Interdisciplinary Analysis of the Lehi Horse: Implications for Early Historic Horse Cultures of the North American West

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Although recognized as one of the most significant cultural transformations in North America, the reintroduction of the horse to the continent after AD 1492 has been rarely addressed by archaeological science. A key contributing factor behind this limited study is the apparent absence of equine skeletal remains from early historic archaeological contexts. Here, we present a multidisciplinary analysis of a horse skeleton recovered in Lehi, Utah, originally attributed to the Pleistocene. Reanalysis of stratigraphic context and radiocarbon dating indicates a historic age for this horse (cal AD 1681–1939), linking it with Ute or other Indigenous groups, whereas osteological features demonstrate its use for mounted horseback riding—perhaps with a nonframe saddle. DNA analysis indicates that the animal was a female domestic horse, which was likely cared for as part of a breeding herd despiteoutlyingitsusefulnessinneffect. Finally, sequentially sampled stable carbon, oxygen, and strontium isotope values from tooth enamel (δ13C, δ18O, and 87Sr/86Sr) suggest that the horse was raised locally. These results show the utility of archaeological science as applied to horse remains in understanding Indigenous horse pastoralism, whereas consideration of the broader archaeological record suggests a pattern of misidentification of horse bones from early historic contexts.

Keywords: horseback riding, equestrianism, zooarchaeology, Utah, Lehi horse, horse pastoralism, Ute archaeology

Pese a ser reconocida como una de las más significativas transformaciones culturales en Norteamérica, la reintroducción del caballo en continente luego del año 1492 raramente ha sido abordada por la arqueología. Una razón de la escasez de tales trabajos es la aparente ausencia de restos esqueletarios de equinos en contextos arqueológicos históricos tempranos. Presentamos un análisis multidisciplinario de los restos de caballo recuperados en Lehi, Utah, los que fueron originalmente identificados como de la “Edad del Hielo” basados sobre el contexto geológico. El reanálisis del contexto estratigráfico junto con la...
The history of the American West is deeply intertwined with the domestic horse. The genus *Equus* first evolved in the Pliocene (Blancan) steppes of the Americas, radiating outward into Eurasia and Africa sometime during the last approximately three million years (Orlando et al. 2013). The ecological landscape of North America evolved with these wild equids, and the first people to move into the continent had a relationship with horses—both as prey species (Bourgeon et al. 2017; Kooyman 2006) and as a source of raw material for bone tools (Webb and Hemmings 2006)—along with other megafauna up to the end of the Pleistocene. Current radiocarbon dates of fossil equids in North American place their disappearance about approximately 10,000 years ago (Haile et al. 2009), and the Americas were without horses until the Spanish introduced the domestic horse (*Equus caballus*) and donkey (*Equus asinus*) after AD 1492.

Horses were first reintroduced to the Americas by Spanish colonists in the sixteenth century, and they were swiftly integrated into the culture, economy, and lifeways of many Indigenous peoples, especially in the North American West (Mitchell 2015). Some Indigenous groups established territorially dominant politics and became “horse nations” rivaling European powers on the continent (Hämäläinen 2008; Mitchell 2015). Although historic records shed some light on the development of riding and breeding horses among Indigenous North Americans, these processes took place largely outside the attention of European observers—limiting the explanatory power of history to address when, why, and how horses were first integrated into Indigenous societies in the Americas (Jones and Gabe 2015). The limitations of the Euro-American historical record in sixteenth- and seventeenth-century North America parallel challenges in understanding early human-horse relations in prehistoric Eurasia, where zooarchaeology has proven to be a valuable source of data about the early stages of horse domestication, management, and use. Osteological study of horse remains in natural history collections and comparison with the archaeological record have revealed characteristic damage patterns to the equine skeleton caused by metal bit use (Anthony et al. 1991; Bendrey 2007), riding with or without a frame saddle (Bartosiewicz and Gal 2013; Levine et al. 2005), heavy exertion (Taylor et al. 2015), and bridle nosebands (Taylor and Tuvshinjargal 2018). Biomolecular study of equid remains can provide sex and species identifications of bone fragments with a high success rate, even in the absence of sufficient DNA for more extensive sequencing (Schubert et al. 2017), whereas stable isotope analysis can shed light on seasonal movements, trade, and supplementary feeding practices (Bendrey et al. 2009; Makarewicz et al. 2018). Such a combination of zooarchaeological and biomolecular analyses of horse remains promises to reveal detailed insights into how horses were managed, ridden, and integrated into ancient lifeways—even in the absence of historical records.

One of the obstacles to the application of a multiproxy zooarchaeological approach to the early historic era in the North American West is the limited sample of horse remains identified in relevant museum collections. For example, in the entire state of Wyoming just two early historic horse specimens plausibly associated with Indigenous cultures have been identified by
Thornhill (2016). It is possible, however, that North America’s rich paleontological record of Pleistocene horses, combined with the well-known significance of horses in historic Euro-American societies, has resulted in the regular misclassification of Indigenous archaeological horses as either Euro-American or Ice Age in origin. Indeed, one of the two early historic horses identified by Thornhill from the site of Black’s Fork, Wyoming, was originally suspected to be the remains of a modern or ancient wild horse that died in that location from a natural death—until it became clear that the horse had been buried as part of a ritual feature, along with three coyote skulls (Eckles et al. 1994). Similarly, Scott and colleagues (2010) identified an early historic horse skull that had remained misidentified for a century, even earning its own Pleistocene taxon—Equus laurentius—before radiocarbon dating and a thorough morphological comparison revealed it to be historic. Furthermore, undated “intrusive” horse burials are sometimes reported from Indigenous archaeological sites after being ascribed to non-native ranchers or settlers (Supplemental Figure 1), despite clear archaeological evidence documenting that horses were sometimes integrated into Indigenous funerary and ritual inhumation (e.g., O’Shea and Ludwickson 1992). These examples suggest that, when confronted with ancient horse assemblages, researchers may often misidentify archaeological horse remains associated with Indigenous cultures.

The Lehi Horse

In the spring of 2018, residents of Lehi, Utah—a city located at the foot of the northern Wasatch Front in the Provo metropolitan area at an elevation of approximately 1,390 m asl (Figure 1)—discovered equine skeletal remains during new construction. The discovery was reported to researchers at the nearby Museum of Ancient Life, who visited the site and determined that the horse was located in sandy lacustrine deposits of Pleistocene Age. Researchers noted several skeletal pathologies, including arthritis, and determined that the animal was small, about “the size of a Shetland pony” (Holson 2018). Although no formal analysis was conducted, the animal was determined to be an elderly Pleistocene horse—perhaps Haringtonhippus francisci (Heintzman et al. 2017)—which had died a
natural death on the Provo shoreline of what was then Lake Bonneville (Holson 2018) and thought to date to approximately 16,000 yr BP (Benson et al. 2011; Oviatt 2015).

Although the evidence for an Ice Age origin was intriguing, given that Pleistocene horse remains have been previously reported from shoreline deposits along the Wasatch Front associated with Lake Bonneville (Milligan and McDonald 2017; Nelson and Madsen 1987), we identified several key aspects of the Lehi horse (reposited at the BYU Museum of Paleontology, specimen #BYU 43000, locality #2428) that raised questions about its age, paleontological associations, and context. The available excavation documentation (Figure 2) suggests a pit-like feature in the likely Pleistocene sands, partially infilled with a dark organic layer and horizontally bedded sands and silts (Supplemental Text 1). The presence of rip-up clasts overlying the horse remains supports at least some fluvial incision and deposition during or following burial. The feature’s irregular morphology led the original investigators to conclude that this pit was not produced by industrial machinery and was thus Pleistocene in origin (i.e., that the horse remains rest at the base of a small channel fill rather than in an anthropogenic pit). During excavations, however, investigators also recovered a human-modified chert flake (specimen BYU 43001) from the “overburden” (Figure 3)—either the fluvial sand and gravel deposits or organic overburden units shown in Figure 2. A large amount of cortex and large flake scars

Figure 2. Stratigraphic profile of the Lehi horse excavation, showing a probable pit incised into Pleistocene sands and filled with dark organic soil below horizontally bedded sands and muds, and overlain by recent fluvial deposits. (Color online)
indicate that this flake was probably produced during early-stage reduction. The precise provenience of this object was not recorded. To us, however, this indicated an Indigenous presence at the location of the Lehi horse after burial. Moreover, our preliminary examination of the horse’s vertebral column revealed osteophytes, cracking, and impinging spinous processes—pathological features that are not commonly observed in wild equids and that are typically associated with mounted horseback riding (Figure 4; Levine 1999; Li et al. 2020). These findings raised the possibility that the Lehi horse was historic or protohistoric rather than Pleistocene in age.

If the Lehi horse was indeed a misidentified, early historic horse managed by Indigenous peoples, it represents a unique opportunity to explore how horses were used, managed, and cared for in Native societies beyond the limited documentation in Euro-American historical records. Here, we apply osteological and biomolecular analyses to evaluate the age and domestic status of the Lehi horse, and we explore the implications of our findings for the sixteenth-century reintroduction of horses to the Americas.

Materials and Methods

Radiocarbon Dating
To assess the age of the Lehi horse, we selected an intact upper molar (the upper right M1, or 109 in the modified Triadan system) for radiocarbon dating at the Oxford Radiocarbon Accelerator Unit (ORAU) in Oxford, UK. The ORAU

![Figure 3. Modified broken chert flake recovered from the “overburden” during initial excavations of the Lehi horse (unprovenienced). (Color online)](image)

![Figure 4. Recovered bones and osteo pathologies: (a) stylized horse skeleton showing missing bones (gray), recovered bones (white), and recovered pathological bones (red); many teeth were recovered, but much of the head/neck was destroyed before excavation. Examples of diagnostic vertebral pathologies (b–d) linked with mounted horseback riding. These include (b) impinging dorsal spinous processes on thoracic vertebra 13, (c, d) undulating, irregular horizontal fissures in the articular surface on the caudal end of the centra, and a proliferation of bony exostoses on thoracic vertebrae (c) 16 and (d) 17.)](image)
performed routine pretreatment and measurement procedures (Brock et al. 2010), extracting dentine and combusting it in an elemental analyzer before graphitizing and measuring results on an HVEE accelerator, alongside blanks and standards, which were used to perform contamination calculations and data corrections. We calibrated our resulting radiocarbon measurement in OxCal using the IntCal13 calibration dataset (Reimer et al. 2013) to produce a calendrical estimate for the horse’s age.

**Osteological Study**

We assessed the age of the Lehi horse using dentition (Evans et al. 2006) and evaluated the skeleton for evidence of pathologies (Bartosiewicz and Gal 2013). We focused particularly on identifying health conditions or osteological changes indicative of—or related to—human activity, including vertebral pathologies associated with mounted horseback riding (Levine et al. 2005) and anthropogenic modifications to the dentition associated with human veterinary care (Taylor et al. 2018). Because the cranium of the horse had been shattered, we were unable to assess premolar changes caused by metal bit use (Bendrey 2007), nasal or premaxillary remodeling associated with bridle use (Taylor and Tuvshinjargal 2018) and heavy exertion (Taylor et al. 2015), or bone formation at the nuchal crest often associated with horseback riding or confinement (Bendrey 2008; Taylor et al. 2015).

**Isotope Analysis**

Analyses of the stable isotope composition of ancient horse remains have the potential to provide important environmental context and reveal aspects of forage, pastoral care, and movement. To shed light on the life history of the Lehi horse, we analyzed the carbon, oxygen, and strontium isotope chemistry of Lehi horse dentition, comparing these data to modern climate and environmental reference data from the region to assess how the horse moved during its lifetime and identify sources of dietary food and water.

\[ \delta^{13}C \]

**Background.** In the terrestrial ecosystems of the Intermountain West, the primary source of variation in stable carbon isotopes (\( \delta^{13}C \)) is the photosynthetic pathway (C₃, C₄, CAM) employed by native plant taxa. The distribution of C₃ versus C₄ taxa in the region is the result of complex interactions between a combination of environmental factors, including temperature, precipitation regime, soil moisture, snow cover, vegetation, and evapotranspiration—all of which vary significantly through space and across time (Notaro et al. 2010). In general, C₄ grasses and forbs are more abundant in the American Southwest, with high summer temperatures and a precipitation regime dominated by the North American Monsoon (NAM), whereas C₃ taxa dominate the Great Basin, with its cooler environments, significant winter precipitation, and associated spring snowmelt events (Cotton et al. 2016; Wer titine et al. 2015). Elevation has a less significant impact on C₃/C₄ abundance in this region than do temperature and seasonality of precipitation (Paruelo and Lauenroth 1996). Within the C₃ pathway, plants respond to reductions in water availability by stomatal closure, leading to a slight increase in leaf \( \delta^{13}C \) values (Farquhar et al. 1989). Environmentally driven shifts in intra-taxonomic plant \( \delta^{13}C \) caused by moisture stress are typically relatively minor—around 2‰ or less (Ehleringer 1989).

Current native grasses in the Lehi region consist primarily of C₃ taxa. Cotton and colleagues (2016) characterize the Lehi region as between 0% and 10% C₄ vegetation. Northern Utah is devoid of C₄ grasses, but C₄ forbs such as Atriplex—also consumed by horses—are present and begin greening in mid-spring, providing a seasonal source of higher \( \delta^{13}C \) values during this time. Although plants expressing the CAM photosynthetic pathway (Lee 2010) can have values overlapping those of C₄ plants, causing interpretive challenges in some settings (e.g., Smith et al. 2014), CAM plants in the study area consist largely of desert succulents unlikely to be consumed by grazers such as horses. Variation in forage \( \delta^{13}C \) is passed up the food web, with isotopic enrichment between diet and tooth enamel carbonates averaging +14.1‰ (±0.5) in a wide range of ungulate mammals (Cerling and Harris 1999:352) and +13.8‰ (±1.9) in horses specifically (Cerling and Harris 1999:349; see also Passey et al. 2005). Prior to fossil fuel depletion of atmospheric CO₂,
Fractionation during photosynthesis led to C3 and C4 grasses expressing an average $\delta^{13}C$ value of approximately $-25\%e$ and $-10\%e$ respectively—about $2\%e$ enriched over modern plant taxa (Dombrosky 2020).

$\delta^{18}O$ Background. Accurate interpretation of archaeological $\delta^{18}O$ relies on contextualization with modern and ancient climate data (e.g., Hamilton et al. 2018). $\delta^{18}O$ of environmental water is influenced by a suite of factors, including temperature, altitude, rainfall amount, aridity, seasonal precipitation patterns, and distance from the coast (Cerling et al. 2010). For example, as water-vapor masses move upslope, H$_2^{18}O$ preferentially rains out, accelerated by declines in temperature. H$_2^{18}O$ also rains out as water-vapor masses move inland, progressively depleting $\delta^{18}O$ values relative to coastal moisture, although this effect is less marked in cold (versus warm) precipitation events. Thus, the $\delta^{18}O$ value of water is influenced by temperature both within and across altitudinal and latitudinal gradients (Daux et al. 2005; Fricke and O’Neil 1999). Generally, colder winter temperatures and precipitation lead to lower water $\delta^{18}O$ values, whereas summer aridity and increased temperatures enrich water $\delta^{18}O$ (Dansgaard 1964). In the Lehi region, most precipitation is concentrated in winter and early spring (Figure 5). Here, deep snowpack with lower $\delta^{18}O$ values accumulates at elevations in the Wasatch range, reaching 3,650 m asl (~12,000 feet). Accordingly, as snowpack melts during late spring and early summer, modern $\delta^{18}O$ data from the Jordan River near Lehi (Coplen and Kendall 2000) exhibit consistently lower $\delta^{18}O$ values due to discharge of snowmelt into numerous alpine streams flowing downslope into Utah Lake and the Jordan River drainage.

Figure 5. (a) monthly temperature at Lehi, Utah (1,391 m asl), and at the nearby peak of Mount Timpanogos, roughly 15 km east of the Lehi site (3,582 m asl), showing warmest temperatures in the summer, June–August; (b) monthly total precipitation at Lehi and Mt. Timpanogos, showing very low precipitation year-round at Lehi itself and higher precipitation with a pronounced decrease in the summer months (June–August) at higher elevations; (c) stable oxygen isotope values for the Jordan River between 1984 and 1987 (Coplen and Kendall 2000), showing $\delta^{18}O$ depletion during spring snowmelt.
Although other processes, such as leaf evapotranspiration, may have minor impacts on the δ¹⁸O value of forage (Pederzani and Britton 2019), horses are obligate drinkers, deriving most of their moisture intake from imbibed water. During summer months, whether drinking from standing or flowing sources, imbibed water will likely express evaporatively enriched δ¹⁸O values relative to snowmelt—the primary source of spring and early-summer drinking water. Thus, in temperate regions, the δ¹⁸O values of obligate drinkers—such as horses—primarily reflect seasonally driven variation in δ¹⁸O values of water bodies (Levin et al. 2006; Pederzani and Britton 2019; Roberts et al. 2018). Significantly, the process of enamel mineralization and maturation allows chronologically ordered isotopic changes to be studied (Balasse 2002; Sponheimer et al. 2006), documenting seasonal patterns of movement or environmental conditions (Balasse 2002; Makarewicz et al. 2018). Although water cisterns may also bias archaeological δ¹⁸O values, there is no evidence of the use of such systems for horse management by Indigenous peoples (Ewers 1955). As a result, δ¹⁸O values from the Lehi horse should assess the input of local water sources and/or associated movements during the animal’s life.

**87Sr/86Sr Background.** Strontium isotope ratios (87Sr/86Sr) of tooth enamel have been used to track mobility based on the principle that bedrock of different ages and compositions have distinctive strontium isotope values that do not fractionate from the bedrock to the biosphere (Graustein and Armstrong 1983; Montgomery 2010). Older lithic formations have relatively higher 87Sr/86Sr in comparison to younger bedrock, while distinctive mineral content leads to characteristic 87Sr/86Sr based on rock “type” (Capo et al. 1998). The surface of Utah Valley is draped by Late Pleistocene Lake Bonneville–related sediments, and the bases of the surrounding mountains are covered with the lake’s shoreline deposits, wave-cut terraces, and related deltaic deposits and alluvial fans. The city of Lehi lies at the north end of Utah Valley, and the burial root juncture and the occlusal surface) of 80–90 mm or more (Levine 1982). Because our sample is derived from the lower portion of this tooth, we infer that our signal derives from the horse’s second year of life. The oblique orientation of enamel formation suggests that each sample taken perpendicular to the vertical axis of a tooth averages a short but thus far undocumented period of mineralization of up to several months (Hoppe et al. 2004). Horses typically complete the process of weaning by the age of roughly 8–9 months, although this process can be as short as 4 months in domestic animals and may last as long as a
year (Waran et al. 2008). Given that mammalian milk is elevated in δ^{18}O and lower in δ^{13}C relative to imbibed water and herbivore forage, respectively, the interpretation of equid stable isotope data can be confounded by enamel values laid down prior to weaning (Zazzo et al. 2002). This bias is particularly problematic with equid M1 data because the tooth mineralizes during the first two years of the animal’s life. The advanced age of the Lehi horse, however, means that the upper 40+ mm or so of tooth has been worn away through occlusal wear. We sampled the remaining 18 mm of enamel on a very worn M1 crown; this vertical section almost certainly mineralized in the second phase of tooth growth after weaning (Hoppe et al. 2004). As a result, although suckling clearly impacts both enamel δ^{13}C and δ^{18}O, and may bias strontium ratios, these processes are unlikely to have impacted our data.

Protocols. The lower right first molar (M1/409 in the modified Triadan system) was sequentially sampled for enamel stable carbon and oxygen isotope values (δ^{13}C and δ^{18}O) at the Stable Isotope Laboratory Facility in the Department of Archaeology, Max Planck Institute for the Science of Human History in Jena, Germany. Sampling was conducted at the base of the tooth root on the lingual margin, moving up at 2 mm intervals to a distance of 18 mm above the root, immediately below the occlusal surface of this heavily worn M1.

Enamel powder was pretreated by a wash in 1.5% sodium hypochlorite for 60 minutes, followed by three rinses in purified H_{2}O, before 0.1 M acetic acid was added for 10 minutes, followed by another three purified H_{2}O rinses (per Lee-Thorp et al. 2012; Roberts et al. 2017; Sponheimer et al. 2005). The resulting residues were freeze-dried overnight before reaction with 100% phosphoric acid. Gases evolved from the samples were analyzed for their stable carbon and oxygen isotopic composition using a Thermo GasBench II connected to a Thermo Delta V Advantage Mass Spectrometer at MPI-SHH. δ^{13}C and δ^{18}O values were compared against International Standards (IAEA-603 [δ^{13}C = 2.5; δ^{18}O = -2.4]; IAEA-CO-8 [δ^{13}C = -5.8; δ^{18}O = -22.7]; USGS44 [δ^{13}C = -42.2]) and in-house standard (MERCK [δ^{13}C = -41.3; δ^{18}O = -14.4]). Replicate analysis of MERCK standards indicates that machine measurement error is c. ± 0.1‰ for δ^{13}C and ± 0.2‰ for δ^{18}O. Overall measurement precision was assessed by measurement of repeat extracts from a bovid tooth enamel standard (n = 20, ± 0.2‰ for δ^{13}C and ± 0.2‰ for δ^{18}O).

We also sampled the same tooth in three locations for strontium isotope analysis—the base of the tooth root (0 mm), the middle of the sampled section (10 mm), and the crown (18 mm). Chemical sample preparation for strontium isotope analysis was performed in the clean chemistry laboratory of the MC-ICP-MS facility, Department of Geological Sciences, University of Cape Town. Strontium isotope ratio analysis was performed using the Nu Instruments Nu Plasma HR MC-ICP-MS in this facility (after Copeland et al. 2008). Twenty milligrams of the sampled tooth enamel powder were dissolved in 2 mL concentrated HNO_{3} in a closed Teflon beaker. The beaker was placed on a hotplate at 140°C for an hour, after which the sample was dried down and redissolved in 1.5 mL 2M HNO_{3} for strontium separation chemistry, following Pin and colleagues (1994). The separated strontium fraction was dried down, dissolved in 2 mL 0.2% HNO_{3} and diluted to 200 ppb strontium concentration prior to MC-ICP-MS isotope ratio analysis. Analyses of NIST SRM987 were used as a bracketing reference standard using a 87Sr/86Sr value of 0.710255. The strontium isotope data was corrected for isobaric rubidium interference at 87 amu using the measurement of 86Rb and the natural 85Rb/87Rb ratio. Instrumental mass fractionation was corrected using the measured 86Sr/88Sr ratio, the exponential law, and a true 86Sr/88Sr ratio of 0.1194. Analysis of in-house carbonate reference material processed and measured with samples from this study (87Sr/86Sr = 0.708908; 2σ = 0.000018) agreed well with long-term results (87Sr/86Sr = 0.708911; 2σ = 0.000040, n = 414).

DNA Analysis

Two independent DNA extractions were carried out at the ancient DNA research facilities of the Laboratoire AMIS CNRS UMR5288, Université de Toulouse III Paul Sabatier. DNA was extracted from 200 and 350 mg of bone powder

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following method Y from Gamba and colleagues (2016), with the slight modifications from Gaunitz and colleagues (2018). Aliquots of DNA extracts were treated with the USER enzyme mix (NEB), following Librado and colleagues (2017), in order to reduce the impact of post-mortem DNA damage in downstream analyses. Triple-indexed DNA libraries were constructed following the methodology from Fages and colleagues (2019), corresponding to the procedure described in Gaunitz and colleagues (2018) applied with the DNA adapters from Rohland and colleagues (2015; which include 7 bp long internal indexes). We amplified DNA libraries in 25 μL reactions using the AccuPrime Pfx DNA polymerase following Gaunitz and colleagues (2018) before concentrating them using Agencourt bead purification, quantifying them on a TapeStation 4200 instrument (Agilent technologies), and pooling them together with other triple-indexed DNA libraries for sequencing on the Illumina MiniSeq. Sequencing reads were demultiplexed based on their internal adapter indexes using AdapterRemoval 2, allowing for a single nucleotide mismatch at best on each internal index, before they were parsed through PALEOMIX v1.2.13 with default parameters, except that seeding was disabled for BWA (Li and Durbin 2009) mapping (version 0.7.17-r1194) in order to identify high-quality reads mapping uniquely against the horse nuclear reference genome EquCab2 and the horse mitochondrial genome (Accession # NC_001640). The resulting read alignment files were then processed with the Zonkey package (Schubert et al. 2017) to identify first-generation equine hybrids and determine the molecular sex, merging the sequence data from both experiments (Supplemental Text 2).

Results

Radiocarbon Dating

AMS radiocarbon dating of dentin from the Lehi horse lower-right first molar returned a calibrated 2σ range of AD 1681–1939 (127 ± 18 14C YBP, OxA-38552), with a median date of AD 1838. The dentine C/N ratio (3.2) and yield (9.5%) fell within standard quality-control parameters, providing good support for the radiocarbon date. These results demonstrate that the Lehi horse is historic, not Pleistocene in age. Although narrowing the timing of the burial is challenging because of the multimodal calibrated probability distribution and shape of the calibration curve over this time period, the co-occurrence of the burial with a modified stone tool suggests that it predates permanent Mormon settlement in the region (ca. 1850) and may fall within the earlier half of this calibrated range.

Osteological Study

The morphology of the incisor grinding table (which lacks enamel spots on the lower but not the upper incisors), the fusion of all identified epiphyseal surfaces, and the crown height measurements of available cheek teeth suggest that the Lehi horse was approximately 12–13 years old at the time of death (Table 1), but it is important to note that the reliability of wear-based age estimates can be influenced by factors such as environment and diet (Allen 2008). Unfortunately, no relevant portions of the cranium or premolars were available for study. Although we identified no canines, the destruction of the cranium makes it impossible to ascribe the absence of these teeth in the recovered assemblage to sex. The presence of marked enamel hypoplasia on the available cheek teeth is far from uncommon in wild or domestic equids, but it indicates some degree of nutritional stress during the animal’s lifetime.

We identified extreme levels of pathological bone formation on the vertebrae. This includes pathologic growth along the articular facets and ventral surface of vertebrae, particularly those of the lower back area most impacted by a seated rider (T16–18 and L1–L2), as well as irregular, undulating, transverse fractures of the caudal

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<tr>
<th>Tooth</th>
<th>Crown Height (mm)</th>
<th>Estimated Age Range (Levine 1982)</th>
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<tr>
<td>Upper right M1</td>
<td>39.91</td>
<td>12–13 years</td>
</tr>
<tr>
<td>Lower right M1</td>
<td>46.42</td>
<td>12–13 years</td>
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<tr>
<td>Lower left M1</td>
<td>45.06</td>
<td>12–13 years</td>
</tr>
<tr>
<td>Lower left M2</td>
<td>47.97</td>
<td>12–13 years</td>
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articulat surface on T17 and T18 (Figure 4). Although much of the vertebral column was destroyed prior to excavation, we found impinging spinous processes on the T13 at the contact with both the T12 and T14.

The specimen exhibits extreme arthritis of the lower limbs and other pathologies that severely impacted its mobility (Holson 2018). The Lehi horse exhibits such severe arthritis of the right rear tibia and tarsal joint that the surface of the astragalus is deeply pitted, and most of the tarsals have fused into a single bony mass (Figure 6). In addition to this rear limb issue, the right radius/metacarpal joint of the forelimb is also impacted by arthritis, and the horse exhibits laminitis of the front left distal phalanx and a fracture to the medial splint bone (the left second metacarpal). These features may be related in that the immobility of the right rear joint may have caused conformation issues (laminitis) and compensation in the forelimbs, adding stress that caused a fracture of the splint bone. Pathological features identified through osteological analysis are summarized in Table 2.

Isotope Analysis
Sequential sampling of M1 $\delta^{13}C$ ($n=10$) yielded an average value of $-10.4 \pm 0.6\%e$ with a $1.9\%e$ range of $-9.9$ to $-11.8\%e$ (Table 3). Assuming a fractionation constant of $+13.8\%e$, as noted above (Cerling and Harris 1999), dietary forage $\delta^{13}C$ ranged between $-25.6$ and $-23.7\%e$ with an average of $-24.2\%e$. Given a fractionation constant of approximately $+14$, a protohistoric horse consuming 100% C3 vegetation would exhibit an enamel $\delta^{13}C$ value of $c. -11\%e$, whereas a 100% C4 diet would yield a carbonate $\delta^{13}C$ value of approximately $+4\%e$ (Lee-Thorp 1989; Lee-Thorp et al. 1989; Levin et al. 2008). Estimates of reliance on C3 forage for the Lehi horse thus range between 100% and 91%, averaging 94.6% C3 intake. Average $\delta^{18}O$ is $-11.7 \pm 0.4\%e$ ($n=10$) with a 1.3\%e range of $-11.2$ to $-12.5\%e$. $\delta^{13}C$ and $\delta^{18}O$ are inversely related, and both datasets suggest seasonal shifts.

Figure 6. Examples of arthritic bone from the right hind leg. Severe arthritic bone formation and pitting of the posterior face of tibia (left) and the astragalus and tarsals (right), resulting in the complete immobility and fusion of many of the tarsal bones. (Color online)
in local forage and the temperature of imbibed water, respectively, across the period of enamel formation (Figure 7; Table 3). Strontium isotope values ($n = 3$; Figure 7) were consistent to the fourth decimal place ($0.710526$–$0.710570$) across the 18 mm sampled region (Figure 8), falling within an expected range for the greater Salt Lake region (Tipple et al. 2018) and in agreement with modern measured values from water sources in the Utah Lake region (Figure 1).

**Dental Analysis**

Combined, our two DNA extractions from the Lehi horse successfully yielded a total of 7,292 reads uniquely mapping to the reference mitochondrial and nuclear genomes of the domestic horse ($E. caballus$). The specimen yielded low but sufficient endogenous DNA to assess the sex, species, and first-generation hybrid status at virtually zero false-positive rates using the Zonkey pipeline (Schubert et al. 2017). These results provide clear indication that the animal is a domestic horse, $E. caballus$, and that the horse is female (Table 4).

### Discussion

**Early Historic Indigenous Horse Riding and Herding in the Great Basin**

Our results provide conclusive evidence that the Lehi horse is early historic rather than Ice Age in origin, and that this animal was used for mounted horseback riding, likely by Native peoples during the eighteenth or early nineteenth centuries. Intact horse teeth have proven to be the most reliable and contamination-resistant biological material for dating archaeological horse remains (Zazzo et al. 2019), and our results place the Lehi horse burial between approximately 1680 and the early twentieth century. The recovery of flaked stone within the excavation points to an association with Indigenous cultures for the Lehi horse, and the deep overburden of sediment may have been generated by

### Table 2. Pathological Features Identified in the Lehi Horse.

<table>
<thead>
<tr>
<th>Element</th>
<th>Pathological Features</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dentition</td>
<td>Enamel hypoplasia</td>
</tr>
<tr>
<td>Lower back (lower thoracic and lumbar vertebrae)</td>
<td>Impinging dorsal spinous processes</td>
</tr>
<tr>
<td>Right rear limb (tibia/tarsals)</td>
<td>Transverse fractures of the caudal articular surface</td>
</tr>
<tr>
<td>Front left limb (distal phalanx, second metacarpal)</td>
<td>Bony exostoses (left biased)</td>
</tr>
<tr>
<td></td>
<td>Arthritic pitting, eburnation, joint fusion</td>
</tr>
<tr>
<td></td>
<td>Healed splint bone fracture</td>
</tr>
<tr>
<td></td>
<td>Laminitis</td>
</tr>
</tbody>
</table>

### Table 3. Normalized Stable Isotope Measurements and Standard Deviation for the Lehi Horse, as Sampled from the Lingual Surface of the Lower-Right First Molar.

<table>
<thead>
<tr>
<th>Sample Number</th>
<th>Distance from Root (mm)</th>
<th>$\delta^{13}C$ ‰ (VPDB)</th>
<th>Standard Deviation</th>
<th>$\delta^{18}O$ ‰ (VPDB)</th>
<th>Standard Deviation</th>
<th>$^{87}Sr/^{86}Sr$</th>
<th>Estimated % C$_3$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>$-11.8$</td>
<td>0.1</td>
<td>$-11.2$</td>
<td>0.1</td>
<td>0.710526</td>
<td>100</td>
</tr>
<tr>
<td>1</td>
<td>2</td>
<td>$-11.1$</td>
<td>0.1</td>
<td>$-11.6$</td>
<td>0.1</td>
<td>—</td>
<td>100</td>
</tr>
<tr>
<td>2</td>
<td>4</td>
<td>$-10.3$</td>
<td>0.1</td>
<td>$-11.7$</td>
<td>0.1</td>
<td>—</td>
<td>94</td>
</tr>
<tr>
<td>3</td>
<td>6</td>
<td>$-10.1$</td>
<td>0.1</td>
<td>$-11.9$</td>
<td>0.1</td>
<td>—</td>
<td>93</td>
</tr>
<tr>
<td>4</td>
<td>8</td>
<td>$-10.0$</td>
<td>0.1</td>
<td>$-12.0$</td>
<td>0.1</td>
<td>—</td>
<td>92</td>
</tr>
<tr>
<td>5</td>
<td>10</td>
<td>$-10.0$</td>
<td>0.1</td>
<td>$-12.5$</td>
<td>0.1</td>
<td>0.710570</td>
<td>92</td>
</tr>
<tr>
<td>6</td>
<td>12</td>
<td>$-10.5$</td>
<td>0.1</td>
<td>$-12.1$</td>
<td>0.1</td>
<td>—</td>
<td>95</td>
</tr>
<tr>
<td>7</td>
<td>14</td>
<td>$-10.6$</td>
<td>0.1</td>
<td>$-11.7$</td>
<td>0.1</td>
<td>—</td>
<td>96</td>
</tr>
<tr>
<td>8</td>
<td>16</td>
<td>$-9.9$</td>
<td>0.1</td>
<td>$-11.5$</td>
<td>0.1</td>
<td>—</td>
<td>91</td>
</tr>
<tr>
<td>9</td>
<td>18</td>
<td>$-10.1$</td>
<td>0.1</td>
<td>$-11.3$</td>
<td>0.1</td>
<td>0.710550</td>
<td>93</td>
</tr>
</tbody>
</table>

**Notes:** Fractionation between enamel and dietary C$_3$ inputs calculated using a value of +13.8‰. Plant CO$_2$ concentrations in the Lehi region during the late prehistoric period were ∼2.0% more positive than today (Dombrosky 2020).
Euro-American farming and land-clearing activities after about 1850 (Morris and Rowe 2014). Based on ethnohistoric reconstructions, the area from which the Lehi horse was recovered would have straddled the boundary between Ute and Shoshone territory (Stewart 1970:209). The Northern Utah Ute acquired the horse as early as the seventeenth century, and they relied on it for their wide-ranging seasonal movements (Duncan 2003). The Shoshone obtained horses by the year 1700, likely from the Ute, and they played a major role in funneling horses to the northern Plains and Plateau (Mitchell 2015). Although our data do not allow us to determine whether this particular horse belonged to the Ute, the Shoshone, or both, our results do articulate with the ethnohistory of the region, given that the horse had become critically important to both Ute and Shoshone societies by the eighteenth century.

Analysis of the animal’s skeleton demonstrates that the horse was ridden, and it reveals health issues that imply a degree of human

Figure 7. Normalized stable isotope measurements for stable oxygen (closed circle), carbon (open circle), and strontium (closed triangle) for the Lehi horse, as sampled from the lingual surface of the lower-right first molar.

Figure 8. Lehi horse’s lower right Molar 1, showing sample locations and location of crown height measurement. (Color online)
protection or care. Damage to the vertebrae of the lower back—osteophytes, impinging spinous processes, and transverse fractures of the caudal articular surface—is characteristically associated with mounted riding, and it is generally absent from wild animals (Levine 1999; Levine et al. 2005). Although this type of damage is found in many ancient assemblages linked to horseback riding, it is particularly pronounced among archaeological horses predating the innovation of the frame saddle because riding bareback entails more direct trauma to the vertebral column (Levine et al. 2005). Among those vertebrae that yielded osteophytes and new bone formation (T5, T13, and T16-L2), this bone growth was universally asymmetric, in favor of more developed bone spurs on the animal’s left side (Figure 9).

We have identified similarly pronounced asymmetry in ridden-horse assemblages from Eurasia (Li et al. 2020). Asymmetric bias in equine skeletal damage may be related to rider handedness or riding technique (Taylor and Tuvshinjargal 2018), but it could also be influenced by consistent mounting from one side. Previously identified vertebral asymmetry in East Asian assemblages is clearly left biased (Levine et al. 2005; Li et al. 2020). Although many Native American horsemen are known historically to have mounted from the right side, it may be that this difference in weight bearing is caused by the handedness of the rider and the dynamics of mounted activities.

The pattern of pathological damage to the lower back observed in the Lehi horse clearly indicates that the animal was used for riding. Archaeological assemblages of animals used only for chariot traction, for example, show increased pathology in the forelimbs and neck region but little damage to the thoracic or lumbar vertebrae (Levine 2005; Weber 2008). This observation does not rule out the animal’s use for other purposes, however. Many Native American groups in western North America employed horses for pack transport (Ewers 1955), either with or without a travois (a pole-based sledge system adapted from existing technology used for dog transport). This system could be, and often was, used in tandem with a saddle and rider, which could have produced an osteological signature similar to that observed in the Lehi horse.

Extreme arthritis and other bone pathologies in the animal’s limbs provide further evidence of its use in transport. The fusion of most of the bones of the lower joint of the right rear limb would have given this horse extreme difficulty in movement. Any horse—wild or domestic—can suffer from these maladies, which are likely to limit an animal’s mobility severely. In the Lehi horse, these pathologies are severe, and they imply that the horse must have been cared for, or at least intentionally retained, even after reaching levels of impaired mobility. The issues with this joint may have radiated throughout the body as the animal experienced continued use. The horse also developed laminitis in a front hoof and fractured a splint bone, both potentially caused by compensation for the impaired hoof. These features suggest that, despite lameness,
the Lehi horse continued to be ridden (or perhaps used in transport in other ways, such as pulling loads in a travois). Indeed, the age of 12 seems remarkably young for a horse to have developed the observed level of pathology through natural aging processes. We argue that one of the only ways an animal with such severe mobility issues could have survived without dying from predation or starvation would be through human management or care.

The genomic identification of the Lehi horse as the domestic *E. caballus* and its identification as a female support its identification as a pastorally managed horse. Although the animal’s injuries would have drastically reduced its usefulness in transport, female horses continue to produce young annually until around 15 years of age (Garrott et al. 1991). As a result, archaeological assemblages related to pastorally managed breeding herds of horses rarely yield breeding-age females. Instead, they consist largely of elderly females (along with young males, who are culled before reaching breeding age; Levine 1999; Taylor 2017). The advanced age of the Lehi horse, and the co-occurrence of mobility-restricting pathological issues in the limbs suggest that the mare was either retained as a breeding animal until she died or that she was culled/sacrificed after reaching the end of her reproductive viability.

**Mobility and Movement**

Our lower stable oxygen isotope values likely correspond to a single, late-spring snow melt event, bracketed by slightly elevated δ¹⁸O values characteristic of lower-elevation precipitation, during the Lehi horse’s second year of life (Figures 4 and 7; Table 3). The lowest Lehi horse δ¹⁸O reading is paired with the highest δ¹³C value, and δ¹³C is loosely inversely related with δ¹⁸O across all but the last two samples taken at 16 mm and 18 mm from the enamel base. These data indicate a slight shift in the type of forage relied upon through winter into late spring—perhaps an increase in consumption of C₄ forbs beginning to green up in mid-spring, incrementally increasing the stable carbon isotope values of the Lehi horse. De Winter and colleagues (2016) also found a similar pattern of higher δ¹³C values with winter δ¹⁸O values in modern horses in Europe, which may be attributable to changes in plant water use efficiency or slight variations in plant choices during the winter season. Maize, also a C₄ domesticate, was farmed by prehistoric Fremont groups along the eastern rim of the Great Basin, but maize agriculture was abandoned by approximately AD 1150 in the Utah and Great Salt Lake Valleys, coincident with the end of the Medieval Climate Anomaly (Coltrain and Leavitt 2002). Prior to pioneer settlement in 1847, maize was not cultivated by indigenous Ute or Shoshone populations. Thus, foddering with maize is unlikely to have made a significant contribution to the Lehi horse’s very slightly elevated spring δ¹³C values.

Historically, the Ute are known to have made fall and winter seasonal movements between
low-elevation wintering grounds and high-elevation summer hunting grounds, as well as between territories to the south and east, ranging as far as eastern Colorado and western Kansas, and hunting in New Mexico, Arizona, and the Texas and Oklahoma panhandles. It is possible that Indigenous pastoralism of equids included moving herds south from Utah Valley during winter months, but neither strontium nor stable oxygen isotope values for the Lehi horse support this interpretation for this particular individual. Strontium signatures do not vary seasonally, but they are consistent at the fourth decimal place (0.710526–0.710570). Moreover, all observed values fall within an expected range for the greater Salt Lake region (Tipple et al. 2018), providing no evidence that carbon and oxygen isotope patterning was the consequence of nonlocal movement.

**Conclusion**

Our results demonstrate that the Lehi horse is indeed an early historic domestic horse, likely managed and used by Indigenous people. Osteological pathologies indicate that the Lehi horse was ridden, perhaps bareback or using a soft pad saddle. The animal’s use continued even after it developed issues with the right rear limb, which caused ancillary effects in other limbs—including arthritis, a fracture, and laminitis. Pathological bone growths on the vertebrae suggest that the horse experienced consistently higher loads on its left-hand side, a pattern that is known from other archaeological assemblages and is characteristic of ridden horses. DNA confirms the identification of the Lehi horse as *Equus caballus*, not an Ice Age wild equid, and reveals that the animal was a mare. This is consistent with expectations for a pastorally managed herd animal, and it explains the apparent care for the horse into advanced age despite severe mobility issues. Stable isotopes sampled from the horse’s enamel point to seasonal shifts in the isotope chemistry of imbibed water and available vegetation, which—in tandem with minimal variance in strontium isotope—suggests that the horse was raised and tended locally within the Wasatch Front near where it was found.

This study provides an example of how to engage data quality in zooarchaeology (e.g., Wolverton 2013) as well as a template for the integration of morphological and biomolecular approaches (Steele 2015)—subjects that have been a key focus of discussion in archaeozoological circles in recent years (e.g., Jones and Gabe 2015; Kansa et al. 2019; LeFebvre et al. 2019; Nims and Butler 2019). Our identification of the Lehi horse as an early domestic rather than an Ice Age horse suggests that prior misclassifications may have influenced museum collection practices and the interpretation of archaeological and paleontological assemblages, leading to gaps in the faunal record of Native horsemanship. Consequently, the reevaluation of horse skeletons in natural history collections, focusing on both osteological markers of human activity and biomolecular analyses, appears warranted—and such studies may significantly change our understanding of the timing and nature of early Indigenous horse use in the Americas.

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**Data Availability Statement.** All data collected during this research are provided in the manuscript and supplemental materials.

**Supplemental Materials.** For supplemental material accompanying this article, visit https://doi.org/10.1017/aaq.2020.109.

Supplemental Text 1. Lehi Horse Stratigraphic Description.


Supplemental Figure 1. Horse burial from the early seventeenth-century Saxman site in Kansas, informally excavated and disposed of during the 1970s, on the assumption that it was modern.

Supplemental Figure 2. Still photo from a video recorded during excavation; photos of the subsurface stratigraphy during excavation; photos of the subsurface stratigraphy directly adjacent to the horse remains during excavation.
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