Exceptional ancient DNA preservation and fibre remains of a Sasanian saltmine sheep mummy in Chehrābād, Iran

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Abstract

Mummified remains have long attracted interest as a potential source of ancient DNA. However, mummification is a rare process that requires an anhydrous environment to rapidly dehydrate and preserve tissue before complete decomposition occurs. We present the whole genome sequences (3.94X) of a ~1600 year old naturally mummified sheep recovered from Chehrābād, a salt mine in northwestern Iran. Comparative analyses of published ancient sequences revealed remarkable DNA integrity of this mummy. Hallmarks of postmortem damage, fragmentation and hydrolytic deamination, are substantially reduced, likely due to the high-salinity of this taphonomic environment. Metagenomic analyses reflect the profound influence of high salt content on decomposition; its microbial profile is predominated by halophilic archaea and bacteria, possibly contributing to the remarkable preservation of this sample. Applying population genomic analyses we find clustering of this sheep with Southwest Asian modern breeds, suggesting ancestry continuity. Genotyping of a locus influencing the woolly phenotype showed the presence of an ancestral “hairy” allele, consistent with hair fibre imaging. This, along with derived alleles associated with the fat-tail phenotype, provides genetic evidence that Sasanian-period Iranians maintained specialized sheep flocks for different uses, with the “hairy”, “fat-tailed”-genotyped sheep likely kept by the rural community of Chehrābād’s miners.

Introduction

In 1993, a remarkably preserved human body dating to the ~1700 years Before Present (BP) was discovered in the Douzlākh salt mine near Chehrābād village in the Zanjan Province of northwest Iran [1–3]. A total of 8 “Salt Men” have been identified at the mine [4,5], several retaining keratinous tissues such as skin, hair, and both endo- and exoparasites, despite dating to the Achaemenid (550-330 BCE, 2500-2280 BP) and Sasanian (224-651 CE, ~1700-1300 BP) periods. The mine, also known as Chehrābād, was active in various periods and its archaeological refilling layers represent an extraction history that ranged from the 6th century BCE to 20th century CE. In addition to the “Salt Men”, textiles, leather objects, and animal remains have been discovered [6,7], likely preserved by high salinity and low moisture content of the mine. Isotopic, genetic, and lipid analyses have been reported for this material [1], and studies have been carried out to characterize genomic DNA survival [8]. These human and animal remains are examples of natural mummification - the spontaneous desiccation of soft tissue by a dry environment that rapidly dehydrates soft tissue before decay begins [9].

Mummification has been suggested as a mechanism that may sufficiently preserve keratinised tissue for ancient DNA (aDNA) sequencing [9]. The effects of age-related damage in aDNA are well documented and include base misincorporation at strand overhangs, fragmentation and low endogenous content [10]. Both deamination and depurination, associated with postmortem transition error and DNA fragmentation, respectively, require water as a substrate [11]. Ancient DNA from Chehrābād, a highly saline, anhydrous environment, presents an
opportunity to investigate potential differences in nucleotide degradation resulting from this unusual taphonomic context.

In this study we sequenced DNA from the ~1600 year old (Sasanian period) mummified sheep leg 4305, recently discovered in a large mining gallery in the northwestern edge of the Douzlākh saltmine of Chehrābād by Iranian-German researchers during archaeological excavations (Figure 1A) [2]. The specimen was likely deposited during refilling activities in the 4-5th centuries CE after the gallery’s reopening in the Early Sasanian period (2nd-3rd centuries CE) and following its initial collapse between 405-380 BCE. The leg was possibly discarded during food preparation activities, as both sheep and goat were likely used as provisioning for Sasanian-period miners; equines may have been used as beasts of burden [12]. By this time, sheep were an established commodity for their meat and secondary products such as wool fibre, which was widespread by 4th millennium BCE and showed regional specialization by the 3rd millennium BCE [13].

We find unusual survival patterns of endogenous DNA given its distance from the equator, implying exceptional preservation of nucleic acid integrity was afforded by the unique salt-rich environment. This enables characterising the mummy skin metagenome and population genomic profiling of this sheep in the context of modern breeds. We also genotyped 51 SNPs within the first intron of the platelet derived growth factor D (PDGFD) that are highly differentiated between fat-tailed and thin-tailed breeds [14], as well as the antisense EIF2S2 retrogene insertion within the 3’ UTR of the IRF2BP2 gene which influences the woolly phenotype and is derived relative to the ancestral coarse “hairy” coat [15], in tandem with fibre analysis using scanning electron microscopy (SEM).

**Materials & Methods**

A sample of the mummified sheep skin (Table 1, MUM2) from sheep leg 4305 was directly radiocarbon dated at the 14CHRONO Centre (Queen’s University Belfast). OxCal 4.3.2 [16] was used to calibrate its age (95.4% confidence interval) using [17].

Sample preparation, extraction and library preparation were performed in a dedicated aDNA laboratory in the Smurfit Institute of Genetics, Trinity College Dublin according to standard protocols (supplementary material). Sequencing of MUM2 and two Iranian sheep bone samples (Khor1 and Azer2) of approximately similar ages (Table 1) for comparison was performed on Illumina MiSeq (50bp SE) and HiSeq 2500 platforms (100bp SE and 100bp PE).

Sequencing reads were aligned to OviAri3.1 and filtered to produce bam files following standard aDNA sequencing pipelines (supplementary material). Damage patterns were assessed using mapDamage2.0 [18].
Filtered reads not aligned to either sheep or human genomes were taxonomically assigned using the metagenomic classifier Kraken2 [19]. Microbial sources were estimated using SourceTracker2 [20] with a custom metagenomic database [21–25] (supplementary material). Bacterial species abundances were generated using MIDAS [26].

Mitochondrial sequences were produced using ANGSD [27] and a Maximum-Likelihood phylogenetic tree was generated using SeaView and phyML [28–30] with the HKY85 substitution model, selected using jmodeltest2 [31,32] and 100 bootstrap repeats.

A SNP dataset of modern breeds [33] was used to investigate genomic affinities (supplementary material). LASER (v2.03) PCA [34], outgroup f₃ statistics [35], TreeMix [36] and ADMIXTURE [37] analyses were completed (supplementary material; Supplementary Table 5).

We investigated the woolly locus located on chromosome 25 [15]. Two modified OviAri3.1 assemblies were produced, one representing the ancestral “hairy” phenotype, the other representing the “woolly” phenotype (supplementary material). Final bam files were visualised using IGV [38]. Hair fibres were examined using scanning electron and light microscopes at USTEM, TU Wien and Austrian Archaeological Institute respectively. We assessed 51 SNPs in the PDGFD gene associated with the derived fat-tail phenotype [14], using the genotype calls of modern fat and thin-tailed breeds to define the derived allele [39] [40]. As the average genome coverage was too low for accurate diploid genotype calls, we report base calls for both alleles.

**Results and Discussion**

The Chehrābād mummy sample (MUM2) was directly dated to the 5-6th century CE (2 sigma 1621-1481 cal BP, uncalibrated 1600 ± 30 BP, Supplementary Figure 3). This aligns with the Sasanian Empire period of Iran, a time when the mine was in active use [1]. Initial DNA screening indicated high endogenous DNA for MUM2, and also the comparative Iranian sheep samples from relatively close time periods (Table 1).

| Name  | Tissue      | Origin            | Period                    | Age              | Endogenous DNA % | Coverage (X) |
|-------|-------------|-------------------|--------------------------|------------------|------------------|--------------|
| MUM2  | Mummified skin | Chehrābād, Iran  | Sasanian Empire period   | *399-539 cal CE  | 31.01            | 3.94         |
| Khor1 | Petrous bone | Nishapur, Iran    | Sasanian - Islamic periods | 600 - 1200 CE    | 58.44            | 0.04         |
| Azer2 | Petrous bone | Tepe Hasanlu, Iran| Iron Age III            | 800-600 BCE      | 31.32            | 0.07         |

*Table 1. Summary information of samples sequenced in this study. * = directly dated.*
Sequencing of the Chehrābād mummy produced a 3.94X genome after quality filtering (Supplementary Table 1), in addition to the low coverage comparative genomes (0.04X and 0.07X). MUM2 differs from the two comparative sheep samples in displaying longer fragment lengths (median 107bp vs 52bp and 56bp; Figure 1B; collapsed reads-only 90bp vs 50bp and 55bp) and substantially lower rates of deamination (Figure 1D) (δS, single strand cytosine deamination probability, mean δS = 0.012 vs. 0.382 and 0.334). Contrasting previously published ancient ovicaprid data from Southwest Asia and Europe (Supplementary Table 3), MUM2 falls outside the ranges of both median fragment length and mean δS values (Figure 1C and 1E), indicating remarkably low fragmentation and deamination of the Chehrābād sheep mummy genomic material given its latitude. Similar length distributions have been reported primarily from high latitude and permafrost environments [41–44]. A low level of thermal fluctuations may also contribute to DNA preservation [45], as comparable fragment lengths have been reported in a human sample from Wezmeh Cave, Iran [46].

Recent models of postmortem DNA fragmentation suggest rate-constant hydrolytic depurination over time [47], or age-independent, driven by environment-dependent biotic and abiotic factors [45]. The depurination rates of MUM2 are similar to the more-fragmented comparative samples (Supplementary Figure 4), implying that other processes in the Chehrābād environment underlie the lower fragmentation rates. The highly alkaline, cool and anhydrous conditions may have contributed to inhibition of cellular nucleases which would otherwise degrade and fragment endogenous DNA [9]. Postmortem DNA deamination via cytosine hydrolysis [48] is thought to be strongly correlated with age [49] and thermal age [45]. The substantially-lower rates of deamination observed in MUM2 is likely due to the scarcity of environmental free water, required for hydrolytic deamination. These results are consistent with Chehrābād providing a taphonomic environment conducive to genome preservation.

DNA preservation may also be influenced by its tissue-of-origin; for example, bone hydroxyapatite rather than keratin fractions are associated with smaller fragment size [50]. As hydrophobic keratinised tissue may provide resistance to environmental water [51], we compared MUM2 to published ancient skins genomes (Figure 1F) to determine if tissue providence was solely responsible for DNA preservation. The mean δS of MUM2 falls outside the range of other ancient skin genomes, including 20th century CE goat skins [52] and leather recovered from the Tyrolean Iceman [53]. While this does not discount keratinized tissue being specifically enriched with longer DNA fragments, the Chehrābād sheep mummy appears to be singular in its DNA integrity among published skin samples.
Figure 1 (A) Mummified sheep leg (4305) after cleaning. Photography: N. Tehrani (B) Read length distributions of MUM2, Khor1, and Azer2, calculated from PE data. MUM2 shows a reduced rate of fragmentation. The median read length of MUM2 (107 bp) exceeds the median read length of Khor1 and Azer2 (52 bp & 56 bp, respectively). (C) Median read lengths of 61 published ancient Ovicaprid samples [54]. The median read length of MUM2 (107 bp; 90 bp among collapsed reads only) exceeds the longest among published ovicaprid genomes (64 bp). (D) Deamination patterns of MUM2, Khor1, Azer2, and other ancient ovicaprims for non UDG-treated libraries. Low levels of base misincorporation at the 5’ ends of reads were observed for MUM2 compared to Khor1 and Azer2. (E) Mean δS of published 182 ancient bone samples [54] [45]. The mean δS of MUM2 (0.012) is singular in its low levels of deamination. (F) Comparison of mean δS of published ancient skins [55] [53] [52]. Lower damage rates are recorded compared to all samples, including some ~50 years old.

Given the distinctive geochemical composition of Chehrābād, we examined if its salt-rich environment was reflected in the metagenomic profile of MUM2. Taxonomic assignment and abundance estimation assigned 57.13% of classified reads to the halophilic Class of Archaea
*Halobacteria* (Supplementary Table 2). Similarly, SourceTracker2 predicted that 0.4725 - 0.7458 of the microbial community originated from a salt-rich environment (Table 2, Supplementary Figure 5). A complementary analysis using MIDAS identified 76 unique bacterial species in the mummified sheep (Supplementary Table 4). The most abundant species is the halophilic bacterium *Actinopolyspora halophila* 58532, accounting for ~29% of identified reads. This signal of a dominant halophilic microbial community is not replicated in comparison samples or controls (Table 2, supplementary material). Rapid colonization by saprophytic microbial communities, with key decomposers being ubiquitous across soil types, is typical for mammalian corpses post-mortem [56]. The halophilic metagenome profile observed in the Chehrābād sheep mummy skin indicates that the typical decomposers may be less abundant in this alkaline, salt-rich setting, which may have contributed to soft tissue and molecular preservation.

| Source Type          | MUM2 (Species) | Azer2 (Species) | Khor1 (Species) | MUM2 (Genus) | Azer2 (Genus) | Khor1 (Genus) |
|----------------------|----------------|----------------|----------------|--------------|--------------|--------------|
| Tissue decomposers   | 0.0212         | 0.1458         | 0.0386         | 0.0426       | 0.3524       | 0.3727       |
| Salt-rich            | 0.4725         | 0.0026         | 0.0036         | 0.7458       | 0.0145       | 0.0151       |
| Laboratory reagents  | 0.0003         | 0.0002         | 0.0002         | N/A          | N/A          | N/A          |
| Sheep skin           | 0.0285         | 0.085          | 0.0463         | 0.0829       | 0.3294       | 0.2244       |
| Soil                 | 0.0009         | 0.0007         | 0.0046         | 0.0          | 0            | 0            |
| Unknown              | 0.4766         | 0.7657         | 0.9067         | 0.1287       | 0.3037       | 0.3878       |

Table 2: Predicted source proportion of metagenomic reads by SourceTracker2.

**Population genomics**

We investigated how the Chehrābād sheep MUM2 relates to modern populations using mitochondrial and autosomal variation. A 664X mitochondrial genome of MUM2 falls within the C haplotype cluster in a Maximum-Likelihood phylogeny of modern sheep mitochondria (Supplementary Figure 7). This clade is found at its highest frequency in southwest and east Asia [57,58], and has been reported in ancient samples from Bronze Age Turkey [59], and is consistent with past and present-day patterns of mitochondrial diversity.

PCA from autosomal variation clusters MUM2 with modern southwest Asian breeds, using both global and Asian reference panels (Supplementary Figure 8). f3 outgroup statistics show MUM2 shares the most genetic drift with southwest Asian breeds, particularly those from Iran (Figure 2A). ADMIXTURE and TreeMix analysis also confirmed the affinity of MUM2 with modern sheep breeds from southwest Asia (supplementary material). Overall, there is genetic continuity between west Iranian sheep populations in Sassanid and modern time periods,
although PCA using Ovine SNP50 genotypes of Asian breeds place MUM2 apart from sampled breeds (Supplementary Figure 9), suggesting a degree of genetic flux during the past 1500-1600 years in Iranian sheep. This is consistent with evidence for genetic exchange across Asia prior to the development of modern breeds [60–62].

**Fibre Genotype and Phenotype and Fat-tail Genotype**

The derived “woolly” coat phenotype is thought to be influenced by a ~1.5 kbp insertion of a *EIF2S2* retrogene into the *IRF2BP2* 3’ UTR, recessive to the ancestral allele associated with “hairy” coat [15]. We exploited the length of the MUM2 DNA fragments to investigate this “woolly” locus by searching for read pairs which either encompassed or overlapped the insertion breakpoint, indicative of a copy of the “hairy” allele. No reads were found to overlap the diagnostic insertion breakpoints of the “woolly” insert, which would indicate a copy of the “woolly” allele (Figure 2B). Five reads were found to uniquely map to the ‘hairy’ allele diagnostic position, with a further two read pairs inferred to overlap this breakpoint (Figure 2C). We therefore infer this animal to be either homozygous or heterozygous for the dominant “hairy” allele. In addition, SEM imaging of the mostly-unpigmented mummified hair fibres revealed mosaic scales typical of sheep [63] with fine lines on the scale surface (Figure 2D, supplementary material), a characteristic of sheep hair fibres and particularly for mouflon and medium-wool breeds [64]. This may reflect MUM2 coming from a herd maintained for meat or milk production rather than wool, consistent with suggestions that ovicaprids were used as food for workers, and that sections of the mine were used as stables [1].
We also find evidence of a fat-tail associated allele (48/51 SNPs) (Supplementary Table 6) at PDGFD, a gene likely controlling tail phenotype [65,66]. This observation, along with MUM2 sharing mitochondrial haplotype C with the majority of modern fat-tailed breeds [62], and the genomic affinity of MUM2 to modern fat-tailed breeds, although based on SNP-chip data, is intriguing. While we cannot determine the MUM2 tail phenotype directly, its genotype is similar to a medium wool or hairy-coated fat-tail breed [1]. Hairy-coated sheep may have lower mortality rates, have higher birth weights, and be more robust than woolly-coated [67], while fat-tail breeds are thought to be better adapted to arid environments [14]. If phenotypically-similar to these sheep breeds, the flock represented by MUM2 could have provided a reliable meat and fat source for Chehrābād’s miners. The faunal assemblage of the mine and paleoparasitological studies, although not very abundant, support the fact that sheep/goat were the most consumed animals by the miners [6,7,12,68].

Both woolly and fat-tailed sheep are depicted in the Early Bronze Age Mesopotamia but the spread of these phenotypes may have been uncoupled, and occurred via distinct processes [69,70]. Fat-tailed breeds were likely introduced from Southwest to East Asia in a period (700 BCE-1000 CE) broadly coinciding with the age of MUM2 [61]; the observed PDGFD genotype supports an ancient origin of this economically important trait. Wider aDNA analysis may elucidate when wool and fat-tailed associated genotypes arose and how they may have influenced sheep breed development, which have their origins in 4th millennium BCE Mesopotamia [[69,70]]. Although the archaeozoological assemblages in the Iranian Plateau from the Antiquity and later Medieval periods are still limited, the diversity of the size of sheep bones is already an indication of the diversification of breeds in these periods [68,71]. Our results are consistent with MUM2 deriving from a herd used for meat and/or milk rather than wool production, and reflect sophisticated Sasanian-period husbandry practices and specialised sheep breeding.

**Data Accessibility:** Sequencing reads and mitochondrial sequences are available under European Nucleotide Archive accession PRJEB43881.

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Authors' contribution: CR, KGD, and MM designed study; CR, KGD, VM, AH, and FM performed laboratory work; CR and MDT performed bioinformatic work; GRP was responsible for SEM analysis; HD, HL, ZL, RK, HF, AA, TS, and MM worked directly with and provided archaeological samples. All authors contributed to writing the manuscript, approve this study, and are accountable for all aspects of the work.

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Supplementary Material

Contextual Information for Archaeological Sites

Chehrābād, Iran

The mummified sheep leg 4305 (Supplementary Figure 1) was discovered during Iranian-German research work in one of the debris layers (feature 31091) in the northwestern edge of the mine during modern archaeological excavations of the Douzlākh saltmine of Chehrābād [2]. At the northwestern edge, part of a large mining gallery was discovered. This part had collapsed between 405 and 380 BCE but was reopened in the Early Sasanian period (2nd-3rd centuries CE). The working area in the northwestern edge was partly refilled most likely between the 4th and 5th century CE, of which the layer 31091 belongs. The dating of the sheep-leg (Table 1, Supplementary Figure 3) is concurrent to the layer dating (reference [2], 43-45, Table 11). It was most likely deposited in the course of refilling activities during a salt extraction phase when mining took place on the northwestern salt rock edge of the mine.

A skin cutting of sheep leg 4305, MUM2 (Supplementary Figure 2), was used for DNA extraction and C\textsubscript{14} date estimation.

Supplementary Figure 1: Sheep leg (4305) in the archaeological context of layer 31091 (left) and recent condition after cleaning (two angles, centre, right), Photos: DBM/RUB/AMZ, Thomas Stöllner (left), N. Tehrani (centre, right).
Supplementary Figure 2. Photograph of the mummified skin of MUM2. MUM2 was sampled from the sheep leg 4305.

Supplementary Figure 3: OxCal 4.3.2 [16] calibration of the radiocarbon age of MUM2 using IntCal13 [17].
Tepe Hasanlu, Western Azerbaijan, Iran

Tepe Hasanlu is one of the key sites of northwestern Iran, due to its long-term occupation and well-defined stratigraphy. It is located in the Solduz valley on the southern shore of Lake Urmia, Western Azerbaijan province of Iran (1043 m ASL, Latitude: 37°0'16.15"N, Longitude: 45°27'31.74"E). Robert H. Dyson Jr. directed ten seasons of excavations at Hasanlu from 1956 to 1977 [72,73]. The site was occupied during ten cultural periods from the Late Neolithic (period X) to the Ilkhanid dynasty (period I) [74]. The most represented periods in the site are the Late Bronze (period V) and Iron Age (period IV-III). The Iron Age II citadel of Hasanlu was destroyed and fired during a battle, probably by the Urartian army, around 800 BC [75,76]. Hasanlu period IIIc and b, are attributed to Iron Age III (Urartian period / 800-600 BCE) and period IIIa allocated to the Achaemenid Empire (550-530 BC). The citadel with fortification wall, a few massive public buildings like garrison quarters, flimsy structures, stables and other architectural remains had been discovered from period III [77]. Extensive excavations in these periods resulted in the discovery of burned buildings, thousands of artifacts in closed contexts, and a large quantity of human and animal bone remains. Animal bones are among the most abundant material recovered in the site. Despite the taxonomic diversity of the remains in Hasanlu, domestic sheep, goat, and cattle were the most exploited animals in all the periods and cattle contributed much less to the diet than sheep and goat during the Bronze Age and Iron I to II, while its contribution increased to as much as Caprini during Iron III and Historical periods [78,79]. The sheep sample from Tepe Hasanlu analysed in this study was a petrous bone recovered from deposits of level IIIb (Iron Age III) in campaign 1974, Op. X32, Stratum 5, Lot 26.

Azer2 Azer2 MM TH3

Nishapur, Khorasan, Iran

Nishapur, one of the major urban centers of Khorasan, is located in Northeast Iran on a plain limited by the Binalud heights in the North and the Kashmar heights in the South (1198 m ASL, Latitude: 36°10'16.72"N, Longitude: 58°50'50.82"E). Nishapur as a major social and economic urban area exists in a cultural landscape in which the economic life of rural and urban entities was based on agriculture, horticulture, and animal husbandry. Nishapur had been an important producing center of silk, wool, cotton and textiles and their export throughout the Islamic world and beyond [80]. Archaeological research in Nishapur began by the Metropolitan Museum of Art during the 1930s [81–85]. Since then, surveys and excavations have been realized by national and international missions. Excavations conducted by the Iranian-French team resulted in the discovery of a significant amount of animal remains in the citadel [86,87]. Recently, a new Iranian excavation that has been carried out in the citadel and the urban area also produced substantial bioarchaeological data [88]. 2470 pieces of animal bones from the first collection found in 2005-2006 were studied in the Archaeozoology section in the Bioarchaeology Laboratory of the Central laboratory of the University of Tehran [71]. The remains could be dated from the Sasanian period to the 12th
century CE. The results indicate that sheep, goats, and cattle were the most exploited animals for both meat and secondary products. The Nishapur petrous bone sample analyzed in the present study was discovered in the campaign 2006, Tr. 15B located in the north part of the citadel, Square: JXV, 6-12th century CE.

Khor1 Khor1 MM NKD1

Supplementary Methods

DNA extraction
Sample preparation, extraction and library preparation were performed in a dedicated ancient DNA laboratory in the Smurfit Institute of Genetics, Trinity College Dublin. A small section of skin (0.454g) was cut using a disposable blade. Prior to the extraction the sample was washed with 1ml of water (lab grade and UVed) then with 600ul of Ethanol (99%) and then again with 1 ml of water. Each time the supernatant was removed after centrifugation. DNA was extracted using a modified version of the classic phenol/chloroform DNA extraction protocol [89]. A modified extraction buffer was used, where 3mM CaCl2 and 30mM DTT were added to the original extraction buffer.

Library preparation and sequencing
Treatment of ancient DNA with Uracil-DNA glycosylase (UDG) and Endonuclease VIII has been shown to remove base misincorporations caused by deamination in aDNA [48]. For UDG-treated libraries, 5 μl USER® Enzyme (1,000U/ml; Uracil-Specific Excision Reagent, New England BioLabs Inc.) was added to 16.25μl purified DNA and incubated in an Eppendorf ThermoMixer for 3 hours at 37°C, prior to library preparation.

Both UDG-treated and not-treated libraries were constructed based on the protocol of [90] with modifications as reported in [91]. Control tubes (21.25μl H2O starting material) were carried out for library construction. Indexing Polymerase Chain Reactions (PCRs) were performed using Accuprime Pfx Supermix (Invitrogen), primer IS4 (10 μM) and a unique indexing primer (5 μM) as detailed in [90]). The concentration of amplified libraries was quantified with a TapeStation 2200 (Agilent).

Three rounds of sequencing were done: single-end and paired-end shotgun sequencing of USER-treated libraries on Illumina HiSeq 2500 (Macrogen), and single-end shotgun sequencing for non USER-treated libraries on Illumina MiSeq (TrinSeq). The UDG-treated library was sequenced on a HiSeq 2500 Illumina platform (100bp SE and 100bp PE) via a commercial sequencing company (Macrogen, Republic of Korea) while the non-UDG-treated library was sequenced on a MiSeq Illumina platform (50bp SE) at TrinSeq (Trinity College Dublin, Ireland) with Phi X control at 1%. 

Khor1 Khor1 MM NKD1
**Raw-read processing**

The quality of fastq files was assessed using FastQC [92]. For single-end libraries, adapters were trimmed from reads of raw fastq files using cutadapt 1.9.1 [93] (cutadapt -a AGATCGGAAGAGCACACGTCTGAACTCCAGTCAC -O 1 -m 30). This removed reads shorter than 30bp and trimmed adapter sequence if an overlap of more than 1 bp was found between the read and adapter sequence.

For paired-end libraries, adapters were trimmed using AdapterRemoval v2.1.1 [94] (AdapterRemoval --file1 reads_1.fq --file2 reads_2.fq --basename output_paired --trimns --trimqualities --minquality 25 --collapse). The --trimns --trimqualities --minquality flags cause stretches of consecutive Ns and/or bases of quality lower than 25 from the 5’ and 3’ termini to be trimmed. The --collapse flag merges mate pairs that overlap by at least 11 bp into a single read. Base quality scores of collapsed reads are then recalculated using the metadata of the paired reads. The collapsed fastq file and the truncated read pairs were used in downstream analysis.

**Read alignment**

All trimmed reads of samples were aligned to OviAri3.1 using BWA [95] 0.7.5a with relaxed parameters (-l 1024 -n 0.01 -o 2). These flags disable seeding, allow for more mismatches with the reference and allow 2 gap openings, respectively.

For single-end reads, the resulting sai files were converted to sam file format using BWA samse [95]. Paired-end reads were handled using BWA sampe [95] to collapse sai pairs into one sam file. Sam files were then converted into the binary bam files using SAMtools (v1.7) view [96]. The -F4 flag was used with SAMtools view to discard unmapped reads for all bam files. For paired-end reads, the -f2 flag was also used to remove pairs of reads where one paired read was aligned to a different chromosome. The bam files were then sorted using SAMtools sort and PCR optical duplicates were removed using SAMtools rmdup. Bam files were then filtered for reads of mapping quality less than 30 using SAMtools view. Read groups were added to each bam file using Picard AddOrReplaceReadGroups (https://broadinstitute.github.io/picard/). Finally, these filtered bam files were merged using SAMtools merge. Genome coverage was calculated using GATK DepthOfCoverage [97]. Alignment statistics are summarised in Supplementary Table 1.
### Supplementary Table 1: Sequencing and alignment statistics.

| Sample | Library type | Sequencing platform | Raw Reads | Trimmed Reads | Aligned Reads | MQ30, rmdup Aligned Reads | Endog. % |
|--------|--------------|---------------------|-----------|---------------|---------------|----------------------------|---------|
| MUM2   | Non-USER treated | Illumina MiSeq 63 SE | 6,371,514 | 6,317,966     | 2,479,430     | 1,959,646                 | 31.02   |
| MUM2   | USER treated  | Illumina HiSeq 100 SE | 63,389,209 | 62,636,422    | 19,172,915    | 15,576,961                | 24.87   |
| MUM2   | USER treated  | Illumina HiSeq 100 PE | 598,909,352 | 348,606,473  | 109,018,213   | 86,945,474                | 24.94   |
| Khor1  | Non-USER treated | Illumina MiSeq 65 SE | 333,116   | 319,697       | 263,477       | 186,846                   | 58.44   |
| Khor1  | USER treated  | Illumina HiSeq 100 PE | 6,230,996  | 3,146,933     | 2,796,988     | 1,934,890                 | 61.48   |
| Azer2  | Non-USER treated | Illumina MiSeq 65 SE | 1,430,246  | 1,379,462     | 645,402       | 432,134                   | 31.33   |
| Azer2  | USER treated  | Illumina HiSeq 100 BP | 20,448,674 | 10,835,702    | 5,029,767     | 3,074,445                 | 28.37   |

**Damage patterns**

Bam files were assessed using mapDamage2.0 [18]. mapDamage2.0 calculates read length distributions and substitution patterns at the 5’ and 3’ ends of read fragments of aligned bam files. mapDamage2.0 also outputs a number of damage parameters estimated in a Bayesian manner including δS, the cytosine deamination probability in single strand context. Published sequencing data of other ancient samples were retrieved from NCBI GenBank in order to contextualise the damage patterns of our sample with published data (Supplementary Table 3).

Depurination levels were inferred using frequencies of purines close to the 5’ end of fragment strand-breaks. The use of T4 DNA polymerase in our library preparation removes 3’ overhangs, therefore only 5’ ends are considered. Only non-UDG libraries were used to calculate these levels as USER excises uracil residues in DNA fragments, creating new strand breaks. Using Qualimap v2.1.3 [98], nucleotide frequencies were calculated on filtered bam files at each position of DNA fragments (Supplementary Figure 4). Following the methods detailed in [49], approximate depurination rates were calculated.
Supplementary Figure 4: Nucleotide frequencies at 5’ end of DNA fragments. The depurination rate was 6.3%, 5.6% and 5.3% for MUM2, Khor1, Azer2, respectively.

Metagenomics

For all metagenomic analysis, we first removed reads originating from the host organism and human contamination. To do this, all trimmed reads were aligned to the sheep genome (oviAri3.1) and the human genome (hg38) using methods detailed above. Unmapped reads were retained using SAMtools view -f 4 and converted to fastq files using SAMtools fastq. The retained reads were then deduplicated using PrinSeq [99]. In total 202,815,624 reads were retained for MUM2 (176,858,319 collapsed/single end reads and 25,957,305 paired end reads). This was repeated for Khor1 and Azer2, resulting in 281,649 and 4,315,523 reads, respectively.

Unmapped, deduplicated reads were searched against the Kraken2’s standard database using default settings [19]. This database includes sequences of bacterial, archaeal and viral genomes. Kraken2 classified 11.96% (24,247,293 reads) of the unmapped, deduplicated reads using this database. To calculate Class abundances, microbial abundances were recalculated using Bracken [100] est_abundance.py, with a minimum threshold of 0.1% of total classified reads. 57.13% (13,499,165) of the classified reads belonged to the salt-adapted *Halobacteria* Class which reflects the high salinity environment of Chehrābād.
Supplementary Table 2: Taxonomically assigned reads of MUM2 by Kraken2, recalculated by Bracken, at Class level. The salt-adapted Class Halobacteria predominates with 57.13% of assigned reads.

Using the same methods, Halobacteria were found in very low abundance in our comparative bone samples, Khor1 and Azer2 (0.7% and 0.67% of assigned reads, respectively with a threshold of 0.1%). As a comparison with other ancient skins, metagenomic analysis of the Tyrolean Iceman’s preserved clothes [53] and ancient parchment [55] also revealed very low levels of Halobacteria present (0-0.02% and 0-0.3% of assigned reads, respectively). The uniquely high levels of halophilic archaea present in MUM2 is suggestive of the large effect the saltmine environment had on the microbial communities.

We used SourceTracker2 to further define the metagenomic composition of MUM2. SourceTracker2 is a Bayesian source-prediction tool that can estimate the proportion of sequencing reads of samples (sinks) originating from selected metagenomes (sources) [20]. DNA sequences of various relevant metagenomes (salt-rich, live sheep skin, soil, tissue decomposers, and laboratory reagents) were downloaded from NCBI (Supplementary Table 4). The sequences were trimmed using AdapterRemoval v2.1.1 and taxonomic assignment for these reference sequences was done by Kraken2, as detailed above. The kraken reports were merged and converted to a BIOM table using kraken-biom (https://github.com/smdabdoub/kraken-biom) and the human taxid (9606) was removed.

Source prediction of MUM2, Azer2 and Khor1 was done at both Species and Genus level using default settings. We were unable to use the “Laboratory reagent” source at Genus level as there was an insufficient number of taxonomically assigned Genuses (27) for our rarefaction level (1,000). At Species level, SourceTracker2 estimates 0.4725 of the microbial community of MUM2 originates from a salt-rich environment, while only 0.003-0.007 of the comparative bone samples were estimated to be from this environment (Table 2;
Supplementary Figure 5A). This estimation of salt rich increases to 0.746 when estimated at Genus level (Table 2; Supplementary Figure 5B).

**Supplementary Figure 5: Prediction of metagenomic source proportions by SourceTracker2 at (A) Species level and (B) Genus level.**

In order to cross-validate our results, and to investigate individual bacterial species, MIDAS [26] was selected as it has been reported to be the amongst most accurate for aDNA microbial communities [101]. This tool quantifies bacterial species abundance from shotgun metagenomes. MIDAS clusters more than 31,000 bacterial reference genomes into nearly 6,000 species groups based on sequence identity of 30 universal marker genes.

We observe a large proportion of detected microbial species in MUM2 (8 of the 9 most abundant species) as being salt adapted or salt tolerant (Supplementary Table 4). Of these, just one species is identified in ancient sample Azer2. This species, *Streptomyces sp 59965*, is a generalist. Similarly, none of the defined salt adapted bacteria in MUM2 were detected in shotgun data of leather from the Tyrolean Iceman ([53], results not shown). Top MIDAS hits for ancient parchment shotgun data ([55], results not shown) largely comprised of bacteria associated with animal skins (e.g. *Erysipelothrix rhusiopathiae*, *Propionibacterium acnes*, *Staphylococcus equorum*, *Brevibacterium linens*). Some salt-tolerant species (Table 2) were found in low abundance (<2%) of several ancient parchment shotgun data except for *Saccharomonospora glauca 62535* which was found at 4% abundance in sample SAMEA104143134. Overall, these results support the observation that MUM2’s microbial profile reflects its unique taphonomic context.
To validate the authenticity of the most abundant bacterium detected, all sequencing data was aligned to GCF_00371785.1, an *Actinopolyspora halophila* reference genome (*Actinopolyspora halophila* DSM 43834, retrieved from NCBI GenBank). A 0.33X genome was generated (std per bp = 1.13X). Using Qualimap v2.1.3 [98], the coverage of this bam file was visualised. The coverage of the bam file was even, as is recommended as a validation in [102]. The median read length of the aligned reads was 74 bp.

**Controls**

Controls that were done throughout the sample preparation, extraction and library preparation were also screened for halophilic signatures. The number of reads taxonomically assigned by Kraken2 was low (643 - 4648 assignments) and in all the presence of the *Halobacteria* clade was low (0.009-0.01). Metagenomic source prediction was done at a Species level using the same SourceTracker2 dataset as above (Supplementary Figure 6). The salt-rich environment was not predicted to contribute highly to any of the controls (0.003-0.024). After “Unknown”, “Laboratory reagents” were predicted to be the highest contribution to all controls except the Library Control. Here, SourceTracker2 predicted 0.66 of its metagenome to belong to “Sheep skin”, suggesting possible skin contamination during library preparation.

![Supplementary Figure 6: Prediction of metagenomic source proportions of controls by SourceTracker2 at Species level.](image-url)
Mitochondrial tree
All UDG-treated sequencing data was aligned to the domestic sheep mitochondrial reference, AF010406.1 (retrieved from NCBI GenBank) using an identical method as to the nuclear genome alignment. Consensus fasta sequences were generated using ANGSD [27] (angsd -doFasta 2 -doCounts 1 -setMinDepth 3 -minQ 20 -minMapQ 30). With these flags, base calls must have a minimum quality of 20 and a minimum MAQ of 30 to be considered. In the event that there are more than one base at a given site, the most common base is chosen. For a site to be called a minimum depth of 3 is required.

Ten modern sheep whole genome sequences and one urial sheep whole genome sequence published in [103] were retrieved from NCBI GenBank. Multiple sequence alignments were performed with MUSCLE [28]. A maximum-likelihood phylogeny (100 bootstraps) was generated. The HKY85 substitution model was selected for tree-building using jmodeltest2 [31,32]. Alignments and maximum-likelihood phylogeny constructions were implemented in SeaView v. 5.0.1 [28–30].

Supplementary Figure 7: Whole mitochondrion maximum-likelihood phylogeny of domestic sheep (100 bootstraps). MUM2 falls within Clade C. All clades of the phylogenetic tree are well-supported by bootstrap values (>0.97).

SNP Calling
The ovine SNP50 HapMap dataset used for the analyses described was provided by the International Sheep Genomics Consortium and obtained from www.sheephapmap.org in agreement with the ISGC Terms of Access. This dataset is genotyped on the Illumina OvineSNP50 Genotyping BeadChip (Illumina), covering >54,000 SNPs evenly spaced over the sheep genome. 1503 genotyped individuals are included in this dataset, representing modern breeds from Africa, the Americas, Europe, SW Asia and Asia [33].The genomic
positions of the OvineSNP50 were updated for the OviAri3.1 genome build (https://doi.org/10.6084/m9.figshare.8424935.v2) and SNPs flipped to the forward strand. Using PLINK v1.9 [104], these SNPs were filtered down to 44,223 sites using a MAF filter of ≤ 0.5.

The coverage of our sample was too low (approx. 4X) for accurate diploid genotype calling. To circumvent this issue, the pileup tool in GATK [97] v3.7 was used to report sample base calls for each read at each position of the filtered SNP list (44,223 SNP positions). A minimum base quality of 30 was required to be considered and sites with three or more different bases present were removed. A single base call was then chosen at random from each site and duplicated to create a pseudohaploid homozygous genotype at that position for each individual. In total 43,027 SNPs were called for MUM2.

Using PLINK v1.90, the pseudohaploidised MUM2 genotype calls were merged with the modern sheep breeds dataset. In cases where the allele of MUM2 matched neither of the alleles given in the sheep breed position, the SNP position was flipped using PLINK v1.90 and attempted to be merged again. Any remaining triallelic SNPs were removed. The final merged dataset was used for f3 statistics, ADMIXTURE and TreeMix analysis.

**PCA**

Projection PCA using Procrustes analysis was performed using LASER v2.03 [34]. The PCA reference space and projection transformation were constructed using a subsample of our modern breeds dataset. This subsample contained representation from all modern breeds, chosen randomly. Using the OvineSNP50 reference panel, pileup files were generated for MUM2, Khor1 and Azer2 using SAMtools mpileup (-q 30 -Q 20). Each sample was projected onto this PCA space and averaged across ten independent runs. The uncertainty based on the standard deviation * 2 for each sample was plotted on the map (Supplementary Figure 8).

The plot of PC1 vs PC2 shows that PC1 differentiates modern European breeds from breeds of the rest of the world. PC2 differentiates non-European breeds. A cline from African, Asian and South West Asian appears. These breeds do not create discrete clusters, however, and some individuals from different locations can be seen to cluster together in geographically-heterogeneous groups. MUM2 falls within the main cluster of SW Asian breeds. Khor1 and Azer2 both project within the diversity of SW Asian breeds but fall away from this main cluster on PC1. However we note that the uncertainty of PC1 of Khor1 and Azer2 is large, likely due to the lower SNPs called while the coordinates of MUM2 had a low uncertainty.
Supplementary Figure 8: PCA of global sheep affinity and Procrustes projection of MUM2, Khor1 and Azer2. Error bars represent the standard deviation of each PC coordinate between independent runs. PC1 separates breeds of European origin and the rest of the world, while PC2 separates Asian and African breeds into an approximate cline of Central/East Asian, SW Asian, African [33].

Local affinity
To investigate local affinity, a PCA was produced using only breeds of SW Asia and Asia using LASER v2.03. Here, PC1 differentiates Asian and SW Asian breeds. PC2 then separates one SW Asian (Turkish) breed from all other SW Asian breeds. When MUM2, Khor1 were projected onto this reference space using LASER, they clustered with SW Asian breeds rather than those from SE Asia (Supplementary Figure 9). Azer2 falls away from this cluster of SW Asian breeds, however, there is a larger amount of uncertainty again due to lower SNPs called.
Supplementary Figure 9: PCA containing only Asian breeds. Error bars represent the standard deviation of each PC coordinate between independent runs * 2. MUM2 clusters with Southwest Asian breeds. Some amount of genetic flux in local breeds in the past 1600 years is inferred by MUM2 falling slightly away from the main cluster of Southwest Asian breeds. Azer2 falls further away from the main cluster of SW Asian breeds, however, we note increased relative uncertainty of its PC1 position due to the lower number of called SNPs.

$f_3$ Statistics

$f_3$ statistics were computed to quantify the amount of genetic drift shared between MUM2 and modern sheep breeds. Three Asiatic mouflon individuals that were genotyped on the OvineSNP50 Genotyping BeadChip were used as an outgroup. Using PLINK v1.9, these were merged with our full dataset of modern breeds and MUM2. Using PLINK v1.9, a filter of 0 for missingness was applied to this merged dataset. The dataset was then filtered for linkage disequilibrium (LD) between markers, with the command --indep-pairwise 50 5 0.5. Outgroup $f_3$ values were not estimated for Khor1 and Azer2 due to the lower number of called SNPs.

Using this complete dataset, we calculated the $f_3$ statistics in the form of $f_3$(MUM2, modern breed; Asiatic Mouflon) with ADMIXTOOLS [35] (Supplementary Table 5). $f_3$ statistics were plotted using approximate geographic origins of breeds with R using the sf package [105] (Figure 2A).
TreeMix
The complete dataset used was converted into TreeMix format with the plink2treemix.py utility [36]. TreeMix [36] was used to construct a model of population splits with no migration events. The tree was rooted with Asiatic moufflon and blocks of 500 SNPs were selected (-k 500). No sample size correction was used (-noss). Clades emerge based on the geographic location of the sheep breeds. MUM2 falls in the SW Asian clade.

Supplementary Figure 10: TreeMix plot with 0 migration edges. MUM2 falls into the SW Asian breed clade. European breed labels were removed for clarity.

ADMIXTURE
The genotyped breed dataset was downsampling using PLINK v1.9 to include approximately 90 individuals from each Asia, Europe and Africa. Using PLINK v1.9, a filter of 0 for missingness was applied to this merged dataset. The dataset was then filtered for linkage disequilibrium (LD) between markers, with the command --indep-pairwise 50 5 0.5. Finally, transitions were removed using PLINK v1.9 and VCFtools [106]. This fully filtered dataset contained 31,261 sites.
Unsupervised ADMIXTURE was explored with a range of hypothetical ancestral populations, $K$s (2-20). ADMIXTURE runs were replicated three times for each $K$ and set a different random seed for each. The three replicated runs for each $K$ were averaged, and the cross-validation values which describe the approximate error of each ADMIXTURE run were inspected [37]. The run with the lowest cross-validation error was selected for presentation, $K=13$. In this ADMIXTURE analysis, the predicted ancestral profile of MUM2 most resembles Qezel, a sheep breed local to Iran.

**Supplementary Figure 11**: Average cross-validation values of unsupervised ADMIXTURE for each $K$, hypothetical ancestral population.
Supplementary Figure 12: ADMIXTURE plot with K=13 ancestral components. MUM2 displays a similar ancestral profile to modern Iranian sheep, represented by the Qezel breed.

Modified Genome Assembly
To investigate the woolly locus, two modified OviAri3.1 genome assemblies were produced. To exactly reproduce the locus detailed in [15], the sequence: ‘ACTTCTGTAATGGAATAGATAATTTAAAGTTATTATAACGATGTATTCTTTCAAAA AATAAAAACACCCTAGA’ was inserted at position chr25:7,452,231/7,452,232. This modified genome represents the woolly locus. To construct the ancestral locus (i.e. with no asEIFS insertion in chromosome 25), this assembly was edited to replace sequence chr25:7,450,861-7,452,231 with ‘TAAAAACACCCTAGA’, as per [15].

All UDG-treated sequencing data was aligned to these modified assemblies in an identical manner as above to produce two bam files. The coverage at the woolly locus was visualised using Integrative Genomics Viewer (IGV) [38], before and after a MAQ filter of 30 was applied. Since there are three copies of EIF2S2 genes annotated on the OviAri3.1 genome assembly (One genuine EIF2S2 gene located on chromosome 13 and two EIF2S2 pseudogenes located on chromosome 7 and chromosome 25), the mapping quality of reads aligned to the insertion are 0 [15].

The ‘breakpoints’ of the woolly locus were defined at positions chr25:7,450,874/5 and chr25:7,452,217/8 of the modified oviAri3.1 assembly. Any reads which straddled either of these sites were deemed evidence of a woolly allele. The ‘breakpoint’ of the ancestral locus was chr25:7,450,861/2. Similarly any read that straddled this point was deemed evidence of
the hairy/ancestral allele. Any read attributed to one genotype must only align to one modified assembly and not the other.

**Microscopy analysis**
Images produced by reflected light and transmitted light microscopy reveal a fat layer between the grain and corium, which is characteristic of domestic sheep skin [107,108] (Supplementary Figure 13). The fibres appear mostly unpigmented, with sporadic dark fibres visible (Supplementary Figure 14).

![Supplementary Figure 13: Cross section of the sheep skin. Fat is deposited between the corium layer and the grain. Images by G. Ruß-Popa.](image-url)
Supplementary Figure 14: Microscopy of fibres reveal a mostly unpigmented phenotype. Image by G. Ruß-Popa.

The leg of the sheep is dominated by coarse fibres (primary fibres) (Supplementary Figure 15). Scanning Electron Microscope (SEM) imagery of these hair fibres showed typical structure of sheep hair fibres, with a mosaic type scale and the fine lines on the scale’s surface (Supplementary Figure 16A). This is characteristic of sheep hair fibres, and particularly for mouflon and medium-wool breeds. Light microscopy of these coarse fibres revealed an amorphous medulla (Supplementary Figure 16B). Thin unmedullated fibres (underwool) were also detected (Supplementary Figure 17).

Overall, these results are consistent with MUM2 having a hairy/medium-wool phenotype, although results are not conclusive as the fibre types of the lower leg do not reflect the fleece type from the sheep body. Therefore, comparative analysis like wool measurements [109] of different sheep types are planned, in which samples from all parts of the body, including the lower leg, are to be examined.
Supplementary Figure 15: SEM image of different mummy hair fibres. Image by A. Steiger-Thirsfeld and G. Ruß-Popa.

Supplementary Figure 16A: SEM image of MUM2 hair fibre, displaying the typical mosaic scales of a sheep hair shaft and details of the fine lines on the scale surface. Image by A. Steiger-Thirsfeld and G. Ruß-Popa. 16B Light microscopy of the coarse hair fibre displaying the amorphous medulla. Image by G. Ruß-Popa.
**Supplementary Figure 17A:** Microscopy displaying the thin fibres in front. Image by G. Ruß-Popa. **17B:** Light microscopy revealing the unmedullated underwool. Image by G. Ruß-Popa.

SEM images were generated by G. Ruß-Popa and Andreas Steiger-Thirsfeld at USTEM (Universitäre Service Einrichtung für Transmissions-Elektronenmikroskopie), TU Vienna. FEI Quanta 250 FEGSEM with AFM for Structural investigations (EBSD) was used for SEM imagery. Microscopy was done by G. Ruß-Popa at the Austrian Academy of Sciences, Austrian Archaeological Institute, Archaeological Sciences, Hollandstraße 11-13, 1020 Vienna, Austria. Zeiss SteREO Discovery.V20 (Magnification: 1–150 X) was used for Stereomicroscope (reflected light) analysis. Zeiss Axio Scope.A1 Pol. (Magnification: 1 X – 2.5 X – 5 X – 10 X – 20 X – 40 X) was used for the polarized light microscope analysis for the thin section. Axiocam 305 color was the microscope camera.

**Genotyping the derived fat-tail allele**

The pileup tool in GATK [97] v3.7 was used to report sample base calls at 51 SNP sites defined that are highly differentiated between fat-tailed and thin-tailed breeds, and thought to underlie the fat-tail phenotype [14]. The derived fat-tail alleles were determined using genotypes of the 29 diverse breeds of fat and thin-tailed breeds. These breeds were genotyped as part of the International Sheep Genomics Consortium (ISGC). Genotypic data were retrieved from [39], with methods detailed therein. The average genome coverage of MUM2 (~4X) was too low for accurate diploid genotype calling at these sites. Instead, we report a simple presence/absence of alleles based on base calls (Supplementary Table 6).

14 sites contained both the ancestral and derived allele base calls. Of these sites, 11 are susceptible to misdiagnosis due to deamination-induced base transitions. However, we also note the very low postmortem deamination levels of MUM2. One site (ss1139270994,ss1215832245,ss1198779144) contained one base call of a third allele (G) but was otherwise homozygous for the derived fat-tail allele (T) with four base calls.

**Supplementary Table 3**
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