ancient samples. After 100,000 repetitions the fraction of values < 0 was calculated. Thereby we were able to approximate the probability of obtaining a smaller F_{ST} in internal comparisons of the ancient samples, compared to a modern population.

Inferring global inbreeding coefficient F

We implemented a method to quantify the Wahlund Effect with the inbreeding coefficient *F*. *Model*

Let *F* be defined as the global deficit in heterozygotes in a population of individuals i = 1, ..., I at all loci l = 1, ..., L. The genotype of an individual (*i*) at a locus (*l*) is denoted by g_{il} and the alternative allele's frequency in the population by p_l .

We define two sub-models for $F: M_F$, where $0 \le F \le 1$ and the genotype are distributed according to Hardy-Weinberg Equilibrium with inbreeding (Equation 1), and M_{HWE} , where the genotype frequencies are determined by the allele frequencies according to Hardy-Weinberg Equilibrium (Equation 2).

$$P(g_{il}|M_F, F, p_l) = \begin{cases} (1-F)(1-p_l)^2 + F(1-p_l) & \text{if } g_{il} = 0\\ (1-F)2p_l(1-p_l) & \text{if } g_{il} = 1\\ (1-F)p_l^2 + Fp_l & \text{if } g_{il} = 2 \end{cases}$$
Equation 1

$$P(g_{ii}|M_{HWE}, F, p_i) = \begin{cases} (1 - p_i)^2 & \text{if } g_{ii} = 0\\ 2p_i(1 - p_i) & \text{if } g_{ii} = 1\\ p_i^2 & \text{if } g_{ii} = 2 \end{cases}$$
 Equation 2

In order to infer F and p from the sequencing data (d) we assume independence of loci and individuals.

We define two submodules for the allele frequency: M_p for polymorphic loci, where $p \sim B(\gamma)$, and M_0 for monomorphic loci. The likelihood function of our model is shown in Equations 3 and 4, where the products runs over all loci in model M_p .

$$P(g_{ii}|M_F, F, p_i, \gamma) = \prod_{M_p^{(i)}} P(p_i|\gamma) \left[\prod_i \sum_g P(d_{ii}|g) P(g|F, p_i) \right]$$
Equation 3

$$P(g_{ii}|M_{HWE}, p_{l}, \gamma) = \prod_{M_{p}^{(l)}} P(p_{l}|\gamma) \left[\prod_{i} \sum_{g} P(d_{ii}|g) P(g|p_{l}) \right]$$
Equation 4

We estimate posterior distributions of the parameters $\theta = \{F, p, \gamma, \pi\}$ with a reversible-jump Metropolis-Hastings algorithm. **Priors**

We assume $P(M_F) = P(M_{HWE})$ and $F \sim U(0, 1)$. We define $P(M_p) = \pi$ and $P(M_0) = 1 - \pi$ and $\pi \sim U(0, 1)$. The polymorphic loci are modeled according to a symmetric beta distribution: $p_l \sim B(\gamma)$. We assume extreme allele frequencies to be the most probable,



i.e., γ to be small. For ease of implementation we update $c = log(\gamma)$, where $c \sim U(-\infty, \infty)$. This variable transformation has the benefit of allowing the use of the symmetric uniform prior for γ instead of the asymmetric exponential distribution.

Inference

We used ATLAS ([70], commit 52a2276, task = majorMinor) to identify the major and minor allele and to create a VCF containing genotype likelihoods of all Tollense > 4x from the GLF files (see above). We then used ATLAS ([70], commit 52a2276, task = inbreeding) to quantify population structure among the Tollense > 4x sample by estimating the inbreeding coefficient *F*, running the MCMC for 10^6 iterations with the parameters minVariantQuality = 40, numBurnins = 10 burninLength = 500. We reran this analysis also with argument probMovingToModelNoF = 0 to restrict the analysis to a model with *F* > 0.

LP allele frequency estimation for NGS data

We estimated the allele frequencies in Table 1 using ATLAS ([70], commit 52a2276, task alleleFreq, doBayesian). Whole genome sequencing was available for Tollense, CEU and Mokrin and we used the genotype likelihoods in the VCFs created as described in section Allele count estimation as input. For the other NGS populations in Table 1, we downloaded the published BAM files and analyzed them with the standard GATK pipeline [77] to obtain genotype likelihoods for rs4988235-A. Frequencies for rs4988235-A for East European Steppe: 3600-2300 BCE and Area of Corded Ware Culture: 2900-2300 BCE are based on pseudo-haploid calls, taken from the V42.4 version of the downloadable genotypes of present-day and ancient DNA data (compiled from published papers), provided by the Reich Lab (https://reich.hms.harvard.edu/downloadable-genotypes-present-day-and-ancient-dna-data-compiled-published-papers).

We used ATLAS ([70], task = compareAlleleFreq, commit 8af19b9) to obtain posterior probabilities of the allele frequency of Tollense to be smaller than the allele frequency of other populations. For this, we assumed the allele frequencies to follow a Beta distribution with parameters alpha and beta set to 0.7. The results only changed minimally when using an uninformative prior with alpha and beta set to 1 and did not change the fact that Medieval populations had higher frequencies for rs4988235-A than the Bronze Age populations.

LP allele frequency estimation for PCR data

In Europe, LP is caused by a single C to T transition 13,910 bp upstream of the LCT gene (-13,910*T, rs4988235). The LCT data was obtained through single or multiplex PCR amplification and Sanger and/or 454 sequencing (see Table S3 for details). The laboratory work was performed in dedicated ancient DNA facilities of the Palaeogenetics Group at Johannes-Gutenberg University Mainz. Methods for sample preparation, extraction, amplification and sequencing, including decontamination methods and criteria for replication and validation, followed Wilde et al. [30]. The primer sequences used in the single and multiplex PCR setups described therein are as follows: LCTa_U 5'CTGCGCTGGCAATACAGATAAGA3' and LCTa_L 5'CAAATGCAACCTAAGGAGAGAGTT3'.

In brief, at least four amplifications of two independent extractions were required to consider a genotype as fully replicated. For homozygous individuals, all four amplifications need to show the same allele, for heterozygous individuals, the alternate allele or a heterozygous signal has to be present at least twice and once in each extraction. Only nine of the 37 samples presented here do not entirely fulfill our own strict criteria but miss them by one amplification (see Table S3). We include them because the associated blank controls and additional mitochondrial and nuclear marker amplifications do not show any sign of contamination (see [30]).

We obtained the allele frequencies by dividing the derived allele count by the total amount of alleles. The Credible Intervals were determined as the 5th/95th percentile of a beta distribution, defined by $\beta(n_p + 1, n_q + 1)$, where n_p and n_q were the number of ancestral and derived alleles.

Estimating LP allele age

Consider two alleles A and a segregating in a population of size 2N. The frequency of Allele A in generation t is a random variable X(t) taking values between 0 and 2N. Under the Wright-Fisher model with selection, these frequencies form a time-homogeneous Markov chain with transition probabilities

$$P_{ij}: = P(X(t) = j | X(t-1) = i) = {\binom{2N}{k}} p_i^j (1-p_i)^{2N-j}$$
 Equation 5

where

$$p_{i} = \frac{w_{AA}i^{2} + w_{Aa}i(2N - i)}{w_{AA}i^{2} + 2w_{Aa}i(2N - i) + w_{aa}(2N - i)^{2}},$$

and $w_{AA} = 1 + s$, $w_{Aa} = 1 + hs$, and $w_{aa} = 1$ are defined by the selection coefficient s and the dominance coefficient h.

Let us denote by t_0 the time at which genomic data $D = (D_1, ..., D_n)$ is available for *n* samples. Further, let *T* denote the age of the selected allele *A*, i.e., $t_0 - T$ denotes the time at which a mutation changed its frequency from 0 to 1. We aim at inferring the posterior probability

$$P(T|D, 2N, s, h) \propto P(D|T, 2N, s, h)P(T).$$





Given known parameters 2*N*, *s* and *h*. We assume here an improper prior $P(T) \propto 1$. Since the true allele frequency at t_0 is unknown, we have

$$P(D|T,2N,s,h) = \sum_{x=1}^{2N} P(D|X(t_0) = x) P(X(t_0) = x|T,2N,s,h).$$

We use ATLAS to determine

$$P(D|X(t_0) = x) = \prod_{i=1}^{n} P(D_i|X(t_0) = x)$$

with

$$P(D_i|X(t_0) = x) = \sum_{g=0}^{2} P(D_i|g)P(g|X(t_0) = x),$$

where $P(D_i|g)$ are the genotype likelihoods calculated by ATLAS and $P(g|X(t_0) = x)$ are the Hardy-Weinberg proportions.

To determine $P(X(t_0) = x | T, 2N, s, h)$ we will resort to a Monte-Carlo scheme. Specifically, we will sample k = 1, ..., K allele frequency trajectories x_k by repeating the following steps:

- Set $x_k(0) = 1$ and time t = 0.
- Simulate the next frequency $x_k(t+1)$ according to the transition probabilities in Equation 1. Increment t.
- Repeat Step 2 for t_{max} generations or until the chain is absorbed at 0.

Let T be a matrix with elements T_{xt} representing the number of simulated trajectories that had frequency x at time t. From this matrix, we get the estimate

$$P(X(t_0) = x | T, 2N, s, h) \approx \frac{T_{xT}}{\sum_{y=1}^{2N} T_{yT}}$$

Inference

We set the parameter to: $t_{max} = 100000$, s = 0.06, $n_0 = 1$, $N \in \{100, 200, 500, 1000, 2000, 5000, 10000, 50000, 100000\}$, $h \in \{0.5, 1\}$ and ran the simulations for 10000 iterations. The allele age was transformed from generations to the Before Present (BP) timescale in the following way:

$$age = 25 * generations + 3150,$$

where 25 is the number of years per generation and 3150 is the age of Tollense in BP.