## Current Biology

## The Arrival of Siberian Ancestry Connecting the Eastern Baltic to Uralic Speakers further East

## Graphical Abstract



## Highlights

- Increase in hunter-gatherer ancestry in Bronze Age Eastern Baltic genomes
- Genetic input from Siberia to the Eastern Baltic during the transition to Iron Age
- Arrival of Siberian ancestry coincides with proposed arrival of Uralic languages
- Light eyes, hair, and skin and lactose tolerance become frequent in the Bronze Age


## Authors

Lehti Saag, Margot Laneman, Liivi Varul, ..., Valter Lang, Mait Metspalu, Kristiina Tambets

## Correspondence

lehti.saag@ut.ee (L.S.), kristiina.tambets@ut.ee (K.T.)

## In Brief

Saag et al. present aDNA from the teeth of 33 individuals from the Eastern Baltic dating to 3,200-400 years ago. They find that the Bronze Age in the Eastern Baltic brings an increase in hunter-gatherer ancestry and the Iron Age delivers genetic input from Siberia that also connects modern European Uralic speakers to those living much further east.

# The Arrival of Siberian Ancestry Connecting the Eastern Baltic to Uralic Speakers further East 

 Valeri I. Khartanovich, ${ }^{6}$ Elena R. Mikhaylova, ${ }^{7}$ Alena Kushniarevich, ${ }^{1}$ Christiana Lyn Scheib, ${ }^{1}$ Anu Solnik, ${ }^{1}$ Tuuli Reisberg, ${ }^{1}$ Jüri Parik, ${ }^{1,2}$ Lauri Saag, ${ }^{1}$ Ene Metspalu, ${ }^{1}$ Siiri Rootsi, ${ }^{1}$ Francesco Montinaro, ${ }^{1}$ Maido Remm, ${ }^{8}$ Reedik Mägi, ${ }^{9}$<br>Eugenia D'Atanasio, ${ }^{10}$ Enrico Ryunosuke Crema, ${ }^{11}$ David Díez-del-Molino, ${ }^{12,13}$ Mark G. Thomas, ${ }^{14,15}$ Aivar Kriiska, ${ }^{3}$ Toomas Kivisild, ${ }^{1,2,10}$ Richard Villems, ${ }^{1,2}$ Valter Lang, ${ }^{3}$ Mait Metspalu, ${ }^{1}$ and Kristiina Tambets ${ }^{1, *}$<br>${ }^{1}$ Estonian Biocentre, Institute of Genomics, University of Tartu, Tartu 51010, Estonia<br>${ }^{2}$ Department of Evolutionary Biology, Institute of Cell and Molecular Biology, University of Tartu, Tartu 51010, Estonia<br>${ }^{3}$ Department of Archaeology, Institute of History and Archaeology, University of Tartu, Tartu 51014, Estonia<br>${ }^{4}$ School of Humanities, Tallinn University, Tallinn 10120, Estonia<br>${ }^{5}$ Department of Slavic and Finnic Archaeology, Institute for the History of Material Culture, Russian Academy of Sciences, St. Petersburg 191186, Russia<br>${ }^{6}$ Museum of Anthropology and Ethnography (Kunstkamera), Russian Academy of Sciences, St. Petersburg 199034, Russia<br>${ }^{7}$ Saint Petersburg State University, St. Petersburg 199034, Russia<br>${ }^{8}$ Department of Bioinformatics, Institute of Cell and Molecular Biology, University of Tartu, Tartu 51010, Estonia<br>${ }^{9}$ Estonian Genome Center, Institute of Genomics, University of Tartu, Tartu 51010, Estonia<br>${ }^{10}$ Department of Human Genetics, KU Leuven, Leuven 3000, Belgium<br>${ }^{11}$ Department of Archaeology, University of Cambridge, Cambridge CB2 3ER, UK<br>${ }^{12}$ Department of Bioinformatics and Genetics, Swedish Museum of Natural History, Stockholm 104 05, Sweden<br>${ }^{13}$ Department of Archaeology and Classical Studies, Stockholm University, Stockholm 106 91, Sweden<br>${ }^{14}$ Research Department of Genetics, Evolution and Environment, University College London, London WC1E 6BT, UK<br>${ }^{15}$ UCL Genetics Institute, University College London, London WC1E 6BT, UK<br>${ }^{16}$ Lead Contact<br>*Correspondence: lehti.saag@ut.ee (L.S.), kristiina.tambets@ut.ee (K.T.)<br>https://doi.org/10.1016/j.cub.2019.04.026

## SUMMARY

In this study, we compare the genetic ancestry of individuals from two as yet genetically unstudied cultural traditions in Estonia in the context of available modern and ancient datasets: 15 from the Late Bronze Age stone-cist graves (1200-400 BC) (EstBA) and 6 from the Pre-Roman Iron Age tarand cemeteries ( $800 / 500$ BC-50 AD) (EstIA). We also included 5 Pre-Roman to Roman Iron Age Ingrian ( 500 BC450 AD) (InglA) and 7 Middle Age Estonian (12001600 AD) (EstMA) individuals to build a dataset for studying the demographic history of the northern parts of the Eastern Baltic from the earliest layer of Mesolithic to modern times. Our findings are consistent with EstBA receiving gene flow from regions with strong Western hunter-gatherer (WHG) affinities and EstIA from populations related to modern Siberians. The latter inference is in accordance with Y chromosome (chrY) distributions in present day populations of the Eastern Baltic, as well as patterns of autosomal variation in the majority of the westernmost Uralic speakers [1-5]. This ancestry reached the coasts of the Baltic Sea no later than the mid-first millennium $B C$; i.e., in the same time window as the diversification of west Uralic (Finnic) languages [6]. Furthermore, phenotypic traits often associated
with modern Northern Europeans, like light eyes, hair, and skin, as well as lactose tolerance, can be traced back to the Bronze Age in the Eastern Baltic.

## RESULTS AND DISCUSSION

The Eastern Baltic has witnessed several population shifts since people reached its southern part during the Final Paleolithic ~11,000-10,000 BC [7, 8] and its northern part during the Mesolithic $\sim 9000 \mathrm{BC}$ [9]. No genetic information is available from Paleolithic populations, but Mesolithic hunter-gatherers of the Kunda and Narva cultures were genetically most similar to Western hunter-gatherers (WHGs) widespread in Europe [10-12]. A genetic shift toward Eastern hunter-gatherer (EHG) genetic ancestry occurred with the arrival of the Neolithic Comb Ceramic culture (CCC) people $\sim 3900$ BC [10-13]. The Late Neolithic (LN) Corded Ware culture (CWC) people of Ponto-Caspian steppe origin [10-13] brought farming into the Eastern Baltic ~2800 BC, contrary to most of Europe, where the Neolithic transition was mediated by Aegean early farmers [14-19]. Human remains radiocarbon dated to the Early Bronze Age (ca. 1800-1200 BC) are rare from this region, and no ancient DNA (aDNA) data are currently available. Genetic data from succeeding Bronze Age (BA) layers in Latvia and Lithuania indicate some genetic affinities with modern Eastern Baltic populations but also notable differences [11].

In this study, we present new genomic data from Estonian Late Bronze Age stone-cist graves (1200-400 BC) (EstBA) and

Pre-Roman Iron Age tarand cemeteries ( $800 / 500$ BC-50 AD) (EstIA). The cultural background of stone-cist graves indicates strong connections both to the west and the east [20, 21]. The Iron Age (IA) tarands have been proposed to mirror "houses of the dead" found among Uralic peoples of the Volga-Kama region [22]. As this time window matches the proposed diversification period of western Uralic languages [6] and the arrival of ProtoFinnic language in the Eastern Baltic from the east [23, 24], our study considers linguistic, archaeological, and genetic data to inform on this.

One of the most notable genetic features of Eastern Baltic populations is a high frequency of Y chromosome (chrY) haplogroup (hg) N3a (nomenclature of Karmin et al. [25]), a characteristic shared mostly with Finno-Ugric-speaking groups in Europe and several populations all over Siberia [1-5]. The rapid expansion of people carrying these lineages likely took place within the last 5,000 years [1], but their arrival time in the Eastern Baltic remains unresolved. The gene flow from Siberia to west-ern-Uralic-speaking populations has also recently been inferred using autosomal data [5, 26]. However, available aDNA data have not revealed chr Y hg N lineages in Eastern Baltic individuals [10-13].

To characterize the genetic ancestry of people from the so-far-unstudied cultural layers, we extracted DNA from the tooth roots of 56 individuals (Figure 1A; Table S1; STAR Methods). No individuals were included from later IA in Estonia because people were mostly cremated during that period. Individuals morphologically sexed as males were prioritized in sampling to make comparisons using autosomal and both sex chromosomes. We shotgun sequenced all samples and they formed 3 groups: (1) 15 with low endogenous DNA content and resulting coverage, which were excluded from further analyses; (2) 8 with sufficient mtDNA (and in some cases, chrY) coverage for determining hgs, but not for informative autosomal analyses; and (3) 33 that yielded sufficient autosomal data for informative analyses. The 33 individuals included 15 from EstBA, 6 from EstIA, 5 from Pre-Roman to Roman Iron Age Ingria ( 500 BC 450 AD) (IngIA), and 7 from Middle Age Estonia (1200-1600 AD) (EstMA) and yielded endogenous DNA ~4\%-88\%, average genomic coverages $\sim 0.017-0.734 \times$, and contamination estimates $<4 \%$ (Table S1). We analyzed the data in the context of modern and other ancient individuals, including from Neolithic Estonia [13].

## Temporal Dynamics of Maternal and Paternal Lineages in Estonia

We identified mtDNA hgs for 41 individuals (Table 1). We then compared these with over 2,000 present-day Estonian whole mtDNA sequences (unpublished data; cohort [29]) and found that all the hgs are also present in modern Estonia and are not restricted to a particular region.

We identified chrY hgs for 30 male individuals (Tables 1 and S2; STAR Methods). All 16 successfully haplogrouped EstBA males belonged to hg R1a, showing no change from the CWC period, when this was also the only chrY lineage detected in the Eastern Baltic [11, 13, 30, 31]. Three EstIA and two IngIA individuals also belonged to hg R1a, but three EstIA males belonged to hg N3a, the earliest so far observed in the Eastern Baltic. Three EstMA individuals belonged to hg N3a, two to hg

R1a, and one to hg J2b. ChrY lineages found in the Baltic Sea region before the CWC belong to hgs I, R1b, R1a5, and Q [10-13, 17, 32]. Thus, it appears that these lineages were substantially replaced in the Eastern Baltic by hg R1a [10-13], most likely through steppe migrations from the east [30, 31]. Although we did not detect N3a chrYs in our BA sample, unlike in BA Fennoscandia [26], we cannot rule out its presence due to small sample size. However, the frequency should not exceed 0.17 with $95 \%$ and 0.25 with $99 \%$ confidence [33]. The frequency of hg N3a was significantly higher in our IA than our BA group (Fisher's exact test $p$ value 0.013 ). Our results enable us to conclude that, although the expansion time for R1a1 and N3a3'5 in Eastern Europe is similar [25], hg N3a likely reached Estonia or at least became comparably frequent to modern Estonia [1] only during the BA-IA transition.

## Autosomal Ancestries in Estonia from the Bronze Age Onward

To assess whether the Eastern Eurasian influence indicated by chrY hg N3a is apparent elsewhere in the genome, we first applied principal-component analysis (PCA). We projected ancient genomes from previous studies (Table S3) and this study on two axes inferred using Estonian Biocenter Illumina genotyping array data (EBC-chipDB) of modern Western Eurasian individuals (Table S3) (Figure 1C). A clear shift toward West Eurasian hunter-gatherers is visible between European LN and BA (including Baltic CWC) and EstBA individuals, the latter clustering together with Latvian and Lithuanian BA individuals [11]. EstIA, IngIA, and EstMA individuals project between BA individuals and modern Estonians, partially overlapping with both.

We performed ADMIXTURE analysis by projecting aDNA data on worldwide EBC-chipDB modern data (Figures S1C and S1D; Table S3) and present results at K = 9 (Figures 1B, S1A, and S1B; STAR Methods). EstBA individuals are clearly distinguishable from Estonian CWC individuals as the former have more of the blue component most frequent in WHGs and less of the brown and yellow components maximized in Caucasus hunter-gatherers and modern Khanty, respectively. The individuals of EstBA, EstIA, IngIA, EstMA, and modern Estonia are quite similar to each other on average, indicating that the relatively high proportion of WHG ancestry in modern Eastern Baltic populations compared to other present-day Europeans [15] traces back to the BA.

When comparing Estonian CWC and EstBA using autosomal outgroup f3 and Patterson's D statistics (Table S3), the latter is more similar to other Baltic BA populations, to Baltic IA and Middle Age (MA) populations, and also to populations similar to WHGs and Scandinavian hunter-gatherers (SHGs), but not to Estonian CCC (Figures 2A and S2A; Data S1). The increase in WHG or SHG ancestry could be connected to western influences seen in material culture [20, 21] and facilitated by a decline in local population after the CCC-CWC period [20]. A slight trend of bigger similarity of Estonian CWC to forest or steppe zone populations and of EstBA to European early farmer populations can also be seen. These differences remain when over 900,000 positions of the '1240k' capture [16] are used instead of $\sim 500,000$ positions of the EBC-chipDB (Figure S2B; Data S1). When comparing to modern populations, Estonian CWC is


Figure 1. Geographical Locations, ADMIXTURE, and Principal-Component Analyses Results
(A) Map of the geographical locations of the individuals of this study.
(B) ADMIXTURE analysis results for a selection of ancient population averages at K9 with ancient individuals projected onto the modern genetic structure. The $x$ axis shows the proportions of the ancestral components.
(C) Principal-component analysis results of modern West Eurasians with ancient individuals projected onto the first two components (PC1 and PC2).

BA, Bronze Age; EF, early farmers; HG, hunter-gatherers; IA, Iron Age; IMA, Iron/Middle Ages; LN, Late Neolithic; LNBA, Late Neolithic/Bronze Age; MA, Middle Ages. See also Figure S1 and Table S3.
slightly more similar to Caucasus individuals but EstBA to Baltic populations and Finnic speakers (Figure 2B; Data S1). Outgroup f 3 and D statistics do not reveal apparent differences when comparing EstBA to EstIA, EstIA to InglA, and EstIA to EstMA (Data S1). These results highlight how uniparental and autosomal data can lead to different demographic inferences-the genetic
change between CWC and BA not seen in uniparental lineages is clear in autosomal data and the appearance of chrY hg N in the IA is not matched by a clear shift in autosomal profiles.

We also tested for sex biases by comparing outgroup f3 statistics calculated on autosomal $(A)$ and $X$ chromosomal $(X)$ data. The high X-to-A ratio of European-early-farmer-related ancestry

Table 1. Archaeological Information, Genetic Sex, mtDNA and Y Chromosome Haplogroups, and Average Coverage of the Individuals of This Study

| Individual | Location | Period | Date | Sex |  | MT hg | Y hg | Av. cov. |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  | Morph. | Gen. |  |  |  |
| X02 | Iru, Harju, EST | BA | 1090-910 BC ${ }^{\text {a }}$ | M | XY | T1a1b | R1a | 0.031 |
| OLS11 | Jõelähtme, Harju, EST | BA | 1060-850 BC ${ }^{\text {a }}$ | M | XY | H1c | R1a1 | 0.214 |
| V9 | Jõelähtme, Harju, EST | BA | 1220-1010 BC ${ }^{\text {a }}$ | M | XY | K1c1h | R1a1'2 | 0.474 |
| V14 | Muuksi, Harju, EST | BA | 1280-1050 BC ${ }^{\text {a }}$ | M | $X Y$ | U2e2a1 | R1a1'2 | 0.443 |
| X05 | Muuksi, Harju, EST | BA | 1210-1010 BC ${ }^{\text {a }}$ | M | XY | T2a1b1a1 | R1a1'2 | 0.029 |
| X08 | Muuksi, Harju, EST | BA | 930-810 BC ${ }^{\text {a }}$ | M | XY | T2a1b1a2 | R1a1c | 0.306 |
| X09 | Muuksi, Harju, EST | BA | 820-770 BC ${ }^{\text {a }}$ | M | XY | J1b1a | R1a | <0.017 |
| X10 | Muuksi, Harju, EST | BA | 1220-1020 BC ${ }^{\text {a }}$ | M | XY | U5a2a1 | R1a1'2 | 0.22 |
| X11 | Napa, Ida-Viru, EST | BA | 1030-890 BC ${ }^{\text {a }}$ | M | XY | J1c2k | R1a | 0.224 |
| X12 | Napa, Ida-Viru, EST | BA | 900-790 BC ${ }^{\text {a }}$ | M | XY | W6 | R1a1'6 | 0.023 |
| X13 | Rebala, Harju, EST | BA | 780-480 BC ${ }^{\text {a }}$ | M | $?$ | K1b2a | - | <0.017 |
| X14 | Rebala, Harju, EST | BA | 780-430 BC ${ }^{\text {a }}$ | M | XY | H1b2 | R1a1c | 0.307 |
| V16 | Väo, Harju, EST | BA | 730-390 BC ${ }^{\text {a }}$ | M | XY | H1b2 | R1a1'2 | 0.22 |
| X16 | Väo, Harju, EST | BA | 1080-910 BC ${ }^{\text {a }}$ | M ? | XY | J1c4 | R1a | 0.018 |
| X17 | Väo, Harju, EST | BA | 930-810 BC ${ }^{\text {a }}$ | M | XY | U4a2b | R1a1c | 0.387 |
| X18 | Väo, Harju, EST | BA? | 1200 BC-... ${ }^{\text {b }}$ | M | XY | U3b2a | ? | <0.017 |
| X19 | Väo, Harju, EST | BA | $1200-400 \mathrm{BC}^{\text {b }}$ | ? | XX | U | - | <0.017 |
| X20 | Väo, Harju, EST | BA | 900-800 BC ${ }^{\text {a }}$ | ? | XY | U4a2b | R1a | 0.085 |
| X15 | Vehendi, Tartu, EST | BA | 1210-1000 BC ${ }^{\text {a }}$ | M ? | XY | U5b1b1 | R1a1c | 0.339 |
| OLS09 | Ilmandu, Harju, EST | IA | 540-380 BC ${ }^{\text {a }}$ | F | XX | H6a1a | - | <0.017 |
| V7 | Ilmandu, Harju, EST | IA | 790-430 BC ${ }^{\text {a }}$ | M | XY | T2a1b1a1 | R1a | $<0.017$ |
| V8 | Ilmandu, Harju, EST | IA | $730-400 B C^{\text {c }}$ | M ? | XX | HVO | - | $<0.017$ |
| OLS10 | Kunda, Lääne-Viru, EST | IA | $770-430 \mathrm{BC}^{\text {c }}$ | M | XY | H13a1a1a | N3a3'5 | 0.319 |
| V10 | Kunda, Lääne-Viru, EST | IA | $790-430 \mathrm{BC}^{\text {a }}$ | M | XY | H1a | R1a1c | 0.403 |
| V11 | Kurevere, Saare, EST | IA | 390-200 BC ${ }^{\text {a }}$ | M ? | XX | W3a1d | - | 0.277 |
| V12 | Kurevere, Saare, EST | IA | 360-40 BC ${ }^{\text {a }}$ | M ? | XY | 11a1c | N3a3a | 0.245 |
| X04 | Loona, Saare, EST | IA | $480-360 \mathrm{BC}^{\text {a }}$ | M | XY | H1c | R1a1'2 | 0.256 |
| VII3 | Poanse, Pärnu, EST | IA | 380-180 BC ${ }^{\text {a }}$ | M | XY | U5a1d | ? | <0.017 |
| VII4 | Võhma, Lääne-Viru, EST | IA | $760-400 B^{\text {a }}$ | M | XY | T1a1b | N3a3a | 0.342 |
| VII15 | Kerstovo, Ingria, RUS | IA | $45 \mathrm{BC}-77 \mathrm{AD}^{\text {a }}$ | ? | XY | U5a2a1 | R1a | 0.244 |
| VIII7 | Kerstovo, Ingria, RUS | IA | 75-200 AD ${ }^{\text {b }}$ | ? | XX | H2a1a | - | 0.062 |
| VIII8 | Kerstovo, Ingria, RUS | IA | 75-200 AD ${ }^{\text {b }}$ | ? | XY | H3h | R1a1c | 0.0517 |
| VIII9 | Kerstovo, Ingria, RUS | IA | $75-200 A D^{\text {b }}$ | ? | XX | U4a2 | - | 0.3 |
| VIII5 | Malli, Ingria, RUS | IA | 75-300 AD ${ }^{\text {b }}$ | ? | XX | T1a1b | - | 0.398 |
| Ila | Karja, Saare, EST | MA | 1230-1300 AD ${ }^{\text {b }}$ | M | XY | H3h1 | N3a3a | 0.734 |
| OLS03 | Kukruse, Ida-Viru, EST | MA | 1180-1220/1240 AD ${ }^{\text {b }}$ | M | XY | U4d1 | R1a1a'b | 0.0696 |
| IVLS09KT | Mäletjärve, Tartu, EST | MA | 1570-1600 AD ${ }^{\text {b }}$ | M | XY | H2a1 | J2b2 | 0.332 |
| Ilf | Otepää, Valga, EST | MA | 1360-1390 AD ${ }^{\text {b }}$ | M | XY | T2b | N3a3a | 0.206 |
| Ilg | Pada, Lääne-Viru, EST | MA | 1210-1230/1240 AD ${ }^{\text {b }}$ | M | XY | U4a2b | N3a3a | 0.102 |
| IIIt | Vaabina, Võru, EST | MA | 1250-1450 AD ${ }^{\text {b }}$ | F | XX | U5a2a1 | - | 0.0413 |
| ILS01 | Vana-Kuuste, Tartu, EST | MA | 1500-1625 AD ${ }^{\text {b }}$ | M | XY | H11a1 | R1a | 0.0827 |

[^0]

Figure 2. Outgroup f3 Statistics Results
Estonian Corded Ware culture (Estonia CWC; blue axis) and Estonian Bronze Age (Est BA; red axis) plotted against each other.
(A) Outgroup f3 statistics' values of form f3 (Yorubas; Estonia CWC/Est BA, ancient). EMBA, Early/Middle Bronze Age; MLBA, Middle/Late Bronze Age; NCA, Neolithic/Copper Age.
(B) Outgroup f 3 statistics' values of form f 3 (Yorubas; Estonia CWC/Est BA, modern).
See also Figure S2, Table S3, and Data S1.
observed in Estonian CWC [13] decreases over time and disappears by the MA (Figures S2C-S2F; Data S1).

We used ChromoPainter-non-negative least squares (NNLS) in the unlinked mode and qpAdm to infer mixing proportions of proxy source populations forming the genetic structure of the study populations. The best model for both analyses included WHG, Yamnaya, Central European Middle Neolithic (Central MN), and modern Nganasans as sources (STAR Methods; Data S1). The study populations have on average 36\% or 20\% WHG-, $42 \%$ or $51 \%$ Yamnaya-, and $21 \%$ or $26 \%$ Central-MNrelated ancestry as estimated by ChromoPainter or qpAdm (Figure 3; Data S1). The differences in WHG- and Yamnaya-related ancestry of the two methods could be due to the large amount of shared ancestry between those populations. Importantly, both analyses differentiate EstBA from other study popula-tions-EstBA individuals have no Nganasan-related ancestry and EstIA, IngIA, and EstMA individuals on average have 2\% or 4\% (Figure 3; Data S1). The differentiation remains when using BA or IA Fennoscandian populations [26] instead of Nganasans (Data S1). Notably, the proportion of Nganasan-related ancestry varies between 0\% and 12\% among sampled EstIA, IngIA, and EstMA individuals (Data S1), which may suggest its relatively recent admixture into the target population. Moreover, two individuals from Kunda (OLS10 and V10) have the highest proportions of Nganasan ancestry among EstIA (6\% and 8\%), one of them has chrY hg N3a, and isotopic analysis suggests neither individual being born in Kunda [34].

To consolidate the previously described evidence of genetic input from Siberia, we applied f 4 statistics (Data S1). A direct comparison between EstBA and EstIA suggests a closer affinity of EstIA to Siberian proxy Nganasan, but the result is non-significant $(|Z|=2.6)$. However, modern Estonians are significantly closer to Nganasan than EstBA $(|Z|=5.6)$, and there is no significant difference between modern Estonians and EstIA in that regard ( $|Z|=1.2$ ). Tests where Nganasans are replaced with Koryaks yield similar results, consistent with the signal relating to Siberian ancestry in general (Data S1). Additionally, the difference between EstBA and EstIA in their affinity to Nganasan can be seen through comparisons with preceding Central European $L N$ and $B A(|Z|=0.2$ and 3.2 , respectively). Furthermore, EstBA had a significantly higher affinity to WHGs than preceding CWC $(|Z|=8.7)$ or modern Estonians $(|Z|=5.1)$. We also tested the in-
than either EstBA or EstIA $(|Z|=4.9$ or 3.9). We then replaced Syrians with Yamnaya Kalmykia ( $|Z|=1.2$ or 0.6 ) and Central $\mathrm{MN}(|Z|=3$ or 2.6$)$. This indicates a slight increase in early farmer ancestry from EstBA and EstIA to modern Estonians.

Finally, we performed formal tests of continuity between individual genomes of this study and modern Estonians. We found that population continuity can be rejected for most scenarios (Data S1; p < 0.05; colored gray; Figure S3E). Taking into account modern Estonian effective population size (STAR Methods), continuity cannot be fully rejected only if the ancient sampling populations had an effective size of a few hundreds (Data S1; p>0.05; colored yellow to red; Figure S3E).

## A Case of Close Genetic Relatedness between Two Stone-Cist Grave Groups

We screened the BA, IA, and MA individuals for the presence of closely related pairs using relationship estimation from ancient DNA (READ) and discovered that two BA individuals, X14 and V16, were ${ }^{\text {nd }}$ degree relatives (Figures S3A-S3C). These individuals also shared mtDNA hg H1b2 and-like all EstBA maleschrY hg R1a (Table 1). Although chrY coverage is not sufficiently high to determine how closely related these individuals are patrilineally, their haplotypes matching across the entire mtDNA genome suggests that they were half-brothers sharing their mother or an uncle and his sister's son. Notably, the two related individuals were not buried in the same site but 13 km apart. Given the small number of just sixteen stone-cist burials available for kinship analyses from a time span of $\sim 500$ years, the finding of cross-site relatedness supports the notion that these structures were built for a limited circle of people [35], possibly the elite.
The plateau in the calibration curve hinders precisely estimating the chronological separation between the radiocarbon dates of X14 ( $2,481 \pm 30$ before present [BP]) and V16 ( $2,399 \pm$ 27 BP), with a $95 \%$ highest posterior density (HPD) -76 to 344 years (V16 dying 76 years earlier to 344 years later than X14). Given the estimated ages at death (35-45 for X14; 30-40 for V16; Table S1), female reproductive age 13-40, and assuming X14 to be the uncle of V16, the biologically plausible difference in time of the two individuals dying is -29 to 72 years (STAR Methods). This interval is associated with a probability of 0.15 and is within the 95\% HPD; hence, the radiocarbon dates do not reject the relatedness inferred from aDNA. The plausible

A


B


Figure 3. ChromoPainter-NNLS and qpAdm Results
(A) ChromoPainter-NNLS unlinked mode summarized results.
(B) qpAdm results. Error bars indicate one SE.

Central MN, Central European Middle Neolithic; EstBA, Estonian Bronze Age; EstIA, Estonian Iron Age; IngIA, Ingrian Iron Age; EstMA, Estonian Middle Ages; WHG, western hunter-gatherers. See also Table S3 and Data S1.
context of the individuals, this seems to have followed the so-called southwestern route from the Volga-Ural region [20, 21]. Notably, the Bronze to Iron Age transition period also coincides with the hypothesized
range of difference in time of deaths in case of V 16 being the uncle is -82 to 19 years, and in case V16 and X14 were halfbrothers, this becomes -42 to 32 years. Both temporal intervals are less likely than the scenario described above (probability 0.08 ).

## Frequency Changes of Phenotype Informative Alleles in the Eastern Baltic

We imputed the genotypes of 37 phenotype informative SNVs from the HlrisPlex-S system, two from TLR1, and one from MCM6 gene and a 32-bp deletion (rs333) in the CCR5 gene for Mesolithic and Neolithic individuals from Latvia and Estonia [10, 13] and the individuals of this study. We inferred a sharp increase to $>50 \%$ in the frequency of the lactase persistence variant (MCM6/rs4988235) in the Baltic area after the LN (Data S 2 ), in line with previous indications of this variant becoming common in Europe in the last 4,000-3,500 years [31, 36] and of its fast increase in populations with steppe ancestry due to local adaptation [37]. In contrast, the rs333, responsible for HIV resistance, which we first detect in a CWC individual, remains at $10 \%-25 \%$ frequency since then (Data S2), comparable to its present-day $14.8 \%$ frequency in Estonia [38]. Both TLR1 variants involved in the protection against leprosy were already present in Europe at medium-high frequencies since the Mesolithic [16, 39] (Data S2). Notably, we infer a high proportion ( $40 \%-60 \%$ ) of dark skin pigmentation in the hunter-gatherers and CWC farmers (Data S2). We infer dark skin and blue eyes for two individuals, similarly to another European Mesolithic individual [39]. However, from BA onward, we infer pale or intermediate skin pigmentation for all individuals and an increase in the proportion of blue eyes and lighter shades of hair (Data S2). This is in line with previous suggestions that light skin pigmentation alleles reached high frequencies in Europe only recently [40].

## Conclusions

We show that a component of possibly Siberian ancestry was added to the gene pool of the Eastern Baltic during the Bronze to Iron Age transition at the latest. This component is present in the autosomes and chrY of many northeastern European Uralic-speaking populations today [5, 26] but arrived in the Eastern Baltic probably later than 3,500 years ago (ya), when it reached Fennoscandia [26]. Considering the archaeological
arrival of westernmost Uralic (Finnic) languages [6] in the Eastern Baltic, supporting the idea that the spread of these languages was mediated by IA migrants from the east.

The EstBA individuals of this study, as other Baltic BA individuals [11], display more WHG ancestry compared to both earlier CWC and modern Estonians. Interestingly, we do not detect this change in their uniparental lineages. However, half of the admittedly small EstIA sample and over one-third of modern Estonian men [1] share a hg N3a chrY-common in other Uralic-speaking populations living much further east [1-5] and not found in the Eastern Baltic earlier-although the autosomes of EstIA individuals only show 3\%-5\% Siberian ancestry on average.

Furthermore, phenotypic characteristics often associated with modern Northern Europeans (light eyes, hair, and skin pigmentation and lactose tolerance) can be traced back to the Bronze Age in the Eastern Baltic.

## STAR $\star$ METHODS

Detailed methods are provided in the online version of this paper and include the following:

- KEY RESOURCES TABLE
- CONTACT FOR REAGENT AND RESOURCE SHARING
- EXPERIMENTAL MODEL AND SUBJECT DETAILS
- Information about the archaeological time periods, sites and individuals of this study
- Late Bronze Age stone-cist graves
- Pre-Roman Iron Age early tarand and other cemeteries
- Pre-Roman and Roman Iron Age cemeteries in Ingria, Russia
- Medieval rural cemeteries in Estonia
- METHOD DETAILS
- DNA extraction
- Library preparation
- DNA sequencing
- QUANTIFICATION AND STATISTICAL ANALYSIS
- Mapping

○ aDNA authentication

- Calculating general statistics and determining genetic sex
- Variant calling

○ Determining mtDNA haplogroups<br>- Y chromosome variant calling and haplogrouping<br>- Preparing the datasets for autosomal analyses<br>- Principal component analysis<br>- Outgroup f3 statistics<br>O D statistics<br>O Admixture analysis<br>- ChromoPainter/NNLS<br>- qpAdm<br>O f4 statistics<br>- Population continuity tests<br>O Kinship analysis<br>- Radiocarbon date difference probability estimation Phenotyping<br>- DATA AND SOFTWARE AVAILABILITY

## SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j. cub.2019.04.026.
A video abstract is available at $10.1016 / \mathrm{j}$. .cub.2019.04.026\#mmc7.

## ACKNOWLEDGMENTS

This work was funded by research projects of the Estonian Research Council IUT24-1 (R.V., T.K., E.M., M. Metspalu, A. Kushniarevich, Lauri Saag, and K.T.), IUT20-7 (V.L., A. Kriiska, H.V., M.L., and M. Malve), IUT34-11 (M.R.), PUT1217 (K.T., Lauri Saag, and Lehti Saag), PRG243 (M. Metspalu, Lauri Saag, C.L.S., R.M., Lehti Saag, and A.S.), and PUT1339 (A. Kushniarevich); of the EU European Regional Development Fund 2014-2020.4.01.16-0030, 2014-2020.4.01.16-0125, and 2014-2020.4.01.15-0012; European Research Council Starting Investigator Grant FP7-261213 (T.K.); Wellcome Trust Senior Research Fellowship Grant 100719/Z/12/Z (M.G.T.); Sapienza University of Rome fellowship "borsa di studio per attività di perfezionamento all'estero 2017" (E.D.); and Arheograator Ltd. (L.V. and A. Kriiska). The authors would like to thank the University of Tartu Development Fund for support to the Collegium for Transdisciplinary Studies in Archaeology, Genetics and Linguistics. We thank Ajai Kumar Pathak and Tarmo Puurand for help with analyses. Analyses were carried out using the facilities of the High Performance Computing Center of the University of Tartu.

## AUTHOR CONTRIBUTIONS

Lehti Saag, A. Kriiska, R.V., V.L., M. Metspalu, and K.T. conceived the study. M.L., L.V., M. Malve, H.V., I.G.S., V.I.K., E.R.M., A. Kriiska, and V.L. assembled skeletal samples and performed osteological analyses. Lehti Saag, A. Kushniarevich, C.L.S., A.S., T.R., J.P., and K.T. performed aDNA extraction and sequencing. Lehti Saag, Lauri Saag, E.M., S.R., F.M., M.R., R.M., E.D., E.R.C., D.D.-d.-M., M.G.T., T.K., and K.T. analyzed data. Lehti Saag, M.L., H.V., M.A.R., A. Kriiska, T.K., V.L., and K.T. wrote the manuscript with input from remaining authors.

## DECLARATION OF INTERESTS

The authors declare no competing interests.
Received: February 5, 2019
Revised: March 18, 2019
Accepted: April 9, 2019
Published: May 9, 2019

## REFERENCES

1. Ilumäe, A.M., Reidla, M., Chukhryaeva, M., Järve, M., Post, H., Karmin, M., Saag, L., Agdzhoyan, A., Kushniarevich, A., Litvinov, S., et al.
(2016). Human Y chromosome haplogroup N: a non-trivial time-resolved phylogeography that cuts across language families. Am. J. Hum. Genet. 99, 163-173.
2. Tambets, K., Rootsi, S., Kivisild, T., Help, H., Serk, P., Loogväli, E.L., Tolk, H.V., Reidla, M., Metspalu, E., Pliss, L., et al. (2004). The western and eastern roots of the Saami-the story of genetic "outliers" told by mitochondrial DNA and Y chromosomes. Am. J. Hum. Genet. 74, 661-682.
3. Pliss, L., Tambets, K., Loogväli, E.L., Pronina, N., Lazdins, M., Krumina, A., Baumanis, V., and Villems, R. (2006). Mitochondrial DNA portrait of Latvians: towards the understanding of the genetic structure of Balticspeaking populations. Ann. Hum. Genet. 70, 439-458.
4. Rootsi, S., Zhivotovsky, L.A., Baldovic, M., Kayser, M., Kutuev, I.A., Khusainova, R., Bermisheva, M.A., Gubina, M., Fedorova, S.A., Ilumäe, A.M., et al. (2007). A counter-clockwise northern route of the Y-chromosome haplogroup N from Southeast Asia towards Europe. Eur. J. Hum. Genet. 15, 204-211.
5. Tambets, K., Yunusbayev, B., Hudjashov, G., Ilumäe, A.M., Rootsi, S., Honkola, T., Vesakoski, O., Atkinson, Q., Skoglund, P., Kushniarevich, A., et al. (2018). Genes reveal traces of common recent demographic history for most of the Uralic-speaking populations. Genome Biol. 19, 139.
6. Honkola, T., Vesakoski, O., Korhonen, K., Lehtinen, J., Syrjänen, K., and Wahlberg, N. (2013). Cultural and climatic changes shape the evolutionary history of the Uralic languages. J. Evol. Biol. 26, 1244-1253.
7. Zagorska, I. (1999). The earliest settlement of Latvia. In Environmental and Cultural History of the Eastern Baltic Region, U. Miller, ed. (European Council), pp. 131-156.
8. Šatavičius, E. (2016). The first Palaeolithic inhabitants and the Mesolithic in Lithuanian territory. In A Hundred Years of Archaeological Discoveries in Lithuania, G. Zabiela, Z. Baubonis, and E. Marcinkevičiūte, eds. (Society of the Lithuanian Archaeology), pp. 8-39.
9. Kriiska, A., and Lõugas, L. (2009). Stone Age settlement sites on an environmentally sensitive coastal area along the lower reaches of the River Pärnu (south-western Estonia), as indicators of changing settlement patterns, technologies and economies. In Mesolithic Horizons, S. McCartan, R. Schulting, G. Warren, and P. Woodman, eds. (Oxford-Oakville: Oxbow Books), pp. 167-175.
10. Jones, E.R., Zarina, G., Moiseyev, V., Lightfoot, E., Nigst, P.R., Manica, A., Pinhasi, R., and Bradley, D.G. (2017). The Neolithic transition in the Baltic was not driven by admixture with early European farmers. Curr. Biol. 27, 576-582.
11. Mittnik, A., Wang, C.C., Pfrengle, S., Daubaras, M., Zarin,a, G., Hallgren, F., Allmäe, R., Khartanovich, V., Moiseyev, V., Tõrv, M., et al. (2018). The genetic prehistory of the Baltic Sea region. Nat. Commun. 9, 442.
12. Mathieson, I., Alpaslan-Roodenberg, S., Posth, C., Szécsényi-Nagy, A., Rohland, N., Mallick, S., Olalde, I., Broomandkhoshbacht, N., Candilio, F., Cheronet, O., et al. (2018). The genomic history of southeastern Europe. Nature 555, 197-203.
13. Saag, L., Varul, L., Scheib, C.L., Stenderup, J., Allentoft, M.E., Saag, L., Pagani, L., Reidla, M., Tambets, K., Metspalu, E., et al. (2017). Extensive farming in Estonia started through a sex-biased migration from the Steppe. Curr. Biol. 27, 2185-2193.e6.
14. Lazaridis, I., Nadel, D., Rollefson, G., Merrett, D.C., Rohland, N., Mallick, S., Fernandes, D., Novak, M., Gamarra, B., Sirak, K., et al. (2016). Genomic insights into the origin of farming in the ancient Near East. Nature 536, 419-424.
15. Lazaridis, I., Patterson, N., Mittnik, A., Renaud, G., Mallick, S., Kirsanow, K., Sudmant, P.H., Schraiber, J.G., Castellano, S., Lipson, M., et al. (2014). Ancient human genomes suggest three ancestral populations for present-day Europeans. Nature 513, 409-413.
16. Mathieson, I., Lazaridis, I., Rohland, N., Mallick, S., Patterson, N., Roodenberg, S.A., Harney, E., Stewardson, K., Fernandes, D., Novak, M., et al. (2015). Genome-wide patterns of selection in 230 ancient Eurasians. Nature 528, 499-503.
17. Skoglund, P., Malmström, H., Omrak, A., Raghavan, M., Valdiosera, C., Günther, T., Hall, P., Tambets, K., Parik, J., Sjögren, K.G., et al. (2014). Genomic diversity and admixture differs for Stone-Age Scandinavian foragers and farmers. Science 344, 747-750.
18. Olalde, I., Schroeder, H., Sandoval-Velasco, M., Vinner, L., Lobón, I., Ramirez, O., Civit, S., García Borja, P., Salazar-García, D.C., Talamo, S., et al. (2015). A common genetic origin for early farmers from Mediterranean Cardial and Central European LBK cultures. Mol. Biol. Evol. 32, 3132-3142.
19. Hofmanová, Z., Kreutzer, S., Hellenthal, G., Sell, C., Diekmann, Y., Díez-Del-Molino, D., van Dorp, L., López, S., Kousathanas, A., Link, V., et al. (2016). Early farmers from across Europe directly descended from Neolithic Aegeans. Proc. NatI. Acad. Sci. USA 113, 6886-6891.
20. Lang, V. (2018). Läänemeresoome Tulemised (Tartu: University of Tartu Press).
21. Lang, V. (2015). Formation of Proto-Finnic - an archaeological scenario from the Bronze Age/Early Iron Age. In Congressus Duodecimus Internationalis Fenno-Ugristarum, Oulu 2015. Plenary papers, H. Mantila, K. Leinonen, S. Brunni, S. Palviainen, and J. Sivonen, eds. (Oulu: University of Oulu), pp. 63-84.
22. Patrushev, V. (2000). The Early History of the Finno-Ugric Peoples of European Russia (Societas Historiae Fenno-Ugricae).
23. Kallio, P. (2006). Suomen kantakielten absoluuttista kronologiaa. Virittäjä 110, 2-25.
24. Häkkinen, J.P. (2009). Kantauralin ajoitus ja paikannus: perustelut puntarissa. Suom.-Ugr. Seuran Aikakauskirja 92, 9-56.
25. Karmin, M., Saag, L., Vicente, M., Wilson Sayres, M.A., Järve, M., Talas, U.G., Rootsi, S., llumäe, A.M., Mägi, R., Mitt, M., et al. (2015). A recent bottleneck of Y chromosome diversity coincides with a global change in culture. Genome Res. 25, 459-466.
26. Lamnidis, T.C., Majander, K., Jeong, C., Salmela, E., Wessman, A., Moiseyev, V., Khartanovich, V., Balanovsky, O., Ongyerth, M., Weihmann, A., et al. (2018). Ancient Fennoscandian genomes reveal origin and spread of Siberian ancestry in Europe. Nat. Commun. 9, 5018.
27. Bronk Ramsey, C. (2009). Bayesian analysis of radiocarbon dates. Radiocarbon 51, 337-360.
28. Reimer, P.J., Bard, E., Bayliss, A., Beck, J.W., Blackwell, P.G., Bronk Ramsey, C., Buck, C.E., Cheng, H., Edwards, R.L., Friedrich, M., et al. (2013). IntCal13 and Marine13 radiocarbon age calibration curves 0-50,000 years cal BP. Radiocarbon 55, 1869-1887.
29. Mitt, M., Kals, M., Pärn, K., Gabriel, S.B., Lander, E.S., Palotie, A., Ripatti, S., Morris, A.P., Metspalu, A., Esko, T., et al. (2017). Improved imputation accuracy of rare and low-frequency variants using population-specific high-coverage WGS-based imputation reference panel. Eur. J. Hum. Genet. 25, 869-876.
30. Haak, W., Lazaridis, I., Patterson, N., Rohland, N., Mallick, S., Llamas, B., Brandt, G., Nordenfelt, S., Harney, E., Stewardson, K., et al. (2015). Massive migration from the steppe was a source for Indo-European languages in Europe. Nature 522, 207-211.
31. Allentoft, M.E., Sikora, M., Sjögren, K.G., Rasmussen, S., Rasmussen, M., Stenderup, J., Damgaard, P.B., Schroeder, H., Ahlström, T., Vinner, L., et al. (2015). Population genomics of Bronze Age Eurasia. Nature 522, 167-172.
32. Günther, T., Malmström, H., Svensson, E.M., Omrak, A., SánchezQuinto, F., Kılınç, G.M., Krzewińska, M., Eriksson, G., Fraser, M., Edlund, H., et al. (2018). Population genomics of Mesolithic Scandinavia: Investigating early postglacial migration routes and highlatitude adaptation. PLoS Biol. 16, e2003703.
33. Burger, J., Kirchner, M., Bramanti, B., Haak, W., and Thomas, M.G. (2007). Absence of the lactase-persistence-associated allele in early Neolithic Europeans. Proc. NatI. Acad. Sci. USA 104, 3736-3741.
34. Oras, E., Lang, V., Rannamäe, E., Varul, L., Konsa, M., Limbo-Simovart, J., Vedru, G., Laneman, M., Malve, M., and Price, T.D. (2016). Tracing
prehistoric migration: isotope analysis of Bronze and Pre-Roman Iron Age coastal burials in Estonia. Est. J. Archaeol. 20, 3-32.
35. Lang, V. (2011). Traceless death. Missing burials in Bronze and Iron Age Estonia. Est. J. Archaeol. 15, 109-129.
36. Olalde, I., Brace, S., Allentoft, M.E., Armit, I., Kristiansen, K., Booth, T., Rohland, N., Mallick, S., Szécsényi-Nagy, A., Mittnik, A., et al. (2018). The Beaker phenomenon and the genomic transformation of northwest Europe. Nature 555, 190-196.
37. Mathieson, S., and Mathieson, I. (2018). FADS1 and the timing of human adaptation to agriculture. Mol. Biol. Evol. 35, 2957-2970.
38. Adojaan, M., Mölder, T., Männik, A., Kivisild, T., Villems, R., Krispin, T., and Ustav, M. (2007). High prevalence of the CCR5Delta32 HIV-resistance mutation among Estonian HIV type 1-infected individuals. AIDS Res. Hum. Retroviruses 23, 193-197.
39. Olalde, I., Allentoft, M.E., Sánchez-Quinto, F., Santpere, G., Chiang, C.W.K., DeGiorgio, M., Prado-Martinez, J., Rodríguez, J.A., Rasmussen, S., Quilez, J., et al. (2014). Derived immune and ancestral pigmentation alleles in a 7,000-year-old Mesolithic European. Nature 507, 225-228.
40. Skoglund, P., and Mathieson, I. (2018). Ancient genomics of modern humans: the first decade. Annu. Rev. Genomics Hum. Genet. 19, 381-404.
41. Li, J.Z., Absher, D.M., Tang, H., Southwick, A.M., Casto, A.M., Ramachandran, S., Cann, H.M., Barsh, G.S., Feldman, M., CavalliSforza, L.L., and Myers, R.M. (2008). Worldwide human relationships inferred from genome-wide patterns of variation. Science 319, 1100-1104.
42. Behar, D.M., Yunusbayev, B., Metspalu, M., Metspalu, E., Rosset, S., Parik, J., Rootsi, S., Chaubey, G., Kutuev, I., Yudkovsky, G., et al. (2010). The genome-wide structure of the Jewish people. Nature 466, 238-242.
43. Yunusbayev, B., Metspalu, M., Järve, M., Kutuev, I., Rootsi, S., Metspalu, E., Behar, D.M., Varendi, K., Sahakyan, H., Khusainova, R., et al. (2012). The Caucasus as an asymmetric semipermeable barrier to ancient human migrations. Mol. Biol. Evol. 29, 359-365.
44. Yunusbayev, B., Metspalu, M., Metspalu, E., Valeev, A., Litvinov, S., Valiev, R., Akhmetova, V., Balanovska, E., Balanovsky, O., Turdikulova, S., et al. (2015). The genetic legacy of the expansion of Turkic-speaking nomads across Eurasia. PLoS Genet. 11, e1005068.
45. Raghavan, M., Skoglund, P., Graf, K.E., Metspalu, M., Albrechtsen, A., Moltke, I., Rasmussen, S., Stafford, T.W., Jr., Orlando, L., Metspalu, E., et al. (2014). Upper Palaeolithic Siberian genome reveals dual ancestry of Native Americans. Nature 505, 87-91.
46. Rasmussen, M., Li, Y., Lindgreen, S., Pedersen, J.S., Albrechtsen, A., Moltke, I., Metspalu, M., Metspalu, E., Kivisild, T., Gupta, R., et al. (2010). Ancient human genome sequence of an extinct Palaeo-Eskimo. Nature 463, 757-762.
47. Kushniarevich, A., Utevska, O., Chuhryaeva, M., Agdzhoyan, A., Dibirova, K., Uktveryte, I., Möls, M., Mulahasanovic, L., Pshenichnov, A., Frolova, S., et al.; Genographic Consortium (2015). Genetic heritage of the Balto-Slavic speaking populations: a synthesis of autosomal, mitochondrial and Y-chromosomal data. PLoS ONE 10, e0135820.
48. Behar, D.M., Metspalu, M., Baran, Y., Kopelman, N.M., Yunusbayev, B., Gladstein, A., Tzur, S., Sahakyan, H., Bahmanimehr, A., Yepiskoposyan, L., et al. (2013). No evidence from genome-wide data of a Khazar origin for the Ashkenazi Jews. Hum. Biol. 85, 859-900.
49. Fedorova, S.A., Reidla, M., Metspalu, E., Metspalu, M., Rootsi, S., Tambets, K., Trofimova, N., Zhadanov, S.I., Hooshiar Kashani, B., Olivieri, A., et al. (2013). Autosomal and uniparental portraits of the native populations of Sakha (Yakutia): implications for the peopling of Northeast Eurasia. BMC Evol. Biol. 13, 127.
50. Unterländer, M., Palstra, F., Lazaridis, I., Pilipenko, A., Hofmanová, Z., Groß, M., Sell, C., Blöcher, J., Kirsanow, K., Rohland, N., et al. (2017). Ancestry and demography and descendants of Iron Age nomads of the Eurasian Steppe. Nat. Commun. 8, 14615.
51. Damgaard, P.B., Marchi, N., Rasmussen, S., Peyrot, M., Renaud, G., Korneliussen, T., Moreno-Mayar, J.V., Pedersen, M.W., Goldberg, A.,

Usmanova, E., et al. (2018). 137 ancient human genomes from across the Eurasian steppes. Nature 557, 369-374.
52. de Barros Damgaard, P., Martiniano, R., Kamm, J., Moreno-Mayar, J.V., Kroonen, G., Peyrot, M., Barjamovic, G., Rasmussen, S., Zacho, C., Baimukhanov, N., et al. (2018). The first horse herders and the impact of early Bronze Age steppe expansions into Asia. Science 360, eaar7711.
53. Narasimhan, V.M., Patterson, N.J., Moorjani, P., Lazaridis, I., Mark, L., Mallick, S., Rohland, N., Bernardos, R., Kim, A.M., Nakatsuka, N., et al. (2018). The genomic formation of South and Central Asia. bioRxiv. https://doi.org/10.1101/292581.
54. Martin, M. (2011). Cutadapt removes adapter sequences from highthroughput sequencing reads. EMBnet.journal 17, 10-12.
55. Li, H., and Durbin, R. (2009). Fast and accurate short read alignment with Burrows-Wheeler transform. Bioinformatics 25, 1754-1760.
56. Li, H., Handsaker, B., Wysoker, A., Fennell, T., Ruan, J., Homer, N., Marth, G., Abecasis, G., and Durbin, R.; 1000 Genome Project Data Processing Subgroup (2009). The Sequence Alignment/Map format and SAMtools. Bioinformatics 25, 2078-2079.
57. McKenna, A., Hanna, M., Banks, E., Sivachenko, A., Cibulskis, K., Kernytsky, A., Garimella, K., Altshuler, D., Gabriel, S., Daly, M., and DePristo, M.A. (2010). The Genome Analysis Toolkit: a MapReduce framework for analyzing next-generation DNA sequencing data. Genome Res. 20, 1297-1303.
58. Jónsson, H., Ginolhac, A., Schubert, M., Johnson, P.L.F., and Orlando, L. (2013). mapDamage2.0: fast approximate Bayesian estimates of ancient DNA damage parameters. Bioinformatics 29, 1682-1684.
59. Fu, Q., Mittnik, A., Johnson, P.L.F., Bos, K., Lari, M., Bollongino, R., Sun, C., Giemsch, L., Schmitz, R., Burger, J., et al. (2013). A revised timescale for human evolution based on ancient mitochondrial genomes. Curr. Biol. 23, 553-559.
60. Korneliussen, T.S., Albrechtsen, A., and Nielsen, R. (2014). ANGSD: analysis of next generation sequencing data. BMC Bioinformatics 15, 356.
61. Skoglund, P., Storå, J., Götherström, A., and Jakobsson, M. (2013). Accurate sex identification of ancient human remains using DNA shotgun sequencing. J. Archaeol. Sci. 40, 4477-4482.
62. Weissensteiner, H., Forer, L., Fuchsberger, C., Schöpf, B., KlossBrandstätter, A., Specht, G., Kronenberg, F., and Schönherr, S. (2016). mtDNA-Server: next-generation sequencing data analysis of human mitochondrial DNA in the cloud. Nucleic Acids Res. 44 (W1), W64-W69.
63. Weissensteiner, H., Pacher, D., Kloss-Brandstätter, A., Forer, L., Specht, G., Bandelt, H.J., Kronenberg, F., Salas, A., and Schönherr, S. (2016). HaploGrep 2: mitochondrial haplogroup classification in the era of high-throughput sequencing. Nucleic Acids Res. 44 (W1), W58-W63.
64. van Oven, M., and Kayser, M. (2009). Updated comprehensive phylogenetic tree of global human mitochondrial DNA variation. Hum. Mutat. 30, E386-E394.
65. Quinlan, A.R. (2014). BEDTools: the Swiss-army tool for genome feature analysis. Curr. Protoc. Bioinformatics 47, 11.12.1-11.12.34.
66. Purcell, S., Neale, B., Todd-Brown, K., Thomas, L., Ferreira, M.A.R., Bender, D., Maller, J., Sklar, P., de Bakker, P.I.W., Daly, M.J., and Sham, P.C. (2007). PLINK: a tool set for whole-genome association and population-based linkage analyses. Am. J. Hum. Genet. 81, 559-575.
67. Patterson, N., Price, A.L., and Reich, D. (2006). Population structure and eigenanalysis. PLoS Genet. 2, e190.
68. Patterson, N., Moorjani, P., Luo, Y., Mallick, S., Rohland, N., Zhan, Y., Genschoreck, T., Webster, T., and Reich, D. (2012). Ancient admixture in human history. Genetics 192, 1065-1093.
69. Alexander, D.H., Novembre, J., and Lange, K. (2009). Fast model-based estimation of ancestry in unrelated individuals. Genome Res. 19, 16551664.
70. Lawson, D.J., Hellenthal, G., Myers, S., and Falush, D. (2012). Inference of population structure using dense haplotype data. PLoS Genet. 8, e1002453.
71. Leslie, S., Winney, B., Hellenthal, G., Davison, D., Boumertit, A., Day, T., Hutnik, K., Royrvik, E.C., Cunliffe, B., Lawson, D.J., et al.; Wellcome Trust Case Control Consortium 2; International Multiple Sclerosis Genetics Consortium (2015). The fine-scale genetic structure of the British population. Nature 519, 309-314.
72. Montinaro, F., Busby, G.B.J., Pascali, V.L., Myers, S., Hellenthal, G., and Capelli, C. (2015). Unravelling the hidden ancestry of American admixed populations. Nat. Commun. 6, 6596.
73. Monroy Kuhn, J.M., Jakobsson, M., and Günther, T. (2018). Estimating genetic kin relationships in prehistoric populations. PLoS ONE 13, e0195491.
74. Danecek, P., Auton, A., Abecasis, G., Albers, C.A., Banks, E., DePristo, M.A., Handsaker, R.E., Lunter, G., Marth, G.T., Sherry, S.T., et al.; 1000 Genomes Project Analysis Group (2011). The variant call format and VCFtools. Bioinformatics 27, 2156-2158.
75. Browning, B.L., and Browning, S.R. (2016). Genotype imputation with millions of reference samples. Am. J. Hum. Genet. 98, 116-126.
76. Gamba, C., Jones, E.R., Teasdale, M.D., McLaughlin, R.L., GonzalezFortes, G., Mattiangeli, V., Domboróczki, L., Kővári, I., Pap, I., Anders, A., et al. (2014). Genome flux and stasis in a five millennium transect of European prehistory. Nat. Commun. 5, 5257.
77. Walsh, S., Chaitanya, L., Clarisse, L., Wirken, L., Draus-Barini, J., Kovatsi, L., Maeda, H., Ishikawa, T., Sijen, T., de Knijff, P., et al. (2014). Developmental validation of the HIrisPlex system: DNA-based eye and hair colour prediction for forensic and anthropological usage. Forensic Sci. Int. Genet. 9, 150-161.
78. Walsh, S., Chaitanya, L., Breslin, K., Muralidharan, C., Bronikowska, A., Pospiech, E., Koller, J., Kovatsi, L., Wollstein, A., Branicki, W., et al. (2017). Global skin colour prediction from DNA. Hum. Genet. 136, 847-863.
79. Tõnisson, E., and Selirand, J. (1964). Nõukogude Eesti arheoloogide välitööd aastail 1958-1962. Proc. Acad. Sci. Est. SSR Soc. Sci. 13, 225-246.
80. Jaanits, K., and Lavi, A. (1978). Über die Ausgrabungen eines Steinkistengrabes in Väo. Proc. Acad. Sci. Est. SSR Soc. Sci. 27, 330-333.
81. Lõugas, V. (1981). Archäologische Rettungsgrabungen im neuen Wohngebiet Lasnamäe in Tallinn. Proc. Acad. Sci. Est. SSR Soc. Sci. 30, 390-393.
82. Lang, V., and Ligi, P. (1991). Muistsed kalmed ajaloolise demograafia allikana. In Arheoloogiline Kogumik, L. Jaanits, and V. Lang, eds. (Muinasaja teadus, 1. (Tallinn: Eesti Arheoloogiaselts)), pp. 216-238.
83. Lang, V. (1983). Ein neues Steinschiffsgrab in Nordestland. Proc. Acad. Sci. Est. SSR Soc. Sci. 32, 293-295.
84. Laneman, M., Lang, V., Malve, M., and Rannamäe, E. (2015). New data on Jaani stone graves at Väo. Northern Estonia. Est. J. Archaeol. 19, 110-137.
85. Howen, A. (1900). Ausgrabungen in Estland. Beitr. Zur Kunde Est- LivKurlands, 92-96.
86. Lõugas, V. (1975). Über die Entstehnung ortsgebundene Bodenbaukultur in Westestland. Proc. Acad. Sci. Est. SSR Soc. Sci. 24, 85-88.
87. Vassar, A. (1936). Kivikalme Nehatus. Eesti Rahvuslaste Klubid 7/8, 190-195.
88. Lõugas, V. (1976). Ausgrabungen der Steingräber und Flurrelikte in Iru. Proc. Acad. Sci. Est. SSR Soc. Sci. 25, 48-52.
89. Lõugas, V. (1983). Über die Steingräbergruppe Lastekangrud in Rebala. Proc. Acad. Sci. Est. SSR Soc. Sci. 32, 295-297.
90. Lang, V., Laneman, M., Ilves, K., and Kalman, J. (2001). Fossil fields and stone-cist graves of Rebala revisited. Archaeol. Fieldwork Est. 2000, 34-47.
91. Kalman, J. (1999). Human remains from the stone-cist graves of Rebala Lastekangrud. North Estonia. Est. J. Archaeol. 3, 19-34.
92. Kraut, A. (1985). Die Steinkistengräber von Jõelähtme. Proc. Acad. Sci. Est. SSR Soc. Sci. 34, 348-350.
93. Varul, L. (2016). Jõelähtme kivikirstkalmete 1-9, 12-24, 34-36 inimluude analüüs (Tartu: Manuscript in the archive of the archaeology department of the University of Tartu).
94. Laneman, M., and Lang, V. (2013). New radiocarbon dates for two stonecist graves at Muuksi, northern Estonia. Est. J. Archaeol. 17, 89-122.
95. Friedenthal, A. (1927). Ein gräberfeld der bronzezeit in Estland. Beitr. Zur Kunde Est. XIII, 1-2, 47-52.
96. Vassar, A. (1938). Drei Steinkistengräber aus Nordestland. Sitzungsberichte Gelehrt. Estnischen Ges. 1937, 304-364.
97. Lõugas, V., and Selirand, J. (1989). Arheoloogiga Eestimaa teedel. Second (Tallinn: Valgus).
98. Vedru, G. (1997). New settlement sites in the surroundings of Lake Kahala and revision excavations of stone-cist grave. In Stilus : Eesti Arheoloogiaseltsi teated (Eesti Arheoloogiaselts), pp. 62-67.
99. Friedenthal, A. (1932). Ein Beitrag zur vorgeschichtlichen Anthropologie Estlands. Z. Ethnol. 63, 1-42.
100. Kalman, J. (1998). Skeletal report. In Aruanne kivikirstkalmete kaevamistest Muuksi Hundikangrutes 1996-1997, G. Vedru, ed. (Tartu: Manuscript in the archive of the archaeology department of the University of Tartu).
101. Spreckelsen, A. (1926). Ausgrabungen in Neuenhof, Kirchsp. Kusal, Dorf Muuksi, Lõokese-Gesinde. Beitr. Zur Kunde Est. XI, 38-42.
102. Friedenthal, A. (1927). Bericht über die im Auftrage der Estländischen Literärischen Gesellschaft im Sommer 1927 vorgenommenen archäologischen Untersuchungen (Manuscript in the archive of the archaeology department of the University of Tartu).
103. Friedenthal, A. (1928). Bericht über die im Auftrage der Estländischen Literärischen Gesellschaft von Dr. A. Friedenthal im Juli 1928 vorgenommenen archäologischen Ausgrabungen (Manuscript in the archive of the archaeology department of the University of Tartu).
104. Schmiedehelm, М.Н. (1955). Археологические памятники периода разложения родового строя на северо-востоке Эстонии. (Archaeological Sites of the Period of the Disintegration of the Tribal System in North-eastern Estonia) (Estonian State Publishing House).
105. von Schroeder, L. (1895). Die Steinhügel-Gräber von Randen. Sitzungsberichte Gelehrt. Estnischen Ges. 1894, 75-81.
106. Laul, S. (1978). Die Steinkistengräber von Vehendi. Proc. Acad. Sci. Est. SSR Soc. Sci. 27, 76-77.
107. Lang, V. (2007). The Bronze and Early Iron Ages in Estonia. Estonian Archaeology, 3 (Tartu: University of Tartu Press).
108. Jaanits, L., Laul, S., Lõugas, V., and Tõnisson, E. (1982). Eesti Esiajalugu (Tallinn: ENSV Teaduste Akadeemia Ajaloo Instituut; Eesti Raamat).
109. Moora, T. (1974). Раскопки каменного могильника у с. Выхма в Северной Эстонии (Excavations of a stone burial site in Võhma in Northern Estonia). Proc. Acad. Sci. Est. SSR Soc. Sci. 23, 84-87.
110. Lang, V. (2000). Keskusest ääremaaks. Viljelusmajandusliku Asustuse Kujunemine ja Areng Vihasoo-Palmse Piirkonnas Virumaal. Muinasaja teadus, 7 (Tallinn).
111. Lang, V. (2003). From centre to periphery. Establishment and history of the agricultural settlement in the Vihasoo-Palmsee area (Virumaa, north Estonia). Acta Archaeol. 74, 123-188.
112. Kalman, J. (2000). Tandemägi stone grave - osteological report. In Keskusest Ääremaaks. Viljelusmajandusliku Asustuse Kujunemine ja Areng Vihasoo-Palmse Piirkonnas Virumaal, V. Lang, ed. (Muinasaja teadus, 7. (Tallinn)), pp. 423-436.
113. Lõugas, V. (1977). Ausgrabungen eines Steingräberfeldes von Kurevere. Proc. Acad. Sci. Est. SSR Soc. Sci. 26, 48-52.
114. Vaab, H. (2003). Kurevere kivikalme Saaremaal. Peaseminaritöö. (Manuscript in the archive of the archaeology department of the University of Tartu (University of Tartu), Bachelor's degree thesis.
115. Lang, V. (1995). A pre-Roman tarand-grave and Late Medieval fossil fields of Ilmandu, NW Estonia. Proc. Est. Acad. Sci. Humanit. Soc. Sci. 44, 429-436.
116. Kalman, J. (2000). Stone grave II of Tõugu - skeletal report. In Keskusest Ääremaaks. Viljelusmajandusliku Asustuse Kujunemine ja Areng Vihasoo-Palmse Piirkonnas Virumaal, V. Lang, ed. (Muinasaja teadus, 7. (Tallinn)), pp. 387-407.
117. Mandel, M. (1978). Über die Ausgrabungen der Tarandgräber von Poanse. Proc. Acad. Sci. Est. SSR Soc. Sci. 27, 78-81.
118. Mandel, M. (2000). Poanse tarandkalmed. In Töid Arheoloogia Alalt II, H. Pauts, and T. Tamla, eds. (Eesti Ajaloomuuseum), pp. 89-111.
119. Kalman, J. (2000). Skeletal analysis of the graves of Kaseküla, Poanse I and Poanse II. In Töid Ajaloo Alalt II, H. Pauts, and T. Tamla, eds. (Eesti Ajaloomuuseum), pp. 17-40.
120. Laneman, M., Lang, V., and Saage, R. (2016). Burial site hidden in a clearance cairn at Alu. Raplamaa. Archaeol. Fieldwork Est. 2015, 35-46.
121. Yushkova, M.A. (2016). New group of sites of the 1st to 7th centuries $A D$ in the south-west of Leningrad Oblast. In New Sites, New Methods. The 14th Finnish-Russian Archaeological Symposium. ISKOS 21 (Helsinki), pp. 143-159.
122. Juškova, M., and Kulešov, V. (2011). Kyorstovo 1: a new burial ground of the period of Roman influences in North-Western Russia. Archaeol. Litu. 12, 99-121.
123. Yushkova, M.A. (2015). Раскопки могильника Малли. (Excavations of the burial ground of Malli). In Археологические открытия 2010-2013 гг. (Archaeological discoveries of 2010-2013), P.G. Gaidukov, E.G. Devlet, D.S. Korobov, S.V. Kuzminykh, N.V. Lopatin, A.A. Maslennikov, S.Z. Chernov, and A.V. Engovatova, eds. (Moskow), pp. 100-102.
124. Shirobokov, I.G., and Yushkova, M.A. (2014). Антропологические материалы из коллективных захоронений по обряду кремации и ингумации каменного могильника с оградками Малли (по результатам раскопок 2010 г.). (Anthropological materials from collective burials according to the rite of cremation and inhumation of a stone fence burial ground in Malli (according to the results of excavations in 2010)). Bull. Archeol. Anthropol. Ethnogr. 2, 71-79.
125. Shirobokov, I.G., and Yushkova, M.A. (2015). Результаты планиграфического и макроскопического анализа антропологических материалов из могильника с каменными оградками Малли. (Results of a planigraphic and macroscopic analysis of anthropological materials from a stone fence burial ground in Malli). Bull. Anthropol. New Ser. 3, 93-109.
126. Yushkova, M.A., Grigorieva, O.V., and Grigorieva, N.V. (2015). Раскопки могильника Малли в 2013 г. (Excavations of the burial ground of Malli in 2013). In Археология и история Шскова и Шсковской земли, 60. (Archeology and History of Pskov and Pskov region, 60), P.G. Gaidukov, T.Y. Zakurina, L.Y. Kostyuchuk, E.V. Koroleva, I.K. Labutina, N.V. Lopatin, E.V. Salmina, V.V. Sedov, B.N. Kharlashov, and E.A. Yakovleva, eds. (Moskow), pp. 245-258.
127. Mikhaylova, E.R. (2016). The population of the south-eastern coast of the Gulf of Finland and its contacts with the regions of the Baltic Sea in the 1st millennium AD. Archaeol. Balt. 23, 181-198.
128. Mikhaylova, E.R. (2015). Древности Западной Ингрии I тыс. н.э.: Новые материалы. (Antiquities of Western Ingria I millennium BC: New materials) (Археологические вести. Вып. 21. (Archaeological News. Issue 21)), pp. 176-186.
129. Valk, H. (2001). Rural Cemeteries of Southern Estonia 1225-1800 AD. CCC Papers, 3. Second (Visby, Tartu: Gotland University College, Centre for Baltic Studies, University of Tartu Archaeology Centre).
130. Muižnieks, V. (2015). Bēru tradicijas Latvijā pēc arheologiski pētito 14.18. gadsimta apbedišanas vietu materiāla. Latvijas Nacionāla Vēstures Muzeja Raksti, 21. (Riga).
131. Palli, H. (1996). Eesti rahvastiku ajalugu aastani 1712. Third (Tallinn: Eesti Teaduste Akadeemia Kirjastus).
132. Pajusalu, K., Hennoste, T., Niit, E., Päll, P., and Viikberg, J. (2018). Eesti murded ja kohanimed. Third (Tallinn: Eesti Keele Sihtasutus).
133. Viires, A. (2008). Tagasivaade. In Eesti rahvakultuur, A. Viires, and E. Vunder, eds. (Tallinn: Eesti Entsüklopeediakirjastus), pp. 449-455.
134. Remmel, M.A., and Valk, H. (2014). Muistised, pärimuspaigad ja kohapärimus: ajalised ning ruumilised aspektid. In Muistis, koht ja pärimus 2. Pärimus ja paigad. Muinasaja teadus, 26: 2 (Tartu: Tartu Ülikooli ajaloo ja arheoloogia instituut), pp. 305-398.
135. Nelis, M., Esko, T., Mägi, R., Zimprich, F., Zimprich, A., Toncheva, D., Karachanak, S., Piskácková, T., Balascák, I., Peltonen, L., et al. (2009). Genetic structure of Europeans: a view from the North-East. PLoS ONE 4, e5472.
136. Kustin, A. (1958). Kalmistu XIII-XIV sajandist Karjas, Saaremaal. Proc. Acad. Sci. Est. SSR Soc. Sci. 7, 47-57.
137. Tamla, T. (1998). Zum Grabraub in vor- und frühgeschichtlichen Gräbern Estlands. In Studien zur Archäologie des Ostseeraumes. Von der Eisenzeit zum Mittelalter. Festschrift für Michael Müller-Wille, A. Wesse, ed. (Mainz, Stuttgart: Akademie der Wissenschaften und der Litteratur, Franz Steiner Verlag), pp. 291-297.
138. Tamla, T. (1989). Aruanne Pada maa-aluse kalmistu kaevamistest Rakvere rajoonis (Viru-Nigula kihelkond, Pada mõis; tänapäeval Viru-Nigula vald, Pada küla) 1989. aastal. (Manuscript in the archive of the archaeology department of the University of Tartu) (Tartu: University of Tartu).
139. Lõhmus, M., Jonuks, T., and Malve, M. (2011). Archaeological salvage excavations at Kukruse: a Modern Age road, cremation field and 12th13th century inhumation cemetery. Preliminary results. Archaeol. Fieldwork Est. 2010, 103-114.
140. Jonuks, T., Oras, E., Best, J., Demarchi, B., Mänd, R., Presslee, S., and Vahur, S. (2018). Multi-method analysis of avian eggs as grave goods: revealing symbolism in conversion period burials at Kukruse, NE Estonia. Environ. Archaeol. 23, 109-122.
141. Oras, E., Tõrv, M., Jonuks, T., Malve, M., Radini, A., Isaksson, S., Gledhill, A., Kekišev, O., Vahur, S., and Leito, I. (2018). Social food here and hereafter: multiproxy analysis of gender-specific food consumption in conversion period inhumation cemetery at Kukruse, NE-Estonia. J. Archaeol. Sci. 97, 90-101.
142. Kurisoo, T. (2014). Pada kalmistu rinnakeed. In Ajast ja ruumist. Uurimusi Mare Auna auks. Muinasaja teadus, 25, Ü. Tamla, and V. Lang, eds. (Ajaloo Instituut), pp. 79-92.
143. Schmiedehelm, M.H. (1928). Aruanne kaevamisest külakalmel Otepääl 1.VI-5.VI 1928. (Manuscript in the archive of Archaeological Research Collection of the University of Tallinn) (Tallinn University).
144. Schmiedehelm, M.H. (1929). Aruanne kaevamisest Otepääl 17.VI-21.VI 1929. a. (Manuscript in the archive of Archaeological Research Collection of the University of Tallinn) (Tallinn University).
145. Saadre, O. (1938). Aruanne kaevamistest Otepää külakalmel 13.-15. oktoobrini 1938. (Manuscript in the archive of Archaeological Research Collection of the University of Tallinn) (Tallinn University).
146. Valk, H. (1997). Archaeological investigations in Otepää and its Surroundings in 1996. In Stilus. Eesti Arheoloogiaseltsi väljaanne, Ü. Tamla, ed. (Eesti Arheoloogiaselts), pp. 124-129.
147. Raudkivi, P. (2010). Maa meie ema, ilm meie isa. Märkmeid looduse rollist Liivimaa 14. sajandi ajaloos. Acta Hist. Tallinn. 15, 3-23.
148. Napiersky, C.E. (1846). Beitrag zur Geschichte des ehemaligen Bisthums Dorpat (Müllersche Buchdruckerei).
149. Valk, H. (1983). Aruanne Vana-Kuuste 15.-17. sajandi külakalme arheoloogilistest kaevamistest 1982. a. (Manuscript in the archive of the archaeology department of the University of Tartu) (University of Tartu).
150. Valk, H. (1985). Der Dorffriedhof von Mäletjärve. Proc. Acad. Sci. Est. SSR Soc. Sci. 34, 376-379.
151. Valk, H. (1986). Der Dorffriedhof von Vaabina. Proc. Acad. Sci. Est. SSR Soc. Sci. 35, 389-393.
152. Damgaard, P.B., Margaryan, A., Schroeder, H., Orlando, L., Willerslev, E., and Allentoft, M.E. (2015). Improving access to endogenous DNA in ancient bones and teeth. Sci. Rep. 5, 11184.
153. Meyer, M., and Kircher, M. (2010). Illumina sequencing library preparation for highly multiplexed target capture and sequencing. Cold Spring Harb. Protoc. 2010, pdb. prot5448.
154. Orlando, L., Ginolhac, A., Zhang, G., Froese, D., Albrechtsen, A., Stiller, M., Schubert, M., Cappellini, E., Petersen, B., Moltke, I., et al. (2013). Recalibrating Equus evolution using the genome sequence of an early Middle Pleistocene horse. Nature 499, 74-78.
155. Malaspinas, A.S., Lao, O., Schroeder, H., Rasmussen, M., Raghavan, M., Moltke, I., Campos, P.F., Sagredo, F.S., Rasmussen, S., Gonçalves, V.F., et al. (2014). Two ancient human genomes reveal Polynesian ancestry among the indigenous Botocudos of Brazil. Curr. Biol. 24, R1035-R1037.
156. Rasmussen, M., Guo, X., Wang, Y., Lohmueller, K.E., Rasmussen, S., Albrechtsen, A., Skotte, L., Lindgreen, S., Metspalu, M., Jombart, T., et al. (2011). An Aboriginal Australian genome reveals separate human dispersals into Asia. Science 334, 94-98.
157. Andrews, R.M., Kubacka, I., Chinnery, P.F., Lightowlers, R.N., Turnbull, D.M., and Howell, N. (1999). Reanalysis and revision of the Cambridge reference sequence for human mitochondrial DNA. Nat. Genet. 23, 147.
158. Poznik, G.D., Xue, Y., Mendez, F.L., Willems, T.F., Massaia, A., Wilson Sayres, M.A., Ayub, Q., McCarthy, S.A., Narechania, A., Kashin, S., et al.; 1000 Genomes Project Consortium (2016). Punctuated bursts in human male demography inferred from 1,244 worldwide Y-chromosome sequences. Nat. Genet. 48, 593-599.
159. ISOGG. (2019). Y-DNA haplogroup tree. https://isogg.org/tree/.
160. YFull. (2019). YFull web site. https://www.yfull.com/.
161. Sudmant, P.H., Mallick, S., Nelson, B.J., Hormozdiari, F., Krumm, N., Huddleston, J., Coe, B.P., Baker, C., Nordenfelt, S., Bamshad, M., et al. (2015). Global diversity, population stratification, and selection of human copy-number variation. Science 349, aab3761.
162. R Core Team (2017). R: a language and environment for statistical computing (Vienna, Austria: R Foundation for Statistical Computing). https://www.R-project.org/.
163. Hellenthal, G., Busby, G.B.J., Band, G., Wilson, J.F., Capelli, C., Falush, D., and Myers, S. (2014). A genetic atlas of human admixture history. Science 343, 747-751.
164. Raveane, A., Aneli, S., Montinaro, F., Athanasiadis, G., Barlera, S., Birolo, G., Boncoraglio, G., Blasio, A.M.D., Gaetano, C.D., Pagani, L., et al. (2018). Population structure of modern-day Italians reveals patterns of ancient and archaic ancestries in Southern Europe. bioRxiv. https://doi. org/10.1101/494898.
165. Jeffreys, H. (1946). An invariant form for the prior probability in estimation problems. Proc. R. Soc. Lond. A Math. Phys. Sci. 186, 453-461.
166. Voight, B.F., Adams, A.M., Frisse, L.A., Qian, Y., Hudson, R.R., and Di Rienzo, A. (2005). Interrogating multiple aspects of variation in a full resequencing data set to infer human population size changes. Proc. Natl. Acad. Sci. USA 102, 18508-18513.
167. Browning, S.R., and Browning, B.L. (2015). Accurate non-parametric estimation of recent effective population size from segments of identity by descent. Am. J. Hum. Genet. 97, 404-418.
168. Bevan, A., Crema, E., and Silva, F. (2018). rcarbon v1.2.0: calibration and analysis of radiocarbon dates. https://CRAN.R-project. org/package=rcarbon.
169. Auton, A., Brooks, L.D., Durbin, R.M., Garrison, E.P., Kang, H.M., Korbel, J.O., Marchini, J.L., McCarthy, S., McVean, G.A., and Abecasis, G.R.; The 1000 Genomes Project Consortium (2015). A global reference for human genetic variation. Nature 526, 68-74.
170. Li, H. (2011). A statistical framework for SNP calling, mutation discovery, association mapping and population genetical parameter estimation from sequencing data. Bioinformatics 27, 2987-2993.

## STAR $\star$ METHODS

## KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
| :---: | :---: | :---: |
| Biological Samples |  |  |
| Human archaeological remains | This paper | N/A |
| Critical Commercial Assays |  |  |
| MinElute PCR Purification Kit | QIAGEN | Cat No./ID: 28006 |
| NEBNext DNA Library Prep Master Mix Set for 454 | New England Biolabs | E6070 |
| Deposited Data |  |  |
| Human reference genome NCBI build 37, GRCh37 | Genome Reference Consortium | http://www.ncbi.nlm.nih.gov/projects/genome/ assembly/grc/human/ |
| Modern comparison dataset | [5] | https://www.ncbi.nlm.nih.gov/geo/query/acc. cgi?acc=GSE108646; GEO: GSE108646 |
| Modern comparison dataset | [41] | http://hagsc.org/hgdp/files.html |
| Modern comparison dataset | [42] | https://www.ncbi.nlm.nih.gov/geo/query/acc. cgi?acc=GSE21478; GEO: GSE21478 |
| Modern comparison dataset | [43] | http://evolbio.ut.ee/caucasus/ |
| Modern comparison dataset | [44] | http://evolbio.ut.ee/turkic/ |
| Modern comparison dataset | [45] | https://www.ncbi.nlm.nih.gov/geo/query/acc. <br> cgi?acc=GSE50727; GEO: GSE50727 |
| Modern comparison dataset | [46] | https://www.ncbi.nlm.nih.gov/sra/?term= SRA010102; SRA: SRA010102 |
| Modern comparison dataset | [47] | http://evolbio.ut.ee/slavic/ |
| Modern comparison dataset | [48] | http://evolbio.ut.ee/khazar/ |
| Modern comparison dataset | [49] | https://www.ncbi.nlm.nih.gov/geo/query/ acc.cgi?acc=GSE46828; GEO: GSE46828 |
| Modern and ancient comparison dataset (including restricted access samples) | [14] | reich@genetics.med.harvard.edu |
| Ancient comparison dataset | [10] | erj35@cam.ac.uk |
| Ancient comparison dataset | [50] | https://www.ebi.ac.uk/ena/data/view/PRJEB18686; ENA: PRJEB18686 |
| Ancient comparison dataset | [13] | http://www.ebi.ac.uk/ena/data/view/PRJEB21037; <br> ENA: PRJEB21037 |
| Ancient comparison dataset | [11] | mittnik@shh.mpg.de |
| Ancient comparison dataset | [12] | https://reich.hms.harvard.edu/datasets |
| Ancient comparison dataset | [51] | https://www.ebi.ac.uk/ena/data/search?query= PRJEB20658; ENA: PRJEB20658 |
| Ancient comparison dataset | [52] | https://www.ebi.ac.uk/ena/data/view/PRJEB26349; https://www.ebi.ac.uk/ena/data/view/PRJEB25389; ENA: PRJEB26349; ENA: PRJEB25389 |
| Ancient comparison dataset | [53] | https://reich.hms.harvard.edu/datasets |
| Eastern Baltic aDNA data | This paper | http://evolbio.ut.ee/; http://www.ebi.ac.uk/ena/data/ view/PRJEB31893; ENA: PRJEB31893 |
| Oligonucleotides |  |  |
| NEBNext Multiplex Oligos for Illumina | New England Biolabs | E7335 |
| Software and Algorithms |  |  |
| cutadapt | [54] | https://cutadapt.readthedocs.io/en/stable/\# |
| Burrows-Wheeler Aligner (BWA) | [55] | http://bio-bwa.sourceforge.net/ |
| samtools | [56] | http://samtools.sourceforge.net/ |
| picard | N/A | http://broadinstitute.github.io/picard/index.html |

Continued on next page)

| Continued |  |  |
| :---: | :---: | :---: |
| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
| GATK | [57] | https://software.broadinstitute.org/gatk/ |
| mapDamage | [58] | https://ginolhac.github.io/mapDamage/ |
| contamMix | [59] | N/A |
| ANGSD | [60] | http://www.popgen.dk/angsd/index.php/ANGSD |
| sex identification algorithm | [61] | http://www.sciencedirect.com/science/article/pii/ S0305440313002495 |
| mtDNA-Server | [62] | https://mtdna-server.uibk.ac.at/index.html |
| HaploGrep2 | [63, 64] | http://haplogrep.uibk.ac.at/ |
| BEDTools | [65] | http://bedtools.readthedocs.io/en/latest/ |
| PLINK | [66] | http://pngu.mgh.harvard.edu/purcell/plink/ |
| EIGENSOFT | [67] | https://github.com/DReichLab/EIG |
| ADMIXTOOLS | [68] | https://github.com/DReichLab/AdmixTools |
| ADMIXTURE | [69] | https://www.genetics.ucla.edu/software/admixture/ |
| ChromoPainter | [70] | https://people.maths.bris.ac.uk/~madj/ finestructure-old/chromopainter_info.html |
| ChromoPainter/NNLS pipeline | [19, 71, 72] | N/A |
| Continuity simulation | [19] | N/A |
| READ | [73] | https://bitbucket.org/tguenther/read |
| VCFtools | [74] | http://vcftools.sourceforge.net/ |
| Beagle | [75] | https://faculty.washington.edu/browning/ beagle/b4_1.html |
| HlrisPlex-S webtool | [76-78] | https://hirisplex.erasmusmc.nl/ |

## CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Lehti
Saag (lehti.saag@ut.ee).

## EXPERIMENTAL MODEL AND SUBJECT DETAILS

The teeth used for DNA extraction were obtained with relevant institutional permissions from the University of Tartu, Institute of History and Archaeology; Tallinn University Archaeological Research Collection; the Museum of Anthropology and Ethnography (Kunstkamera) in St. Petersburg.

DNA was extracted from the teeth of 56 individuals - 23 from Late Bronze Age Estonia (EstBA; 1200-400 BC), 14 from Pre-Roman Iron Age Estonia (EstIA; between 800 and 500 BC-50 AD), 12 from Pre-Roman to Roman Iron Age Ingria, Russia (IngIA; 500 BC-450 AD) and 7 from Middle Age Estonia (EstMA; 1200-1600 AD) (Figure 1; Tables 1 and S1). More detailed information about the archeological periods and the specific sites and burials of this study is given below.

## Information about the archaeological time periods, sites and individuals of this study

In the archaeological record of Estonia, inhumation burials, which make the extraction of aDNA possible with current methods, date mainly from three major periods: 1) the Stone Age (9000-1800 BC), 2) the Bronze Age and Pre-Roman Iron Age (1800 BC-50 AD), 3) 2nd millennium AD. Thereby, inhumations from the Stone Age are presently known from the 7 th millennium to the 3rd millennium BC, and those from the Bronze and Pre-Roman Iron Age from ca. 1200 BC until the beginning of CE , with a few exceptions from later Iron Age. From the third major period, individuals from ca. 1200-1600 AD, conventionally regarded as 'medieval', have been involved in this study.

If the information in question was lacking beforehand, the preliminary estimation of the age at death of the individuals of this study was made at sample collection and the Bronze and Iron Age individuals who were included in autosomal analyses (having at least 10,000 overlapping SNPs with the EBC-chipDB) were radiocarbon dated.

## Late Bronze Age stone-cist graves

Late Bronze Age ( $1100-500 \mathrm{BC}$ ) in Estonia is a period where first stone graves, extensive permanent field systems and cup-marked stones appear in the near-coastal alvar areas. Settlement sites of the period are small and poor in both finds and construction
remains. The main settlement units were probably single farms, inhabited by small family or kin groups who subsisted on agriculture. Around 900-800 BC a few so-called fortified settlements were established on the island of Saaremaa and on the northern coast. These probably functioned as centers of (bronze) trade and were inhabited by larger groups. Late Bronze Age inland Estonia, on the other hand, is considerably poorer in archaeological sites, apart from settlement sites in open landscape and, in smaller numbers, on hilltops. Some hilltop settlements may have been similar to the coastal fortified settlements in function and nature.

Stone-cist graves are above-ground burial/mortuary structures, built of limestone or granite stones, or a combination thereof. They are round in shape, with a diameter from a few up to a few dozen meters, and their height rarely exceeds a meter. The graves feature one or more stone circles or drystone walls, which surround one or more human-length stone cists. A cist usually encloses several inhumations, both adults and children, with no clear patterning in age or sex categories. Some cists contain also or only cremated bones. Burials (or secondary bone deposits), both burnt and unburnt, are also common outside the cists. Grave goods are usually few and can rarely be associated with a particular skeleton. The most characteristic finds are bone pins; pottery is also commonplace (except for the earliest stone-cist graves); metal, amber and stone items are infrequent. While the grave type and one group of grave goods (imported bronzes) refer to dense contacts with Scandinavia, the other items (e.g., bone pins, temple ornaments, some of the pottery) witness the contacts with people in the East-European Forest Belt.

Stone-cist graves are distributed along the near-coastal zone of northern and western Estonia, including the largest islands. The graves usually come in groups. One such group is believed to have served a single family or kin group for several centuries. It is possible that only selected members of a household or kin group were accorded a burial in a stone-cist grave. Radiocarbon dating of bones have shown that the stone-cist graves appeared in what today is Estonia between 1200 and 1100 BC and were built until ca 400 BC at the latest. Many of them, however, contain occasional burials from the Iron and even the Middle Ages.

Besides stone-cist graves, cairn graves may have been occasionally built. The main difference of this grave type from stone-cist graves is the absence of cist(s) and, in some cases, ring wall(s). At the end of the period, early tarand graves appeared (see below). Most probably other types of burial sites existed, for instance burials in pits, but information on such sites is very limited.

In view of the hypothesis that a group of stone-cist graves represents a single kin group, aDNA samples were collected from as wide variety of such groups as possible. Males were targeted, since the aim was to focus on Y chromosome diversity. In this article, twenty-three burials from Late Bronze Age in Estonia are analyzed.

## Kangru at Väo

Location: Väo, Harjumaa, Estonia
Excavations: 1959 [79], 1976-1977 [80], 1980 [81]
Cemetery: A minimum of nine stone-cist graves (numeration of graves differs in publications; we follow the numeration as in Lõugas 1981 [81]). Beside inhumations within cists, occasional cremations and inhumations outside cists were also present. No proper osteological analysis has been performed (but see Lang and Ligi 1991 [82]). Bronze Age artifact finds were rare and included a bronze razor, dated to the IV or V period of the Nordic Bronze Age, and a few bone pins. Some Iron Age objects were also uncovered. The scarce evidence for dating suggests that the grave group was established around 900 BC at the latest.

DNA-analyzed individuals:
X16: Male(?) from the cist of grave 1 (Al 4303), age unknown (the bones were too fragmented and intermingled for a preliminary age estimation). Sampled tooth right lower second molar ( $\left(\mathrm{M}_{2}\right)$, date $2834 \pm 26$ BP (SUERC-80019 (GU47830); 1080-910 cal BC). The cist contained remains of at least one other individual.

X17: Male from grave 8 (skeleton 1; Al 4939), age 25-35 years. In the absence of excavation records, location of the skeleton within the grave is indeterminate; position in one of the grave's two cists is likely. Sampled tooth $r M_{2}$, date $2732 \pm 28$ (SUERC-80020 (GU47831); 930-810 cal BC).

X18: Male from grave 8 (skeleton 3; AI 4939), age 17-22 years. Location of the skeleton within the grave cannot be established, but there are grounds to suggest that it was a burial outside the cists. The excavators dated the burial outside the cists to the Middle Ages [80], but an earlier date cannot be excluded.Sampled tooth left upper canine (I C ${ }^{1}$ ).

## Jaani at Väo

Location: Väo, Harjumaa, Estonia
Excavations: 1982 [83]
Cemetery: Two stone-cist graves and a ship grave, attached to one another. A minimum of thirty-eight individuals, predominantly inhumations, had been interred in to the stone-cist graves; original cremation deposit of the ship grave has been lost since excavations [84]. The bones were heavily intermingled. Radiocarbon dates of the bones [84] show that the first of the stone-cist graves (B) was probably erected between 800 and 500 BC ; grave A was attached in the $5^{\text {th }}$ century BC at the latest. The graves were used for burial also in the Pre-Roman Iron Age and even later, until at least the $7^{\text {th }}$ century AD. Artifact finds comprise pottery, a bone pin, and several poorly datable metal objects from the Iron Age and even later periods. The distance from the Pärna graves (see below) was ca 190 m , which means that the separation of the grave groups may be artificial.

DNA-analyzed individual:
V16: Male from the cist of grave A (skeleton 1; AI 5220), age 30-40 years [84], date $2399 \pm 27$ BP (UBA-24124; 730-390 cal BC) [84]. Sampled tooth $r \mathrm{M}^{1}$. The cist contained an iron knife and, most probably, an indeterminate number of other skeletons.
Pärna at Väo
Location: Väo, Harjumaa, Estonia
Excavations: 1895 [85], 1972-1973 [86]

Cemetery: The original number of the stone graves is unknown. At least four graves have been excavated, but the information on the results is poor. No osteological analysis has been carried out. Artifact finds include pottery and a bone pin; a few Roman-period metal objects were also uncovered. The artifacts and the radiocarbon date suggest that the graves were present before 800 BC . There is a possibility that the Pärna and Jaani graves (see above) were built and used by the same community.

DNA-analyzed individuals:
X19: Individual from the cist of grave 1 (AI 4620: L44), sex and age unknown (the bones were too poorly preserved for a preliminary estimation during sample collection). Sampled tooth $r \mathrm{M}_{1}$.

X20: Individual from the cist of grave 1 (AI 4620: L46), sex and age unknown (the bones were too poorly preserved for a preliminary estimation during sample collection). Sampled tooth $r \mathrm{M}_{2}$, date $2677 \pm 26$ BP (SUERC-80018 (GU47829); 900-800 cal BC).

The cist also contained a clay vessel and remains of at least one sub-adult [82].
Iru
Location: Iru, Harjumaa, Estonia
Excavations: 1936 [87], 1974 [88]
Cemetery: Nine dispersed stone-cist graves, all excavated. The original number of graves was greater, and the graves possibly formed several (sub-)groups. Excavations yielded more than twenty inhumations [82], predominantly but not exclusively from the cists. A few deposits of cremated bone were also recorded. A proper osteological analysis is still to be done. The most characteristic grave inclusions were bone pins and pottery. The finds and a few radiocarbon dates (unpublished) suggest that the cemetery was established around 900 BC at the latest. Some burials or bone deposits outside cists may be later than Bronze Age in date.

DNA-analyzed individuals:
X01: Male from the cist of grave 6 (AI 4808: L10), age 17-25 years. Sampled tooth I M ${ }^{2}$. The cist also contained remains of at least two sub-adults, a bone pin and pottery.

X02: Male from the cist of grave 14 (AI 4810: L5), age 17-25 years. Sampled tooth I M ${ }^{3}$, date $2834 \pm 28$ (SUERC-80017 (GU47828); 1090-910 cal BC). The cist also enclosed skeletons of at least two children and a bone pin.

X03: Male from the cist of grave 18 (Al 4811: L11), age 35-45 years, date $2595 \pm 30$ BP (HeLa-2413; 830-590 cal BC [Laneman, unpublished]). Sampled tooth $1 \mathrm{M}_{3}$. The cist contained a skeleton of another adult and two bone pins.

## Lastekangrud at Rebala

Location: Rebala, Harjumaa, Estonia
Excavations: 1982 [89], 2000 [90]
Cemetery: Six stone-cist graves, one of them almost completely destroyed before excavations. The five remaining graves contained at least 40 inhumations, both inside and outside cists; cremated human bones were also present in almost each grave [ 90,91 ]. A quarter of the inhumations were infants, interred in grave 2 in the $15^{\text {th }}$ century AD. The cist burials date from ca 800400 BC , and a few individuals outside cists from the following centuries (unpublished radiocarbon data). Bronze Age artifact finds include clay vessels and bone pins, mostly in cists. Other areas of the graves contained occasional poorly datable metal items from various periods of the Iron Age and even beyond.

DNA-analyzed individuals:
X13: Male from the cist of grave 2 (AI 5229), age 18-22 years [91], date $2485 \pm 30 \mathrm{BP}$ (HeLa-2127; 780-480 cal BC) [Laneman, unpublished]. Sampled tooth I $\mathrm{M}_{1}$.

X14: Male from the cist of grave 2 (AI 5229), age 35-45 years [91], date $2481 \pm 30 \mathrm{BP}$ (HeLa-2061; 780-430 cal BC) [Laneman, unpublished]. Sampled tooth I M ${ }^{2}$.

The cist also contained inhumed remains of an infant, cremated human bones, and a poorly preserved iron object.

## Jõelähtme

Location: Jõelähtme, Harjumaa, Estonia
Excavations: 1982-1984 [92]
Cemetery: A dense cluster of thirty-six stone-cist graves with the remains of roughly a hundred inhumations (osteological analysis is incomplete, see Varul 2016 [93]). The cemetery was originally even bigger, as part of it has been destroyed by road construction and was in use between ca 1200/1100 and 800 BC (unpublished radiocarbon data). Grave goods include small bronze items, mostly of Scandinavian origin (razors, tweezers, buttons), bone pins and a few amber beads.

DNA-analyzed individuals:
OLS11: Male from grave 34 (Al 5306), age 30-50 years [34], date $2815 \pm 33$ BP (HeLa-2361; 1060-850 cal BC) [34]. The bones were commingled with the remains of at least one other adult and were located both inside and outside the cist. The DNA-analyzed tooth was found outside the cist, but it is likely that the original location of the skeleton was in the cist. Fragments of two bone pins were found alongside. Isotope (Sr and O) analysis showed that the man had been born locally [34]. Sampled tooth left upper second premolar (I P²).

V9: Male from the cist of grave 7 (AI 5306), age 30+ years [34], date $2924 \pm 32$ BP (HeLa-2365; 1220-1010 cal BC) [94]. Bronze tweezers and a bronze razor were found together with the skeleton. Isotope ( Sr and O ) analysis showed that the man had been born locally [34]. Sampled tooth $r \mathrm{P}_{1}$.
Toomani at Muuksi (Hundikangrud)
Location: Muuksi, Harjumaa, Estonia
Excavations: 1924-1926 [95], 1936 [96], 1976-1983 [97], 1995-1996 [98]

Cemetery: About forty closely spaced stone-cist graves, five of which have been excavated in their entirety and twelve partially [94]. Inhumations occur both inside and outside of cists; the same applies to the few cremation deposits. The number of excavated inhumations is well over thirty. Artifact finds comprise a few items of flint, quartz and bone. Radiocarbon data from the completely excavated grave 5 shows that burial began around 1100 BC at the latest, and ceased around 900 BC at the latest [94]. No such data is available for other graves, but in view of their uniform characteristics it is likely that the whole group dates from ca 1200-800 BC. Distance from the Lõokese graves (see below) is ca 1 km .

DNA-analyzed individuals:
V14: Male from cist 1 of grave 5 (AM 365: T4), age 50-60 years [99], date $2966 \pm 29$ BP (SUERC-44064 (GU29245); 1280-1050 cal $B C$ ) [94]. Isotope (Sr and O) analysis showed that the man had been born locally [34]. Sampled tooth I $P_{1}$. The cist also housed remains of an adult female.

X05: Male from cist 2 of grave 5 (AI 6320: L135), age 20-25 years [100], date $2908 \pm 26$ BP (SUERC-44069 (GU29247); 1210-1010 cal BC) [94]. Sampled tooth I M ${ }^{3}$. A tooth of a dog was found nearby.

X06: Male from cist 4 of grave 5 (AI 6320: L176), age 25-35 years [100], date $2906 \pm 25$ BP (SUERC-44070 (GU29248); 1200-1010 cal BC) [94]. Sampled tooth I $M_{3}$. The cist also housed remains of a child.

X07: Male from cist 2 of grave 12 (AM 365: T15), age 30-40 years [99]. Sampled tooth I M ${ }^{3}$.

## Lõokese at Muuksi

Location: Muuksi, Harjumaa, Estonia
Excavations: 1921 [101]
Cemetery: Six stone graves, only one (partially) excavated. The grave had three parallel cists (A, B, C) built crosswise over the fourth (D). The upper cists housed a single skeleton each; cist D housed two inhumations. Remains of a child were uncovered outside the cists. Artifact finds comprise only three potsherds. Distance from the Toomani graves (see above) is ca 1 km .

DNA-analyzed individuals:
X08: Male from cist A (AM ?: L1), age 50 years [99]. Sampled tooth I M ${ }^{3}$, date $2733 \pm 26$ BP (SUERC-80021 (GU47832); 930-810 cal BC).

X09: Male from cist B (AM ?: L2), age 18-20 years [99]. Sampled tooth I M ${ }_{2}$, date $2606 \pm 28$ BP (SUERC-80025 (GU47833); 820770 cal BC ).

X10: Male from cist C (AM ?: L3), age 60 years [99]. Sampled tooth I M ${ }_{1}$, date $2926 \pm 28$ BP (SUERC-80026 (GU47834); 12201020 cal BC). A potsherd and a tooth of a dog were reported nearby.
Napa
Location: Napa, Ida-Virumaa, Estonia
Excavations: 1927-1928 [102-104]
Cemetery: Around fifteen or twenty stone graves, of which partially excavated were at least five stone-cist graves and a probable tarand grave. Numeration of graves differs in publications; in this paper we generally follow Friedenthal 1932 [99]. The excavated cists housed a minimum of fourteen inhumations, and some cists had an assemblage of cremated bones beneath the cist floor. Both inhumations and cremations were observed outside the cists. Osteological analysis is available for only the cist inhumations [99]. Grave goods included a few bone pins and items of flint, bronze, and iron. The finds and radiocarbon dates show that the cemetery was present in the $9^{\text {th }}$ century BC at the latest; it may also contain a few centuries older as well as a few centuries younger burials.

DNA-analyzed individuals:
X11: Male from the cist of grave 3 (5 in other referred sources) (AM 331: N10), age 50 years [99]. Sampled tooth r $\mathrm{M}_{2}$, date $2805 \pm$ 26 BP (SUERC-80010 (GU47824); 1030-890 cal BC). A single potsherd was found nearby.

X12: Male from the cist of grave 4 ( 6 in other referred sources) (AM 331: N11), age 40-50 years [99]. Sampled tooth $r M_{2}$, date $2652 \pm$ 26 BP (SUERC-80011 (GU47825); 900-790 cal BC). The cist also contained an infant. Burnt bones and a fragment of a bone pin were found under the cist floor.

## Vehendi

Location: Vehendi, Tartumaa, Estonia
Excavations: 1894 [105], 1975-1976 [106]
Cemetery: Eleven stone mounds distributed within a one kilometre long stretch along the coast of Lake Võrtsjärv. Two mounds, nos 11 and probably 1 , have been excavated, but information on the $19^{\text {th }}$-century digs is poor. The available evidence suggests that the graves are probably cairn and not stone-cist graves, i.e., their structure includes a stone circle but no cists. Grave 11 contained an inhumation in the center (not available for analysis) and a few other bone deposits, both burnt and unburnt, in other parts of the cairn. The burials were poorly preserved, and no osteological analysis has been applied to the bones. No artifact finds were uncovered. The radiocarbon date obtained for the current project shows that the grave(s) must have been present around 1000 BC at the latest.

DNA-analyzed individual:
X15: Male(?) from the eastern periphery of grave 11 (skeleton 3; AI 6950). The teeth indicate a relatively aged person. Sampled tooth $1 \mathrm{M}^{1}$, date $2899 \pm 28 \mathrm{BP}$ (SUERC-80016 (GU47827); 1210-1000 cal BC).

## Pre-Roman Iron Age early tarand and other cemeteries

In the Pre-Roman Iron Age ( $500 \mathrm{BC}-50 \mathrm{AD}$ ), new developments took place in the culture and settlement pattern in what today is Estonia. The fortified sites were abandoned around 500 BC and an open settlement pattern (most likely in the form of single
households) spread everywhere, both in coastal and interior regions. In the later Pre-Roman Iron Age, a new short-lived fortification wave can be observed all over the country. The building of new stone-cist graves was terminated around 400 BC at the latest. At some point of time within the period of ca 800-500 BC (due to difficulties in calibration of radiocarbon dates of that period it is not known when exactly), a new form of burying sites was introduced in coastal zone - the so-called early tarand cemeteries. Some of the earliest tarands were erected side by side with, or in close proximity to, stone-cist graves, the rest of them were built separately from other burial sites. In addition, burial sites of other forms are known, such as cairn graves, pit graves with either inhumations or cremations, and burial sites where cremated bones have been scattered over an open surface of the ground.

Early tarand cemeteries form a peculiar and diverse group of burial sites that were spread in Estonia, northern and western Latvia, south-western Finland, Ingria, and eastern central Sweden. Tarands are quadrangular stone enclosures for individual or collective burials built on the ground, with the straight flat sides of the walls facing outward. The number of tarands in a cemetery can vary from one to a few dozens, and if there is more than a single tarand they are joined together. Inhumation was the original and most common burial custom in the earliest cemeteries during the early Pre-Roman Iron Age; cremation was introduced later, at the end of this period, but inhumation did not disappear. The number of burials in one tarand can vary greatly: in earlier cemeteries with smaller tarands this number rarely exceeds two or three; in later cemeteries, one tarand can house up to a dozen or even more individuals.

Grave goods were quite rare in the earliest graves that can be dated to the period of ca 800-500 BC by the radiocarbon method. The only grave goods of that time were clay pots of Ilmandu type, a new style in Estonian Final Bronze Age pottery, which was formed under the influences from the Oka and Moscow rivers' region [20]. During the $5^{\text {th }}-3^{\text {rd }}$ centuries $B C$, many metal artifacts appeared among the grave goods, such as neck-rings and bracelets of bronze, massive bracelets of iron, temple ornaments with spoonshaped ends, a variety of decorative pins (of bronze and iron, and bimetallic), etc. A distinguished group of grave goods originates in the East-European Forest Belt [107] but artifacts imported from central and northern Europe were not unique either. During the last centuries BC and the first century AD, the finds in tarand graves became more numerous: ornaments (shepherd's crook pins, bracelets, finger-rings, etc.), small-sized tools (knives), and pottery (incl. cord- and comb-decorated vessels).

In this article, fourteen burials from Pre-Roman Iron Age in Estonia are analyzed.

## Loona

Location: Loona, Saaremaa, Estonia
Excavations: 1958-1959 [108]
Cemetery: Two stone graves, one of them excavated; four other stone graves at a distance of 300 m [97]. The excavated grave was a stone-cist grave which contained at least seventeen inhumations outside the empty cist in generally lower layers and numerous deposits of cremated bone in upper layers. No osteological analysis has been performed. Artifact finds include various bone and amber objects (probably ornaments), iron and bronze bracelets, temple ornaments, and pottery. The finds, the majority of which have close parallels in early tarand graves, suggest that the grave was built in the Bronze Age, and was used for burial also in the Pre-Roman Iron Age. More precise dates are difficult to establish.

## DNA-analyzed individual:

X04: Male in the south-western part of grave 1 (skull 10; Al 4210). In preliminary examination, teeth and bones yielded contradictory evidence on age at death estimate (17-25 and 40+ years, respectively). Sampled tooth $r M_{1}$, date $2331 \pm 26$ BP (SUERC-80015 (GU47826); 480-360 cal BC).

## Tandemägi IV at Võhma

Location: Võhma, Lääne-Virumaa, Estonia
Excavations: 1969-1972 [109-111]
Cemetery: Tandemägi is a long ridge with seven stone settings. In the north-western part of this ridge there were three stone-cist graves (I-III) of the Late Bronze Age. The tarand cemetery (IV), dated from the Pre-Roman Iron Age, had been built on the south-eastern end of the ridge, 76 m apart from the stone-cist graves. It consisted of three quadrangular enclosures with altogether at least fifty inhumations and five cremations [112]. The cemetery was rather rich in grave goods, which mostly belonged to the late Pre-Roman Iron Age: ceramics, shepherd's crook pins of iron, bracelets of bronze, knives and an axe of iron, etc. In contrast to generally very fragmentary and intermingled skeletons there was a well-preserved triple burial in tarand 2 . It consisted of a $30-35$ years old male, a 20-25 years old (fe)male, and a 6-7 years old child. The adults were richly furnished with grave goods: the older male had a neck-ring and a decorative pin of bronze, two bracelets of iron and one more of bronze; the other adult had a similar neck-ring and three bronze bracelets whereas the child had only a bronze temple ornament [110, 111]. All these grave goods have an early Pre-Roman Iron Age date.

DNA-analyzed individual:
VII4: Male from the triple burial (AI 5074: L64), age 30-35 years [112]. Sampled tooth $r$ M $^{3}$, date $2425 \pm 35$ BP (Poz-98210; 760-400 cal BC).

## Hiiemägi at Kunda

Location: Kunda, Lääne-Virumaa, Estonia
Excavations: 2004-2006 [34]
Cemetery: The cemetery is located on a ridge called Hiiemägi in the outskirts of the town of Kunda. The cemetery has been ca 50 m long but was largely destroyed by quarrying. Only a small part of the cemetery was excavated but the results are not properly published as yet [34]. There were eleven small cist-like tarands distinguished in the excavated area, each of them contained
one or more inhumation burials (altogether 32). Grave goods were very poor: a few potsherds, animal bones, a knife and three small decorative pins of iron from the early Pre-Roman Iron Age.

DNA-analyzed individuals:
OLS10: Male from tarand III (burial 9; TÜ 1325: L777), age 17-25 years [34]. He had a fragment of a sheep/goat bone and ceramics as grave goods. This burial has two radiocarbon dates: $2430 \pm 35$ BP (Poz-10801; 760-400 cal BC) and $2530 \pm 41$ BP (UBA-26114; $800-530 \mathrm{cal} \mathrm{BC}$ ) [34]. According to the isotopic analysis, the person was not born in the vicinity of Kunda; his place of birth is still unknown (but south-western Finland and Sweden are excluded) [34]. Sampled tooth $r P_{1}$.

V10: Male from tarand XI (burial 24; TÜ 1325: L1925), age 25-35 years [34], date $2484 \pm 40$ BP (UBA-26115; 790-430 cal BC) [34]. He had a few potsherds near the skull. Likewise, this person was not locally born [34]. Sampled tooth I $P_{1}$.

## Kurevere

Location: Kurevere, Saaremaa, Estonia
Excavations: 1974-1975 [113, 114]
Cemetery: It was one of the stone settings in a larger group and consisted of three structural parts: (1) a round-shaped grave surrounded with two concentric stone circles (but no cist in the center), (2) a much larger stone circle around the former, and (3) ca 20 tarand-like enclosures by the northern, southern and western sides of the large stone circle. The majority of burials were inhumations, but the bones were rather fragmentary and intermingled. Cremated bones occurred sporadically and can be connected with the latest stage in the use of this burial site. Osteological material has not been analyzed so far, however. Grave goods were quite numerous consisting mainly of pottery, various ornaments of bronze and iron (shepherd's crook pins, a pin with a spiral-shaped head, bracelets, various temple ornaments, decorative mounts, etc.), tools (knives, awls, an axe, and a grinding stone), and a few weapons (fragments of a spearhead and a battle knife). The earliest part of the cemetery (the two concentric circles) was already built in the Late Bronze Age, while the rest of the cemetery belongs to the Pre-Roman Iron Age.

DNA-analyzed individuals:
V11: Male(?) buried in the northern portion of the large stone circle (Al 4780: L17), age 25-35 years. Sampled tooth $\mathrm{r} \mathrm{M}_{2}$, date $2220 \pm$ 35 BP (Poz-98256; 390-200 cal BC).

V12: Male(?) buried in tarand VII (AI 4780: L118), age 25-35 years. Close to the bones there were also pieces of a clay pot with cord decoration found. Sampled tooth $r \mathrm{M}_{3}$, date $2125 \pm 35$ BP (Poz-98257; 360-40 cal BC).

## IImandu III

Location: Ilmandu, Harjumaa, Estonia
Excavations: 1994 [115]
Cemetery: The cemetery belongs to a larger group of burial sites (stone-cist graves and early tarand cemeteries), which are dispersed over the lands of IImandu and Rannamõisa villages close to northern Estonian limestone cliff. Cemetery III of IImandu was partially destroyed by building a house. Altogether six tarands and two cist-like constructions were distinguished in the preserved part of the cemetery. All burials were inhumations, except a few cremated bones that were of later date. Osteological material is properly not analyzed but during excavations at least seventeen adult individuals were distinguished. Grave goods were very poor consisting of pottery of Ilmandu type and a temple ornament.

DNA-analyzed individuals:
OLS09: Female from cist I (AI 6009: L180), age 19-25 years [34], date $2361 \pm 29 \mathrm{BP}$ (SUERC-44060 (GU29241); 540-380 cal BC) [94], most likely locally born [34]. Sampled tooth $r P_{1}$.

V7: Male from tarand IV (burial 1; AI 6009: L166), age 35-45 years [34], date $2484 \pm 41$ BP (UBA-26113; 790-430 cal BC) [34]. According to isotopic analyze, this person was most likely locally born [34]. Sampled tooth I M3.

V8: Male(?) from tarand IV (burial 9; AI 6009: L184), age 17-25 years, date of right femur $2413 \pm 29$ BP (SUERC-44062 (GU29243); $750-400$ cal BC) [94]. Furnished with a clay pot of Ilmandu type and a bronze temple ornament (fragment). Sampled tooth I M $\mathrm{M}_{1}$, date $2405 \pm 35$ BP (Poz-98215; 750-390 cal BC).

## Tôugu II

Location: Tõugu, Lääne-Virumaa, Estonia
Excavations: 1993-1995 [110, 111]
Cemetery: There is a group of at least eleven stone settings at Tõugu but only one of them is excavated. Cemetery II consisted of three separate parts: a stone-cist grave from the Bronze Age (IIA), topped with a large single tarand (IIB) of the Pre-Roman Iron Age, and a chain of five tarands (IIC) that was erected next to the latter structures also in the Pre-Roman Iron Age. According to Jonathan Kalman [116], there were altogether at least twenty-five inhumations excavated from the Tõugu II cemetery, sixteen of them from the series of five interconnected tarands IIC. Grave goods were rather poor, including pottery, iron knives, some bracelets of bronze, pieces of quartz and a few grinding stones.

DNA-analyzed individual:
V15: Male from tarand 1 of the cemetery IIC (AI 6003: L637), age 25-35 years [116]. Sampled tooth I M $\mathrm{M}_{2}$.
Poanse I
Location: Poanse, Läänemaa, Estonia
Excavations: 1975-1976 [117, 118]
Cemetery: There were two Pre-Roman tarand cemeteries close to each other. Cemetery I consisted of seven enclosures. Kalman [119] identified forty-four burials in this burial site, whereas most remains were commingled and fragmentary. In some cases, the
skeletons were preserved well enough to make the identification of individual burials possible. The majority of burials were without grave goods, but some were furnished quite remarkably with bracelets of iron and bronze, shepherd's crook pins, temple ornaments with spoon-shaped ends, and cord-decorated pottery; as an extraordinary find for tarands also a sickle should be mentioned. Cemetery II was smaller than cemetery I, it consisted of two tarands and housed altogether thirty-four burials. Judging from grave goods - a spearhead, bracelets, shepherd's crook pins, knives, and pottery - cemetery II was at least partly contemporary with cemetery I in the mid- and late Pre-Roman Iron Age.

DNA-analyzed individuals (cemetery I):
VII2: Male(?) buried in tarand 1 (AM A483: L18), age 17-25 years. Sampled tooth $r \mathrm{M}_{3}$, date $2275 \pm 35$ BP (Poz-98208; 410-200 cal BC). Tarand 1 was built as the first enclosure in this cemetery. Together with this male person there were also two juveniles (14-18 and 16-18 years old) and a 50+ years old male, and a few subadults buried.

VII3: Male from tarand 4 (AM A483: L30), age 30-40 years [119]. Sampled tooth r M 3 , date $2205 \pm 35$ BP (Poz-98209; 380-180 cal BC). Buried together with four adults and two children. Tarand 4 was built some time (perhaps a few generations) later than tarand 1. Alu
Location: Alu/Kalevi, Raplamaa, Estonia
Excavations: 2015 [120]
Cemetery: The site, a low moraine hump covered in field clearance stones, contained only two inhumations, a few meters apart from each other. The clearance cairn, which was of a later date, made it difficult to determine the original appearance and type of the burial site. One individual had been interred in a shallow earth-cut grave which, possibly, may have been surrounded and/or covered with stones, including sizeable boulders. The other burial structure possibly also included a shallow pit grave, and most certainly boulders and smaller rocks had been used in its construction. The stone structure had been disturbed and the bones were scattered. Both of the skeletons belonged to adults, perhaps mature adults, but a more precise age-at-death estimation was impossible due to poor preservation of bones; sex determination by osteological methods was not possible. No grave goods were found, though some of the pottery, scattered over the site, may have been contemporary with the burials.

DNA-analyzed individuals:
OLS07: Individual in the earth-cut grave (TÜ 2525: L264), adult [120], dates $2209 \pm 33$ BP (SUERC-63659 (GU38997); 380-190 cal BC), $2213 \pm 33$ BP (SUERC-63660 (GU38998); 380-190 cal BC) [120]. Sampled tooth I P ${ }_{2 \text { ? }}$.

OLS08: Individual in the stone structure (TÜ 2525: L291), adult [120], dates $2162 \pm 31 \mathrm{BP}$ (SUERC-63661 (GU38999); 360-110 cal BC), $2166 \pm 33$ BP (SUERC-63665 (GU39000); 360-110 cal BC), $2145 \pm 31$ BP (SUERC-63666 (GU39002); 360-50 cal BC) [120]. Sampled tooth $\mathrm{r} \mathrm{M}_{2}$.

## Pre-Roman and Roman Iron Age cemeteries in Ingria, Russia

Archaeological material from Pre-Roman (500 BC-50 AD) and Roman Iron Age (50-450 AD) in Ingria, south-western part of Leningrad district in Russia, are quite limited and studied only a little more than 30 years [121]. Most common type of archaeological sites is socalled tarand cemeteries. The tarand cemeteries have been excavated more widely at the burial sites of Kerstovo 1 and Malli, but similar structures are found also at the cemetery of Valgovitsy and Velikino. Isolated finds, possibly originating from disturbed burials, were found in the villages of Ratchino, Georgiyevsky, Voynosolovo and Ropsha. The walls of tarands were built of granite stones and limestone, while the inner space was filled with smaller stones and limestone gravel.

The overwhelming number of finds from Ingrian tarands is dated to the Early Roman Period, that is, to the time span from ca 75 to 200 AD. The grave goods included different types of fibulas, bracelets, rings, temple rings, weapons and iron tools for everyday life (spearheads and javelin heads, socketed axes, razors, awls, needles, scythes, knives). In Kerstovo 1 and Malli plaques imported from more eastern regions of the East-European Forest Belt (basins of the Upper Volga, Mologa, Middle Volga and the Kama region) were found. The tarand cemeteries in Ingria represent a local variant that finds its closest parallels at sites in north-eastern Estonia. The easternmost site in Estonia - the cemetery of Utria - is located some 40 km to the west of the sites on the Izhora Heights. The tarands in Ingria have a distinctive difference compared to those in Estonia by the presence of numerous weapons (spearheads, javelin heads, axes) and objects imported from the more eastern areas.

Among other findings there are three hoards of Roman coins that were discovered near the village of Koporye worth mentioning [121].

In this study, twelve burials from Pre-Roman and Roman Iron Age in Ingria are analyzed.

## Kerstovo I

Location: Kerstovo, district of Kingisepp, north-western Russia
Excavations: 2008-2009 [122], 2016 [121]
Cemetery: The burial ground is situated on an arable field and its upper level was partly disturbed. A funerary installation, elongated along the west-east line, consisting of a chain of four tarands was investigated. Numerous skeletal remains were discovered - altogether ca. 19 kg of bones, mostly calcined. The bones were found within the structures both as isolated pieces and in associations. The rite of an outside cremation prevailed and the skeletal remains are predominantly represented by small calcined fragments. Also fragments of unburned bones were found; these were lying in no anatomical order. At least 38 persons were interred at the site. The grave goods from the excavations - 155 items altogether - included parts of garments and bronze ornaments, among others different types of fibula. Other ornaments include bronze bracelets, rings, temple rings, large beads, with a lug, and an iron clasp. In addition, weapons and iron tools for everyday life were found (spearheads and javelin heads, half-moon-shaped razors, a scythe, a needle,
awls, and knives), as well as a gold-glass bead, a bronze needle, and fragments of ceramic vessels. The surface finds ( 150 metal objects) included bronze ornaments - eye brooches, profile fibulae, rings, and a plaque in the form of a rosette, as well as iron javelin heads and spearheads, socketed axes, and knives. The materials from the excavations and the surface finds suggest that also other tarand cemeteries can be found here.

DNA-analyzed individuals:
VII15: Adult from tarand 3 (horizon 3). Sampled tooth I C ${ }^{1}$, date $1980 \pm 30$ BP (Poz-103328; 45 cal BC-77 cal AD).
VIII7: Adult from tarand 3(horizon 2), No. 219. Sampled tooth I P . $^{\text {. }}$
VIII8: Adult from tarand 2 (horizon above bedrock), No. 2979. Sampled tooth r C ${ }^{1}$.
VIII9: Adult from tarand 2 (horizon 3). Sampled tooth r $\mathrm{P}^{2}$.

## Malli

Location: Malli, district of Kingisepp, north-western Russia
Excavations: 2010-2011, 2013 [121, 123-126]
Cemetery: The burial structure was consisting of two tarands and stone pavements. The westernmost tarand (NNE-SSW) was evidently built first. After destroying its eastern wall, a new tarand was constructed there in a slightly different orientation. The walls were joined by a lateral mound constructed of limestone gravel and granite pavement. The lateral mound was well preserved along the western wall of the western tarand. To the south and east of the tarands, a stone pavement was discovered.

The calcined and unburned bones (ca 116 kg ) were deposited within the structures both dispersed and in accumulations but with no anatomical order.

The grave goods - 850 artifacts in total - are distinctly subdivided into two chronologically different groups. The first group is dated to the time of construction of the tarand cemeteries, i.e., the Roman Iron Age; the second group derives from the third quarter of the 1st millennium AD. The finds of the Roman Iron Age are represented by bronze and iron ornaments (fibulae and their parts), as well as ceramics. The discovered bronze ornaments also included closed and spiral finger rings, bracelets, spiral beads, spiral temple rings, possible fragments of neck rings and some other rare specimens. These objects are typical of the Pyanobor archaeological culture and were evidently imported from the Kama River region. The weaponry and tools included iron spearheads, scythes and knives with a curved back, awls, a miniature pick-axe, and a miniature knife. Fragments of ceramics with striated and smoothed surfaces belong to the same period.

DNA-analyzed individuals:
VII14: Adult from pit No. 8, No. 2479. Sampled tooth r P ${ }^{1}$.
VIII4: Adult from the stone pavement, 2011, No. 2348 (horizon 4). Sampled tooth ? P?.
VIII5: Adult from the eastern tarand, 2011, No. 1622 (horizon 2). Sampled tooth r C ${ }^{1}$.
VIII6: Adult from the eastern tarand, 2013 (horizon 2). Sampled tooth $\mathrm{II}^{2}$.

## Udosolovo

Location: Udosolovo, district of Kingisepp, north-western Russia
Excavations: 2013 [127, 128], 2016-2017 [Stasyuk, unpublished]
Cemetery: The cemetery was originally a low flat stone mound of approximately rectangular shape (oriented NW-SE), badly damaged. The lower layer of burials in this mound reveals some inhumations in single stone cists, six of which were investigated. The skeletons were lying stretched on the back, head to the north. The cists were fragmentarily preserved, the bones were crushed into pieces by the weight of the stones and soil, some of the bones were displaced. Only a few items were found in graves: a narrow bronze bracelet, a javelin head and a fragment of an iron plate. Numerous small fragments of pottery (including those with striated surfaces) were found in this layer. The lower layer of burials in Udosolovo cemetery should be dated by the late Pre-Roman Iron Age ( $1^{\text {st }}$ century $B C$ - the first half of the $1^{\text {st }}$ century $A D$ ).

The upper layer of burials in the cemetery contained some scatterings of cremated bones mixed with gravel and soil, lying directly under the present turf. Between the two stages of the use of this cemetery there was a chronological gap, during which the stone cists were destroyed. No stone structures were found in the upper layer of the mound. There were almost no ceramics in the upper layer, but there were numerous metal items, often melted: an iron razor, iron knives, spirals of bronze wire, pieces of narrow bronze bracelets, a fragment of a silver neck-ring, etc. Finds from the upper layer with cremations are similar to those of tarand cemeteries in Northern Estonia and allow to date the assemblage to the 3rd century or even later, to the $5^{\text {th }}-7^{\text {th }}$ centuries.

DNA-analyzed individuals:
VII16: Male from burial 1, age 25-35 years [127]. Sampled tooth I M ${ }^{3}$.
VIII10: Male from burial 5, age 20-40 years [127]. Sampled tooth I M ${ }^{1}$.
VIII11: Adult [127] from square 4 (upper horizon). Sampled tooth I $\mathrm{M}_{1}$.
VIII12: Adult [127] from square 4 (horizon on stone layer). Sampled tooth $r \mathrm{M}_{2}$.

## Medieval rural cemeteries in Estonia

During the entire first millennium AD cremation burials were practised in Estonia. Inhumations with potential for aDNA analysis reappear in the late $10^{\text {th }} / 11^{\text {th }}$ century. The $11^{\text {th }}$ and $12^{\text {th }}$ centuries belong to the High Middle Ages in the historical chronology of western and central Europe, but Iron Age societies and culture still continued in the eastern Baltic area in that time.

The territory of Estonia was gradually conquered by German and Danish crusaders in the wars of 1208-1227. This conquest and forced Christianization mark the end of the Iron Age and the birth of medieval Livonia - a confederation of small states: the bishoprics
of Tartu and Ösel-Wiek in Estonia, those of Riga and Couronia in Latvia, and the Livonian branch of the Teutonic Order in a part of both countries. Northern Estonia belonged to Denmark until 1346, then it was sold to the Order. Although the end of the Middle Ages is usually dated around 1500 AD in Western Europe, for the area of medieval Livonia it is defined by the war with Russia (1558-1561).

In the rural archaeology of Estonia, the borders of the medieval period are, however, flowing and conventional. Burials from preconquest decades cannot clearly be distinguished from post-conquest ones. Until the transition of the country to Lutheran Sweden (since 1583 in Northern Estonia, since 1625 in Southern Estonia), the archaeological record of native Estonian population preserves features characteristic for medieval times. Thus, in the context of present research, the Middle Ages are regarded in a long-term perspective and individuals from ca 1200-1600 AD are conventionally regarded as 'medieval'.

As the Christianization of Estonia took place in a forced and violent way, the acceptance of Christian practices remained limited and a lot of pre-Christian traditions survived in medieval times. While in medieval Christian Europe people were normally buried in consecrated churchyards, in Livonia the dead were often buried at the home place, near villages and hamlets until the early $18^{\text {th }}$ century [129, 130]. Although cremation as a pagan practice was banned and greatly abandoned together with Christianization, the nonchurchyard village cemeteries existed parallel to churchyards. As the Livonian nobility of German origin was buried in churches and churchyards, individuals from Estonian village cemeteries represent the native Estonian population.

The village cemeteries lie usually 200/300-600/700 m from medieval village centers. If the landscape allows, they are located on low hummocks with the diameter usually from 15-20 to 40-60 m, sometimes more. In Estonia, there were usually ca 20-30 village cemeteries per parish. The number of people buried there depends on the local situation and duration of use, but it usually comprises several hundreds. Most of rural people were probably buried in village cemeteries in medieval time. The hinterlands of a local cemetery may have comprised from one to $2-4$ villages/hamlets, the number probably increasing in time, in parallel to population growth and settlement expansion. In Northern Estonia, the size of a village was mostly between 5-15 ploughlands in the mid-13 ${ }^{\text {th }}$ century, whereby each unit might roughly correspond to the number of farms, probably inhabited on the average by 5-8 people (incl. children) [131]. Villages of Southern Estonia were often of similar size in the $16^{\text {th }}$ century (earlier data are missing) but in areas with dispersed settlement there were small hamlets based on a few farms only.

Culturally, Estonia can be divided into coastal (sea-oriented) and inland (southern and eastern) areas. This distinction is clearly expressed in Estonian dialects [132], ethnography [133], folklore and traditional popular culture [134], as well as in present-day population genetic data [135]. The difference between the two macro-regions distinctly appears in the archaeological record also in the medieval period.

In the present study, Estonia's coastal areas are represented by the cemeteries of Karja, Pada and Kukruse, the inland areas by those of Otepää, Vana-Kuuste, Mäletjärve and Vaabina. In coastal Northern and Western Estonia, inhumations appear on some of the village cemeteries (e.g., Pada and Kukruse) some decades before the crusades, as a sign of transition to Christian religion and burial traditions. Some of these sites may have been deserted already soon after the conquest in the $13^{\text {th }}$ century. In that region, grave goods almost disappeared on rural cemeteries since the $2^{\text {nd }}$ half of the $13^{\text {th }}$ century but re-appeared again in the $16^{\text {th }}$ century. In inland Estonia, the pre-Christian practice of burying the dead dressed, together with jewelry items (brooches, rings, necklaces) and furnished with some minor grave goods - coins, knives, needles and other small utensil, survived continuously until the early $18^{\text {th }}$ century. The dead were buried mostly with the head toward west or south-west, according to medieval Christian practices, but in south-eastern Estonia the opposed orientation of men and women, a tradition of pre-Christian origin, lasted until the $17^{\text {th }}$ century.

Considering the presence of well-datable grave goods and coins, as well as relative chronology - in case of cemeteries of longterm use, earlier graves are often cut by later ones - the dates of $2^{\text {nd }}$ millennium AD inhumation burials are not based on radiocarbon samples which often provide a vague and wide date range, but on artifact chronology.

## Karja

Location: Karja, Saaremaa, Estonia
Excavations: 1955 [136]
Cemetery: Village cemetery on flat land, studied with rescue excavations (ca $150 \mathrm{~m} 2,32$ burials). The cemetery (full number of graves estimated as ca 70 ) with graves mainly from the $13^{\text {th }}$ century was probably founded soon after the Christianization (1227) and seems to have been deserted in the early $14^{\text {th }}$ century or by its middle. Burials of both sexes were oriented with the head toward W or SW. Some graves were furnished with jewelry (brooches, bracelets, rings), knives and belt accessories, some were unfurnished. DNA-analyzed individual:
Ila: Male (burial 16; Al 4115), 45+ years old, orientation WSW, furnished with a knife sheath. Sampled tooth r M $\mathrm{M}_{1}$, date 12301300 AD.

## Pada

Location: Pada, Lääne-Virumaa, Estonia
Excavations: 1987-1989 [137, 138]
Cemetery: Cemetery on flat land beside large $12^{\text {th }}$ and $13^{\text {th }} \mathrm{cc}$. Pada hill fort, a Final Iron Age district center, separated from it by a deep valley. The cemetery (investigated 171 burials and $253 \mathrm{~m}^{2}$ ) which dates from ca 1180-1250 probably belonged to the inhabitants of the hill fort and was deserted when the churchyard of Viru-Nigula was founded. Burials of both sexes were irregularly oriented with the head toward W, SW, E and NE. Graves were rich furnished with jewelry (brooches, bracelets, neck rings, breast chains with
pins, rings, necklaces), tools (axes, senses, knives), weapons (spears) and belt accessories. In four graves Gotlandic coins from 1140-1210/1220 were found.

DNA-analyzed individual:
IIg: Male (burial 151; Al 5366), 25-35 years old, WSW-oriented, richly furnished - horse harness, 4 silver coins (1140/60-1210/20), knife, belt accessories. Sampled tooth I M ${ }_{3}$, date 1210-1230/1240 AD.

## Kukruse

Location: Kukruse, Ida-Virumaa, Estonia
Excavations: 2009-2010 [139]
Cemetery: Cemetery on flat land, ca 300 m SE of Kukruse manor center. Rescue excavations (ca 600 m 2 ) revealed 44 inhumations mainly from the late $12^{\text {th }}$ and $13^{\text {th }}$ century and traces of earlier cremations. Burials of both sexes were of diverse orientation W, NW, SW, SSW, S, SE, E, N. Until Christianization (in 1220), and maybe also somewhat later, burials were rich in grave goods. A group of W-oriented graves (inc. grave 9) was most richly furnished with jewelry (brooches, bracelets, neck rings, breast chains with pins, rings, necklaces with silver sheet pendants), tools (axes, senses, knives), weapons (spears, a sword), and metal accessories of the costume. Special publications relate to burial rites [140, 141] and artifacts [142].

DNA-analyzed individual:
OLSO3: Male (burial 9; TÜ 1977), 25-30 years old, oriented toward W, richly furnished (clay vessel, sense, spearhead, knife, fire steel, neck rings, bracelets, brooch etc). Sampled tooth I M ${ }^{3}$, date 1180-1220/1240 AD.
Otepää
Location: Otepää, Tartumaa, Estonia
Excavations: 1928 [143], 1929 [144], 1938 [145], 1996 [146]
Cemetery: Located on flat land, studied with rescue excavations (ca $330 \mathrm{~m}^{2} ; 136$ burials). Otepää was a main castle of Tartu bishopric, with a big urban settlement at its foot in the $13^{\text {th }}$ and $14^{\text {th }} \mathrm{cc}$., the cemetery belongs to its inhabitants. Graves in parallel irregular rows were oriented with the head between W and SW. Judging by the almost total lack of disturbed graves, the site was of short-time use, dated by coin finds to the last third of the $14^{\text {th }}$ century. As most graves contained $2-4$ skeletons, the site seems to relate to some epidemic, maybe the plague of 1378 in which $5 / 6$ [147] or even about $9 / 10$ [148] of the people of the bishopric died. Judging by finds typical for the village cemeteries of the region - jewelry (brooches, rings, necklaces of cowry shells, glass beads, bells), knives, and belt accessories, the cemetery belongs to Estonian population.

DNA-analyzed individual:
IIf: Male (burial 1; AI 3680), 25-35 years old, oriented toward SW, finds: belt buckle, belt ring, knife. Sampled tooth $r \mathrm{M}_{3}$, date 1360-1390 AD.
Vana-Kuuste
Location: Vana-Kuuste, Tartumaa, Estonia
Excavations: 1982 [149]
Cemetery: Village cemetery on a low hummock in a forest, excavated (ca $75 \mathrm{~m} 2,99$ burials) to identify the character of the site. Investigated burials from the late $13^{\text {th }}$ or $14^{\text {th }}$ to the late $17^{\text {th }}$ century were oriented with the head toward W and SW, furnished with jewelry (brooches, rings, necklaces), knives, coins and belt accessories.

DNA-analyzed individual:
ILS01: Male (burial 73; TM A 153), 25-35 years old, oriented toward WSW, finds: knife, penannular brooch. Sampled tooth I M $\mathrm{M}_{1}$, date 1500-1625 AD.

## Mäletjärve

Location: Mäletjärve, Tartumaa, Estonia
Excavations: 1984 [150]
Cemetery: Village cemetery on flat land, founded beside a Roman Iron Age tarand cemetery. Trial excavations ( $50 \mathrm{~m} 2,50$ burials) in 1984 to establish the preservation/destruction state of the cemetery. Investigated graves from the late $14^{\text {th }}$ to the early $17^{\text {th }}$ century were oriented toward W and SW, furnished with jewelry (brooches, rings, necklaces), knives, coins and belt accessories.

DNA-analyzed individual:
IVLS09KT: Male (burial 18; TM A 155), 30-40 years old, oriented toward SSW, finds: coin from 157?, penannular brooch, knife. Sampled tooth I M ${ }_{3}$, date 1570-1600 AD.

## Vaabina

Location: Vaabina, Võrumaa, Estonia
Excavations: 1985 [151]
Cemetery: Village cemetery on top of a high hummock, studied with rescue excavations (ca 350 m 2 , remains of 64 skeletons), dates from the mid- $13^{\text {th }}$-late $17^{\text {th }}$ century. Male graves were oriented with the head toward W , female, according to local regional tradition, toward E. Burials were furnished with jewelry (brooches, rings, necklaces), knives, coins and belt accessories.

DNA-analyzed individual:
IIIt: Female (burial 43; AI 5354), 40+ years old, oriented toward E, finds: knife, $13^{\text {th }}-14^{\text {th }} \mathrm{cc}$. brooch. Sampled tooth $r \mathrm{M}^{1}$, date 1250-1450 AD.

## METHOD DETAILS

All of the laboratory work was performed in dedicated ancient DNA laboratories of the Institute of Ecology and Earth Sciences, University of Tartu or the Department of Archaeology and Anthropology, University of Cambridge. The library quantification and sequencing were performed at the Estonian Biocenter Core Laboratory. The main steps of the laboratory work are detailed below.

## DNA extraction

The teeth of 56 individuals were used to extract DNA.
Tooth roots were broken off and used for extraction since root cementum has been shown to contain more endogenous DNA than crown dentine [152]. The roots were used whole to avoid heat damage during powdering with a drill and to reduce the risk of crosscontamination between samples. Contaminants were removed from the surface of tooth roots by soaking in $6 \%$ bleach for 15 mi nutes, then rinsing twice with water and lastly soaking in $70 \%$ ethanol for 2 minutes, shaking the tubes during each round to dislodge particles. Finally, the samples were left to dry under a UV light for 30 minutes on both sides.

Next, the samples were weighed, [20 * sample mass (mg)] $\mu$ l of EDTA and [sample mass ( mg ) / 2] $\mu$ l of proteinase K was added and the samples were left to digest for 72 hours on a slow shaker at $20^{\circ} \mathrm{C}$ to compensate for the smaller surface area of the whole root compared to powder. Undigested material was stored for a second DNA extraction if need be.

The DNA solution was concentrated to $250 \mu \mathrm{l}$ (Amicon Ultra-15 30 kDa , Merck Millipore) and purified in large volume columns (High Pure Viral Nucleic Acid Large Volume Kit, Roche) using 2.5 mL of PB buffer, 1 mL of PE buffer and $50 \mu \mathrm{~L}$ of EB buffer (MinElute PCR Purification Kit, QIAGEN).

## Library preparation

Sequencing libraries were built using NEBNext DNA Library Prep Master Mix Set for 454 (E6070, New England Biolabs) and Illuminaspecific adaptors [153] following established protocols [153-155]. The end repair module was implemented using $18.75 \mu \mathrm{~L}$ of water, $7.5 \mu \mathrm{~L}$ of buffer and $3.75 \mu \mathrm{~L}$ of enzyme mix, incubating at $20^{\circ} \mathrm{C}$ for 30 minutes. The samples were purified using $500 \mu \mathrm{LPB}$ and $650 \mu \mathrm{~L}$ of PE buffer and eluted in $30 \mu \mathrm{~L}$ EB buffer (MinElute PCR Purification Kit, QIAGEN). The adaptor ligation module was implemented using $10 \mu \mathrm{~L}$ of buffer, $5 \mu \mathrm{~L}$ of T 4 ligase and $5 \mu \mathrm{~L}$ of adaptor mix [153], incubating at $20^{\circ} \mathrm{C}$ for 15 minutes. The samples were purified as in the previous step and eluted in $30 \mu \mathrm{~L}$ of EB buffer (MinElute PCR Purification Kit, QIAGEN). The adaptor fill-in module was implemented using $13 \mu \mathrm{~L}$ of water, $5 \mu \mathrm{~L}$ of buffer and $2 \mu \mathrm{~L}$ of Bst DNA polymerase, incubating at $37^{\circ} \mathrm{C}$ for 30 and at $80^{\circ} \mathrm{C}$ for 20 minutes. The libraries were amplified and both the indexed and universal primer (NEBNext Multiplex Oligos for Illumina, New England Biolabs) were added by PCR using HGS Diamond Taq DNA polymerase (Eurogentec). The samples were purified and eluted in $35 \mu \mathrm{~L}$ of EB buffer (MinElute PCR Purification Kit, QIAGEN). Three verification steps were implemented to make sure library preparation was successful and to measure the concentration of dsDNA/sequencing libraries - fluorometric quantitation (Qubit, Thermo Fisher Scientific), parallel capillary electrophoresis (Fragment Analyzer, Agilent Technologies) and qPCR.

## DNA sequencing

DNA was sequenced using the Illumina NextSeq 500 platform with the 75 bp single- or paired-end method. As a norm, 12 samples were sequenced together on one flow cell; additional data was generated for 6 samples on one flow cell to increase coverage.

## QUANTIFICATION AND STATISTICAL ANALYSIS

## Mapping

Before mapping, the sequences of adaptors and indexes and poly-G tales occurring due to the specifics of the NextSeq 500 technology were cut from the ends of DNA sequences using cutadapt 1.11 [54]. Sequences shorter than 30 bp were also removed with the same program to avoid random mapping of sequences from other species.

The sequences were mapped to reference sequence GRCh37 (hs37d5) using Burrows-Wheeler Aligner (BWA 0.7.12) [55] and command mem with re-seeding disabled.

After mapping, the sequences were converted to BAM format and only sequences that mapped to the human genome were kept with samtools 1.3 [56]. Next, data from different flow cell lanes was merged and duplicates were removed with picard 2.12 (http:// broadinstitute.github.io/picard/index.html). Indels were realigned with GATK 3.5 [57] and lastly, reads with mapping quality under 10 were filtered out with samtools 1.3 [56].

The average endogenous DNA content (proportion of reads mapping to the human genome) for EstBA samples was $21 \%$, for EstIA samples $23 \%$, for IngIA samples $15 \%$ and for EstMA samples $36 \%$ (Table S1). The variation in the endogenous DNA content was variable as is common in aDNA studies, ranging from under $1 \%$ in most sample groups to at least over $60 \%$ in all four groups (Table S1).

## aDNA authentication

As a result of degrading over time, aDNA can be distinguished from modern DNA by certain characteristics: short fragments and a high frequency of $\mathrm{C}=>\mathrm{T}$ substitutions at the $5^{\prime}$ ends of sequences due to cytosine deamination. The program mapDamage 2.0 [58] was used to estimate the frequency of $5^{\prime} \mathrm{C}=>\mathrm{T}$ transitions.
mtDNA contamination was estimated using the method from Fu et al. 2013 [59]. This included calling an mtDNA consensus sequence based on reads with mapping quality at least 30 and positions with at least $5 x$ coverage, aligning the consensus with 311 other human mtDNA sequences from Fu et al. 2013 [59], mapping the original mtDNA reads to the consensus sequence and running contamMix 1.0-10 with the reads mapping to the consensus and the 312 aligned mtDNA sequences while trimming 7 bases from the ends of reads with the option trimBases.

For the male individuals, contamination was also estimated based on X chromosome using the two contamination estimation methods first described in Rasmussen et al. 2011 [156] and incorporated in the ANGSD software [60] in the script contamination.R.

The Bronze and Iron Age samples on average showed $14 \% \mathrm{C}=>\mathrm{T}$ substitutions at the $5^{\prime}$ ends while for the considerably more recent Middle Age samples this result was $7 \%$ (Table S1). The mtDNA contamination point estimate for samples with > $6 x \mathrm{mtDNA}$ coverage ranged from $0.05 \%$ to $3.65 \%$ with an average of $0.6 \%$ (Table S1). The average of the two X chromosome contamination methods of male individuals with average $X$ chromosome coverage $>0.1 x$ was between $0.07 \%$ and $3.02 \%$ with an average of $1.07 \%$ (Table S1).

## Calculating general statistics and determining genetic sex

Samtools 1.3 [56] option stats was used to determine the number of final reads, average read length, average coverage etc.
Genetic sex was calculated using the script sexing.py from Skoglund et al. 2013 [61], estimating the fraction of reads mapping to Y chromosome out of all reads mapping to either X or Y chromosome.

The average coverage of the whole genome for the samples was between 0.0001 x and 0.7 x (Table S1). Genetic sexing confirmed morphological sex estimates or provided additional information about the sex of the individuals involved in the study. The sex of 12 of the samples could not be reliably estimated due to low coverage. Apart from those samples, the study involves 12 females and 32 males (Table 1) since a focal point of the study is chromosome Y .

## Variant calling

Variants were called with the ANGSD software [60] command doHaploCall, sampling a random base for the positions that are present in the EBC-chipDB [5, 41-49] (Table S3).

## Determining mtDNA haplogroups

mtDNA haplogroups were determined by submitting mtDNA BAM files to mtDNA-Server [62] which uses HaploGrep2 [63, 64] for assigning haplogroups. Subsequently, the results were checked visually by aligning mapped sequences to reference sequence rCRS [157] with samtools 0.1.19 [56] command tview and confirming the haplogroup assignments in PhyloTree [64].

41 of the 56 individuals were successfully haplogrouped (Table 1).

## Y chromosome variant calling and haplogrouping

Y chromosome variants were called from the BAM files of the samples using ANGSD [60] doHaploCall. The resulting VCF files were filtered for regions of a total length of 8.8 Mbp of sequence that uniquely maps to human Y chromosome when using short read sequencing technology [25]. Variants called within this 8.8 Mbp region were further filtered for 113,217 haplogroup informative positions [25, 29, 158-160] using BEDTools 2.19 .0 [65] intersect option. Haplogroup assignments of each individual sample were made by determining the haplogroup with the highest proportion of informative positions called in derived state in the given sample. Y chromosome haplogrouping was performed on all samples to check if any of the samples estimated to be female also give a result.

None of the female samples were successfully haplogrouped as expected. 30 out of the 32 males were successfully haplogrouped (Table 1).

## Preparing the datasets for autosomal analyses

The EBC-chipDB [5, 41-49] was used as the modern DNA background. Individuals from Lazaridis et al. 2016 [14], Jones et al. 2017 [10], Unterländer et al. 2017 [50], Saag et al. 2017 [13], Mittnik et al. 2018 [11], Mathieson et al. 2018 [12], two Damgaard et al. 2018 [51, 52] papers, Narasimhan et al. 2018 [53] and Lamnidis et al. 2018 [26] were used as the ancient DNA background. The full genome sequencing data of the aDNA background dataset $[10,13,51,52]$ in the form of FASTQ files was called as described in the Variant calling section. The 1240k capture data of the aDNA background dataset [11, 12, 14, 26, 50,53] was downloaded in EIGENSTRAT format. The data of the two comparison datasets and of the individuals of this study was converted to BED format using PLINK 1.90 (http://pngu.mgh.harvard.edu/purcell/plink/) [66], the datasets were merged and the 503,714 SNPs of the modern comparison dataset were kept. Due to low coverage ( $<0.017 x$ ) resulting in a low number of SNPs ( $<10,000$ of the 503,714 ), 23 of the individuals of this study were removed from further autosomal analyses, leaving 15 individuals from Bronze Age Estonia, 6 from Iron Age Estonia, 5 from Iron Age Ingria and 7 from Middle Age Estonia to be used in autosomal analyses (Table S1).

## Principal component analysis

To prepare for principal component analysis (PCA), a reduced comparison sample-set composed of 817 modern individuals from 46 populations of Europe, Caucasus and Near East and 645 ancient individuals from 97 populations was assembled (Table S3). The data was converted to EIGENSTRAT format using the program convertf from the EIGENSOFT 7.2 .0 package [67]. PCA was performed with the program smartpca from the same package, projecting ancient individuals onto the components constructed based on
the modern genotypes using the option Isqproject and trying to account for the shrinkage problem introduced by projecting by using the option autoshrink.

## Outgroup f3 statistics

For calculating autosomal outgroup f3 statistics, the same ancient sample-set as for PCA was used and the modern sample-set was increased to 1490 individuals from 92 populations from Europe, Caucasus, Near East, Siberia, Central Asia and East Asia and Yorubas as outgroup (Table S3). Heterozygous positions were converted to homozygous by randomly choosing one of the alleles at each position to enable comparison between pseudo-haploid ancient samples and diploid modern samples. The data was converted to EIGENSTRAT format using the program convertf from the EIGENSOFT 5.0.2 package [67]. Outgroup f3 statistics of the form f3(Yorubas; EstBA/EstIA/IngIA/EstMA, modern/ancient) were computed using the ADMIXTOOLS 1.1 [68] program qp3Pop.

To allow for X chromosome versus autosomes comparison, outgroup f3 statistics using X chromosome SNPs were computed. However, the overlap between the X chromosome positions of the EBC-chipDB [5, 41-49] and the 1240k capture data of the ancient comparison sample-set was only 17,852 SNPs. To be able to use the whole ancient comparison dataset for this analysis, the full genome sequencing data of that dataset and the individuals of this study were called as described in the Variant calling section but using the positions of the Lazaridis et al. [14] ancient dataset. To allow for the use of the bigger number of positions in the ancient over the modern dataset from Lazaridis et al. [14], Mbuti from Panel C of the Simons Genome Diversity Project [161] was used as the outgroup. The outgroup f 3 analyses of the form $\mathrm{f} 3(\mathrm{Mbuti}$; EstBA/EstIA/IngIA/EstMA, ancient) were run both using 991,166 autosomal SNPs and also 40,185 X chromosome positions available in the Lazaridis et al. [14] ancient dataset. Since all children inherit half of their autosomal material from their father but only female children inherit their $X$ chromosome from their father then in this comparison X chromosome data gives more information about the female and autosomal data about the male ancestors of a population.

The autosomal outgroup f3 results of the two different SNP sets were compared to see whether the SNPs used affect the trends seen.

## D statistics

D statistics of the form D(Yorubas, EstBA/EstIA/IngIA/EstMA; Estonians, modern/ancient) were calculated on the same EBC-chipDB [5, 41-49] as outgroup f3 statistics (Table S3). The ADMIXTOOLS 1.1 [68] package program qpDstat was used.

## Admixture analysis

Three Paleolithic individuals were added to the ancient sample-set used for previous analyses and the modern sample-set was increased to 1799 individuals from 115 populations from all over the world for Admixture analysis [69] (Table S3). The analysis was carried out using ADMIXTURE 1.3 [69] with the $P$ option, projecting ancient individuals into the genetic structure calculated on the modern dataset due to missing data in the ancient samples. The dataset of modern individuals was pruned to decrease linkage disequilibrium using the option indep-pairwise with parameters 10002500.4 in PLINK 1.90 (http://pngu.mgh.harvard.edu/purcell/ plink/) [66]. This resulted in a set of 216,398 SNPs. Admixture was run on this set using $\mathrm{K}=3$ to $\mathrm{K}=18$ in 100 replicates. This enabled us to assess convergence of the different models. $\mathrm{K}=10$ and $\mathrm{K}=9$ were the models with the largest number of inferred genetic clusters for which $>10 \%$ of the runs that reached the highest Log Likelihood values yielded very similar results. This was used as a proxy to assume that the global Likelihood maximum for this particular model was indeed reached. Then the inferred genetic cluster proportions and allele frequencies of the best run at $\mathrm{K}=9$ were used to run Admixture to project the aDNA individuals on the inferred clusters. The same projecting approach was taken for all models for which there is good indication that the global Likelihood maximum was reached (K3-18). We present all individuals on Figure S1 but only population averages of those aDNA samples on Figure 1 for which the intersection with the LD pruned modern dataset yielded data for more than 10,000 SNPs. The resulting membership proportions to K genetic clusters are sometimes called "ancestry components" which can lead to over-interpretation of the results. The clustering itself is, however, an objective description of genetic structure and as such a valuable tool in population comparisons.

## ChromoPainter/NNLS

In order to infer the admixture proportions of ancient individuals, the ChromoPainter/NNLS pipeline [19, 71, 72] was applied. Due to the low coverage of the ancient data, it is not possible to infer haplotypes and the analysis was performed in unlinked mode (option -u). Only samples with more than 20,000 SNPs were used in the analyses. Since ChromoPainter [70] does not tolerate missing data, every ancient target individual was iteratively painted together with one representative individual from potential source populations as recipients. All the remaining modern individuals from the sample-set used for Admixture analysis were used as donors (Table S3). Subsequently, we reconstructed the profile of each target individual as a combination of three or more ancient individuals, using the non-negative least square approach. Let $X g$ and $Y p$ be vectors summarizing the proportion of DNA that source and target individuals copy from each of the modern donor groups as inferred by ChromoPainter. $Y p=\beta_{1} X_{1}+\beta_{2} X_{2}+\beta_{3} X_{3}+\ldots+\beta_{z} X_{z}$ was reconstructed using a slight modification of the nnls function in R [162] and implemented in GlobeTrotter [163] under the conditions $\beta_{g} \geq 0$ and $\sum \beta_{g}=1$. In order to evaluate the fitness of the NNLS estimation, we inferred the sum of the squared residual for every tested model and reported the one with the lowest value [164]. The model with the smallest residual values included WHG (Loschbour
[15]), Yamnaya (Yamnaya [52]), Central MN (I0172 [16]) and modern Nganasans (Nganassan11 [46]) as sources (see other models in Data S1). The resulting painting profiles, which summarize the fraction of the individual's DNA inherited by each donor individual, were summed over individuals from the same population.

## qpAdm

The ADMIXTOOLS 1.1 [68] package programs qpWave and qpAdm were used to estimate which populations and in which proportions are suitable proxies of admixture to form the populations of this study. Only samples with more than 100,000 SNPs were used in the analyses. The best model tested (taking into account $p$ values, standard errors and the presence of negative values for proportions) included EstBA/EstIA/IngIA/EstMA, WHG, Yamnaya Kalmykia, Central MN and Nganasans as left populations and Yorubas, Ust-Ishim, Mal'ta, Kostenki, SHG and Han as right populations (see other models in Data S1).

## f4 statistics

$f 4$ statistics of the form $f 4$ (Yorubas, Nganasans; period in Estonia/Central LNBA, period in Baltics), f 4 (Yorubas, Koryaks; period in Estonia, period in Baltics), 44 (Yorubas, WHG; period in Estonia, period in Baltics) and $f 4$ (Yorubas, Syrians/Yamnaya Kalmykia/Central MN; period in Estonia, period in Baltics) were calculated on the same EBC-chipDB [5, 41-49] as outgroup f3 statistics (Table S3). The ADMIXTOOLS 1.1 [68] package program qpDstat and the option f4mode: YES was used.

## Population continuity tests

We applied the forward simulation method described in Hofmanová et al. 2016 [19] to test whether individual genomes from the ancient Estonian populations can be considered as sampled from a population directly ancestral to modern Estonians under a model of full continuity. We used the overlapping positions between the pseudo-haploid calls of our ancient genomes and the biallelic calls of ten modern Estonian genomes extracted from the Human Origins dataset [14] to estimate their population allele frequencies and infer the site frequency spectrum (SFS). In order to preserve the SFS shape, we only tested ancient genomes for which more than 100,000 SNPs overlapped with the modern dataset (Data S1). Alleles were then polarized into ancestral and derived by comparing them with the alleles in the chimpanzee to obtain the derived folded SFS.

For each combination of an ancient genome and the ten modern genomes we performed the steps described in Hofmanová et al. 2016 [19]. Briefly, we first incorporated uncertainty on the allele frequencies by sampling 100 frequency vectors using a beta distribution and the Jeffreys' prior [165] from the distribution of allele frequencies of the SFS of the modern Estonian individuals. We then use binomial sampling in forward simulations to emulate a genetic drift process and generate possible allele trajectories given the age of the ancient sample in generations. We explored two parameters, ancient $\left(\mathrm{Ne}_{\mathrm{a}}\right)$ and modern $\left(\mathrm{Ne}_{\mathrm{m}}\right)$ effective population sizes, assuming a model of exponential growth between them. For each simulation we sampled a haploid genome from the initial frequency vector and another one from each simulated final frequency vector. We compared the observed calls with the simulated ones using an allelic sharing classification consisting of six possible classes formed by all possible combinations of haploid calls of an ancient genome ( t 0 ) and the biallelic calls of each modern genome (tn) for the same position: 1) match A: t0 ancestral (A) and tn AA, 2) match D: t0 derived ( D ) and tn DD, 3) mismatch AD: to $A$ and tn DD, 4) mismatch DA: t0 $D$ and tn AA, 5) half match A: t0 A and tn AD, and 6) half match D: t0 D and tn AD (see Hofmanova et al. 2016 [19]). Allele sharing fraction values are calculated for both the observed and simulated data as the proportion of all analyzed positions that fall into each one of these six classes. Finally, we calculated an overall $p$ value for the null-hypothesis of rejection of population continuity for each combination of parameters by combining the individual p values for each allelic sharing fraction using Fisher's and Voight's methods [166] as described in Hofmanová et al. 2016 [19].

We explored a wide parameter space of $\mathrm{Ne}_{\mathrm{a}}$ and $\mathrm{Ne}_{\mathrm{m}}$ for each ancient genome and the modern individuals by performing the test in a $50 \times 50$ grid composed for values of these effective population sizes ranging from 10 to 10 million individuals on a log scale. For each combination of parameters, we performed 1,000 simulations, thus the total number of simulations per test (each ancient genome versus modern Estonians) was 2.5 million. The ranges of realistic effective population sizes in which continuity could not be rejected were examined by slicing the parameter grid by the mean, upper and lower Cl of the effective population size of modern Estonians, estimated on over 2,000 modern Estonian full genomes (unpublished; cohort [29]) using the program IBDNe [167]. The two-tail $p$ values of the test for each ancient effective size are reported.

## Kinship analysis

A total of $4,375,438$ biallelic single nucleotide variant sites, with MAF $>0.1$ in a set of over 2,000 high coverage genomes of EGC (unpublished; cohort [29]) were identified and called with ANGSD-0.916 [60] command doHaploCall from the BAM files of 12 Bronze Age, 11 Iron Age and 6 medieval individuals with coverage $>0.03 x$. The ANGSD output files were converted to .tped format as an input for the analyses with READ script to infer pairs with $1^{\text {st }}$ and $2^{\text {nd }}$ degree relatedness [73].

## Radiocarbon date difference probability estimation

Probabilistic estimates of the temporal distance between the radiocarbon dates associated with X14 (2481 $\pm 30 \mathrm{BP})$ and V16 (2399 $\pm$ 27 BP) have been obtained by: 1) calibrating both dates using the IntCal13 calibration curve [28] (using the rcarbon R package [168]); and 2) sampling one million pairs of random dates from each distribution and calculating their differences. The resulting distribution of differences had a 95\% HPD between between - 76 (i.e., V16 earlier) and 344 years (i.e., V16 later). We then calculated the expected difference in time between the date of death of X14 and V16, assuming that: 1) the former was the uncle, the latter the nephew; 2 ) an
age of death between 35 and 45 for X 14 [91] and between 30 and 40 for V 16 [84]; 3) a reproductive age span between 13 and 40 years old; and 4) a maximum age difference between X14 and his sister of 27 years (i.e., 40-13). The difference in the date of death can then be calculated using the following formula $(a+b)-(c+d)$, where $a$ is the age at death of $V 16, b$ is the age at which X 14 's sister gave birth to V16, $c$ is the age at death of X14 and $d$ is the difference in age between X14's sister and X14 (i.e., negative if X14 is assumed to be older, positive if his sister was born first). It follows that the difference in time between the date of death of X14 and V16 could range between - 29 (i.e., V16 dying before X14) and 72 years (V16 dying later). We then computed that the probability that difference in the age of the radiocarbon dates is within this interval computing the proportion of dates within -29 to 72 , which was equivalent to 0.15 . We also calculated the ranges and probabilities if V16 was X14's uncle and if the two were half brothers sharing a mother.

## Phenotyping

The phenotype prediction was performed only on the samples with an average genomic coverage greater than 0.1 x , for a total of 23 subjects (Data S2).

In order to predict eye, hair and skin color in the ancient individuals (Data S2), we selected all the 41 variants from 19 genes in 9 autosomes in the HIrisPlex-S system [76] and, for each autosome, we selected the region to be analyzed adding 5 Mb at each side of the chromosomal segment delimited by the first and the last SNP. We analyzed the three genes on chromosome 15 in two different regions (OCA2-HERC2 region and SLC24A5 region), because the distance between the two nearest SNPs of the two chromosomal segments was greater than 20 Mb . We obtained a total of 10 regions ranging from about 10 Mb to about 15 Mb . We chose as reference panel a set of 606 modern individuals, from all the European (EUR) populations and one Asian (CHB) population of the 1000 Genomes Phase 3 [169]. The Chinese outgroup was added to include also the variants that are very rare in Europe. The variant sites in the 10 chromosomal regions were extracted from the phased VCF files of the modern individuals with VCFtools [74], discarding the indels. The resulting VCF files were filtered using bcftools [170] to keep only the biallelic SNPs with a minor allele frequency (MAF) above a chosen threshold. We set the MAF threshold to $1 \%$ for all the genes, with the exception of the region on chromosome 16 , for which the MAF threshold was set to $0.1 \%$ to retain as many rare SNPs as possible from the MC1R gene. These settings allowed us to exclude only 3 SNPs and one indel out of the 41 HIrisPlex-S informative markers. The final VCFs were manipulated with PLINK 1.9 (http://pngu.mgh.harvard.edu/purcell/plink/) [66] to obtain a list of variant sites and a map file for each region.

We calculated the genotype likelihoods for the variant sites in the ancient individuals using the ANGSD [60] -GL command, with the -dopost 1 option and a reference sequence in the FASTA format. We then performed the imputation step using the Beagle 4.1 and Beagle 5.0 software [75]. First, we loaded on Beagle the ANGSD VCF output and we used the -gl command to infer genotype probabilities (GP). We obtained a VCF output including a GT value in the FORMAT field and filtered it to discard all variants with a GP $\leq 0.99$. The filtered VCF was loaded again on Beagle 5.0 for a second run with the -gt command to impute at ungenotyped sites. The resulting VCF files were filtered again to keep only variants with a GP $\geq 0.85$, with the exception of the HERC2 rs12913832 variant: since the lack of this SNP will not produce an eye color prediction result, we set the GP threshold to 0.6 . The resulting VCF files were subset with VCFtools [74] to extract the SNPs relevant for the phenotype prediction. The SNPs were recoded and organized with PLINK 1.9 (http://pngu.mgh.harvard.edu/purcell/plink/) [66] and R [162] and the missing variants were coded as "NA" to produce a csv input file for the HlrisPlex-S webtool (https://hirisplex.erasmusmc.nl/), that was used to perform the phenotype prediction [76-78].

We used the same approach to extract the allele information for the lactase persistence variant (rs4988235 in the MCM6 gene) and two variants involved in the protection against leprosy (rs5743618 and rs4833095 in the TLR1 gene). For the TLR1 variants, a more relaxed GP threshold of 0.60 was used. We also used the same approach to impute rs333 (CCR5-32bp deletion) but used a larger local Estonian reference panel of over 2,000 EGC high coverage genomes (unpublished; cohort [29]).

We tested the accuracy of our imputation pipeline by downsampling a high-coverage sample and comparing the variants imputed in the downsampled samples to the variants in the original one. To this aim, we selected the high-coverage (20x) NE1 individual from Gamba et al. 2014 [76] and randomly downsampled it to a coverage of $0.05 x$ and $0.1 x$ using SAMtools [56]. We applied the same ANGSD commands described above to calculate the genotype likelihoods for the variants in the chromosome 20 region in both the high-coverage and low-coverage NE1 bams. We then followed the same pipeline described above, obtaining an overall concordance rate of about $95 \%$.

## DATA AND SOFTWARE AVAILABILITY

The DNA sequences are available through the data depository of the EBC (http://evolbio.ut.ee) and the accession number for the sequences reported in this paper is ENA: PRJEB31893.

## Supplemental Information

## The Arrival of Siberian Ancestry Connecting

## the Eastern Baltic to Uralic Speakers further East

Lehti Saag, Margot Laneman, Liivi Varul, Martin Malve, Heiki Valk, Maria A. Razzak, Ivan G. Shirobokov, Valeri I. Khartanovich, Elena R. Mikhaylova, Alena Kushniarevich, Christiana Lyn Scheib, Anu Solnik, Tuuli Reisberg, Jüri Parik, Lauri Saag, Ene Metspalu, Siiri Rootsi, Francesco Montinaro, Maido Remm, Reedik Mägi, Eugenia D'Atanasio, Enrico Ryunosuke Crema, David Díez-del-Molino, Mark G. Thomas, Aivar Kriiska, Toomas Kivisild, Richard Villems, Valter Lang, Mait Metspalu, and Kristiina Tambets


Figure S1. ADMIXTURE analysis results with ancient data projected onto the modern genetic structure. Related to Figure 1. A. Ancient individuals at K9. B. Modern population averages at K9. C. Ancient individuals at K3 to K18. D. Modern population averages at K3 to K18.


Figure S2. D and outgroup f3 statistics' results. Related to Figure 2. EF - early farmers; HG - huntergatherers; LNBA - Late Neolithic/Bronze Age; IMA - Iron/Middle Ages; LN - Late Neolithic; BA Bronze Age; IA - Iron Age; MA - Middle Ages. A. D statistics’ values of form D(Yorubas, Estonian Corded Ware culture/Bronze Age; Estonians, ancient). B. Outgroup f3 statistics' values of form f3(Mbuti; Estonian Corded Ware culture/Bronze Age, ancient) using 1240k capture SNPs. C-F. Outgroup f3 statistics' values of form f (Mbuti; autosomal SNPs/chr X SNPs, modern) of Estonian Corded Ware culture, Bronze Age, Iron Age and Middle Ages respectively using 1240k capture SNPs.


Figure S3. Kinship test results, date difference probability estimation for found relatives and continuity test results. Related to Table 1. A. Bronze Age Estonian kinship test results. B. Iron Age Estonian and Ingrian kinship test results. C. Middle Age Estonian kinship test results. D. Probabilistic estimate for the distance in time between the radiocarbon dates associated with X14 and V16 obtained via one million MC simulations of calendar dates. E. Continuity test results between a Middle Age Estonian genome and modern Estonians.

| Modern population | References | PCA | $\begin{aligned} & \mathrm{f3}, \\ & \mathrm{D} \end{aligned}$ | ADM, CP | Ancient population | References | PCA | $\begin{aligned} & \mathrm{f} 3, \\ & \mathrm{D} \end{aligned}$ | ADM |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Abkhasians | [S15] | + | + | + | Afanasievo | [S16,S17] | + | + | + |
| Adygei | [S18] | + | + | + | Aktogai_MLBA | [S17] | + | + | + |
| Altaians | [S19,S20] |  | + | + | Alan | [S21] | + | + | + |
| Armenians | [S15] | + | + | + | Anatolia_ChL | [S16] | + | + | + |
| Azeris | [S22] | + | + | + | Anatolia_N | [S16] | + | + | + |
| Balkars | [S15] | + | + | + | Andronovo | [S16,S21] | + | + | + |
| Balochi | [S18] |  |  | + | Armenia_ChL | [S16] | + | + | + |
| Bantus | [S18] |  |  | + | Armenia_EBA | [S16] | + | + | + |
| Bashkirs | [S22] |  | + | + | Armenia_MLBA | [S16] | + | + | + |
| Basques | [S18] | + | + | + | Bell_Beaker | [S16] | + | + | + |
| Belorussians | [S23,S24] | + | + | + | Bichon | [S16] | + | + | + |
| Brahui | [S18] |  |  | + | Bolshoy Oleni Ostrov | [S25] | + | + | + |
| Bulgarians | [S15] | + | + | + | Botai | [S26] | + | + | + |
| Burusho | [S18] |  |  | + | Cardial_EN | [S16] | + | + | + |
| Buryats | [S19] |  | + | + | Central_LNBA | [S16] | + | + | + |
| Chechens | [S15] | + | + | + | Central_MN | [S16] | + | + | + |
| Chukchis | [S19,S27] |  | + | + | CentralSteppe_EMBA | [S26] | + | + | + |
| Chuvash | [S23] |  | + | + | Chalmny Varre | [S25] | + | + | + |
| Cypriots | [S23] | + | + | + | CHG | [S16] | + | + | + |
| Daur | [S18] |  | + | + | Dali_MLBA | [S17] | + | + | + |
| Dolgans | [S19,S27] |  | + | + | Dashti_Kozy_BA | [S17] | + | + | + |
| Erza | [S15] |  |  | + | EHG | [S16,S28] | + | + | + |
| Estonians | [S20,S24,S29] | + | + | + | Estonia_CCC | [S28,S30] | + | + | + |
| Ethiopians | [S23] |  |  | + | Estonia_CWC | [S28,S30] | + | + | + |
| Evenkis | [S19] |  | + | + | Estonia_HG | [S28,S30] | + | + | + |
| Evens | [S20,S27] |  | + | + | Globular_Amphora | [S31] | + | + | + |
| Finns | [S29] | + | + | + | HallstattBylany | [S21] | + | + | + |
| French | [S18] | + | + | + | Hungarian_Med | [S21] | + | + | + |
| Gagauzes | [S15,S29] | + | + | + | HungarianScythian | [S21] | + | + | + |
| Germans | [S15,S29] | + | + | + | Hungary_BA | [S16] | + | + | + |
| Han | [S18] |  | + | + | Hungary_EN | [S16] | + | + | + |
| Hazara | [S18] |  |  | + | Iberia_BA | [S16] | + | + | + |
| Hezhen/Nanaj | [S18] |  | + | + | lberia_Chl | [S16] | + | + | + |
| Hungarians | [S23] | + | + | + | Iberia_EN | [S16] | + | + | + |
| Ingrian_Finns | [S29] | + | + | + | lberia_MN | [S16] | + | + | + |
| Iranians | [S23] |  | + | + | Iran_ChL | [S16] | + | + | + |
| Japanese | [S18] |  | + | + | Iran_Hotulllb | [S16] | + | + | + |
| Jordanians | [S23] | + | + | + | Iran_LN | [S16] | + | + | + |
| Kalmyks | [S22] |  | + | + | Iran_N | [S16] | + | + | + |
| Karelians | [S15,S29] | + | + | + | Iron_Gates_HG | [S31] | + | + | + |
| Kazakhs | [S20] |  | + | + | Kairan_MLBA | [S17] | + | + | + |
| Kets | [S19,S27] |  | + | + | Kangju | [S21] | + | + | + |
| Khakases | [S27] |  | + | + | Karagash_MLBA | [S17] | + | + | + |
| Khanty | [S29] |  | + | + | Kashkarchi_BA | [S17] | + | + | + |
| Komis | [S22] |  | + | + | Kazakh_His | [S21] | + | + | + |
| Koryaks | [S19,S27] |  | + | + | Kazakh_Mys_MLBA | [S17] | + | + | + |
| Kumyks | [S15] | + | + | + | Kostenki | [S16] |  |  | + |
| Kurds | [S15] |  | + | + | Krasnoyarsk_MLBA | [S17] | + | + | + |
| Kyrgyzians | [S20] |  | + | + | Latvia_BA | [S28] | + | + | + |
| Latvians | [S24,S29] | + | + | + | Latvia_CCC_EHG | [S31,S32] | + | + | + |
| Lebanese | [S23] | + | + | + | Latvia_CCC_WHG | [S31,S32] | + | + | + |
| Lezgins | [S23] | + | + | + | Latvia_CWC | [S32] | + | + | + |
| Lithuanians | [S23] | + | + | + | Latvia_HG | [S31,S32] | + | + | + |


| Makrani | [S18] |  |  | + | LBK_EN | [S16] | + | + | + |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Malayan | [S23] |  |  | + | LchashenMetsamor | [S21] | + | + | + |
| Mandenkas | [S18] |  |  | + | Levant_BA | [S16] | + | + | + |
| Mansis | [S29] |  | + | + | Levant_N | [S16] | + | + | + |
| Maris | [S20,S29] |  | + | + | Lisakovskiy_MLBA_Alakul | [S17] | + | + | + |
| Moksha | [S15] |  |  | + | Lithuania_BA | [S28] | + | + | + |
| Mongola | [S18] |  | + | + | Lithuania_CWC | [S28] | + | + | + |
| Mongolians | [S19] |  | + | + | Lithuania_HG | [S28] | + | + | + |
| Mordovians | [S15] | + | + | + | Levänluhta | [S25] | + | + | + |
| Moroccans | [S23] |  |  | + | Maitan_MLBA_Alakul | [S17] | + | + | + |
| Mozabites | [S18] |  |  | + | Mal'ta | [S16] |  |  | + |
| Nenets | [S22] |  | + | + | Natufian | [S16] | + | + | + |
| Nganasans | [S19] |  | + | + | NLithuania | [S21] | + | + | + |
| Nivkhs | [S27] |  | + | + | Northern_LNBA | [S16] | + | + | + |
| Nogais | [S15] |  | + | + | Okunevo_EMBA | [S26] | + | + | + |
| North_Italians | [S18] | + | + | + | Oy_Dzhaylau_MLBA | [S17] | + | + | + |
| North_Kannadi | [S19] |  |  | + | Petrovka | [S17] | + | + | + |
| North_Ossetians | [S15] | + | + | + | Poltavka | [S16,S17] | + | + | + |
| northwest_Mingrelians | [S23] | + | + | + | Poprad | [S21] | + | + | + |
| Orcadians | [S18] | + | + | + | Potapovka | [S16,S17] | + | + | + |
| Oroqens | [S18] |  | + | + | Remedello | [S16] | + | + | + |
| Palestinians | [S18] | + | + | + | Romania_HG | [S31] | + | + | + |
| Paniya | [S23] |  |  | + | Russia_EBA | [S16] | + | + | + |
| Pathan | [S18] |  |  | + | SaltovoMayaki | [S21] | + | + | + |
| Poles | [S33] | + | + | + | Samara_Eneolithic | [S16] | + | + | + |
| Portugese | [S29] | + | + | + | Sarmatian | [S21] | + | + | + |
| Romanians | [S23] | + | + | + | Sarmatian_SU | [S34] | + | + | + |
| Russians_Central | [S22,S23] | + | + | + | Scythian_East | [S34] | + | + | + |
| Russians_North | [S18,S22-S24] | + | + | + | Scythian_IA | [S16] | + | + | + |
| Russians_South | [S22] | + | + | + | SHG | [S16,S28] | + | + | + |
| Saami_Kola | [S29] |  | + | + | SidelkinoEHG_ML | [S26] | + | + | + |
| Saami_SWE | [S29] |  | + | + | Sintashta | [S16,S17] | + | + | + |
| Sakilli | [S23] |  |  | + | Srubnaya | [S16,S17] | + | + | + |
| Sardinians | [S18] | + | + | + | Sweden_LNBA | [S28] | + | + | + |
| Saudis | [S23] |  |  | + | Sweden_TRB | [S28] | + | + | + |
| Selkups | [S19,S20] |  | + | + | Tagar | [S21] | + | + | + |
| Shors | [S27] |  | + | + | Trypillia | [S31] | + | + | + |
| Sindhi | [S18] |  |  | + | Ukraine_Eneolithic | [S31] | + | + | + |
| Slovaks | [S24] | + | + | + | Ukraine_Mesolithic | [S31,S32] | + | + | + |
| Spaniards | [S23] | + | + | + | Ukraine_Neolithic | [S31,S32] | + | + | + |
| Swedes | [S29,S33] | + | + | + | Ust-Ishim | [S16] |  |  | + |
| Syrians | [S23] | + | + | + | West_Siberia_N | [S17] | + | + | + |
| Tajiks | [S15] |  | + | + | WHG | [S16,S31] | + | + | + |
| Tatars | [S15,S29] |  | + | + | Wusun | [S21] | + | + | + |
| Tu | [S18] |  | + | + | Yamnaya | [S26] | + | + | + |
| Turkmens | [S15] |  | + | + | Yamnaya_Bulgaria | [S31] | + | + | + |
| Turks | [S23] | + | + | + | Yamnaya_Kalmykia | [S16] | + | + | + |
| Tuscans | [S18] | + | + | + | Yamnaya_Samara | [S16] | + | + | + |
| Tuvinians | [S19,S22] |  | + | + | Yamnaya_Ukraine | [S31] | + | + | + |
| Udmurts | [S22] |  | + | + | Zevakinskiy_LBA | [S17] | + | + | + |
| Ukrainians | [S15] | + | + | + |  |  |  |  |  |
| Uygurs | [S18] |  | + | + |  |  |  |  |  |
| Uzbeks | [S20,S23] |  | + | + |  |  |  |  |  |
| Vepsas | [S15,S29] | + | + | + |  |  |  |  |  |
| Welsh | [S29] | + | + | + |  |  |  |  |  |


| Xibo | $[S 18]$ | + | + |
| :--- | :---: | :---: | :---: |
| Yakuts | $[S 18, S 27]$ | + | + |
| Yemenese | $[S 23]$ |  | + |
| Yorubas | $[S 18]$ | + | + |
| Yukaghirs | $[S 19]$ | + | + |

Table S3. Modern and ancient comparison populations for principal component (PCA), outgroup f3 and D statistics' (f3, D), ADMIXTURE (ADM) and ChromoPainter (CP) analyses. Related to Figures 1, 2, 3, S1 and S2.

## Supplemental References

S1. Oras, E., Lang, V., Rannamäe, E., Varul, L., Konsa, M., Limbo-Simovart, J., Vedru, G., Laneman, M., Malve, M., and Price, T.D. (2016). Tracing prehistoric migration: isotope analysis of Bronze and Pre-Roman Iron Age coastal burials in Estonia. Est. J. Archaeol. 20: 1, 3-32.

S2. Laneman, M., and Lang, V. (2013). New radiocarbon dates for two stone-cist graves at Muuksi, northern Estonia. Est. J. Archaeol. 17: 2, 89-122.

S3. Friedenthal, A. (1932). Ein Beitrag zur vorgeschichtlichen Anthropologie Estlands. Z. Für Ethnol. 63, 1-42.

S4. Kalman, J. (1998). Skeletal report. In Aruanne kivikirstkalmete kaevamistest Muuksi Hundikangrutes 1996-1997, G. Vedru. (Manuscript in the archive of the archaeology department of the University of Tartu). (Tartu).

S5. Kalman, J. (1999). Human remains from the stone-cist graves of Rebala Lastekangrud, North Estonia. Est. J. Archaeol. 3: 1, 19-34.

S6. Laneman, M., Lang, V., Malve, M., and Rannamäe, E. (2015). New data on Jaani stone graves at Väo, Northern Estonia. Est. J. Archaeol. 19: 2, 110-137.

S7. Laneman, M., Lang, V., and Saage, R. (2016). Burial site hidden in a clearance cairn at Alu, Raplamaa. Archaeol. Fieldwork Est. 2015, 35-46.

S8. Kalman, J. (2000). Skeletal analysis of the graves of Kaseküla, Poanse I and Poanse II. In Töid ajaloo alalt II. (Tallinn: Eesti Ajaloomuuseum), pp. 17-40.

S9. Kalman, J. (2000). Stone grave II of Tõugu - skeletal report. In Keskusest ääremaaks. Viljelusmajandusliku asustuse kujunemine ja areng Vihasoo-Palmse piirkonnas Virumaal, V. Lang. Muinasaja teadus, 7. (Tallinn), pp. 387-407.

S10. Lang, V. (2000). Keskusest ääremaaks. Viljelusmajandusliku asustuse kujunemine ja areng Vihasoo-Palmse piirkonnas Virumaal. Muinasaja teadus, 7. (Tallinn).

S11. Kalman, J. (2000). Tandemägi stone grave - osteological report. In Keskusest ääremaaks. Viljelusmajandusliku asustuse kujunemine ja areng Vihasoo-Palmse piirkonnas Virumaal, V. Lang. Muinasaja teadus, 7. (Tallinn), pp. 423-436.

S12. Yushkova, M.A., and Kulešov, V.S. (2011). Kyorstovo 1: a new burial ground of the period of Roman influences in North-Western Russia. Archaeol. Litu. 12, 99-121.

S13. Shirobokov, I.G., and Yushkova, M.A. (2014). Антропологические материалы из коллективных захоронений по обряду кремации и ингумации каменного могильника с оградками Малли (по результатам раскопок 2010 г.). (Anthropological materials from collective burials according to the rite of cremation and inhumation of a stone fence burial ground in Malli (according to the results of excavations in 2010)). Bull. Archeol. Anthropol. Ethnogr. 2, 71-79. (In Russian).

S14. Mikhaylova, E.R. (2016). The population of the south-eastern coast of the Gulf of Finland and its contacts with the regions of the Baltic Sea in the 1st millennium AD. Archaeol. Balt. 23, 181-198.

S15. Yunusbayev, B., Metspalu, M., Järve, M., Kutuev, I., Rootsi, S., Metspalu, E., Behar, D.M., Varendi, K., Sahakyan, H., Khusainova, R., et al. (2012). The Caucasus as an asymmetric semipermeable barrier to ancient human migrations. Mol. Biol. Evol. 29, 359-365.

S16. Lazaridis, I., Nadel, D., Rollefson, G., Merrett, D.C., Rohland, N., Mallick, S., Fernandes, D., Novak, M., Gamarra, B., Sirak, K., et al. (2016). Genomic insights into the origin of farming in the ancient Near East. Nature 536, 419-424.

S17. Narasimhan, V.M., Patterson, N.J., Moorjani, P., Lazaridis, I., Mark, L., Mallick, S., Rohland, N., Bernardos, R., Kim, A.M., Nakatsuka, N., et al. (2018). The genomic formation of South and Central Asia. bioRxiv, 292581.

S18. Li, J.Z., Absher, D.M., Tang, H., Southwick, A.M., Casto, A.M., Ramachandran, S., Cann, H.M., Barsh, G.S., Feldman, M., Cavalli-Sforza, L.L., et al. (2008). Worldwide human relationships inferred from genome-wide patterns of variation. Science 319, 1100-1104.

S19. Rasmussen, M., Li, Y., Lindgreen, S., Pedersen, J.S., Albrechtsen, A., Moltke, I., Metspalu, M., Metspalu, E., Kivisild, T., Gupta, R., et al. (2010). Ancient human genome sequence of an extinct Palaeo-Eskimo. Nature 463, 757-762.

S20. Raghavan, M., Skoglund, P., Graf, K.E., Metspalu, M., Albrechtsen, A., Moltke, I., Rasmussen, S., Stafford, T.W., Orlando, L., Metspalu, E., et al. (2014). Upper Palaeolithic Siberian genome reveals dual ancestry of Native Americans. Nature 505, 87-91.

S21. Damgaard, P. de B., Marchi, N., Rasmussen, S., Peyrot, M., Renaud, G., Korneliussen, T., Moreno-Mayar, J.V., Pedersen, M.W., Goldberg, A., Usmanova, E., et al. (2018). 137 ancient human genomes from across the Eurasian steppes. Nature 557, 369-374.

S22. Yunusbayev, B., Metspalu, M., Metspalu, E., Valeev, A., Litvinov, S., Valiev, R., Akhmetova, V., Balanovska, E., Balanovsky, O., Turdikulova, S., et al. (2015). The genetic legacy of the expansion of Turkic-speaking nomads across Eurasia. PLoS Genet. 11, e1005068.

S23. Behar, D.M., Yunusbayev, B., Metspalu, M., Metspalu, E., Rosset, S., Parik, J., Rootsi, S., Chaubey, G., Kutuev, I., Yudkovsky, G., et al. (2010). The genome-wide structure of the Jewish people. Nature 466, 238-242.

S24. Kushniarevich, A., Utevska, O., Chuhryaeva, M., Agdzhoyan, A., Dibirova, K., Uktveryte, I., Möls, M., Mulahasanovic, L., Pshenichnov, A., Frolova, S., et al. (2015). Genetic heritage of the Balto-Slavic speaking populations: a synthesis of autosomal, mitochondrial and Y-chromosomal data. PloS One 10, e0135820.

S25. Lamnidis, T.C., Majander, K., Jeong, C., Salmela, E., Wessman, A., Moiseyev, V., Khartanovich, V., Balanovsky, O., Ongyerth, M., Weihmann, A., et al. (2018). Ancient Fennoscandian genomes reveal origin and spread of Siberian ancestry in Europe. Nat. Commun. 9, 5018.

S26. de Barros Damgaard, P., Martiniano, R., Kamm, J., Moreno-Mayar, J.V., Kroonen, G., Peyrot, M., Barjamovic, G., Rasmussen, S., Zacho, C., Baimukhanov, N., et al. (2018). The first horse herders and the impact of early Bronze Age steppe expansions into Asia. Science 360.

S27. Fedorova, S.A., Reidla, M., Metspalu, E., Metspalu, M., Rootsi, S., Tambets, K., Trofimova, N., Zhadanov, S.I., Hooshiar Kashani, B., Olivieri, A., et al. (2013). Autosomal and uniparental portraits of the native populations of Sakha (Yakutia): implications for the peopling of Northeast Eurasia. BMC Evol. Biol. 13, 127.

S28. Mittnik, A., Wang, C.-C., Pfrengle, S., Daubaras, M., Zariṇa, G., Hallgren, F., Allmäe, R., Khartanovich, V., Moiseyev, V., Tõrv, M., et al. (2018). The genetic prehistory of the Baltic Sea region. Nat. Commun. 9, 442.

S29. Tambets, K., Yunusbayev, B., Hudjashov, G., Ilumäe, A.-M., Rootsi, S., Honkola, T., Vesakoski, O., Atkinson, Q., Skoglund, P., Kushniarevich, A., et al. (2018). Genes reveal traces of common recent demographic history for most of the Uralic-speaking populations. Genome Biol. 19, 139.

S30. Saag, L., Varul, L., Scheib, C.L., Stenderup, J., Allentoft, M.E., Saag, L., Pagani, L., Reidla, M., Tambets, K., Metspalu, E., et al. (2017). Extensive farming in Estonia started through a sex-biased migration from the Steppe. Curr. Biol. CB 27, 2185-2193.e6.

S31. Mathieson, I., Alpaslan-Roodenberg, S., Posth, C., Szécsényi-Nagy, A., Rohland, N., Mallick, S., Olalde, I., Broomandkhoshbacht, N., Candilio, F., Cheronet, O., et al. (2018). The genomic history of southeastern Europe. Nature 555, 197-203.

S32. Jones, E.R., Zarina, G., Moiseyev, V., Lightfoot, E., Nigst, P.R., Manica, A., Pinhasi, R., and Bradley, D.G. (2017). The Neolithic transition in the Baltic was not driven by admixture with early European farmers. Curr. Biol. CB 27, 576-582.

S33. Behar, D.M., Metspalu, M., Baran, Y., Kopelman, N.M., Yunusbayev, B., Gladstein, A., Tzur, S., Sahakyan, H., Bahmanimehr, A., Yepiskoposyan, L., et al. (2013). No evidence from genome-wide data of a Khazar origin for the Ashkenazi Jews. Hum. Biol. 85, 859-900.

S34. Unterländer, M., Palstra, F., Lazaridis, I., Pilipenko, A., Hofmanová, Z., Groß, M., Sell, C., Blöcher, J., Kirsanow, K., Rohland, N., et al. (2017). Ancestry and demography and descendants of Iron Age nomads of the Eurasian Steppe. Nat. Commun. 8, 14615.


[^0]:    See also Figure S3, Tables S1 and S2, and Data S2. <0.017, not included in autosomal analyses; Av. cov., average genomic coverage; BA, Bronze Age;
    EST, Estonia; F, female; Gen., genetic; IA, Iron Age; M, male; MA, Middle Ages; Morph., morphological; MThg, mtDNA haplogroup; RUS, Russia; Y hg,
    Y chromosome haplogroup.
    ${ }^{\text {a14 }} \mathrm{C}$ date; calibrated using OxCal v4.2.4 [27] and IntCal13 atmospheric curve [28]
    ${ }^{\text {b }}$ Typo-chronological date
    ${ }^{\mathrm{c}}$ Combined ${ }^{14} \mathrm{C}$ date of multiple dates using OxCal v4.2.4 [27] R_combine; calibrated using OxCal v4.2.4 [27] and IntCal13 atmospheric curve [28]

