Deglacial landscapes and the Late Upper Palaeolithic of Switzerland

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Abstract:
The presence of people in Switzerland in recently deglaciated landscapes after the Last Glacial Maximum represents human utilisation of newly available environments. Understanding these landscapes and the resources available to the people who exploited them is key to understanding not only Late Upper Palaeolithic settlement in Switzerland, but more broadly human behavioural ecology in newly inhabited environmental settings. By applying bone collagen stable isotope analysis ($\delta^{13}$C, $\delta^{15}$N and $\delta^{34}$S) to faunal remains from Late Upper Palaeolithic localities in Switzerland, we investigate animal ecology and environmental conditions during periods of human occupation. High and relatively uniform $\delta^{34}$S values indicate that landscapes north of the Jura Mountains provided comparatively stable environmental conditions, while lower and more variable $\delta^{34}$S values on the Swiss Plateau suggest a dynamic landscape with diverse hydrological and pedological conditions, potentially linked to regionally different patterns of permafrost thaw. This contrasts with the archaeological record that appears relatively uniform between the two regions, suggesting people were employing similar subsistence behaviours across a range of environmental settings. The pattern of change in $\delta^{15}$N across the deglacial period appears consistent between areas that remained ice-free throughout the LGM and those that were glaciated. Most notable is a period of exclusively low $\delta^{15}$N values between 15,200 and 14,800 cal. BP, which could relate a regional expansion of floral biomass in response to environmental change.

Keywords: Magdalenian, collagen, sulphur isotopes, nitrogen isotopes, carbon isotopes, Pleistocene, palaeogeography, Europe, horse, reindeer
1. Introduction

The period after the Last Glacial Maximum (LGM) in Switzerland saw significant expansion of human settlement into previously ice-covered landscapes. This was facilitated by the rapid development of pioneer floral and faunal communities, and most likely by changes in the range dynamics of key prey species, such as horse (Equus sp.) and reindeer (Rangifer tarandus) (Leesch et al., 2012; Cupillard et al., 2015). Understanding the environmental developments that drove such changes is key to interpreting the subsistence and settlement patterns of the human populations in this region and, more broadly, to investigating human behavioural ecology in newly available landscapes. Here we examine post-LGM paleoenvironmental conditions at two archaeological localities; Kastelhöhle-Nord (Figure 1), which is situated north of the Jura Mountains and remained ice-free throughout the LGM, and Monruz and Champréveyres (Figure 1), which are situated on the Swiss Plateau and became ice-free before 17,500 cal. BP (Ivy-Ochs et al., 2004). Bone collagen stable isotope analyses (δ¹³C, δ¹⁵N, and δ³⁴S) are applied to the archaeological faunal assemblages from these sites to provide a direct record of past prey species ecology and environmental conditions during the period of human activity. New radiocarbon dates are also obtained from the fauna to better contextualise the chronology of human occupation. By combining this data with other published results, the pattern of post-LGM environmental change is compared between regions that remained ice-free throughout the LGM and those that were glaciated.

1.1 Post-LGM environment and archaeology in Switzerland

Switzerland during the LGM was almost entirely covered by ice with only a small region north of the Jura Mountains remaining unglaciated (Ivy-Ochs et al., 2004; Bini et al., 2009; Ivy-Ochs 2015). While a brief phase of human occupation during the LGM occurred in the ice-free region (Terberger and Street, 2002; Reade et al., 2020), it is not until after the LGM that evidence of widespread and sustained Late Upper Palaeolithic human activity becomes apparent (Leesch et al., 2012). Post-LGM ice sheet decay in Switzerland was rapid, and the entire central Swiss Plateau was ice free before the onset of Greenland Stadial 2.1a (GS-2.1a, c.17,450 – 14,650 BP, Figure 2; Ivy-Ochs et al., 2004; Rasmussen et al., 2014). The exposure of new landscapes during Greenland Stadial 2.1b (GS-2.1b, c. 20,850 to 17,450 BP, Figure 2; Rasmussen et al., 2014) was quickly followed by the development of pioneer floral communities. Vegetation on the northern margins of the Jura Mountains was dominated by Poaceae, Artemisia, Juniperus and Hippophae (Cupillard et al., 2015), while species-rich treeless steppe tundra had developed on the Swiss Plateau by around 18,700 cal. BP (Rey et al., 2017). This was followed by an increase in herbaceous vegetation (e.g. Ammann and Lotter, 1989; Lotter, 1999; Wehrli et al., 2007) that allowed
large herbivores to recolonise deglaciated regions by around 17,000 cal. BP (Morel and Hug, 1996; Hajdas et al., 2007).

There is some evidence for a relatively early phase of post-LGM human activity early in GS-2.1a, but the main expansion of settlement in Switzerland did not occur until the latter part of GS-2.1a (Weniger, 1989; Napierala, 2008; Leesch et al., 2012; Maier, 2015). These sites, associated with the Magdalenian culture, are characterised by a dominance of horse and reindeer in their faunal assemblages (Leesch et al., 2012; Nielsen, 2013; Maier, 2015). Attributed to Magdalenian techno-complexes D and E, the sites cover a wide geographic distribution across the Swiss Plateau and Jura region, indicating the exploitation of a variety of landscapes (Leesch et al., 2012). Archaeological evidence comes from both caves and rockshelters, such as at Kesslerloch and Kastelhöhle-Nord, and open-air localities, such as the sites of Monruz and Champréveyres (Leesch et al., 2019). Coleoptera-based temperature estimates suggest summer and winter mean air temperatures on the Swiss Plateau during GS-2.1a were around 9°C and –20°C respectively (Coope et al., 2000; Thew et al., 2009), while summer temperatures of around 12°C are estimated for the region north of the Jura mountains (Cupillard et al., 2015). Pollen and plant macrofossil evidence attest to vegetation rapidly increasing in diversity, but still dominated by cold-tolerant herbaceous species (Thew et al., 2009; Cupillard et al., 2015).

The onset of Greenland Interstadial 1 (GI-1, c. 14,650 to 12,850 BP, Figure 2; Rasmussen et al., 2014) corresponds to a rapid rise in temperatures on the Swiss Plateau and the expansion of juniper and birch vegetation into open shrub and grasslands (Thew et al., 2009). It is currently not clear whether Magdalenian activity continued into this early phase of GI-1e or ended with the onset of this warm period, but by c. 14,400 cal. BP Azilian occupation of the Swiss Plateau was established (Leesch et al., 2012). During this time, mean summer and winter temperatures are estimated to have been around 15°C and 0°C respectively, with vegetation composed of a mosaic of open birch woodland, patches of dwarf birch and shrubs, and areas of grasses and sedges (Leesch, 1997; Thew et al., 2009). These more temperate conditions are reflected in the change in subsistence focus, with red deer (Cervus elaphus) and horse being important prey species (Leesch et al., 2004; Nielsen, 2013; Maier, 2015).

Considering the availability of different prey species and the ecologies and environments they represent is key to understanding the landscapes past people would have experienced. Whilst there is a wealth of Swiss Lateglacial palaeoenvironmental data from numerous lake and mire archives (e.g. Lotter, 1999; Coope et al., 2000; Wehrli et al. 2007; Lotter et al., 2012; Cupillard et al., 2015; Rey et al., 2017), analysis of the archaeological faunal assemblage provides the means to directly link inferences on habitat, ecology, and palaeoenvironmental conditions, to human hunting, settlement and subsistence practices.
1.2 Paleoenvironmental records from faunal stable isotopes

Palaeoenvironmental and ecological data can be obtained directly from skeletal remains of prey species from archaeological contexts through stable isotope analyses (e.g. Stevens and Hedges, 2004; Stevens et al., 2008, 2014; Drucker et al., 2011a; 2011b; 2012; Bocherens et al., 2015; Jones et al., 2018; Reade et al., 2020). In this study we use carbon (δ\(^{13}\)C), nitrogen (δ\(^{15}\)N) and sulphur (δ\(^{34}\)S) isotope ratios in bone collagen to explore post-LGM environments and prey species ecology in Switzerland. The measured isotopic signals are underpinned by dietary specialisation, animal behaviour and environmental conditions.

Bone collagen δ\(^{13}\)C values are largely determined by species-specific dietary behaviours, such as grazing versus browsing, or specific dietary specialisations, such as the consumption of lichens by reindeer, which leads to systematically higher δ\(^{13}\)C values in comparison to other herbivore species (Drucker et al., 2010; Bocherens et al., 2015). However, dietary δ\(^{13}\)C values also reflect atmospheric CO\(_2\) concentration and δ\(^{13}\)C, environmental variables such as temperature and moisture availability, and vegetation density and type (Heaton, 1999; Stevens and Hedges, 2004; Drucker et al., 2008; Kohn, 2010). Faunal δ\(^{15}\)N values are linked to both dietary specialisation/niche position and to climatic parameters, such as temperature and precipitation, mediated through soil processes (Amundson et al., 2003; Stevens et al., 2008; 2014; Drucker et al., 2011b; 2012; 2018 Craine et al 2015; Rabanus-Wallace et al., 2017). In particular, nutrient availability and microbial activity may be reflected in herbivore bone collagen δ\(^{15}\)N values, parameters that were likely strongly influenced by the presence of permafrost and ice sheets in the European Lateglacial (Stevens and Hedges, 2004; Stevens et al., 2008; Drucker et al., 2011b). Collagen δ\(^{34}\)S values relate to the soil environment upon which the animal fed. Bioavailable sulphur can be derived from sulphates in groundwater and rain, atmospheric sulphur, and from mineral weathering of the underlying geology (Nehlich, 2015). As such, bone collagen δ\(^{34}\)S values are spatially variable and often considered a tool for exploring mobility and landscape utilisation (e.g. Drucker et al., 2012; 2018; Jones et al., 2018; Wißing et al., 2019).

However, bone collagen δ\(^{34}\)S values may also hold significant promise as a palaeoenvironmental proxy, as soil-bedrock interactions, mineral weathering, and sulphur cycling in the soil are driven by hydrological and microbial processes (Thode, 1991). These are dynamic systems influenced by climatic and environmental conditions, such that δ\(^{34}\)S values in the local landscape are unlikely to have remained static across major environmental transitions, for example the last deglaciation.

2. Materials and Methods
In this study we present $\delta^{13}$C, $\delta^{15}$N and $\delta^{34}$S data generated from horse, reindeer, red deer and $Bos/Bison$ bone collagen samples from the Swiss sites of Kastelhöhle-Nord, Monruz and Champréveyres to explore animal ecology and environmental conditions after the LGM.

2.1 Archaeological samples

The faunal assemblage from Kastelhöhle-Nord provides a record from the ice-free region of Switzerland, on the north edge of the Jura Mountains. Excavation of the cave between 1948 and 1954 revealed an ‘intermediate’ horizon associated with a Badegoulian phase of occupation, dated to the latter part of the LGM (Terberger and Street, 2002; Reade et al., 2020), and a 25cm-thick ‘upper’ horizon associated with the post-LGM Magdalenian (Leesch et al., 2012). This upper horizon contained a rich lithic assemblage that certainly represents more than one phase of Magdalenian activity at the site and most likely comprises more than one techno-complex (Magdalenian D-a and E; Leesch et al., 2012). A relatively long duration of accumulation for this horizon was confirmed by three faunal radiocarbon dates, which range from 16,350–15,965 cal. BP (ETH-45024) to 14,265–13,967 cal. BP (ETH-45026) (Leesch et al., 2012; Figure 2, Table 1). We sampled the large herbivore species found within the upper horizon (Schweizer, 1959): reindeer ($n=10$), horse ($n=5$) and $Bos/Bison$ ($n=6$). Of these, one reindeer bone displayed evidence of anthropogenic impact (cut marks). To investigate the chronology of this horizon, 3 specimens (two horse and the cut-marked reindeer bone) were selected for radiocarbon dating. The three specimens that had previously been dated (two reindeer, one $Bos/Bison$) were also re-dated to ensure methodological consistency between laboratories.

Comparative post-LGM samples from the Swiss Plateau, which was glaciated during the LGM, come from the open-air localities of Monruz and Champréveyres. Excavated between 1984 and 1992, both sites produced Late Upper Palaeolithic Magdalenian and Azilian occupation horizons, with rich faunal, botanical and lithic assemblages. Phases of Magdalenian activity at the sites date to the later part of GS-2.1a and were focused primarily on the exploitation of horse, together with a broad spectrum of smaller mammals, birds and fish, particularly in spring and summer (Müller et al., 2006; Müller, 2013). Three horse bones from these assemblages have previously been radiocarbon dated, producing age determinations between 15,874–15,349 cal. BP (OxA-20699) and 15,585–15,053 cal. BP (OxA-20701) (Figure 2, Table 1). We selected five reindeer and 12 horse samples from Monruz (all from Magdalenian sector 1) and 7 red deer samples from Champréveyres (four from Magdalenian sector 2, two from Azilian sector 1, and one from sector 2 where the association to the Magdalenian or Azilian was uncertain) for stable isotope analysis. While the specimens do not bear direct traces of anthropogenic action, the large herbivore faunal
assemblage from the sites is confidently interpreted as the product of human action (Müller et al., 2006; Müller, 2013). While Magdalenian activity has been dated both from faunal and charcoal remains, dating of the Azilian phase has so far relied solely on charcoal samples. Therefore, one bone from the Azilian concentration at Champréveyres was selected for radiocarbon dating as part of this study.

2.2 Sample pre-treatment

A small sample of bone (0.2 to 1.3g) was collected from each specimen using a dental drill with either a small cutting wheel or tungsten burr attachment. Samples were prepared at University College London (UCL) using a modified version of the Oxford Radiocarbon Accelerator Unit (ORAU) collagen extraction procedures (AF and AG methods; Brock et al., 2010), which is based on a modified version of the Longin (1971) method. All samples were treated with 0.5M hydrochloric acid (HCl) at 4°C until fully demineralised and then thoroughly rinsed with ultrapure water. Some samples were then also treated with 0.1M sodium hydroxide (30mins), and 0.5M HCl (1hr) to remove humic contaminants (Szpak et al., 2017), again being thoroughly rinsed with ultrapure water between reagents. All samples were then gelatinised in pH3 HCl solution at 75°C for 48hrs and filtered using a pre-cleaned Ezee-filter. For most samples, including all those to be radiocarbon dated, the filtrate was then passed through a pre-cleaned 15–30 kD ultrafilter, with the >30 kD fraction collected and freeze-dried (AF method). For some samples the ultrafiltration step was omitted (AG method); while ultrafiltration has been shown to successfully improve the removal of contaminants that can influence radiocarbon determinations (Higham et al., 2006), it also significantly reduces collagen yield, while at the same time producing little difference in measured stable isotope compositions (Sealy et al., 2014; Szpak et al., 2017). Details of pre-treatment methodology are given for each sample in the supplementary information S1.

2.3 Stable isotope analysis

Collagen yields from Kastelhöhle-Nord ranged from 2.3 to 16.2%. Collagen preservation at Monruz and Champréveyres was poorer; 12 out of 24 samples failed to produce enough collagen for stable isotope analysis (yields ≤0.7%), while the other 12 samples had collagen yields ranging from 0.8 to 3.6% (supplementary information S1). Samples with adequate collagen were analysed for their nitrogen (δ¹⁵N), carbon (δ¹³C), and sulphur (δ³⁴S) isotopic ratios at the Scottish Universities Environmental Research Centre (SUERC). 1.2–1.5mg aliquots of freeze-dried collagen were weighed into tin capsules and analysed using a Delta V Advantage continuous-flow isotope ratio mass spectrometer coupled via a ConfloIV to an EA IsoLink elemental analyser (Thermo Fisher Scientific, Bremen). For every ten archaeological samples, three in-house standards, calibrated to the
International Atomic Energy Agency (IAEA) reference materials, were analysed (Sayle et al., 2019). Results are reported as per mil (‰) relative to the internationally accepted standards VPDB, AIR and VCDT. Measurement uncertainty was determined to be ±0.1‰ for δ^{13}C, ±0.2‰ for δ^{15}N, and ±0.3‰ for δ^{34}S, on the basis of repeated measurements of an in-house bone collagen standard and a certified fish gelatin standard (Elemental Microanalysis, UK). Standard quality control criteria were used to assess the δ^{13}C, δ^{15}N and δ^{34}S data (DeNiro, 1985; Ambrose, 1990; Nehlich and Richards, 2009). Each sample was analysed in duplicate and reproducibility was better than ±0.1‰ for δ^{13}C, ±0.2‰ for δ^{15}N and ±0.3‰ for δ^{34}S. All analysed samples had C:N atomic ratios between 3.2–3.6, and %C and %N between 35–46% and 12-16%, respectively, indicating good bone collagen preservation (DeNiro, 1985; Ambrose, 1990). All analysed samples except UPN-240 had C:S and N:S atomic ratios within the recommended ranges of 600 ± 300 and 200 ± 100, and %S content between 0.14 and 0.30% (Nehlich and Richards, 2009).

### 2.4 Radiocarbon analysis and background corrections

Radiocarbon dating was performed at ORAU using their standard procedures, as described by Brock et al. (2010). For each sample, approximately 5mg of dry collagen, which had been weighed into tin capsules baked at 500°C for 12 hours, was combusted using an elemental analyser coupled to an isotope ratio mass spectrometer, employing a splitter to allow for collection of the CO₂ (Bronk Ramsey and Humm, 2000; Brock et al., 2010). Samples were graphitised by reduction of collected CO₂ over an iron catalyst in an excess H₂ atmosphere at 560°C (Bronk Ramsey and Hedges, 1997; Dee et al., 2010). ¹⁴C dates were measured on the Oxford AMS system using a cesium ion source for ionisation of the solid graphite sample (Bronk Ramsey et al., 2004). To denote the bone pretreatment at UCL rather than at ORAU, all measured dates were given “OxA-V-wwwww-pp” numbers, where “wwwww” indicates the wheel number, and “pp” is the position of the sample on the wheel (Brock et al., 2010). As collagen extraction was performed at UCL according to the ORAU pretreatment protocol, background corrections were applied to our dates based on repeat AMS measurements at ORAU of known-aged reference samples prepared in the UCL laboratory, following the method outlined by Wood et al. (2010). A full description of our correction methodology is detailed in Reade et al. (2020). Corrected dates are denoted by adding a “C” to the end of the date code assigned by ORAU. Uncorrected measured date values as well as further details of the correction calculations are provided in the supplemental information S2.

### 3. Results and Discussion
3.1 Chronology of the Lateglacial assemblage at Kastelhöhle-Nord and Champréveyres

Six new radiocarbon determinations were made on fauna from the upper horizon at Kastelhöhle-Nord (Figure 2, Table 1). Three of these were undertaken on previously dated specimens for inter-laboratory comparison, and the results reflect those previously obtained (Leesch and Müller, 2012) (Table 1). The new dates for Kastelhöhle-Nord’s upper horizon range from 13,550 ± 60 14C BP (OxA-V-2794-25C) to 12,200 ± 50 14C BP (OxA-V-2793-56C), giving a range of calibrated ages between 16,350 – 15,965 cal. BP and 14,260 – 13,935 cal. BP, further confirming an extended period of bone accumulation in the horizon. There appears to be a species-based chronological pattern to the results (Figure 2);

*Bos/Bison* and horse date to GS-2.1a and reindeer to GI-1. However, this finding is likely coincidental as reindeer dating to GS-2.1a are known at other sites within the same valley system (e.g. Hollenberg-Höhle 3; Müller and Leesch, 2011), as are horse dating to GI-1 (e.g. Kohlerhöhle; Leesch and Müller, 2012). Thus, both species were present within the local landscape at the same time as one another.

Archaeologically, the dating of a cut-marked reindeer bone found in association with Magdalenian artefacts to the later, colder part of GI-1ed is problematic when considered against our current understanding of the chronology of Magdalenian/Azilian development and their subsistence systems in the region. While it is recognised that the Kastelhöhle-Nord upper horizon lithic assemblage most probably represents more than one Magdalenian techno-assemblage (Leesch et al., 2012), the reindeer date is significantly later than those at other Magdalenian sites in the region and also later than dates associated with the Early Azilian assemblages at Monruz and Champréveyres. Therefore, it appears reindeer hunting at Kastelhöhle-Nord continued or resumed after the Magdalenian disappeared from the archaeological record in adjacent regions. As the bone was found in the mixed Magdalenian techno-assemblage the question arises whether this lithic tradition survived or was revived with the hunting of reindeer during GI-1ed, or whether there is a yet unrecognised lithic assemblage type admixed in the upper horizon. Comparable late GI-1ed radiocarbon dates on faunal remains (horse, ptarmigans, red deer and dog) have come from the nearby cave of Büttenloch, and from other northern Swiss localities, namely Rislisberghöhle and Kesslerloch (Napierala, 2008; Leesch and Müller, 2012). Chronologically, the dates are compatible with the Azilian phase in Switzerland, although the dated fauna cannot be certainly attributed to the subsistence activities of this culture.

One new radiocarbon determination was made on a red deer bone from the Azilian horizon at Champréveyres, which had so far been chronologically constrained by radiocarbon determinations made on charcoal and macrobotanical remains (Leesch, 1997). The date, 12,480 ± 50 14C years BP (OxA-V-2754-49C, Figure 2, Table 1), falls within the
range of dates on charcoal from hearth deposits, confirming the contemporaneity of the faunal assemblage with human activity at the site.

3.2 Lateglacial ecology and environment at Kastelhöhle-Nord and Monruz/Champréveyres

Kastelhöhle-Nord upper horizon $\delta^{13}C$ values overlap for reindeer and Bos/Bison, (–20.3‰ to –19.4‰ and –20.1‰ to –19.7‰, respectively) while horse $\delta^{13}C$ values differ (–21.3‰ to –20.2‰) (Figure 3, Table 2). A similar offset between reindeer and Bos/Bison, and horse is also observed in the nitrogen isotopic data, which ranges from 0.7‰ to 2.3‰ for horse, compared to 2.3‰ to 3.5‰ and 2.6‰ to 4.0‰ for reindeer and Bos/Bison respectively (Figure 3, Table 2). The significant species-based differences between $\delta^{13}C$ and $\delta^{15}N$ values (Figure 3, supplementary information S3.1 and S3.2) can largely be explained by species ecology and dietary specialisation. The comparable Bos/Bison and reindeer $\delta^{13}C$ values indicate ecological overlap. Reindeer typically display higher $\delta^{13}C$ values than other herbivore species due to lichen consumption; this dietary behaviour has also been observed in some modern bison populations and has previously been suggested for the species in other Late Pleistocene contexts (Larter and Gates, 1991; Julien et al., 2012; Bocherens et al., 2015). The two species also display similar $\delta^{15}N$ and $\delta^{34}S$ values (Figure 3, Table 2), further supporting the interpretation of overlapping habitat preferences. Lower $\delta^{13}C$ and $\delta^{15}N$ values in horse compared to reindeer is a pattern that is observed across Pleistocene Europe and indicates occupation of a different ecological niche (Stevens and Hedges, 2004; Stevens et al., 2008; Bocherens et al., 2015). The $\delta^{34}S$ values observed for horse also differ from those of reindeer and Bos/Bison (Figure 3, Table 2), further suggesting that the animals were not only occupying different niches, but also possibly different landscapes and/or different topographical features/locations within a given area. However, it should be noted that the species-dependent chronological pattern observed in the radiocarbon dates (Figure 2) means that it cannot be certainly demonstrated that any of the individuals analysed here overlap in their chronology, and as such, the differences observed may partly be influenced by temporally changing underlying environmental parameters. Pollen records indicate the vegetation north of the Jura mountains was dominated by a mosaic of Poaceae, Artemisia, Juniperus and Hippophae species during GS-2.1a, with Juniperus and Betula expanding at the start of GI-1e, while climatic proxies suggest both an increase in temperature and precipitation across this time interval (Cupillard et al., 2015). Thus, the observed isotopic data likely represents a combined signal of environmental change and species-specific patterns of habitat utilisation.

A similar pattern of variation between horse and reindeer is seen in the Monruz results (Figure 4). At this site average horse $\delta^{15}N$ values (1.8 ± 0.6‰) and $\delta^{13}C$ values (–21.3‰ to –20.2‰) indicate ecological overlap. Reindeer typically display higher $\delta^{13}C$ values than other herbivore species due to lichen consumption; this dietary behaviour has also been observed in some modern bison populations and has previously been suggested for the species in other Late Pleistocene contexts (Larter and Gates, 1991; Julien et al., 2012; Bocherens et al., 2015). The two species also display similar $\delta^{15}N$ and $\delta^{34}S$ values (Figure 3, Table 2), further supporting the interpretation of overlapping habitat preferences. Lower $\delta^{13}C$ and $\delta^{15}N$ values in horse compared to reindeer is a pattern that is observed across Pleistocene Europe and indicates occupation of a different ecological niche (Stevens and Hedges, 2004; Stevens et al., 2008; Bocherens et al., 2015). The $\delta^{34}S$ values observed for horse also differ from those of reindeer and Bos/Bison (Figure 3, Table 2), further suggesting that the animals were not only occupying different niches, but also possibly different landscapes and/or different topographical features/locations within a given area. However, it should be noted that the species-dependent chronological pattern observed in the radiocarbon dates (Figure 2) means that it cannot be certainly demonstrated that any of the individuals analysed here overlap in their chronology, and as such, the differences observed may partly be influenced by temporally changing underlying environmental parameters. Pollen records indicate the vegetation north of the Jura mountains was dominated by a mosaic of Poaceae, Artemisia, Juniperus and Hippophae species during GS-2.1a, with Juniperus and Betula expanding at the start of GI-1e, while climatic proxies suggest both an increase in temperature and precipitation across this time interval (Cupillard et al., 2015). Thus, the observed isotopic data likely represents a combined signal of environmental change and species-specific patterns of habitat utilisation.

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21.0 ± 0.2‰) are significantly different to the reindeer values (2.8 ± 0.2‰ and −19.8 ± 0.2‰, respectively) (S3.3). Furthermore, δ^{13}C and δ^{15}N values for horse at Monruz and Kastelhöhle-Nord upper horizon are indistinguishable from one another; the same is also observed for reindeer from the two sites (S3.4 and S3.5). By contrast, horse and reindeer δ^{34}S values at Monruz are statistically indistinguishable from one another (Figure 4; S3.6), but significantly different to those of the horse and reindeer at Kastelhöhle-Nord (S3.7).

Therefore, δ^{13}C and δ^{15}N values cluster by species, indicating that ecological niche and animal behaviour are the primary factors influencing these values, while δ^{34}S values cluster by site, indicating that location-based factors are most strongly represented in the sulphur isotope ratios. Almost twice as much variability is observed in the Monruz δ^{34}S values (16.5‰) than in the Kastelhöhle-Nord δ^{34}S values (8.8‰), despite the Monruz samples representing a significantly shorter time span (Figure 2) and representing only two, rather than three, different animal species (Table 2). Further, unlike at Kastelhöhle-Nord, the horse and reindeer δ^{34}S values from Monruz completely overlap with one another. Thus, while the Monruz δ^{13}C and δ^{15}N values suggest that horse and reindeer were occupying different ecological niches, the sulphur isotopic data suggests this was likely taking place within the same geographical region(s), and therefore under the same range of environmental and climatic conditions. The large range in δ^{34}S values could indicate both species were displaying a number of different long-distance mobility behaviours, or alternatively, could suggest a high level of environmental variability within a relatively small geographical region.

As the Monruz horse produce a significantly greater range of δ^{34}S values (16.2‰) than the reindeer (7.0‰), and unlike reindeer, horse generally do not undertake long-distance seasonal migrations, we suggest the second interpretation is more plausible. However, we recognise mobility behaviours may have differed between different environments in Late Pleistocene Europe (e.g. Bignon et al., 2005; Pelligrini et al., 2008; Pryor et al., 2016).

The isotopic composition of bioavailable sulphur is spatially variable at a range of scales. At the regional level, soil δ^{34}S values are controlled by location-based inputs from underlying bedrock geology, sea spray and the atmosphere (Thode, 1991; Nehlich, 2015). However, Monruz and Kastelhöhle-Nord are situated far from the coast and occupy similar, relatively uniform sedimentary geologies comprised of limestone, sandstone, and clay that are unlikely to account for the range of δ^{34}S values that we observe (Asch, 2005; Figure 1). Other potential sources of spatial variation in δ^{34}S values relate to more local-scale differences in soil microbial activity, influenced by soil temperature, water content and oxygen availability (Orchard and Cook 1983; Liu et al., 2018; Nitsch et al., 2019). Indeed, paleoenvironmental records from the Swiss Plateau document a highly heterogeneous landscape existed during this period; increasing temperatures led to permafrost thaw, terrestrial landscape instability and the development of localised marshy habitats in some
areas, while in other areas it facilitated vegetation development and soil stabilisation (Thew et al., 2009; Rey et al., 2017; 2019). It is these local-scale processes which we suggest are represented in the high level of variation observed in the Monruz δ²⁴S values.

Only one result is available for the Azilian period from Monruz and Champréveyres. This was the sole sample from an Azilian context at Champréveyres to yield adequate collagen for analysis, and the only red deer sample in our data set. This sample produced δ¹³C and δ¹⁵N values of −20.6‰ and 2.4‰, respectively, which are comparable to the red deer previously analysed from the site (−20.6‰ and 3.4‰, Drucker et al., 2009).

Deglaciated versus unglaciated Lateglacial landscape development

In the context of Late Pleistocene Europe, much of the spatial and temporal variation observed in herbivore bone collagen stable isotope values has been linked to soil processes, related to variations in temperature, permafrost extent and proximity to ice sheet margins (Stevens and Hedges, 2004; Stevens et al., 2008; Drucker et al., 2011b; 2012). To explore these possible drivers of the isotopic signatures, and by inference post-LGM environmental change, we compare the temporal patterns observed in fauna from neighbouring regions that had been glaciated at the LGM to those that had remained ice-free during the last glacial cycle. We combine the results of this study with those previously published from regions of Switzerland, the French Jura and Western Alps (Drucker et al., 2003; 2009; 2011a; 2011b; 2012; Stevens et al., 2008; Bocherens et al., 2011; Gröcke et al., 2017; Reade et al., 2020; Figure 5; Supplementary Information 5). Sites are assigned to two groups based on reconstructed maximum ice extents; those in locations that remained ice-free through the LGM, and those that were ice-covered at the LGM (Campy, 1992; Bini et al., 2009; Schlüchter et al., 2010). As this analysis combines data from multiple species (Bos/Bison, horse, red deer, and reindeer), differences in dietary behaviours mask possible environmental interpretations from the δ¹³C values. Dietary ecology is also likely responsible for some of the scatter in the δ¹⁵N data, but temporal trends are nonetheless apparent (Figure 5). Herbivore δ²⁴S values appear to primarily reflect the underlying environment, irrespective of differences in dietary ecology, and the different temporal patterns in the δ²⁴S values between the two location-based groups is striking (Figure 5).

For δ¹⁵N values, the temporal pattern of variation appears consistent between the ice-covered and ice-free areas, suggesting that the presence/absence of ice sheets and processes of deglaciation are not the direct primary influences being recorded in the signal. The most notable aspect of the record is the absence of δ¹⁵N values greater than c. 2.5‰ between around 15,200 and 14,800 cal. BP. As this pattern is present in samples that span a large geographical area, which is topographically and environmentally diverse, a regional-scale explanation should be sought. Low herbivore δ¹⁵N values (<2.5‰) in Lateglacial
Europe have previously been linked to environments with nutrient-poor soils, where low temperatures and the presence of permafrost impeded the soil nitrogen cycle, or to increased environmental moisture; conversely, high herbivore $\delta^{15}N$ values (>5‰) have been considered typical of environments where nitrogen supply is not a limiting factor to plant growth and environmental conditions do not inhibit the soil nutrient cycle (Schulze et al., 1994; Hobbie et al., 1998; Jonasson et al., 1999; Stevens and Hedges 2004; Stark, 2007; Stevens et al., 2008; 2009; Drucker et al., 2011b; 2012; Rabanus-Wallace et al., 2017). Low $\delta^{15}N$ values occur alongside an absence of high $\delta^{15}N$ values at various points in the Swiss/French record shown in Figure 5 (at c. 21,000 – 18,200 cal. BP, c. 15,200 – 14,800 cal. BP, and c. 12,200 cal. BP). However, we suggest that it is only the 15,200 – 14,800 cal. BP interval that can be discussed with a degree of confidence, as the exclusively low $\delta^{15}N$ values during this time interval are clearly bounded by the presence of higher $\delta^{15}N$ values both before and after (Figure 5). The timing of this disappearance of $\delta^{15}N$ values greater than c. 2.5‰ in the Swiss/French record broadly corresponds to the end of Heinrich Event 1 (HE1), a period of climatic cooling following the LGM, during which Alpine ice sheets expanded (Hemming, 2004; Ivy-Ochs et al., 2006). Immediately after HE1, a small but significant climatic warming is evident, although lake level data suggests annual precipitation amounts remained relatively low (Magny et al., 2006; Magny, 2013). On the Swiss Plateau localised permafrost degradation and an increase in insect and plant species diversity is evident (Thew et al., 2009; Rey et al., 2017). A similar increase in vegetation density is also evident northwest of the Jura mountains (e.g. Magny et al., 2006). It is possible that if this period corresponds to the first significant regional increase in vegetation after the LGM, such an increase even if relatively small, occurring in an already nutrient-limited environment would initially deplete nutrient availability even further. The effect of this would be a short-term decline in average plant $\delta^{15}N$ values, followed by a rapid increase, as soils matured and nutrient cycling accelerated (Hobbie et al., 1998; 2005; Ammann et al., 2013).

Unlike the nitrogen record, sulphur isotope ratios display a significantly different pattern of change between ice-covered and ice-free areas. Ice-free areas display low $\delta^{34}S$ values (<–8‰) at around 23,000 cal. BP, but then remain consistently high across the deglacial period (>–8‰, except for one outlier). By contrast, in locations that were covered by ice at the LGM, low $\delta^{34}S$ values (<–8‰) are recorded at around 15,900 to 15,400 cal. BP, increasing to a minimum of –5‰ by 15,000 cal. BP. While some of the scatter in the data can be explained by different patterns of mobility between different species, such influences cannot explain the clear location-specific differences in the temporal sulphur isotopic record. Further, while location-based geological differences may produce different absolute $\delta^{34}S$ values, no relationship is observed between bedrock type, $\delta^{34}S$, and glaciated/ice-free location (Figure S4). As such, spatially and temporally variable environmental parameters...
need to be considered. Temperature-mediated controls on soil mineralisation and
volatilisation, and bacterial reduction of sulphur have been suggested to explain temporal
changes in herbivore δ^{34}S values (Drucker et al., 2011a). However, temperature change
alone cannot explain the location-based differences we observe in the data, unless it was
acting on a sub-regional scale. Soil maturity has also been linked to differences in herbivore
sulphur isotopic ratios (Drucker et al., 2012), but this interpretation is not supported in this
instance by the corresponding δ^{15}N values.

Here, we suggest that the herbivore δ^{34}S values reflect local soil conditions that are
primarily related to hydrology and microbial activity. Plants acquire sulphur from the soil as
sulphate, derived from mineral weathering and atmospheric deposition, and influenced by
soil microbial action (Walker, 1957; Krouse, 1980; Robinson and Bottrell, 1997; Newton and
Bottrell, 2007). Changing soil hydrological dynamics, particularly the development of water-
logged environments that result in anaerobic conditions, can lead to significant differences in
plant δ^{34}S values brought about by changing bacteria-mediated fractionations (Fry et al.,
1982; Trust and Fry, 1992; Groscheová et al., 2000; Nitsch et al., 2019). While the
processes that govern soil δ^{34}S in relation to changing hydrological conditions are complex
and not yet fully understood (Mandernack et al., 2000; Nitsch et al., 2019), we suggest that
the different pattern observed between the ice-free and recently-deglaciated regions in post-
LGM Switzerland could relate to location specific hydrological dynamics, potentially related
ice sheet melt, or more likely, permafrost thaw processes. Indeed, the palaeoenvironmental
record from Lake Neuchâtel on the Swiss Plateau identifies a period of permafrost thaw
coinciding with considerable instability of the terrestrial landscape and localised marshy
conditions, that is largely contemporaneous with the low herbivore δ^{34}S values identified in
this study at Monruz (Thew et al., 2009). By contrast, while there is greater debate about the
distribution and character of permafrost north and west of the Jura, recent studies suggest
that if present, it did not persist into the latter part of GS-2.1a (Renssen and Vandenberge,
2003; Bertran et al., 2014; Vandenberghe et al., 2014). This could explain why low δ^{34}S
values during GS-2.1a are not observed in these regions.

Conclusion

The carbon, nitrogen and sulphur isotopic ratios reported in this study attest to
regionally variable environmental development in post-LGM Switzerland. Significant
differences between areas that remained ice-free throughout the last glacial cycle and those
that were glaciated at the LGM are identified. Deriving such information from archaeological
faunal assemblages allows these environmental records to be directly related to human
presence in these areas, and to subsistence and settlement strategies in such landscapes,
which can more broadly inform on human behavioural ecology in peripheral settings. Our results show the post-LGM period in Switzerland and adjacent regions was characterised by diverse environmental conditions, indicating that a range of habitats and landscapes were available for humans and animals to exploit. Herbivore δ¹³C and δ¹⁵N values are most strongly influenced by species ecology and dietary behaviours, underpinned by environmental influences. δ³⁴S values appear to most strongly correspond to location-specific environmental conditions. Our results suggest that during the period of Magdalenian activity in Switzerland the Swiss Plateau was a dynamic and diverse landscape, while greater environmental stability may have existed north of the Jura Mountains.

The absence of high (>2.5‰) δ¹⁵N values between 15,200 and 14,800 cal. BP, both on the Swiss Plateau and north of the Jura, indicates a regional-scale environmental phenomenon that we suggest is related to the combination of prolonged low temperatures, limited bioavailable soil nutrients, and elevated nutrient demand from increasing vegetation cover. In contrast, low (<–8‰) δ³⁴S values occur at different times in different locations, and we suggest that these reflect locally variable hydrological dynamics, either related to changing rates of mineral weathering and soil-bedrock interactions or to changing soil redox conditions, which govern microorganism-mediated isotopic fractionations. We suggest that either interpretation is congruous with the regionally different patterns of ice sheet melt and permafrost thaw.

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Author contributions
Conceptualization: RES, HR, SG, TH & IB; Sample collection: HR, SC, DL, WM; Formal analysis: HR, JT, AF, & KLS; Investigation: HR, JT & RES; Supervision: RES; Writing - original draft: HR; Writing - review & editing: all authors; Funding acquisition: RES.

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<td>OxA-V-2754-49</td>
<td>12,480 ± 50</td>
<td>15,023−14,284</td>
<td>GS-2.1a/GL-1e</td>
<td>1</td>
</tr>
</tbody>
</table>

Table 1. AMS radiocarbon determinations on bone collagen from the sites and levels discussed in the text, and as shown in Figure 2. \(\delta^{13}C\) and C/N ratio measured by IRMS as part of the radiocarbon dating procedure at the Oxford Radiocarbon Accelerator Unit. Calibration of radiocarbon age determinations was performed using OxCal 4.3 (Bronk Ramsey 2017) and the INTCAL13 dataset (Reimer et al. 2013). References (Ref) for dates: ¹this study; ²Leesch and Müller, 2012; and ³Bodu et al., 2009.
Table 2. Summary statistics for carbon, nitrogen and sulphur isotopic ratios for each species and archaeological site. Each sample was analysed in duplicated by IRMS at the Scottish Universities Environmental Research Centre. Data in () was deemed unreliable based on standard quality control criteria. Full results are presented in the Supplementary Information 1.
Figure Captions

Figure 1: Map showing the location of Monruz, Champréveyres and Kastelhöhle-Nord. Bedrock geology is from the International Geological Map of Europe (IGME 5000; Asch 2005). Black line indicates present day country borders. White hatching indicates reconstructed limits of Last Glacial Maximum (LGM) ice sheet extent from Becker et al. (2015). Symbols indicate archaeological sites from which isotopic data used in our discussion comes (full data in Supplementary Information 5), from areas that were covered by ice at the LGM (pink circles) and those in areas that remained ice free throughout the LGM (black squares). Inset: location of Switzerland in Europe indicated by red box.

Figure 2. Calibrated AMS radiocarbon determinations on faunal bone collagen from sites discussed in the text. Calibration was performed using OxCal 4.3 (Bronk Ramsey 2017) and the INTCAL13 dataset (Reimer et al. 2013) and plotted with the Last Glacial period INTIMATE event stratigraphy and NGRIP ice core δ¹⁸O values values (top; North Greenland Ice Core Project members. 2004; Rasmussen et al., 2014). Purple = Bos/Bison, blue = horse, red = reindeer, green = red deer. Kastelhöhle-Nord dates from this study (OxA codes) and Leesch and Müller (2012; ETH codes); Monruz and Champréveyres Magdalenian dates from Bodu (2009); Champréveyres Azilian dates from this study. Symbols *, ✦, and ▵ indicate date is on same bone specimen.

Figure 3. δ¹³C, δ¹⁵N, and δ³⁴S values of Kastelhöhle-Nord upper horizon samples. Overlap in values can be seen between Bos/Bison (green triangles) and reindeer (Rangifer tarandus, orange circles), while horse (Equus sp., blue squares) are dissimilar in their isotopic values. Enlarged symbols represent directly dated samples reported in Figure 2.

Figure 4. δ¹³C, δ¹⁵N, and δ³⁴S values from reindeer (Rangifer tarandus, circles) and horse (Equus sp, triangles) from Kastelhöhle-Nord upper horizon (yellow) and Monruz (green). δ¹³C and δ¹⁵N values cluster by species, while δ³⁴S values cluster by location.

Figure 5. δ¹⁵N (middle) and δ³⁴S (bottom) isotope values from LGM and Lateglacial fauna (Bos/Bison, horse, red deer, reindeer) from the French and Swiss Jura, western Alps and Swiss Plateau (full data in Supplementary Information 5). INTIMATE event stratigraphy and NGRIP ice core δ¹⁸O values values (top; North Greenland Ice Core Project members. 2004; Rasmussen et al., 2014). Black symbols indicate samples from locations that remained ice-free throughout the LGM, pink symbols indicate samples from locations that were ice-
covered at the LGM, as displayed in Figure 1. Circles indicate directly dated specimens (by radiocarbon), triangles indicate context dated specimens, where age has been inferred from dates on other faunal specimens from the same site/stratigraphic context.
Figure 2

Champréveyres Azilian
OxA-V-2754-49C
Monruz and Champréveyres Magdalenian
OxA-20700
OxA-20701
OxA-20699
Kastelhöhle-Nord Upper Horizon
OxA-V-2793-56C
ETH-45026
OxA-V-2793-54C
ETH-45025
OxA-V-2793-55C
OxA-V-2748-25C
ETH-45024
OxA-V-2794-25C
ETH-45023
OxA-V-2754-48C
ETH-45022

NGRIP δ18O

Calibrated date (calBP)

OxCal v4.3.2 Bronk Ramsey (2017); r:5 IntCal13 atmospheric curve (Reimer et al 2013)
Figure 3
Figure 4

The diagram illustrates the variation in δ¹⁵N_AIR (%o) and δ¹³C_VPBD (%o) for different species and locations. The left side of the figure shows the species-based variation, while the right side demonstrates the location-based variation.

- **Species-based variation**:
  - **Kastelhöhle**: Yellow circles for Equus sp.
  - **Kastelhöhle**: Green triangles for Rangifer tarandus

- **Location-based variation**:
  - **Monruz**: Yellow circles for Rangifer tarandus
  - **Monruz**: Green triangles for Equus sp.
Figure 5

- δ¹⁸O NGRIP
- δ¹⁵N AIR
- δ³⁴S VCDT

Years cal. BP:
23000 21000 19000 17000 15000 13000 11000

- Ice-free at LGM (direct date)
- Ice-free at LGM (context date)
- Ice-covered at LGM (direct date)
- Ice-covered at LGM (context date)