A detailed life history of a Pleistocene steppe bison (*Bison priscus*) skeleton unearthed in Arctic Alaska

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Abstract:

Detailed paleoecological evidence from Arctic Alaska’s past megafauna can help reconstruct paleoenvironmental conditions and can illustrate ecological adaptation to varying environments. We examined a rare, largely articulated and almost complete skeleton of a steppe bison (*Bison priscus*) recently unearthed in Northern Alaska. We used a multi-proxy paleoecological approach to reconstruct the past ecology of an individual representing a key ancient taxon. Radiocarbon dating of horn keratin revealed that the specimen has a finite radiocarbon age ~46,000 +/- 1000 cal yr BP, very close to the limit of radiocarbon dating. We also employed Bayesian age modeling of the mitochondrial genome, which estimated an age of ~33,000-87,000 cal yr BP. Our taphonomic investigations show that the bison was scavenged post-mortem and infested by blowflies before burial. Stable carbon and oxygen isotope (δ¹³C and δ¹⁵N) analyses of sequentially sampled horn keratin reveal a seasonal cycle; furthermore, high δ¹⁵N values during its first few years of life are consistent with patterns observed in modern bison that undertook dispersal. We compared sequential analyses of tooth enamel for strontium isotope ratios (⁸⁷Sr/⁸⁶Sr) to a spatial model of ⁸⁷Sr/⁸⁶Sr ratios providing evidence for dispersal across the landscape. Synthesis of the paleoecological findings indicates the specimen lived during interstadial conditions. Our multi-proxy, paleoecological approach, combining light and heavy isotope ratios along with genetic information, adds to the broader understanding of ancient bison ecology during the Late Pleistocene, indicating that ancient bison adopted different degrees of paleo-mobility according to the prevailing paleoecological conditions and climate.

Keywords:
Quaternary; Beringia, Paleoecology; Stable Isotopes; Strontium; Oxygen; Carbon; Nitrogen; Steppe Bison
1. Introduction:

1.1 Paleoecological context:

Northern Alaska is currently experiencing environmental changes as the result of global warming, which is occurring most rapidly at northern latitudes (Moon et al., 2019). These changes impact the mobility and ecologies of extant megafauna including caribou, moose, and muskoxen (Post and Forchhammer, 2008; Sharma et al., 2009). For example, migratory species are experiencing mismatch in timing of migration and peak resource availability (Post and Forchhammer, 2008), while caribou are losing habitat in the north due to warmer Arctic summers and winters (Sharma et al., 2009). Detailed paleoecological evidence from the remains of past megafauna from this region provides an opportunity to examine how past megafauna lived in this environment (Guthrie, 1989), and therefore help predict responses of living megafauna to present and projected environmental changes.

During the height of the last glaciation (~28-18 thousand years ago (kya)) (Clark et al., 2009), the North Slope of Alaska was part of an expansive land-mass known as Beringia. Sea levels were ~130 meters lower than today (Lambeck et al., 2014), exposing a shallow continental shelf between northeast Asia and North America known as the Bering Land Bridge (BLB). The BLB extended approximately from the Lena River, Russia, in the west and the Mackenzie River, Yukon, Canada, in the east (Elias and Crocker 2008) (Figure 1). The Beringian ecosystem was primarily that of a mammoth steppe, a graminoid-dominated ecosystem that supported a community of large herbivorous mammals, dominated by mammoths (*Mammutthus primigenius*), horses (*Equus* sp.), and steppe bison (*Bison priscus*) (Guthrie, 2001; Mann et al., 2013; Shapiro and Cooper, 2003; Zimov et al., 2012). The mammoth steppe supported large populations of these herbivores, many of which had larger body sizes than their descendants today at similar latitudes (Zimov et al., 2012). Bison, in particular, had larger body sizes and horns than present-day American bison (*Bison bison*) (Martin et al., 2018), and were present throughout most of Eurasia and North America in what has been termed “The Bison Belt” (Guthrie, 1989). Steppe bison first arrived in North America ~195-135 kya (Froese et al., 2017) and their population began to decline around 37 kya (Heintzman et al., 2016; Shapiro et al., 2004).
Modern bison ecology can provide an analog for inferring ancient bison behavior as well as the basis for comparative (paleo) biology and anatomy. Although modern plains bison (*Bison bison bison*) are often considered grassland grazing specialists (Bamforth, 1987), plains bison in northern habitats (Waggoner and Hinkes, 1986), wood bison (*Bison bison athabascae*) (Larter and Gates, 1991), European wisents (*Bison bonasus*) have been observed to regularly utilize browse in their diet (Kowalczyk et al., 2011). Evidence from macro and micro tooth-wear analysis indicates that steppe bison likely had a broader herbivorous diet and ecological niche that included browsing (Rivals et al., 2010, 2007; Saarinen et al., 2016). The long-distance (>100km) (Berger, 2004; Hanson, 2015; Plumb et al., 2009) migrations of American bison (*Bison bison*) across the American Great Plains were legendary and a key component of bison life history (Bamforth, 1987; Flores, 1991). However, isotopic (strontium) analyses of ancient (~18,500 \(^{14}\)C yr BP) bison (*Bison priscus*) from a site in Ukraine found no evidence of paleomobility (Julien et al., 2012). Analyses of ancient bison specimens can provide opportunities to flesh out the paleoecological life-history of a taxon that shaped Beringian ecosystems (Zimov et al., 2011).

Fortunately for paleoecologists, bones, teeth, and horns of bison are some of the most numerous fossil remains found in Alaska (Guthrie, 1970; Mann et al., 2013). On rare occasions, high sediment deposition rates along with freezing temperatures can result in preservation of virtually complete carcasses or skeletons of past Beringian fauna, revealing vivid paleoecological snap-shots of life in Beringia (Boeskorov et al., 2016; Kirillova et al., 2015; Van Geel et al., 2014; Zazula et al., 2017; Zazula et al., 2009). These rare and well-preserved glimpses of past megafaunal paleoecology can emerge from eroding river-banks (Mann et al., 2013; Zazula et al., 2009), during mining operations (Guthrie, 1968), and during construction activities (Zazula et al., 2017). In some instances, soft tissue such as hair, skin, organs, and stomach contents are preserved (Kirillova et al., 2015; Van Geel et al., 2014), as well as associated insects. These remains, along with bones and teeth, can retain chemical clues about an individual’s paleoecology. Isotopic analyses (Kirillova et al., 2015) and analyses of ancient DNA (aDNA) (Zazula et al., 2017) can reveal past mobility patterns and population interconnectivity, contributing to an understanding of past ecosystems, landscapes, and evolution (Froese et al., 2017; Haile et al., 2009; Marsolier-Kergoat et al., 2015; Shapiro and Cooper, 2003, Heintzman et al., 2016; Shapiro et al., 2004).
In this study, we conducted a multi-proxy study, combining isotopic and aDNA analyses, with supporting paleo-forensic analyses, to investigate the sex, age, paleoecology and life history of an exceptionally well-preserved and largely articulated steppe bison (*Bison priscus*) found on the North Slope of Alaska (Figure 1 a-c). We assess peri- and post-mortem events together with the taphonomic history of skeletal remains using a taphonomic analysis of the skeleton and of the plant and insect remains present. Our examination of the physical condition of the remains (bones and teeth) from the specimen provides clues about an individual’s age, sex, and appearance (Fuller, 1959). The taphonomy and geology associated with the specimen provides information about the context surrounding an organism’s death and a possible cause of death. We used insect (Elias et al., 2000) and plant (Bigelow et al., 2013) macrofossils associated with the bison’s remains to provide valuable paleoecological information. Our multi-proxy, paleoecological approach adds to the broader understanding of ancient bison ecology during the Late Pleistocene.

### 1.2 Stable Isotopes:

Isotopic analyses have become a popular tool in paleoecology for determining the ecological and life-history traits of ancient fauna (Bocherens, 2003; Britton et al., 2009). Tissues that form in discrete layers over a period of an individual’s life, such as tooth enamel, hair, or horn, can be subsampled to allow inference of inter- and intra-annual paleo-mobility and paleoecology using isotopic analyses of these sample types (Balasse et al., 2001; Britton et al., 2009; Stevens et al., 2011; Zazzo et al., 2012). Intra-tooth strontium isotope ratio ($^{87}\text{Sr}/^{86}\text{Sr}$) analysis has been developed as a useful tool to track the mobility of past animals and humans (Balasse, 2002; Britton, 2009; Britton et al., 2011; Hoppe et al., 1999; Hoppe, 2004; Julien et al., 2012; Koch et al., 1995; Pellegrini et al., 2008; Radloff et al., 2010; Viner et al., 2010; Widga et al., 2010). $^{87}\text{Sr}/^{86}\text{Sr}$ ratios can vary across landscapes depending in part on the age and rock type of the underlying geology (Bataille et al., 2014; Brennan et al., 2014). These landscape $^{87}\text{Sr}/^{86}\text{Sr}$ signatures can enter animals through their diet and drinking water, replacing calcium in tissues such as teeth and bones (Capo et al., 1998). The combination of $^{87}\text{Sr}/^{86}\text{Sr}$ ratios and oxygen isotopic ($\delta^{18}\text{O}$) values from analyses of animal tissues have proved to be a powerful predictor of geographic location (Britton et al., 2009; Gigleux et al., 2017; Knudson et al., 2009). The $\delta^{18}\text{O}$
values from a specimen can indicate location based on latitude and distance from the coast (Hoppe, 2006; Lachniet et al., 2016). The δ18O values from analyses of bison teeth have also been used to determine approximate local climate and seasonal temperature variation, because δ18O values in water can be closely related to temperature (Bernard et al., 2009; Hoppe et al., 2006; Scherler et al., 2014). Examining bison mobility has been one of the more common applications of this isotopic methodology, partly due to the abundance of bison remains in the archaeological and paleontological records (Britton et al., 2012; Julien et al., 2012; Widga et al., 2010).

Stable carbon and nitrogen isotope ratios (expressed as δ13C and δ15N values, respectively) from analyses of sub-fossils of animals can add a dietary dimension to a paleoecological reconstruction (Drucker et al., 2008; Stevens and Hedges, 2004). δ13C, along with δ18O, values can be generated from the analysis of inorganic carbon in bones and teeth (Koch et al., 1997). Bison horns, which grow throughout the life of a bison, are a carbon and nitrogen-rich keratin tissue that allows intra- and inter-annual paleoecological inferences. δ13C values can also be generated from analyses of organic carbon preserved as bone collagen and the horn keratin, these methods also produce δ15N values (Iacumin et al., 2001; Schoeninger and DeNiro, 1984). In order to interpret the δ13C and δ15N values of these analyses, potential sources of variability need to be evaluated. One common source of δ13C variation in the diet of herbivores is variation in the proportional contribution C4 vs. C3 plants. However, C4 plants are exceedingly rare or non-existent in the Arctic (Wooller et al., 2007). Furthermore, a lack of trees and even shrubs in Pleistocene Arctic Alaska, even during some warmer interglacials (Willerslev et al., 2014), also excludes “the canopy effect” caused by the concentration of CO2 in dense forest (Drucker et al., 2008) and differences between herbs and shrubs (Schwartz-Narbonne et al., 2019). Thus, the main source of variability in δ13C values we can expect in Arctic vegetation is between wetter and drier environments (Wooller et al., 2007).

There are several drivers of δ15N variation in animals, which can make it a challenging system to study although δ15N values can demonstrate important relationships. For example, δ15N values in ancient megafauna have been found to have a strong relationship with the amount of precipitation during the Pleistocene (Carlson et al., 2016; Drucker et al., 2003; Graham et al., 2016; Heaton et al., 1986; Rabanus-Wallace et al., 2017), which is likely due in part to major changes in soil ecology in response to climatic change (Hobbie and Hobbie, 2006; Stevens et al.,...
However, for higher resolution $\delta^{15}$N we must consider behavioral and physiological explanations for change found in serially analyzed tissues (Drucker et al., 2010). Behavioral explanations including migration or dispersal could explain some of the differences and $\delta^{15}$N values found in relation to aridity (Barbosa et al., 2009) and altitude (Männel et al., 2007). In northern regions, studies have found consistent difference in plants based on mycorrhizal relationships with the soil (Kristensen et al., 2011). However, these are not sufficient to account for the seasonal differences or major regional differences that would be required to explain some of the magnitude of variation in $\delta^{15}$N values we go on to document in this ancient bison and other modern bison in the region (Funck et al., 2020). Alternatively, physiological changes in $\delta^{15}$N values of animals can occur in animals adapted to extreme conditions. Notably, animals who undergo hibernation (Lee et al., 2012), lack sufficient calories or are fasting (Hobson et al., 1993; Hobson and Clark, 1992; Mekota et al., 2006; Voigt and Matt, 2004) as well as animals managing acute physical stress (Delgiudice et al., 2000; Fuller et al., 2005; Habran et al., 2010; Rode et al., 2016) can exhibit relatively large increases in $\delta^{15}$N values. The effect of dietary stress on $\delta^{15}$N values in wood bison was evident in individuals that had undergone nutritional stress during the winter in Northern Alaska (Funck et al., 2020). For this reason, we will focus our interpretation on nutritional stress as the primary factor of intra-tissue $\delta^{15}$N variation but acknowledge that other behavioral and ecological factors may be involved. From our review of the literature, we have noted that changes in $\delta^{15}$N values in herbivores from the mammoth steppe over time have largely not included the possibility that some of the variation could be driven by physiological stress and even starvation. To some degree this is likely obscured by the fact that a majority of the previous research has focused on $\delta^{15}$N values generated from analyses of bone collagen, which provides a more integrated and essentially a life-time measure. Our analyses of horn-sheaths are providing a new a more detailed temporal perspective, which could be translated to the huge abundance of archived bison horn sheaths available. Using a combination of isotopic approaches on different sequentially grown tissues can provide a multi-proxy perspective of an individual’s life-history and eventually put this in the context of larger herds.

1.3 Ancient DNA:
Ancient DNA (aDNA) from ancient bison specimens from Beringia has been used to monitor gene-flow across the BLB and through the ice-free corridor that connected eastern Beringia to the rest of the Americas (Froese et al., 2017; Heintzman et al., 2016). It has also been used to estimate past changes in effective population size (Lorenzen et al., 2011), which has been used to identify the decline of steppe bison since ~37 kya in Beringia (Shapiro et al., 2004). Analyses using aDNA can also add a further perspective on individual steppe bison specimens. They can provide information on an individual’s genomic sex and population affinity, and help constrain age estimates for specimens potentially outside the range or towards the limits of radiocarbon dating. For the latter analysis, one such approach uses Bayesian analysis of aDNA sequences from dated specimens that lived at different time periods to calibrate a molecular clock, and then use this calibration to estimate the age of the undatable individual (Shapiro et al., 2011). The mitochondrial genealogy of Pleistocene bison in North America has been relatively well sampled, both geographically and temporally (Froese et al., 2017; Heintzman et al., 2016; Shapiro et al., 2004), and is therefore likely to be well suited for estimating the age of ancient bison specimens.

2. Material and methods:

2.1 Specimen and study site:

Dr. Dan Mann and Dr. Pamela Grooves found a nearly complete, and partially articulated specimen of steppe bison (*Bison priscus*) eroding out of a bank of the Ikpikpuk River on the North Slope of Alaska (Figure 1 a-c) located at 69.71563 N, -154.863 W (WGS 84) (Figure 1). They excavated and transported to the University of Alaska Museum of the North (UAMN), where the University of Alaska Museum Earth Sciences collections (UAMES) assigned an identification number UAMES 29458. The North Slope of Alaska is delimited by the Brooks Range of mountains to the south and the Arctic Ocean to the north (Figure 1). The southern portion is predominantly low foothills that become a coastal plain at ~69° north. The Ikpikpuk River cuts through Quaternary-aged aeolian sand deposits, known as the Ikpikpuk Sand Sea (Carter, 1981). These deposits preserve an abundance of Quaternary vertebrate fossils that erode from banks of the Ikpikpuk River, though rarely in such excellent and complete condition (Mann
et al., 2015). The general geology of this region consists of undifferentiated alluvial and aeolian deposits (organic-rich silt, loess and sand) of Holocene to Pleistocene age. The specimen was discovered and excavated from the eroding banks of the Ikpikpuk River in these unconsolidated sediments. The vast majority of vertebrate remains recovered from here are single isolated elements found in reworked sediments while floating the river by small boat. Unfortunately, a quarry map is not available from the site at the time of collection, in part due to the unexpected nature of the find and constraints of rapidly conducting fieldwork in remote Alaska. The sediment and associated microfossils analyzed in this study were collected from matrix directly surrounding or within cavities of the skeleton.

2.2 Fossil Material:

Skeletal elements of specimen UAMES 29458 were removed from the encasing sediment, cleaned, and anatomically re-articulated (Figure 2a). Hair, preserved soft tissue, and associated sediment were stored by individual body element and kept frozen. We rearticulated the specimen at the UAMES (Figure 2 a, b) and examined it for signs of scavenging and other taphonomic processes, using methods outlined elsewhere (Binford, 1981; Domínguez-Rodrigo, 1999; Fisher, 1995; Haynes, 1980; Hudson, 1993; Lyman, 1994, 1987). We found a large number of invertebrate and plant macrofossils within the interstices of the skull, particularly the brain cavity. These were collected and identified to provide taphonomic and paleoenvironmental context. We removed larger macrofossils and examined them using a dissecting microscope. We wet sieved sediments at 150 µm and 250 µm. All diagnostic plant macrofossils were collected from sediments larger than 250 µm and all invertebrate materials were collected and archived.

2.3 Radiocarbon analysis:

Radiocarbon dating was conducted on different materials associated with specimen UAMES 29458. Previously the bone collagen from the cap of a spinous process, extracted without ultrafiltration, was analyzed at Beta Analytic (Miami, Florida), and found to be non-finite in age >43,500 radiocarbon years (Mann et al., 2013). However, collagen is porous and can be subject to contamination, so we selected additional materials for further analysis: keratin from the horn
sheath, along with a fly pupal case and a plant macrofossil (herbaceous stem) that were both recovered from inside the bison skull. W. M. Keck Carbon Cycle Accelerator Mass Spectrometry Laboratory analyzed the samples on a National Electrostatics Corporation (NEC 0.5MV 1.5SDH-2 AMS system). The horn sheath keratin was run alongside three non-finite-\(^{14}\)C aged bison horn sheath samples and the mean blank from these three was subtracted from sample results, with an assumed 30% uncertainty. All of the sample preparation backgrounds were also subtracted, based on measurements of a \(^{14}\)C-free wood standard, with an assumed 30% uncertainty.

2.4 Isotopic analyses:

We removed thirty-three intra-tooth subsamples of enamel from the right mandibular molars 1, 2, and 3 (M1, M2, M3) of UAMES 29458 (Supplemental Figure 1). We collected samples in a laminar flow hood at the Alaska Stable Isotope Facility (ASIF) at the University of Alaska Fairbanks (UAF). We removed the surface layer of the target region from each molar and subsequently used a Dremel diamond-coated circular saw to divide samples along each tooth parallel to the growth plane (Figure S1). Enamel samples were removed as chips, which has been found to reduce the potential for contamination (Diego Fernandez, University of Utah, ICPMS Facility 2018 personal communication).

The University of Utah, Department of Geology and Geophysics, ICPMS facility analyzed half of each sample for \(^{87}\)Sr/\(^{86}\)Sr ratios using a multi-collector inductively coupled plasma mass spectrometer (MC-ICPMS - ThermoFisher Scientific, High Resolution NEPTUNE, Bremen, Germany). Strontium isotope analyses followed previously published protocols (Glassburn et al., 2018; Nelson et al., 2018). Additionally, we sampled whole molars from 14 modern rodent museum specimens (UAM:Mamm: Microtus oeconomus, Microtus mirus, Dicrostonyx groenlandicus) from within a 250-km radius surrounding the UAMES 29458 locality (Figure 1a) to determine the bioavailable \(^{87}\)Sr/\(^{86}\)Sr ratios in the region (Bataille et al., 2020). This method has previously been used to determine local bioavailable \(^{87}\)Sr/\(^{86}\)Sr ratios for comparison to mobile individuals (Hoppe et al., 1999; Kootker et al., 2016; Radloff et al., 2010). The results of this investigation augment existing spatial models of variability in \(^{87}\)Sr/\(^{86}\)Sr ratios in Alaska (Bataille et al., 2014).
We chemically pre-treated the remaining half of each bison enamel sample to remove contaminants from gas exchange using the modified method from Pellegrini and Snoeck (2016, 2015) in order to produce $\delta^{18}O$ and $\delta^{13}C$ values. To compensate for the use of small (~1 × 2 mm) enamel chips rather than powder we increased the soaking time. We added one milliliter of 2% sodium hypochlorite (NaOCl) to each sample, shaken to mix, and soaked for 48 hours to remove organic particulates. We then rinsed samples with deionized water and soaked them in 1 M acetate buffered acetic acid (0.1 M) (CH$_3$CO$_2$H) for 48 hours to removed carbonate portion. Finally, we rinsed the samples three times with deionized water before freezing and then freeze-dried them for approximately 10 hours on a VirTiS benchtop Lyo-Centre lyophilizer to remove any moisture prior to isotopic analysis. We analyzed the carbonate fraction using a Thermo Scientific GasBench II carbonate analyzer attached to a Thermo Scientific DeltaV$^{\text{Plus}}$ Isotope Ratio Mass Spectrometer at ASIF following previously published protocols (Glassburn et al., 2018). Stable carbon and oxygen isotope ratios are reported in $\delta$ notation as parts per thousand (‰) relative to the international standard Vienna Pee Dee Belemnite (VPDB). We ran the samples with laboratory standards of calcium carbonate (Merck, Suprapur 99.95% Lot # B510959 313) every 10 samples to determine analytical precision for $\delta^{13}C$ values (0.3 ‰) and $\delta^{18}O$ values (0.2 ‰) (all errors are expressed as one standard deviations). The $\delta^{18}O$ and $\delta^{13}C$ values were initially determined relative to VPDB for oxygen and carbon, but the $\delta^{18}O$ values were subsequently converted to the Vienna Standard Mean Ocean Water (VSMOW) scale to allow comparison of the values to meteoric water values. The $\delta^{18}O$ values were converted to VSMOW using Formula 1 (below) (Verkouteren and Klinedinst, 2004) and then Formula 2 (below), a conversion developed by Hoppe et al. (2006) and Velivetskaya et al. (2016) to compare bison enamel to meteoric water $\delta^{18}O$ values:

Formula 1: VSMOW = 30.92+1.03092*VPDB

Formula 2: Enamel carbonate = 0.7(±0.12)*VSMOW- 30.06(±1.40)

We removed horn sheath keratin as a wedge perpendicular to the direction of the keratin growth layers (Figure S2). We selected the location to maximize the years of horn growth covered by the core. We then subsampled the core by peeling off thin layers between 0.1-0.3 mm
in thickness with a razor blade, for a total of \( n = 113 \) sub-samples. Sub-samples were analyzed to produce \( \delta^{15}\text{N} \) and \( \delta^{13}\text{C} \) values (vs. AIR for nitrogen and VPDB for carbon) using a Flash 2000 Organic Elemental Analyzer (EA) connected via a Conflo IV to an IRMS (DeltaV Plus) at ASIF. Internal reference checks using peptone (No. P-7750 meat-based protein. Sigma Chemical Company, Lot #76f-0300) were run every 10\(^{th}\) sample, and blanks were run every 20\(^{th}\) sample. Repeated measurements of standards provided the analytical precisions, which were \( \pm 0.1\%\) and \( \pm 0.6\%\) for \( \delta^{15}\text{N} \) and \( \delta^{13}\text{C} \) values, respectively.

### 2.5 aDNA analyses:

The first premolar (PM1) of specimen UAMES 29458 was sent to the specialized Paleogenomics Laboratory at the University of California Santa Cruz (UCSC) for aDNA analysis. We followed standard protocols as outlined in Froese et al. (2017) to extract and analyze ancient DNA unless stated otherwise. Briefly, we extracted aDNA using a silica column method (Dabney et al., 2013), and then converted extracts into an Illumina-compatible double-stranded DNA library (Meyer and Kircher, 2010). We enriched an aliquot of this DNA library for bison mitochondrial genomic fragments using a commercial MyBaits target capture kit (Arbor Biosciences, Ann Arbor, MI). We sequenced both the enriched and remaining unenriched library on separate runs of an Illumina MiSeq using v3 150 cycle chemistry75 cyc, with paired-end 75 bp reads. We then merged the paired-end read data, trimmed adapters, and removed short (<25 base pairs (bp) for the enriched library; <30 bp for the unenriched library) and low-complexity sequences (DUST cutoff: 7) using SeqPrep and PRINSEQ-lite v0.20.4 (Schmieder and Edwards, 2011).

We aligned the filtered reads from the unenriched library to a reference database that included the cow (\textit{Bos taurus}) genome (Genbank: Btau_4.6.1) and an American bison (\textit{Bison bison}) mitochondrial genome (Genbank: NC_012346), using the Burrows Wheeler Aligner (BWA; Li and Durbin, 2009) aln algorithm with the seed disabled (-l 1024). We filtered aligned reads by map quality score (minimum of 20) and removed duplicates using SAMtools v0.1.19 (Li and Durbin, 2009). We assessed aDNA damage patterns using mapDamage v2.0.9 (Jónsson et al., 2013). To infer the genomic sex of UAMES 29458, we followed the method of Heintzman et al. (2017), which compares the relative mapping frequency of the X chromosome to the
autosomes. A male is inferred if this ratio is 0.45-0.55, whereas the expectation for a female is 0.95-1.05.

We mapped the filtered reads from the enriched library to a steppe bison reference mitochondrial genome (Genbank: KX269138) using the multiple iterative assembler (MIA; Briggs et al. 2009) and BWA, as above. For aDNA damage assessment of the enriched data, we used the BWA mapping consensus results as the reference sequence for mapDamage. We calculated a mitochondrial genome consensus sequence from the MIA mapping results using the criteria outlined in Froese et al. (2017), and added this sequence to the full mitochondrial genome alignment of Froese et al. (2017), as modified by Zazula et al. (2017). This alignment consisted of four yak and 47 Siberian or North American bison mitochondrial sequences.

Before estimating the age of UAMES 29458 with Bayesian time-tree methods, we tested for temporal signal in the data set. Tip-dating methods are only valid if this signal is present, as these methods will usually converge on an estimate whether or not there is a temporal signal present (Firth et al., 2010). We used a linear regression of phylogenetic root-to-tip distance against the sampling date to test for temporal signal (following Murray et al., 2016). We estimated a neighbor-joining tree of the data set (excluding the yak and two radiocarbon non-finite bison sequences) using a K80 nucleotide substitution model with pairwise deletion, using the ape package in R (Paradis et al., 2004), in which the root was fit simultaneously with the regression, so as to minimize the residual mean squares, with the resulting root matching the root obtained by using the yak sequences as an outgroup. Analyses of the full alignment and a reduced alignment, with MS022 omitted (see below), returned positive correlations that are significantly different from random permutations over clusters of samples with similar dates (full: \( r = 0.50, p = 0.010 \); reduced: \( r = 0.44, p = 0.025 \); Figure S3). We defined similar-date clusters (n=34-35) as monophyletic clades that had the same date after rounding to the nearest thousand years. A Mantel test suggested that this clustering was sufficient to eliminate a correlation between genetic and temporal distances in the data (without clustering: \( p = 0.001 \), with clustering: \( p = 0.18 \)) (Murray et al., 2016), which can result in a false positive result. Overall, these analyses suggest the presence of temporal signal in the bison mitochondrial genome data set.

We then estimated a time-tree in BEAST (v1.10.4, Drummond et al., 2012) so as to estimate both the age of UAMES 29458 and its placement in the bison mitochondrial genome.
phylogeny. We used a strict molecular clock with either finite radiocarbon dates or stratigraphic data associated with the other sequences in the analysis as priors, following Froese et al. (2017) and Zazula et al. (2017). To estimate the age of UAMES 29458, we used a uniform prior distribution of between 30 kya BP (as this specimen is borderline radiocarbon finite; see results) and 195 kya BP (the earliest mitochondrial estimate for the arrival of bison in North America; Froese et al., 2017), following the method of Shapiro et al. (2011). To test the robustness of the oldest age estimates, we also ran analyses with the minimum age prior set at 0, 20, 40, or 50 kya BP. We ran analyses with each set of priors twice, each time with the Markov chain Monte Carlo (MCMC) chains run for 60 million iterations, sampling every 3,000 iterations, and discarding the first 10% as burn-in. Using Tracer v1.6, we observed that all parameters reached convergence with the exception of the tree likelihood, which swapped between two optima in all analyses. This was due to the shifting placement of sequence MS022 (KX269130; Froese et al., 2017). This sequence was therefore removed from the alignment and all analyses were re-run. Convergence was then observed for all parameters. For each prior and alignment result set, we summarized trees and calculated a maximum clade credibility tree. All ages are reported as 95% highest posterior density (HPD) credibility intervals.

3. Results:

3.1 Physical description:

Overall, we observed specimen UAMES 29458 to be relatively complete and well preserved (Figure 2). Based on the observed number of annual growth cycles visible in the horn sheath, we estimated the bison specimen to be minimally 12 years old, which is considered a mature individual (Fuller, 1959). The specimen’s horn span was 90 cm at the widest point and the comparative angle and length of the horns indicate that it was a male (Guthrie, 1966). Much of the hair had sloughed off the specimen and was found in the surrounding sediment. In some areas, hair and skin had dried to the bone and were still attached. Hair from the front and hind limbs was yellow ochre in color, as were the shorter undercoat hairs from the body. The longer hairs were a reddish-brown or dark brown and tail hairs were black to dark brown. Long hairs
found in the surrounding sediment, which likely came from the cape (hair on the shoulders), beard (long hair on the chin) or cap (long hair on the top of the head), were light to dark brown.

The skull was in excellent condition and the left and right mandibles were fused at the symphysis. The incisors were missing from a pre-burial break. There were no gnaw marks on the surrounding bone to indicate another animal had gnawed them off. Evidence of root etching was observed on the ventral surfaces of the mandible with vivianite deposits observed on the articular surface. Vivianite was identified visually, it is commonly found on fossils found in these environments, and is unique in its blue appearance. We observed that the cervical vertebra 2 (C2) through thoracic vertebra 7 (T7) had the dorsal portion of the spinous process gnawed (Figure 3b) and bitten off (Figure 3a). We also observed T4 and T5 had evidence of damage from a carnivore, including puncture marks (Figure 3 a). Lumbar vertebra 1 (L1) to the sacrum also had evidence of carnivore damage on the ventral surface of the vertebrae (Figure 3 b). The distal ends of four thoracic ribs (towards the anterior portion of the body) also had puncture marks and gnawing. There were several pre- and post-burial breaks, and several medial portions of the ribs were missing. One rib on the right side had a pathology consistent with a healed break (Figure 3c). The left side was missing two thoracic ribs. The costal cartilage had carnivore gnawing damage on two fragments and the manubrium was missing one section.

The right appendicular skeleton was complete and we observed no evidence of carnivore damage. In addition, there was soft tissue (keratinous hoof sheathes, leather-like flesh and connective tissue) and vivianite present. There were gnaw and puncture marks observed on the proximal end of the left humerus (lateral tuberosity) near its articulation with the scapula; the scapula was missing. There was root etching on the medial surface of the humerus where it would have rested on the ground after the specimen’s death and we observed soft tissue adhering to the bone. The left metacarpal, carpals, and proximal, medial, and distal phalanges were missing, which could be consistent with loss during taphonomic processes upon emerging from the riverbank because the bones would have been exposed first and eroded out of the bank. In addition to the lack of carnivore marks near the articulations, they do not possess much food value for a carnivore and would not have been a primary target. On the left femur there was a pre-burial break at the greater trochanter, but no obvious carnivore damage. The left patella, calcaneus, several left tarsals, one sesamoid, one proximal, medial and distal phalanx was
missing, which is also consistent with secondary loss upon the specimen eroding out of the riverbank.

### 3.2 Radiocarbon analysis:

We generated three new radiocarbon dates associated with specimen UAMES 29458 (Table 1). The background-subtracted fraction modern values for both the keratin and insect chitin are different from zero by ~6 standard deviations, which indicates the ages are finite. The age of the keratin from the bison horn sheath was 46,000 ± 1,100 ¹⁴C yr BP (46,962 cal yr BP using CALIB 7.1; Stuiver et al., 2019), which was very similar to the radiocarbon date produced from the keratinous fly puparium (Table 1, Figure 4) taken from inside the bison skull cavity (46,800 ± 1,200 ¹⁴C yr BP, just outside the range for calibration using CALIB 7.1 (Stuiver et al., 2019). The dated plant macrofossil found in the skull, along with the puparia, had a radiocarbon date of >49,900 radiocarbon years, which was outside the range for calibration (Table 1).

### 3.3 Macrofossils:

#### 3.3.1 Fossil plant remains

The contents of the neural canal of the spine and skull from specimen UAMES 29458 included a sand-rich matrix with abundant moss, wood, and we could identify other plant fragments to species in some instances (Figures 4 and 5). The plant macrofossil assemblage from inside the skull is consistent with a mesic tundra or a wet flood plain surrounded by a shrub tundra (Figure 5). The presence of *Daphnia* spp. ephippia indicate that the sediment was also, at least partially, aquatic in origin. Bryophytic material, likely from wet tundra and rivers, was abundant and well preserved in the skull cavity. Wood was very common and possibly came from the shrubs identified from seeds, leaves, and other materials in the skull cavity, including *Salix* spp. (willow), *Betula* spp. (birch), and *Andromeda polifolia* (bog rosemary) (Figure 4 n). These shrubs are typical of wet tundra and tend to be more abundant in more protected areas (Cody, 2000; Hulten, 1968), suggesting an interstadial environment. In addition, *Potentilla palustris* and *Juncus* spp. were also present and are typically associated with riparian habitats and bogs (Cody,
2000). In contrast, the remains of *Papaver* spp. (poppies) and *Polygonum bistorta* (Figure 4 l), which favor drier open slopes (Hulten, 1968), were also present in the bison skull cavity. The largest number of plant macrofossils from the skull cavity was the mustard and *Draba*-type seeds (Figures 4 m and 5), which are from a large and very diverse family making it difficult to use these specimens to characterize paleoecological conditions (Cody, 2000; Hulten, 1968). However, many of the species from this group are typical of open slopes, gravel and/or sandy river banks (Cody, 2000). In addition, there is a portion of grasses (Poaceae) (Figure 4 k) (and *Carex* Figure 4 i and j) that are typical of tundra or steppe ecosystems in the Arctic, and are consistent with an open landscape with no large shrubs (Hulten, 1968). The burial occurred in a time scale that prevented lots of skeletal disarticulation by scavengers but allowed some activities by insects in an energy regime allowed deposition of sand but not coarser sediment.

### 3.3.2 Fossil insect remains

Analysis of the skull cavity of UAMES 29458 revealed the presence of insect remains, including complete or incomplete blowfly puparia (*Calliphoridae*) (*n* = 15, UAMES 52319-52330). From their excellent state of preservation, a determination at the species level was possible and the taxon that colonized the bison at the time of its death was identified as the northern blowfly *Protophormia terraenovae* (Robineau-Desvoidy) (Figure 4 a-e). With the exception of 4 fragmentary specimens, all the puparia were complete and unhatched indicating that the life cycle was interrupted before adult emergence. An attempt to open two fossil puparia (*#29458c, #29458g*), in order to find a possible nymph and to observe its maturity stage, proved unsuccessful since the pupa had disintegrated inside.

Partial abdomen remains of a carabid beetle (*Pterostichinae*: cf. *Pterostichus* sp.) were also found inside the skull (UAMES 52334; Figure 4 f). Thirty-five *Pterostichus* species occur in Alaska (Bousquet et al., 2013). Some representatives of this genus (i.e. *Pterostichus costatus* Ménétriés), live in the damp peaty areas of lowland tundra regions (Lindroth, 1966). Other insect remains belonging to several distinct orders were also recovered from inside the skull: a complete wing of *Psylloidea* (Homoptera) (UAMES 52331; Figure 4 g); a cephalic capsule of a larval *Chironomidae* (Diptera), an incomplete elytron of *Elaphrinae* (Coleoptera, Carabidae) (UAMES 52332; Figure 4 h) and finally, some disarticulated coleopteran remains, too
fragmented to be identified. The presence of a psylloidean species in this context is not surprising since fossil remains of the three main plants *(Salix, Betula, Polygonum)* exploited by these insects were preserved evidenced within the skull cavity of the bison. The larval stages of chironomids develop in almost any aquatic or semiaquatic habitat, both standing and flowing waters, but also occur in tree-holes, rotting vegetation, and damp soils. Finally, the presence of *Elaphrus* sp. (*cf. trossulus* Semenov) (UAMES 52333; Figure 4 h), another ground beetle, is recorded. The species belonging to this genus are representative of riparian communities. Adults of all species of *Elaphrus* live along rivers, small streams, swamps, sloughs, or bogs, which is fully consistent with the bison discovery site. Overall, these taxa are typical of mesic to wet tundra habitats at the time of burial.

**3.4 Isotopic analyses:**

The δ\(^{13}\)C and δ\(^{15}\)N values from the collagen sample were -20.0 \(\pm\) 0.6 and 4.2 \(\pm\) 0.1, respectively. The δ\(^{13}\)C value is consistent with a diet of vegetation consists of plants using C\(_3\) photosynthesis in a relatively open environment (Drucker et al., 2008). The δ\(^{15}\)N value is consistent with values from other bison specimens dated to interstadial conditions, and contrast with stadial conditions, which can have higher values (Hedges et al., 2004; Rabanus-Wallace et al., 2017). Based on an 8-10 year collagen turn-over rate (Hedges et al., 2007), the collagen would exclude the higher δ\(^{15}\)N values from this individual’s early life based on analyses of the horn sheath (described below), and thus the bone collagen and horn can be considered in the same range. The mean horn keratin δ\(^{13}\)C value was slightly lower than the collagen values (mean δ\(^{13}\)C value -21.3 \(\pm\) 1.1; -20.0 \(\pm\) 0.1 respectively), which is consistent with the difference in fractionation factors associated with the two different tissue types (keratin Δ=3.1 \(\pm\) 0.3; collagen Δ=5.1 \(\pm\) 0.3 (Drucker et al., 2008)).

The mean δ\(^{15}\)N value from analyses of the horn keratin (4.6 \(\pm\) 1.1) was similar to the δ\(^{15}\)N value from the collagen value (4.2 \(\pm\) 0.1). The periodicity of the isotopic fluctuations in the horn sheath (Figure 6) appears to reflect annual cycles because the number of oscillations (n = 11) is consistent with the number of observed annual growth layers observed from the horn (11-12). The first two oscillations (~2 years) likely correspond with the period covered by the tooth record (~2.5 years) from the same specimen (described below). These apparently annual
fluctuations would be consistent with the peaks in δ¹⁵N values reflecting a seasonal shift to greater nutritional stress and a more water-limited plant diet during the winter 2020). Shortly before the death of the bison, the δ¹⁵N values from the horn sheath increases (Figure 6), which would correspond with a transition from summer to winter that is observed in modern bison from Alaska (Funck et al., 2020), indicating the bison may have died in the late summer/early fall. Two peaks in the δ¹⁵N values from the horn sheath occurred towards the start of the life of the bison and the second of these corresponds with a decrease in δ¹³C values from the horn sheath. These features are consistent with periods of nutritional stress (Funck et al., 2020). Most notably, this pattern seems consistent with catabolism and the use of lipid reserves respectively (Funck et al., 2020). A marked decrease in δ¹³C values is evident in the year prior to the bison’s death, which also seems consistent with the use of the animal’s lipid reserves (Funck et al., 2020).

The results of the ⁸⁷Sr/⁸⁶Sr ratio analyses (n = 33) from specimen UAMES 29458 (Table 2), with an analytical precision of 0.00001, show a mean ⁸⁷Sr/⁸⁶Sr ratio from the first molar (M1) of 0.71139 (±0.00028). These values are consistent with the rodent teeth values from the Arctic Coastal Plain, which range from 0.70940 to 0.71107 (Table 3, Figure 7). The mean ⁸⁷Sr/⁸⁶Sr ratio from the M2 was higher compared to the values from the M1 (p < 0.000, t.test), averaging at 0.71206 (±0.00025). The mean ⁸⁷Sr/⁸⁶Sr value from the M3 was 0.71308 (± 0.00008) and was higher than the values from both the M1 and M2 (p < 0.000, t.test). The range of the M3 values was similar to the values found from the modern rodents found in the foothills of the Brooks Range mountains (Table 3) (from 0.7137 to 0.7205) (Figure 8). Based on the bedrock ⁸⁷Sr/⁸⁶Sr model (Bataille and Bowen, 2012), the expected ⁸⁷Sr/⁸⁶Sr ratio at the place of death of specimen UAMES 29458 was ~0.70982 (Figure 8). Overall, the M1, M2 and M3 sequential data show a marked increase in ⁸⁷Sr/⁸⁶Sr ratios throughout the tooth sequence (p < 0.000, ANOVA) (Figure 7).

The mean δ¹⁸O value across all teeth samples from specimen UAMES 29458 (n = 30) was −21.1 ‰ (±0.4) vs. VPDB (with an analytical precision of 0.2 ‰) (Table 4). Values were highly variable across the M1-M3 sequence (Figure 7) ranging from −18.4 ‰ to −26.5 ‰, but were not significantly different between teeth (p = 0.34, ANOVA). The values appear to cycle through approximately two full years, with three peaks representing summers, which seasonally have higher δ¹⁸O values (Velivetskaya et al., 2016) and separated by two cooler periods with lower values (Figure 7).
The mean δ¹³C value of the enamel calcium carbonate from the teeth of specimen UAMES 29458 was -11.0 ‰ (±2.0) (with an analytical precision of 0.03) and had a relatively small range of values (-10.3 ‰ to -11.7 ‰) (Table 4), which are consistent with a C₃ diet. Typical values for C₃ plants are ~-27 ‰ (Wooller et al., 2007) and the difference between animal diets and tooth calcium carbonate is Δ+14.6 ‰ (Cerling and Harris 1999; Passey et al. 2005). Overall the δ¹³C values started at their lowest during early life (M1) and gradually became higher in the M2 (-11.1 ‰ ±0.2) and M3 (-10.5 ‰ ±0.1) (p < 0.000, ANOVA), a trend that appears to correlate with the temporal shift from lower to higher ⁸⁷Sr/⁸⁶Sr ratios from the same teeth (Figure 7).

3.5 aDNA analyses:

We generated 1.15 million reads from the unenriched library of UAMES 29458 (raw reads available at the NCBI Short Read Archive under BioProject PRJNA61324), which had an endogenous DNA content of 15.4% based on reads aligning to the cow genome. The relative mapping frequency ratio between the X chromosome and autosomes was 0.528, with a range of 0.493-0.561 across all 29 autosomes (Table 5). This is consistent with UAMES 29458 representing a genomic male individual.

A full mitochondrial genome (JK319) was generated from the enriched library, with an average coverage of 139× (Genbank: MN549280). The phylogenetic placement of the UAMES 29458 mitochondrial genome was marginally impacted by the minimum age prior used, and was found either to fall at the base of Clade 2 (prior of 50 kya BP) or to be sister to MS002 within bison mitochondrial Clade 2 (all other tested priors) (Figure 9, Table 6). The inclusion or exclusion of MS022 did not affect the phylogenetic placement of UAMES 29458. Across all analyses, the age of UAMES 29458 was estimated to be ~33-87 kya BP (Table 6). Analyses with MS022 excluded generally yielded slightly older age estimates, but varying the minimum age prior did not greatly impact this individual’s maximum estimated age (range of ~81-87 kya BP across all analyses). This suggests that the priors did not drive the estimated age results.

Mapped reads from both the unenriched (primarily nuclear) and enriched (mitochondrial) DNA libraries exhibit damage patterns characteristic of authentic aDNA, including short DNA fragments and elevated relative deamination frequencies at the ends of reads (Figure S4).
4. Discussion:

4.1 Constraining a chronological age for UAMES 29458:

Specimen UAMES 29458 provides several lines of evidence that can be used to constrain the individual’s chronological age (Figure 11). The radiocarbon dates from the keratin and the blow fly puparium (chitin) are nearly identical (46,000 ± 1100 \(^{14}\)C yrs and 46,800 ± 1200 \(^{14}\)C yrs – inside and outside the range for calibration, respectively). Keratin is a structurally dense tissue and less susceptible to contamination than collagen (Taylor et al., 1995). The blow flies \(P. terraenovae\), which colonizes a corpse within hours or days following death, seemed a good candidate for precise \(^{14}\)C dating of specimen UAMES 29458. The congruence of the keratin and chitin \(^{14}\)C dates lends support for a finite age of approximately 46,900 \(^{14}\)C yrs. In contrast, the radiocarbon dates from collagen and from plant material are non-finite. Collagen, from porous bone material, can be contaminated or degraded. The sample from the bone of UAMES 29458 was also run on a lower energy \(^{14}\)C instrument, which cannot discriminate low levels of \(^{14}\)C from old samples compared with the chitin and keratin, which were run on a more powerful instrument. The radiocarbon date of the plant material and its apparent older age compared to the puparium and keratin is difficult to interpret. The plant material recovered from the skull was specifically selected because it was a delicate herbaceous structure that would not have experienced substantial reworking or be influenced by variations in carbon sources that can occur in aquatic plants (Marcenko et al., 1989). However, the radiocarbon date of the plant was determined to be non-finite (>49,000) and could have been reworked from older sediments. Given that the dates from keratin and chitin are on the margins of radiocarbon limits we must consider two possibilities; first, that the finite dates are accurate, or second, that these dates are not distinguishable from non-finite dates.

Other evidence supports the hypothesis that UAMES 29458 dates are finite in age. Environmental information from the \(^{18}\)O values from the bison specimen and plant macrofossils add further dimensions that help constrain the chronological age estimation for UAMES 29458. The mean \(^{18}\)O value (as VSMOW) from analyses of the bison’s molars is -3.9 ‰ (±1.2) lower than the mean modern precipitation values. In contrast, full stadial \(^{18}\)O values for precipitation...
tend to be around -8% lower than modern precipitation values (Bowen, 2018; Gaglioti et al., 2017; Meyer et al., 2010; Rasmussen et al., 2014). The $\delta^{18}O$ values from UAMES 29458 therefore indicate that although the climate was slightly colder than modern it was not full stadial (Figure 11). The finite radiocarbon date from the keratin coincides with a period of time that had an offset in $\delta^{18}O$ values of $\Delta -3.9 \%$ ($\pm 1.2$), between 44,380 cal yrs BP and 46,880 cal yrs BP (Rasmussen et al., 2014). This inference that the bison specimen may have lived during a more moderate (interstadial) climate is also supported by the plant macrofossils and insect remains found in the skull, which represent a typical flood plain in tundra ecosystems of the Arctic. However, as stated above, the plant remains may not be exactly contemporaneous with the specimen.

The Bayesian molecular age estimate for the specimen was between ~33-87 kya BP, which is consistent with the radiocarbon ages and corresponds to Marine Isotope Stage (MIS) 3 through to MIS 5b (Lisiecki and Raymo, 2005). Although the results of this analysis are not informative as to whether UAMES 29458 is of older radiocarbon finite or non-finite age, they do suggest that, if the individual were radiocarbon non-finite, it lived after MIS 5e. The finite radiocarbon age from the bison’s keratin is within the range of this molecular-clock derived age estimate, which could narrow the age estimate to within MIS 3 (29-57 kya BP). Our phylogenetic analyses show that UAMES 29458 belonged to bison mitochondrial clade 2 (Heintzman et al., 2016). This lineage did not contribute to extant bison diversity, and instead became extinct during the late Holocene, by as long as ~400 years ago (Heintzman et al., 2016; Shapiro et al., 2004). Given the varied methods used for dating this specimen, it is expected that there might be some disagreement between results. However, the two nearly identical dates from the chemically stable portions associated with the specimen (i.e. keratin and insect chitin, 46,000 ±1100 $^{14}$C yr BP and 46,800 ±1200 $^{14}$C yr BP, respectively) provide support for this as a finite-aged specimen very close to the limit of radiocarbon dating. This age is congruent with an appropriate climatic period suggested by the $\delta^{18}O$ values (Figure 11) and the macrofossil assemblage of flora.

4.2 Life history of an individual steppe bison:
Multiple lines of evidence suggest that specimen UAMES 29458 was a large bull that was 11 to 12 years old at the time of death. Present-day male bison of this age are typically lone bulls, either in or just past their prime (Maher and Byers, 1987; Soper, 1941). During his life, this individual incurred injury to a rib and the bone callous clearly indicates it healed (Figure 3c). This type of injury typically occurs in present-day bison during the rut when bison compete for access to females (Lott, 1971). In addition to this evidence of injury, the early stages of this bison’s life, recorded as the δ¹⁵N values from the horn sheath (Figure 6), could indicate some periods of nutritional stress. Increases in δ¹⁵N values in the tissues of bison can represent stressors, such as starvation, illness, or long-distance movement (Funck et al., 2020). Shifts in δ¹⁵N values are generally used to determine changes in the trophic level (Post, 2002), an interpretation that is not appropriate for a herbivore, except during breast feeding when an juvenile is essentially a trophic level above their mother (Gadbury et al., 2000; Reitsema and Muir, 2015). However, during nutritional stress, such as a hard winter (Funck et al., 2020) or nutritional transition like weaning (Fuller et al., 2003), an animal can break down its muscles to build new proteins, during which tissues fractionate and lead to elevated δ¹⁵N values (Lee et al., 2012). The bison specimen analyzed here exhibited two periods of elevated δ¹⁵N values towards the start of its life (Figure 6). Horn stubs begin developing in utero but do not solidify into horn spikes until the calves are older (Wiener et al., 2015). The two periods of higher δ¹⁵N values likely occurred in the first 2-3 years of life, overlapping with some of the time periods represented by the molar development, which was used to produce the δ¹⁸O values and ⁸⁷Sr/⁸⁶Sr ratios. This allows us several lines of evidence for interpreting what early life was like for this bison.

The interpretation of intra-tooth serial samples as a measure of change over time requires an understanding of how teeth develop and mineralize. For example, the molars of bison have been shown to grow over 2 to 2.5 years (Gadbury et al., 2000; Higgins and MacFadden, 2004; Velivetskaya et al., 2016) and the pattern of molar eruption occurs in a particular sequence and time in an animal’s life. As each molar develops, discrete layers of enamel are laid down and retain the isotopic composition of conditions at each point in time. However, as a tooth develops it continues to mineralize over 6 to 7 months leading to a degree of isotopic averaging (Balasse, 2002; Montgomery et al., 2010). As a result, changes that may appear progressive could in fact occur over much shorter periods of time and the exact timing of tooth development and
mineralization can be subject to some degree of variation between individuals and species. However, Velivetskaya et al. (2016) used high-resolution sampling for analysis of $\delta^{18}$O values to pinpoint the timing of tooth development in two late-Pleistocene steppe bison from the Middle Urals, Russia and found it to be close to those of present-day bison. Seasonal oscillation in $\delta^{18}$O values can also be compared to those of present-day animals to track the speed of tooth development.

The M1 is formed in utero and shortly after birth (Bernard et al., 2009; Widga et al., 2010), and thus represents the mother’s home range and the calving grounds. The M1 formation likely corresponds with the timing of the first $\delta^{15}$N value peak (Figure 6). The $^{87}$Sr/$^{86}$Sr ratios from the M1 had less variation than the M2 suggesting the mother remained in a relatively consistent geological area. When the M1 $^{87}$Sr/$^{86}$Sr ratios are compared to bioavailable $^{87}$Sr/$^{86}$Sr ratios from the region, it appears that this bison began life on the Alaska coastal plain (Figure 8). Contemporary caribou from similar regions of Alaska currently use the coastal plain as calving grounds, taking advantage of the emerging graminoids and open terrain (Fancy et al., 1990; Post and Forchhammer, 2008). Subsequently, they move to higher ground to avoid insects and to take advantage of abundant lichens that sustain them during non-calving seasons (Fancy et al., 1990). Bison calves are dependent on milk for the first ~5 months and gradually stop suckling over 9-21 months (Green et al., 1993). So the first peak in $\delta^{15}$N values from the horn (Figure 6) likely corresponds to this M1 period and could be interpreted as a maternal/weaning signal (Gadbury et al., 2000; Reitsema and Muir, 2015), either attributed to the bison being at a higher trophic level than its mother from which it was nursing, or the effect of weaning and the nutritional stress that it may have imparted (Fuller et al., 2003).

After the period of initial relative geographic stability inferred from the $^{87}$Sr/$^{86}$Sr ratios in the M1, which likely represented the first half-year of life, the bison seems to have dispersed into a new geographic area during its second summer according to the $^{87}$Sr/$^{86}$Sr ratios from the M2 and M3. The M2 development starts at ~2 to 3 months of age and continues for 12-15 months (Gadbury et al., 2000; Velivetskaya et al., 2016). The M3 subsequently develops from the beginning of the second summer after weaning, ~10-11 months until the age of about 2 to 2.5 years (Gadbury et al., 2000; Velivetskaya et al., 2016). Comparisons of the bison’s $^{87}$Sr/$^{86}$Sr ratios from the M2 and M3 to bioavailable $^{87}$Sr/$^{86}$Sr ratios from the region indicate that the bison may have moved into the foothills of the Brooks Range (Figure 1 and Figure 8). The second peak
in δ¹⁵N values from the analyses of the horn sheath (Figure 6) could correspond to mobility on the landscape, which seems to have occurred during this period. A study on present-day wood bison in Alaska (Funck et al., 2020) found that bison that traveled long distances, or experienced nutritional stress, produced elevated δ¹⁵N values due to the energetic costs of travel. The horn δ¹³C record begins relatively low shortly before this point, which could be related to the bison beginning to draw on the bodies lipids which have lower δ¹³C values (DeNiro and Epstein, 1977; Rode et al., 2018). This offset follows the same pattern found in a modern wood bison experiencing a dispersal related dietary stress (Funck et al., 2020). In the present study, the tooth carbonate δ¹³C values show an overall increase (+1 ‰) during this same period (~2.5 years), consistent with a change from a wetter environment to a drier one (Wooller et al., 2007), which also seems to correspond with a shift towards the higher and drier elevations of the foothills compared to the wetter, lower reaches, of the coastal plain. Later in the bison’s life, after the record preserved in the molars is fully mineralized (~2.5 years), we lose track of the specimen’s geolocation based on the ⁸⁷Sr/⁸⁶Sr record. However, this bison must have eventually returned to the coastal plain to the location where he eventually died and was found (Figure 1).

In contrast to the tooth record, the horn sheath continues to develop over approximately the lifetime of the individual. The δ¹⁵N fluctuates through what appear to be seasonal oscillations of slightly more elevated δ¹⁵N values in the winter, likely due to nutritional stress during this season, then lower δ¹⁵N values during more favorable summers, interpreted based on isotopic patterns exhibited in present-day bison from Alaska (Funck et al., 2020). Present-day wood bison south of the Brooks Range in Alaska only had changes in δ¹⁵N values during particularly hard winters (Funck et al., 2020). Thus, these winters were likely harder than the conditions experienced by present-day analogous populations. Applying this seasonal pattern implies that the sample closest to the horn core (i.e. the period leading to the animal’s death) represented the start of the transition from the summer to winter. This indicates that the bison was not experiencing unusual nutritional stress and did not appear to have been in a weakened state prior to death. Other potential causes for changes in δ¹⁵N values in herbivores could include changes in the seasonal δ¹⁵N values of forage and seasonal movement to regions with very different baseline δ¹⁵N values. Our data indicates that this bison individual did move substantially across the landscape. However, the tooth record of paleomobility is not long enough to cover the longer record represented by the horn sheath data. We find that our interpretation of the δ¹⁵N values as
a marker in terms of changes in the degree of nutritional stress is in some ways supported by the 
\( \delta^{13}C \) data from the horn sheath. For example, the low \( \delta^{13}C \) values from the horn sheath in the last 
year of life, relative to the rest of the record from the horn, could indicate that the animal may 
have begun to draw on its lipid reserves (Funck et al., 2020). Although in some animals these 
isotopic shifts occurs only at extreme thresholds of starvation this is a more common strategy in 
animals that have large fat and muscle reserves for winter use, such as bison (Funck et al., 2020).

The degree of mobility of the bison specimen analyzed, based on the \( ^{87}\text{Sr} / ^{86}\text{Sr} \) ratio data 
from the North Slope specimen, is relatively high compared to data from other ancient and 
present-day bison (Figure 10, Table 7; Britton et al., 2011; Glassburn, et al., 2015; Julien et al. 
2012; Widga et al., 2010). Although the range of \( ^{87}\text{Sr} / ^{86}\text{Sr} \) ratios cannot be directly compared, 
because variability is highly dependent on the geological heterogeneity of a landscape that 
individuals inhabit it is worth considering this bison in the context of others analyzed using the 
similar methods. Britton et al. (2011) examined a single bison from western France dating to 
49,000 ±5000 BP and demonstrated high fidelity to an area, with almost no variability in 
\( ^{87}\text{Sr} / ^{86}\text{Sr} \) ratios despite a variable geological environment. Julien et al. (2012) found that steppe 
bison from the Last Glacial Maximum (LGM) (~20.5 kya BP) in Amrosievika in Eastern 
Ukraine had a \( ^{87}\text{Sr} / ^{86}\text{Sr} \) range of 0.00074 (Julien et al., 2012). This range was attributed to the 
low degree of variability to minimal mobility on the landscape because there was no signals from 
mixed geology as close as 20 km to the south-west of the site where the bison were found (Julien 
et al., 2012). A study of bison herds on the Great Plains of North America from the Middle 
Holocene indicated that bison also had limited seasonal mobility (<50 km), while inter-annual 
movement of herds over ~4-5 yrs moved further afield (<500 km) (Widga et al., 2010a). In 
contrast, present-day \textit{Bison bison} from the interior of Alaska, with an observed seasonal and 
regional migration route of about a 100km, had a \( ^{87}\text{Sr} / ^{86}\text{Sr} \) ratio range from 0.71714 to 0.71540 
(0.00174) (Glassburn et al., 2018). Although these regions and studies cannot be quantitatively 
compared because of difference in local geology (Bataille et al., 2020) we can still draw some 
qualitative comparisons. The variability in \( ^{87}\text{Sr} / ^{86}\text{Sr} \) ratio ranges among bison from different 
localities and periods suggests that bison utilize different mobility strategies that are dependent 
on (paleo)ecological conditions. The range of \( ^{87}\text{Sr} / ^{86}\text{Sr} \) ratio variation from the North Slope bison 
specimen analyzed here (0.00237) is relatively large compared to all these previous measures, 
indicating that he was likely a gregarious individual who traveled some distance during his early
years of life. There is a large geological range in this region (Bataille et al., 2020) that this bison appears to be utilizing. Future research could integrate genetic testing of sex from specimens to determine if males and females used different mobility strategies.

4.3 Necrology and Biostratinomy:

Evidence from the specimen UAMES 29458 helps shape a paleoecological picture leading up to the individual’s death, decomposition, and the mechanism of burial. The bones from the bison specimen were overall in a state of excellent preservation as evidenced by the presence of intact horn, hooves, fur, spinal cord, cartilage, and possible brain tissues. Specimens of this level of completeness are extremely rare in Quaternary deposits. There were no obvious signs of the cause of death, such as bite marks around the nose or neck, and all carnivore marks were related to scavenging (Andersson et al., 2011) but predation cannot be ruled out. After death, the specimen came to finally rest on his right side, which protected that side from scavenging, this is further supported by the presence of vivianite. Vivianite deposits tended to be on the bone surfaces that were exposed to the ground surface and root etching also tended to be present on the surfaces of the bones that rested on the ground (Supplemental data).

Carnivore tooth marks are present at various locations on the specimen, indicating scavenging of the bison carcass occurred post mortem (Figure 3 a and b and Supplemental data). There are four major carnivore taxa present contemporaneously that could cause the type of damage seen on UAMES 29458: Arctodus simus (short-faced bear), Canis Lupus (wolf), Homotherium serum (scimitar cat), and Panthera atrox (American lion) (Fox-dobbs et al., 2008; Schubert, 2010). The dentition marks (Figure 3) were consistently narrow punctures, which are not consistent with the wide teeth of A. simus (Sorkin, 2006) or H. serum (Ewald et al., 2018). The incisor punctures indicated that the space between the canines is at least 6 cm apart (Figure 3a, which is consistent with P. atrox (Baryshnikov and Boeskorov, 2001; Christiansen and Harris, 2010). This may be out of the range of the smaller C. lupus (Sorkin, 2006), leaving large felidae as the most likely scavenger. Further analysis of dentition morphometrics in predator species would be useful for this type of taphonomic analysis. Indication of lion scavenging have also been found on other preserved bison specimens from in Alaska (Guthrie, 1989). It is likely that only one individual scavenger caused all of the damage based on the consistency in size of
the tooth marks of the 14 tooth marks with punctures. The 11 instances of gnawing are less
diagnostic but also appear consistent. The limited scavenging also supports a single scavenger, as
more individuals would require less time to remove meat, disarticulate the corpse, and scavenge
from a greater area of the carcass (Blumenschine, 1986). The carnivore appears to have
consumed the bison carcass and ate portions of the backstrap (meat along the top of the spine)
based on observed damage along the spinous processes of several vertebrae (Figure 3 a and b).

The carnivore also seems to have consumed the carcass from the inside of the body and likely ate
the internal organs and tenderloin based on damage to the costal cartilage, the distal ribs, the
transverse processes of the lumbar vertebrae, and the sacrum. In addition, the carnivore gnawed
the left humerus and likely removed the left scapula, in an attempt to get at the flesh of the left
fore-limb. Scavenging only seems to have occurred in high-value areas, supporting an inference
that the carcass was likely buried rapidly. In most contexts a carcass on an open landscape with
carnivores can be consumed and disarticulated within hours to a few days (Blumenschine, 1986).
It is for this reason that complete specimens are extremely rare. Thus, for this specimen to
preserve, something exceptional must have occurred to protect it. During the erosion that re-
exposed the bison it seems the left extremities of the front and hind limbs (phalanxes, carpals,
tarsals, etc.) may have been lost due to erosional processes (Figure 2). These skeletal elements
would be the first to be exposed and vulnerable to gravity, weathering, and would have been
preferentially removed via taphonomic agents (e.g., eroded into the river and lost). These skeletal
elements of the lower extremities are also not associated with high food value flesh and likely
would not have been a target for scavenging.

The presence of the blow fly puparia recovered from inside the skull made it possible to
propose some "paleoforensic" hypotheses associated with postmortem events. Blow flies figure
among the most relevant witnesses in forensic entomology in order to determine the time elapsed
since death, namely the postmortem interval (PMI) because blow flies (Calliphoridae) are the
first and predominant organisms to colonize a body after death. Within hours of death, these
insects are attracted to the smell of the decomposing remains, which are both a site for laying
eggs and a source of protein for larval development. Under normal circumstances, egg laying
occurs soon after death (1–3 days) (Gomes et al., 2006). The favored egg-laying sites are the
natural openings of the body (mouth, eyes, anus) as well as any wounds present on the corpse
(Gomes et al., 2006). The species preserved on the carcass *Protophormia terraenovae* (Robineau
Desvoidy) is a Holarctic, cold-adapted species, present throughout northern Europe and Asia. Very common in the cooler high latitude regions and notably in the Arctic, representatives of this species are found within 890 km of the North Pole (Smith, 1986). Experiments conducted by Marchenko (2001) found that the development period of *P. terraenovae* (from egg to adult) is relatively long. According to Warren and Anderson (2013), egg-carrying females of *P. terraenovae* do not lay eggs at temperatures lower than 10.3°C. In the North Slope region, temperatures only reach above 10°C during the summer period (https://www.weatherbase.com). Insects, like all arthropods, are coldblooded and the duration of their life-cycle is primarily temperature driven. Rate of development is species-specific and influenced by biotic (e.g. maggot mass that can significantly accelerate rate of development) and environmental factors (exposure of the carcass to sunlight or shade, rain, and wind). Following the biological data provided by Marchenko (2001), and focusing on temperatures ranging from 11 to 18°C (respectively the average minimal temperature required for laying eggs and the current maximum temperatures recorded on the paleontological site), *P. terraenovae* development extends from 15 to 50 days (egg to pupal stage) and adult emergence from 24 to 78 days. In the case of UAMES 29458, all of the puparia were intact with no evidence of adult emergence.

The fact that the bison's skeleton was almost complete and articulated, and not fully scavenged by carnivores suggests that the fly pupal stage intervened fairly quickly, indicating relatively high temperatures at the time of death (e.g., at 18°C, the pupal stage would have started on the 15th day). The bison carcass was then quickly covered with sediment or snow precluding further access to predators; however, a few centimeters of soil cover would not prevent the development and emergence of flies (Balme et al., 2012). Trapped in the skull, the pupae would have preserved in situ. The presence of unhatched fly puparia of *P. terraenovae* recovered from the bison individual would suggest that its carcass had been subaerially exposed for at least 2 weeks before the life cycle was interrupted either due to a more complete burial or freezing. The fact that the carcass was little affected by scavengers over this period might be explain by low scavenger density on site, high carcass availability or because the bison’s remains were only accessible to flying insects and not by terrestrial scavengers (e.g. carcass partially exposed in river water). Finally, we can argue that the ambient temperature was very likely over 10°C when the fly infestation occurred, requiring the events to occur during a warm period of the
Fluvial sediments including plant macrofossils likely entered the skull after this initial burial during which the *P. terrae novae* puparia developed. The plant macrofossils recovered from inside the skull and neural canal are unlikely penecontemporaneous with the death of this bison individual, but could indicate an overall glimpse of the surroundings. The macrofossil assemblage of plants is consistent with an Arctic flood plain. The immediate surrounding of the river was probably covered in moss, low shrubs, and bogs (Figure 5). Further away from the banks of the river were likely drier tundra with grasses, sedges, and a variety of tundra flowers including *Papaver* sp., *Polygonum bistorta*, and *Draba* type (Figure 5). Overall, the plant assemblage is more typical of a mesic tundra than a steppe and possibly originated from an interglacial or interstadial environment (Gaglioti et al., 2018).

Overall, we conclude that this bison died on a flood plain during the warm period prior to the onset of winter and fell on to its right side. A carnivore, possibly a large felidae, quickly consumed high-value areas of meat and blow flies swiftly laid eggs within the skull. Alluvial sediments or possibly snow then quickly buried the carcass before the carcass could be further disarticulated. At least two weeks passed before temperatures inside the carcass or the oxygen availability fell too low for the puparia to continue developing.

5. Conclusion:

Our multi-proxy paleoecological evidence from UAMES 29458 fills out the details of an individual bison’s life and death in the Arctic, where bison were once a dominant herbivore. Combining serial isotope analyses with aDNA allowed us to combine mobility information with genealogical information, and to place the individual within bison meta-population dynamics over millennia. This augments a paleoecological picture based on the physical evidence of the skeletal remains alone, providing a more vivid image of a highly mobile individual moving across the North Slope of Alaska. The mitochondrial lineage of this bison is one that thrived in Beringia but ultimately died out. Our multiproxy evaluation indicates that this bison likely lived during an interstadial period, which in some ways may have been somewhat similar to today. Strontium isotope data from the specimen indicates that he dispersed across the Northern Alaska
landscape from possible calving grounds located on the Arctic coastal plain to the foothills of the Brooks Range in early life and ultimately back to his place of death on the Arctic coastal plain. Nitrogen isotopes from the horn sheath indicate that the mobility of his early life may have resulted in significant nutritional stress, and that this individual suffered somewhat harsher winters than those on the North Slope today. The taphonomic analysis revealed that this bison was scavenged in parts of high nutritional value, which was followed by a rapid burial in a riverine environment. In this case, a multiproxy approach has uncovered a life history and detailed examination of a member of a dominant Pleistocene species in the Arctic.

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38


Figure titles:

Figure 1. a) A map on northern Alaska, with the provenance of steppe bison (Bison priscus) specimen UAMES 29458 indicated by the red star. Yellow dots show rodent specimen localities and strontium isotope ratios measured from their teeth. The underlying map is modified from Bataille et al. (2016). b) UAMES 29458 skull in situ during excavation. c) UAMES 29458 articulated thoracic vertebrae attached with connective tissue, in situ during excavation.

Figure 2. a) Specimen UAMES 29458 rearticulated for taphonomic analysis. b) Skeletal elements missing from specimen are indicated in gray and elements with carnivore marks indicated in red. All elements from the right side were present.
Figure 3. Scavenger dentition marks on the a) thoracic vertebral processes of *Bison priscus* specimen UAMES 29458 with bites indicated by arrows, and b) ventral side of lumbar vertebrae with gnaw marks indicated by arrow c) Vertebrae with healed pathology indicated by arrow.

Figure 4. The macrofossil assemblage taken from skull of specimen UAMES 29458: *Protophormia terraenovae* puparium UAMES 52319. a), dorsal view b), lateral view c), ventral view d), posterior view, the posterior spiracles surrounded by a rectangle e), the posterior spiracles in close-up view. (photos: J.-B. Huchet, 2018), f) Partial abdomen sternites of a ground beetle UAMES 52334 (Carabidae: Pterostichinae) (photo: J.-B. Huchet, 2018), g) Psyllid forewing recovered from inside the bison skull UAMES 52331 (photo: J.-B. Huchet, 2018), h) Right elytron of a ripicolous ground beetle of the genus *Elaphrus* Fabricius UAMES 52333 (Carabidae: Elaphrinae) (photo: J.-B. Huchet, 2018), i) Carex seed lenticular, j) Carex seed Trigonal, k) Poaceae caryopsis, l) *Polygonum bistorta*, m) *Draba*-type seed, n) *Andromeda polifolia*, o) *Bryophyta* sp.

Figure 5. The relative abundance of plant macrofossil specimens in the skull and spine of specimen UAMES 29458, categorized by characteristic eco-type.

Figure 6. Carbon and nitrogen stable isotope values of serial horn sheath samples (n=113) with dotted trend-line of weighted moving averaging of the nearest 3 samples and collagen carbon and nitrogen stable isotope values. * Indicate lowest $\delta^{15}N$ values associated with summer. Labels on the bottom indicate approximate timing of identifiable life events.

Figure 7. Isotope data from analyses of molar 1 (M1), molar 2 (M2), and molar 3 (M3) from specimen UAMES 29458. a) Strontium $^{87}$Sr/$^{86}$Sr ratios, b) Oxygen stable isotope ratios expressed as $\delta^{18}O$ values and c) Stable carbon isotope ratios (expressed as $\delta^{13}C$ values). Shading represents estimates of winter, based on the $\delta^{18}O$ values. * Indicate highest $\delta^{18}O$ values associated with summer.
Figure 8. Rodent $^{87}\text{Sr}/^{86}\text{Sr}$ ratios grouped by region, compared to $^{87}\text{Sr}/^{86}\text{Sr}$ ratios from specimen UAMES 29458 over time and a model-predicted $^{87}\text{Sr}/^{86}\text{Sr}$ ratio for the death location of specimen UAMES 29458.

Figure 9. A Bayesian time-calibrated genealogy of bison mitochondrial genomes, with major well-supported Clades (1, 1A, 2, 2A, 2B) highlighted (following Heintzman et al., 2016 and Zazula et al., 2017). All living bison fall within Clade 1A, whereas the specimen UAMES 29458 falls near the base of Clade 2. Purple bars are 95% highest posterior density intervals for node heights and are shown for nodes with posterior probability $>$0.95. This maximum clade credibility tree resulted from the analysis that excluded MS022 and had a minimum age prior of 30 kya BP for UAMES 29458. Results from the other analyses can be found in Table 6. The diverged yak tips have been removed.

Figure 10. Comparison of the ranges ($\Delta$) of $^{87}\text{Sr}/^{86}\text{Sr}$ ratios recorded within the teeth of different bison including, present-day plains bison ($Bison bison$) from interior Alaska (Glassburn et al., 2018); Holocene plains bison ($Bison bison$) from the American Plains (Widga et al., 2010); Late Pleistocene steppe bison ($Bison priscus$) from Ukraine (Julien et al., 2012) and Late Pleistocene steppe bison ($Bison priscus$) from France (Britton et al., 2011).

Figure 11. Summary of all radiocarbon dates ($^{14}\text{C}$), molecular clock estimates, and compatible $\delta^{18}\text{O}$ value periods based on Greenland Ice core values (Rasmussen et al., 2014) and a $\Delta 4\%$ lower values than present. *dates are calibrated radiocarbon dates where possible, otherwise they are before present.
Supplementary materials

Figure S1. Intra tooth sampling of molar 1 (M1), molar 2 (M2) and molar 3 (M3) from *Bison priscus* specimen UAMES 29458, number indicates sample number.

Figure S2. Serial sampling scheme for bison horn sheath from *Bison priscus* specimen UAMES 29458.

Figure S3. Results of the tests for temporal signal in the bison mitochondrial genome data set. a) Full alignment data set, and b) reduced alignment data set (excluding MS022). Left panels: linear regression of phylogenetic root-to-tip distance against sampling date. Center panels: similar-date clusters (rounded to the nearest thousand years). Right panels: testing the results against random permutations over the clusters presented in b).

Figure S4. Assessment of damage patterns in nuclear (A-C) and mitochondrial (D-F) DNA fragments from *Bison priscus* specimen UAMES 29458. (A, D) DNA fragment length distributions, with means of 90 bp (nuclear) and 127 bp (mitochondrial). (B, E) The relative frequency of cytosine to thymine deamination (red line) and observed guanine to adenine misincorporation (blue line) from cytosine deamination on the opposite DNA strand. Deamination is more prevalent at the ends of ancient DNA fragments. (C, F) DNA fragmentation plots showing an elevated occurrence of purines (guanine, adenine) immediately upstream of the ancient DNA fragment (to the left of the grey box) and the opposite signal immediately downstream of the fragment (to the right of the grey box).
## Tables

Table 1. Summary of radiocarbon dates.

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Table 2. Results of isotopic analysis of strontium, carbon and oxygen of UAMES specimen 29458.

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Table 3. Rodents use for characterizing \(^{87}\text{Sr}/^{86}\text{Sr}\) isoscape.

<table>
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<tr>
<th>UAM: Mamm</th>
<th>Species</th>
<th>Locality</th>
<th>Environment</th>
<th>Lat</th>
<th>Long</th>
<th>(^{87}\text{Sr}/^{86}\text{Sr})</th>
</tr>
</thead>
<tbody>
<tr>
<td>4650</td>
<td>Microtus oeconomus</td>
<td>Meade River village</td>
<td>Coastal Plain</td>
<td>70.481</td>
<td>-157.417</td>
<td>0.71007</td>
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<td>8163</td>
<td>Microtus oeconomus</td>
<td>Umiat region</td>
<td>Coastal Plain</td>
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<td>-152.133</td>
<td>0.71309</td>
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<tr>
<td>11149</td>
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<td>Kikitaliorak Lake</td>
<td>Mountain Drainage</td>
<td>68.125</td>
<td>-156.233</td>
<td>0.71721</td>
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<tr>
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<td>Microtus oeconomus</td>
<td>Feniak Lake</td>
<td>Mountain Lake</td>
<td>68.250</td>
<td>-158.333</td>
<td>0.71372</td>
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<td>Barrow, NARL</td>
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</table>
Table 4. Summary of isotopic results ($\delta^{13}\text{C}$ and $\delta^{18}\text{O}$ reported here vs. VPDB in ‰).

<table>
<thead>
<tr>
<th></th>
<th>Mean</th>
<th>Max</th>
<th>Min</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>$^{87}\text{Sr}/^{86}\text{Sr}$</td>
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<td>0.71324</td>
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<td>$\delta^{13}\text{C}$</td>
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<td>$\delta^{18}\text{O}$</td>
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</table>
Table 5. Results of the genomic sex determination analysis. X:A ratio is the ratio of the relative mapping frequency of the X chromosome to that of an autosome (1-29, all). Chr.: chromosome. The minimum and maximum X:A ratios are highlighted in bold.

<table>
<thead>
<tr>
<th>Chr.</th>
<th>Length</th>
<th>Reads mapped</th>
<th>X:A ratio</th>
<th>Chr.</th>
<th>Length</th>
<th>Reads mapped</th>
<th>X:A ratio</th>
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Table 6. Summary of the Bayesian molecular analyses of JK319/UAMES 29458, with overall maximum age ranges highlighted in bold

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<th>Analysis variables</th>
<th>Phylogenetic placement</th>
<th>Estimated age</th>
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<td>Min. age prior</td>
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<tr>
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<td>Excluded</td>
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</table>

Phylogenetic placements are: 1) sister to MS002, and 2) at the base of Clade 2 (see Figure 9).

Minimum and maximum ages are based on 95% highest posterior density credibility intervals.

Note that, for analyses with a minimum age prior of 40 and 50 kya BP, the estimated minimum ages are likely to have been truncated by the prior. ESS: estimated sample size.
Table 7. Comparison to other Bison strontium data. See Figure 10 *Stratigraphically determined to be from MIS4 **Molecular clock.

<table>
<thead>
<tr>
<th>Context</th>
<th>Species</th>
<th>N</th>
<th>Date</th>
<th>Average Range</th>
<th>Citation</th>
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<td></td>
<td></td>
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<td>3</td>
<td>2014</td>
<td>0.00111</td>
<td>Glassburn et al. 2018</td>
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<td>Holocene American</td>
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<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>American Plains</td>
<td><em>Bison bison</em></td>
<td>12</td>
<td>8930-6980 cal yr BP</td>
<td>0.00077</td>
<td>Widga et al. 2010</td>
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<tr>
<td>Late Pleistocene</td>
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<td></td>
</tr>
<tr>
<td>Ukraine</td>
<td><em>Bison priscus</em></td>
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<td>20557-20,491 cal yr BP</td>
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<td>Julien et al. 2012</td>
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<tr>
<td>Late Pleistocene</td>
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<td></td>
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</tr>
<tr>
<td>France</td>
<td><em>Bison priscus</em></td>
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<td>49,000±5 yr BP*</td>
<td>0.00010</td>
<td>Britton et al. 2011</td>
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<tr>
<td>Bison Bob</td>
<td><em>Bison priscus</em></td>
<td>1</td>
<td>33-87,000 yr BP**</td>
<td>0.00237</td>
<td>This paper</td>
</tr>
</tbody>
</table>
Figure 1
Figure 2

(a) A detailed skeleton of an animal, including the skull and ribcage.

(b) A graphical representation of an animal's skeleton, showing the skeletal structure in detail.
Figure 3
Figure 5

Shrubs
- Salix Fragments
- Betula nana leaf

Grasses and sedges
- Carex seed lenticular
- Carex seed trigonal
- Poacea seed
- Drabs type seed
- Mustard type seed

Open tundra flowers
- Polygonum bistort seed
- Papaver sp.
  - Asteracea cf. solidago seed
  - cf. Caryophyllaceae sp. seed
- Potentilla Palustre seed
- Andromeda polifolia leaf

Wet indicators
- Juncus
- Shells
- Daphnea ephippia

Skull
- 1
- 2
- 1
- 3
- 1
- 5

Spine
- 2
- 3
- 3
- 2
- 3
- 3
- 3
Figure 6

Approximate overlap with tooth growth
Figure 7

- **M1**
- **M2**
- **M3**

**Strontium**

**Oxygen**

**Carbon**
Figure 8

A scatter plot showing the distribution of Sr/$^{87}$Sr values and latitude. The plot includes:

- Black triangles: Rodents
- White circle: Bison Bob UAM 29458
- Black dot: Expected Bison Bob

The plot is divided into regions:

- Coastal Plain
- North Slope foothills
- Brooks Range

The $^{{87}}$Sr/$^{86}$Sr values range from 0.705 to 0.725, and the latitude ranges from 67 to 72 degrees north.
Figure 9
Figure 10

Range of $^{87}\text{Sr}/^{86}\text{Sr}$ values within individuals

- Modern Interior Alaska: $N=3$
- Holocene American: $N=24$
- Late Pleistocene Ukraine: $N=12$
- Late Pleistocene France: $N=1$
- UAM 29458: $N=1$
- Bison Bob: $N=1$
Supplemental materials
S1

M1

M2

M3
Supplemental materials
S2
Supplemental materials
S4

Regression Dates

A

R_oot-o_to-tip distance

Dates

\( r = 0.5 \)

Time of MRCA = 174334 yrs BP

B

R_oot-o_to-tip distance

Dates

\( r = 0.44 \)

Time of MRCA = 213001 yrs BP

No clusters = 35

Randomised data sets

\( p\text{-value} = 0.01 \)

\( \text{min } p\text{-value} = 0.001 \)

\( p\text{-value} = 0.025 \)

\( \text{min } p\text{-value} = 0.001 \)