TITLE Gazelle seasonal mobility in the Jordanian Steppe: the use of dental isotopes and microwear as environmental markers, applied to Epipalaeolithic Kharaneh IV **ABSTRACT** The Early/Middle Epipalaeolithic aggregation site of Kharaneh IV in the east Jordan steppe is one of the largest of the period, showing repeated use, high degrees of site investment, and relatively prolonged periods of occupation. Hunters relied heavily on the single prey species, Gazella subgutturosa. This paper reports on isotopic analyses of gazelle teeth, drawn from the Kharaneh IV assemblage, to explore the seasonal and spatial distribution of gazelle in life and examine models of long-distance seasonal gazelle migrations. Dental microwear analysis is also employed to assess hunting seasons. Results from sequential isotope analyses of 11 Epipalaeolithic gazelle molars show that, with one exception, gazelle did not move seasonally from the limestone steppe onto other geologies, nor did they seasonally relocate to areas with different climate regimes. Rather, seasonal herd mobility seems to have been local, meaning animals would have been available in the vicinity of Kharaneh IV year-round. Results from 19 microwear analyses show some gazelle to have died in spring when grass graze was available, whilst others had been browsing around their time of death, indicating non-spring hunting. Microwear evidence thus suggests gazelle were hunted in more than one restricted season. The implications of these new datasets for hunter-gatherer use of the steppe, and potential hunting methods, is highlighted. **KEY WORDS** Early/Middle Epipalaeolithic Oxygen, carbon, strontium isotopes Dental microwear analysis Gazelle seasonal mobility Hunting seasonality

INTRODUCTION

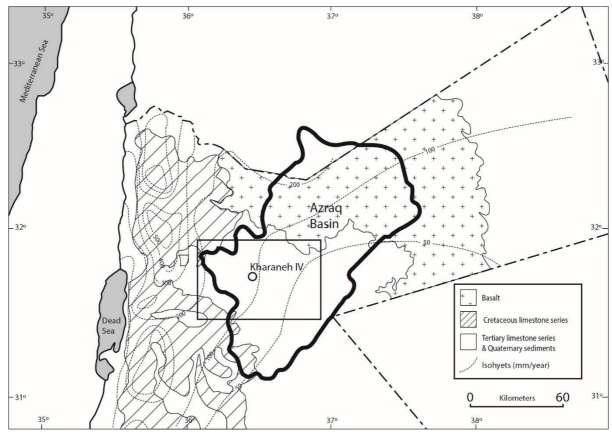


Fig.1 Map showing the Azraq Basin (black line), surrounding areas of Northeast Jordan (dash/dotted line) and area shown in Fig. 2 (rectangle surrounding Kharaneh IV) (after Bender 1974; Jordan National Geographic Centre 1984)

There are compelling reasons for archaeologists to attempt reconstruction of wildlife mobility in past environments. Where zooarchaeological evidence shows hunter-gatherers have relied heavily on single prey species, knowledge of herd seasonal movements is key for better understanding seasonal settlement organization and hunting techniques (Frison 2004; Price et al 2015). The body chemistry of archaeological animal remains can make aspects of the past visible (Feranec et al 2007), more so where multiple lines of evidence are studied in combination (Lubinski 2012). For example, combined evidence from three isotope sequences retrieved from bison dental enamel were employed to determine seasonal location patterns, and thus availability, to Palaeolithic hunters (Julien et al 2012). Signatures of extended and confined seasons of death, determined from dental microwear analysis (DMA) of extant ungulates, were applied to investigating mass cull and attritional hunting techniques from DMA of archaeological assemblages (Rivals et al 2015). The current study utilises both approaches in combination.

This study focuses on the Early/Middle Epipalaeolithic of the Jordan steppe (*Fig. 1*) where evidence points to reliance on goitered gazelle (*Gazella subgutturosa*) to the exclusion of other prey. Specifically, we focus on the site of Kharaneh IV, one of the largest Levantine occupations of the period, in an attempt to understand how gazelle herds may have underpinned large-scale, repeated occupations. The dominant model of Levantine prehistoric *G. subgutturosa* mobility, proposed by Legge and Rowley-Conwy (1987; 2000), is based on

2 draws on ethno-historic observations (18th - 20th C AD) of gazelle movements. Their study posits that the 3 northern migration point of gazelle herds was near Abu Hureyra in spring for fresh graze, and their 4 southernmost over-wintering point was the Jordanian steppe. This long-distance migration model has found 5 widespread acceptance, particularly as an explanation for mass-capture 'kite' structures (e.g. Bar-Oz et al 2011), 6 while some authors have also suggested that large Epipalaeolithic sites such as Kharaneh IV developed on 7 gazelle migration routes (Goring-Morris 1995, 156). 8 9 Gazelles are extinct in the Jordanian steppe today, and historical accounts of recent mobility patterns cannot be 10 assumed for the Epipalaeolithic before millennia of landscape degradation and livestock grazing (Martin 2000). 11 Independent approaches are required for reconstructing the seasonal mobility of prehistoric gazelle herds (Julien 12 et al 2012). We infer mobility and feeding seasonality from oxygen, carbon and strontium stable isotope 13 sequences retrieved from Kharaneh IV gazelle teeth to ask whether, during the Epipalaeolithic, targeted herds 14 undertook long-distance seasonal migrations in and out of the Jordan steppe area. Dental microwear analysis of 15 the same tooth samples augment hunting seasonality evidence, providing indicators of diet, known to vary 16 seasonally, at time of death. This study is the first use of archaeological isotopes to examine Gazella 17 subgutturosa seasonal mobility in the Levantine Epipalaeolithic and the first regional application of a combined 18 isotope/DMA approach. 19 20 KHARANEH IV 21 Kharaneh IV (hereafter Kharaneh) is within the Azraq Basin drainage system (Fig. 2), notable as one of the 22 largest Epipalaeolithic sites (>21,000 m²) of the region, with occupation radiocarbon dated between 19,830 and 23 18,600cal BP. Recent Epipalaeolithic Foragers in Azraq Project excavations have focused on two areas. Area B 24 dates to the earliest millennium whilst Area A overlaps and continues later, 18,850-18,600cal BP (Maher et al 25 2016). Dates sit comfortably within a southern Levantine Early Epipalaeolithic chronology (Belfer-Cohen & 26 Goring-Morris 2013 Table 3.3.1; Goring-Morris et al 2009, Table 10.1), although Area A lithics show clear 27 typo-technological affinities with Middle Epipalaeolithic traditions (Maher & Macdonald 2013; Maher et al 28 2016; Richter et al 2013), thus suggesting an earlier Early-Middle division (Maher et al 2011) and confusing 29 current classification. We use the broad term Early/Middle Epipalaeolithic for Area A, the focus of this study. 30 31 Findings at Kharaneh raise questions about the nature of hunter-gatherer occupation in Levantine steppes. Site 32 size, artefact density, investment in features and variety of lithic traditions (Maher & Macdonald 2013) attest to 33 large aggregations of hunter-gatherer groups, gathering perhaps for social/economic events (Maher et al 2012a). 34 Combined with palaeoenvironmental evidence for lusher, better-watered settings (see Palaeoenvironment), 35 research argues that Kharaneh would have been "...an attractive locale for repeated and prolonged occupation" 36 (Maher et al 2012a, 8). For Area A, >90% of animal remains represent locally hunted gazelle (Gazella 37 subgutturosa) (Spyrou 2014, Table 2 & Fig. 7). This focus on gazelle hunting is typical of other Jordanian 38 prehistoric steppic assemblages and of earlier zooarchaeological studies at Kharaneh (Martin et al 2010) and 39 begs questions about the seasonal spatial distribution of this mainstay herd animal. Did gazelle seasonally

zooarchaeological analysis of late Epipalaeolithic gazelle remains from Abu Hureyra (Middle Euphrates), and

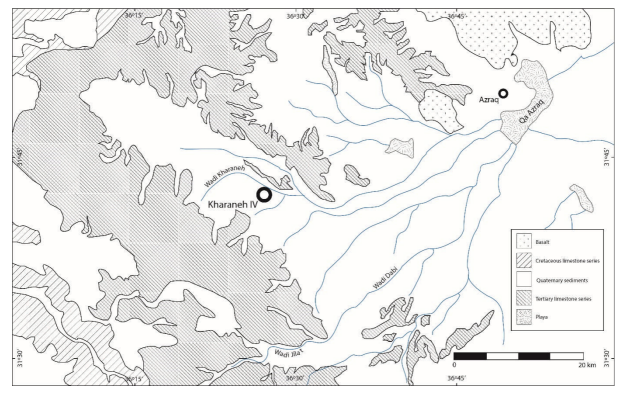


Fig. 2 Map showing the location of Kharaneh and the wadi system draining into the central Azraq Oasis

migrate in and out of the Jordan steppe or could they have maintained a more year-round presence? The latter could have aided the prolonged hunter-gatherer site residence proposed by Kharaneh excavators (Maher et al 2012a).

There are presently two sources of evidence for Kharaneh gazelle hunting seasonality. Fusion data shows targeted winter culling and slight evidence for spring culls (Martin et al 2010), and a tooth cementum pilot study (Jones 2012) finds evidence for both spring/summer and autumn/winter culls. Questions remain as to whether gazelle herds were on northerly migrations in spring (as Legge and Rowley-Conwy predict) and were present in the Jordan steppe mainly in winter, or whether other mobility patterns pertained. Animal behavioural plasticity is an adaptive response to environmental conditions, and Martin (2000) has argued that small-bodied gazelle may not have needed to move far in search of resources. Considering *G. subguttorosa* behavioural and forage requirements, and the environmental parameters of our study region, we propose four potential herd mobility patterns for Epipalaeolithic gazelle herds (*Fig. 3*). Gazelle herds might seasonally aggregate and disperse locally in the steppe (1), or might undergo longer seasonal migrations following better resources, westwards into the Jordanian Highlands in summer (2), northwards towards the Middle Euphrates in spring (3) or south-eastwards along the Wadi Sirhan corridor (4). In cases 2-4, animal presence in the Jordan steppe is predicted to be seasonal.

AZRAQ BASIN PALAEOENVIRONMENT

23 Geology

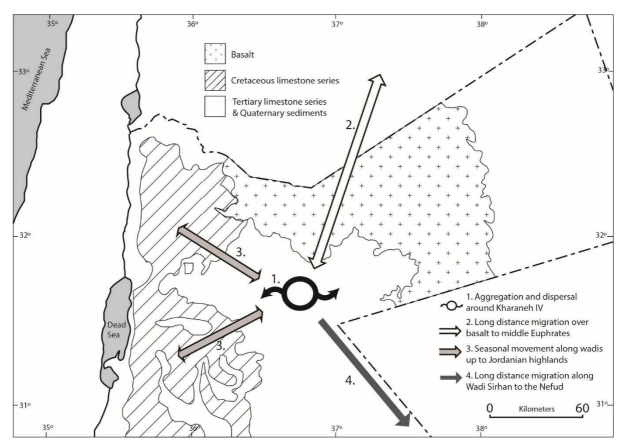


Fig.3 Map showing the four proposed movement patterns of gazelle in the Early/Middle Epipalaeolithic (after Bender 1974; Jordan National Geographic Centre 1984)

The Azraq Basin is a 12,000km² endorheic drainage basin covering north-central Jordan (Garrard & Byrd 2013), with Early Tertiary (tt₁) chalky limestones and marls exposed along western and eastern flanks. In the north there is an extensive cover of Pliocene/Oligocene (B₄) and Pliocene/Miocene (B₅) basalts, whilst in the centre and southeast there is a spread of Quaternary gravels and silts (q₂). Further west, the Jordanian Highlands are composed of Cretaceous dolomitic limestones and marls (c₂) (Bender 1974). Wadi Kharaneh, in the Basin's central-western sector, drains from Early Tertiary uplands (c850m) to the central Qa Azraq (c500m). Kharaneh is situated in the broad wadi c640masl, c40km west of Azraq (*Fig.* 2). Where conditions allow, small pockets of rendzina soils have accumulated.

Climate and hydrology

Present-day Azraq Basin temperatures range from 45°C to -10 °C, and precipitation from 200mm (NW) to <50mm (SE). The Mediterranean controls rainfall distribution; westerly winds bring rain in the cold season but effectively none in the highly evaporative hot season (Enzel et al 2008) (*Fig. 4*). Winter run-off collects in playas and stream channels but evaporates as temperatures rise, becoming highly saline and drying within months; only spring-fed Azraq Oasis marshlands have perennial freshwater.

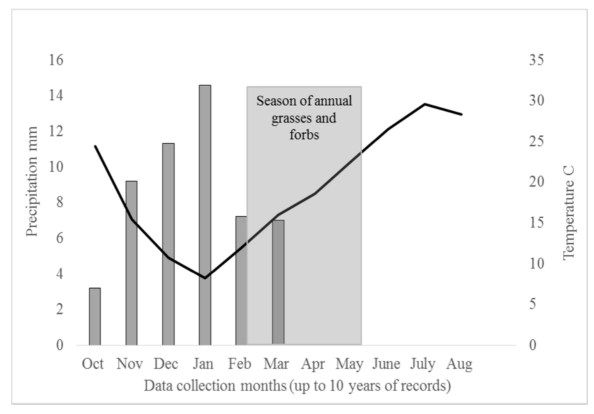


Fig. 4 Mean monthly GNIP temperature and precipitation data for Azraq (WMO/IAEA 2014) with the prime vegetation growth season

Regional and local evidence (Frumkin et al 1999; Hunt and Garrard 2013; Jones & Richter 2011; Jones et al 2016; Ramsey & Rosen 2016) suggest the Last Glacial Maximum (LGM) climate in the Mediterranean zone and regions due east was wetter than now, but with the same atmospheric circulation and seasonality (Enzel et al 2008). However, there is evidence for increasing dryness during the post-LGM period (Hunt and Garrard 2013; Jones et al 2016). Kharaneh was established on the marls of a receding wetland area, possibly 2km maximum, which appears to have dried out during occupation, later overlain by loess (Garrard et al 1985; Jones et al 2016).

<u>Vegetation</u>

Due to precipitation seasonality, opportunistic annual grasses and forbs germinate rapidly in moist, warming spring soils, outcompeting slower growing perennials but soon setting seed and withering as aridity increases, leaving only xeric shrubs and species tolerant of soil salinity (Batanouny 2001; Hillman 1996). Vegetation zones grade according to isohyets; the Highlands have higher rainfall and lower temperatures supporting oak woodland (>400mm) and park woodland/ woodland steppe (400-200mm). Within the Azraq Basin, this grades into the steppe plant community described above (200-100mm), and desert vegetation (<100mm) (Hillman 1996; Hillman in Moore et al 2000). Each zone extends further east along wadi edges than on interfluvial plains, and in the Oasis permanent springs support marshland plants. Today, Kharaneh is located on the Irano-Turanian steppe/desert 100mm isohyet boundary.

Modelled palaeovegetation distribution (Byrd et al 2016; Colledge 2001; Hillman 1996) shows an eastward shift and greater species abundance, such that xeric woody thickets encroached along wadis and perennial pools;

1 annual grasses blanketed interfluvial areas for an extended season; halophytic species were less dominant in arid

seasons. The zooarchaeological spectrum offers support to this model with water dependent species by large

wadis and in the Oasis, numerous gazelle associated with dry steppic grasslands and an absence of woodland

habitat species such as deer (Edwards pers. comm; Martin et al 2010).

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PRINCIPLES UNDERLYING METHODS

- 7 Isotope analysis – tooth enamel
- 8 Gazelle tooth enamel apatite is chosen for isotope analyses because developmental morphology of hypsodont
- 9 mandibular molars allows bi-monthly sampling resolution along the growth axis (Balasse et al 2003; Fricke &
- 10 O'Neil, 1996; Henton 2012; Kohn et al 1998; Zazzo et al 2002) (Fig. 5). A sampling sequence spans an annual
- 11 cycle as enamel precipitation proceeds during the first year in the second molar and between nine and eighteen
- 12 months in the third (Davis 1980; Munro et al 2009). Intra-species developmental consistency pertains,
- 13 particularly in M₂s (Blaise & Balasse 2011; Tornero et al 2013). In this study, trajectories of change during the
- 14 gazelle's annual cycle are most pertinent to research questions. Each data point retrieved from the tooth growth
- 15 column is not considered in isolation but as a fixed-position contributor to an annual curve. The research goal
- 16 does not require precise conversions of stand-alone data-point values, diminishing problems arising from sample
- 17 resolution and enamel formation time-lags (Britton et al 2011; Hoppe et al 2005).

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- 19 Within the tooth enamel time capsule, where seasonality exists, the sequence of retrieved oxygen (δ^{18} O) isotope
- 20 signatures identify (via ingested water) patterns of seasonal change (Bryant & Froelich 1995; Dansgaard 1964;
- 21 Fricke et al 1998; Iacumin & Longinelli 2002); where geological substrata variation allows, strontium (87Sr/86Sr)
- 22 isotope signatures identify (via ingested forage) changing location (Bentley 2006; Faure & Powell 1972;
- 23 Graustein 1989; Sillen et al 1998); and where plant species have varied morphological or physiological stress
- responses, carbon (δ^{13} C) isotope signatures identify (via ingested forage) changing feeding habits (Bender 1971; 25 Cerling & Harris 1999; Ehleringer et al 1997; Heaton 1999; de Niro & Epstein 1978; O'Leary 1988; Tieszen
- 26 1991; Vogel et al 1986). By analysing all three isotopes in the same sample, seasonality of carbon signatures can
- 27 be inferred from oxygen signatures, but strontium incorporates into enamel more slowly introducing an
- 28 unknown time-lag (Montgomery et al 2010).

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Isotope analysis – the palaeoenvironment

- 31 The Azraq Basin palaeoenvironment is eminently suitable for investigation through these isotopic signatures, as
- 32 its climate, geology and vegetation can be related to modern environmental parameters, and as isotopic variation
- 33 potentially discriminates season, location and forage. Global Natural Isotopes in Precipitation records (Fig. 6)
- 34 show clear summer peaks and winter troughs in monthly δ^{18} O; >0.57 to -6.32 for Azraq, and >-3.32 to -7.28 for
- 35 the Jordanian Highlands (IAEA/WMO 2014). Labile 87Sr/86Sr signatures retrieved from modern Jordanian
- 36 plants and animals (Shewan 2004) fall within accepted bedrock parameters (Bentley 2006; Hartman & Richards
- 37 2014), discriminating the Cretaceous limestone series in the Jordanian Highlands, the Tertiary limestone series
- 38 in the steppe, and the younger basalt shield. Short-lived spring annuals constitute 80% of Azraq Basin plant
- 39 species (Zohary 1973) and have C₃ photosynthetic pathways (Bocherens et al 2001; Ramsey & Rosen 2016), as
- 40 do slower growing woody shrubs and trees; these would return δ^{13} C that group around -27% (Bender 1971;

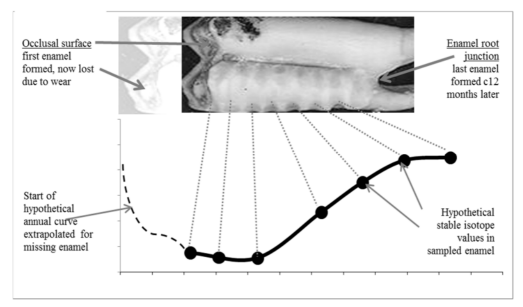


Fig. 5 Diagram showing a hypothetical annual graph retrieved from the sequential drilling of enamel apatite in an ungulate second mandibular molar formed over 12 months

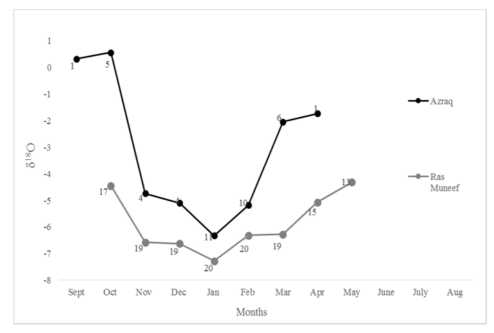


Fig 6. Mean monthly GNIP $\delta 18O_{VSMOW}$ data (WMO/IAEA2014) comparing highland Ras Muneef to Azraq (number of modern records

Ehleringer et al 1997; O'Leary 1988). Whereas halophytic chenopods, more predominant throughout the arid season, have C_4 photosynthetic pathways that enrich $\delta^{13}C$ to c.-12‰ (Akhani et al 1997; Shomer-Ilan et al 1981).

<u>Isotope analysis – gazelle ethology</u>

This research focuses on seasonal changes experienced by gazelle, and their ethology constrains likely environmental situations. Gazelle are non-obligate drinkers with water needs met by plant tissues, where arid-

1 season δ^{18} O is considerably enriched by evapo-transpiration (Ayliff & Chivas 1990), enhancing, but not 2 masking, the summer/winter range in values (Kohn et al 1996, 1998). However, should gazelle migrate to 3 cooler, less-evaporative summer feeding grounds (Poage & Chamberlain 2001), the sinusoidal annual δ^{18} O 4 curve would be disrupted due to loss of the extreme arid season signature. 5 6 Gazelle ingest strontium from plants which take up only the labile fraction. In more arid regimes, soils largely 7 derive from underlying bedrock (Bentley 2006), but where geologies intersect, where upland sediments are 8 washed downstream (Graustein 1989; Sillen et al 1998), or where windborne dust settles, the resulting soil 9 ⁸⁷Sr/⁸⁶Sr reflects the contributory mix. For this research, key location soil ⁸⁷Sr/⁸⁶Sr signatures are refined with 10 the aid of a small modern plant baseline; results for locations deep within major geologies conform to 11 established ⁸⁷Sr/⁸⁶Sr ranges; local to Kharaneh, Tertiary limestone series 0.70807-0.70819, Highlands 12 Cretaceous limestone series 0.70845-0.70854 and basalt lava flows 0.70764-0.70778 (Supplement 1, Table 1 & 13 Fig. 1; Henton et al submitted to Environmental Archaeology). Curves constructed from sequential 87Sr/86Sr 14 sampling are predicted to be flat in gazelle remaining on a single bedrock type but where they might have 15 moved between different geologies within a yearly cycle, the curves would, respectively, progressively rise or fall in gazelle progressing onto bedrock with higher or lower 87Sr/86Sr signatures. Climate-induced variation in 16 17 aeolian dust input (Hartman & Richards 2014; Stein et al 2007) would not register within an annual 87Sr/86Sr 18 cycle in a gazelle tooth. 19 20 Gazelle primarily ingest carbon from forage; grazing or browsing as seasonal mixed feeders on many Irano-21 Turanic species, but preferentially eating high-nutrient grasses, particularly during late gestation and lactation 22 (Kingswood & Blank 1996). Seasonal δ^{13} C change in archaeological gazelle enamel is predicted due to the 23 combined effects of seasonal availability of C₃ and C₄ plants and feeding preferences. In spring/early summer, when water-stress is minimal, enamel δ^{13} C would be most depleted as gazelle graze C₃ grasses, whereas high 24 25 summer to mid-winter browsing on water-stressed C_3 shrubs might enrich δ^{13} C by as much as 6% (Farquhar et 26 al 1989), and the inclusion of C4 halophytes would further raise values. After accounting for a +14.1% 27 fractionation factor between plant and animal tissue (Cerling & Harris 1999), δ^{13} C of -12% might be 28 approached in spring/early summer, becoming enriched by >-8% in the rest of the year. 29 30 Dental microwear analysis (DMA) 31 Isotope analyses reveal the seasonal forage landscape experienced by gazelle throughout a year of life as a sub-32 adult, whereas DMA elucidates forage conditions in the final weeks before death. Microwear describes marks 33 on the enamel surface that last only a few weeks. Their interpretation draws on correlation with diets in modern 34 animals (Teaford & Walker 1984). A 2-feature discrimination of pits (length:breadth <4:1) and striations 35 (length:breadth >4:1) usefully identifies grazers, with more striations, from browsers, with more pits (Solounias 36 & Hayek 1993; Solounias & Moelleken 1992). This holds true across mammalian orders and between 37 geographical zones (Mainland 1998; Merceron et al 2005; Wilkie et al 2007), however tooth type introduces 38 DMA variation, where pit percentage increases towards the third molar (Gordon 1982). The rapidity of turnover 39 allows a bimodial distribution of features in seasonal mixed feeders such as gazelle (Merceron et al 2007; Rivals

et al 2011; Solounias and Hayek 1993; Solounias & Moelleken 1992).

1 2 As Jordanian steppe gazelle are seasonal mixed feeders on seasonally available food types, a strong seasonal 3 signature might be revealed through DMA. Striation-rich microwear can be associated with spring/early summer 4 as the only time when fibrous grasses flourish, pits with later summer, autumn and early winter when soft 5 dicotyledonous plants are the sole dietary contributors, and an intermediate signal with the brief transitional 6 periods where one diet replaces the other. 7 8 **Summary** 9 This research asks simple questions of each dataset; the strength of the approach becomes apparent in their 10 combined interpretation constrained by gazelle ethology and palaeoenvironmental reconstruction. In considering 11 the seasonal mobility represented by each gazelle tooth sample throughout a year of early life, the δ^{18} O sequence 12 enables identification of four seasons, ${}^{87}\text{Sr}/{}^{86}\text{Sr}$ identifies movement over different geologies, and the $\delta^{13}\text{C}$ 13 sequence identifies dietary shifts between spring/early summer and other seasons. Seasonal mobility patterns for

sampled individuals are assumed to be representative of those of whole herds; sub-herds of *G. subgutturosa* may briefly form throughout the year, but unite for long-distance movements (Blank et al 2012). We also assume that

patterns are likely to represent annually repeated movements if environmental pressures are largely unchanged

(Cunningham & Wronski 2011). Microwear data provide insights on forage conditions around the time of death,

when the gazelle's lifecycle was intercepted by hunters; a single tightly clustered pattern can be associated with

a short, single hunting season, a bimodal pattern with two hunting seasons, and a non-modal, diffuse pattern

with year-round hunting.

MATERIALS, PREPARATION PROTOCOLS AND ANALYSES

Fully developed third mandibular molars (M₃), just in wear and with closed roots, are most identifiable in the

fragmented Kharaneh assemblage, but some securely identifiable second molars (M₂) were also sampled,

meeting recommended sample sizes of nine teeth for isotope analyses (Hoppe et al 2005; Pearson & Grove

26 2013) and fourteen for microwear analyses (Mainland pers. comm.). Table 1 provides the context and

description of fourteen M₃ and eight M₂ archaeological specimens suitable for analyses; these represent twenty

gazelle as two individuals had both M₂ and M₃ sampled.

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After taking morphometric and photographic records, impression and resin casts were made following Mainland (1998) protocols. Four images of the infundibular enamel of the paraconid cusp were captured under a Hitachi S–3400N scanning electron microscope (SEM) at x500 magnification. The clearest and most representative was imported into Ungar's (2002) image analysis software (Microware 4.02) where the dimensions and orientation of each microwear feature in a surface area of 0.0404mm² were quantified and categorised.

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For isotope analyses, the mesial tooth-column buccal surface of each specimen was cleaned and six sequential samples of >20mg drilled at measured distances from the enamel:root junction (see *Fig. 5*); contamination and diagenetic carbonates were removed following established protocols (Balasse 2002). The carbonate fraction from 10mg of each was analysed for δ^{13} C and δ^{18} O by Isotope Ratio Mass Spectrometry at UCL Bloomsbury Environment Isotope Facility (BEIF). Each remaining 10mg was processed for 87 Sr/ 86 Sr analysis under clean

laboratory conditions in the Isotope Geochemistry Laboratories at Royal Holloway College, UL. The sample was digested in concentrated HNO_3 and the soluble residue passed through columns that collected the strontium on Sr-resin. This was eluted with triple-reduced water, evaporated, and mounted on Rhenium filaments for analysis in a VG354 Thermal Ionisation Mass Spectrometer.

RESULTS

The Kharaneh depositional environment resulted in gazelle tooth specimens being considerably abraded, concreted and fractured; of the 22 originally sampled, 11 yielded strontium data, 10 carbonate data and 19 DMA data (*Table 1*). Each dataset sample size is adequate for robust analysis although variation due to tooth type requires discussion.

<u>Isotope results</u>

Looking first at the isotope results ($Table\ 2$), external reproducibility of laboratory analyses is $\pm 0.04\%$ and $\pm 0.08\%$ for $\delta^{13}C$ and $\delta^{18}O$ respectively, and $\pm 0.000014\ (2\sigma)$ for $^{87}Sr/^{86}Sr$. Measured on the PDB scale, maximum $\delta^{18}O$ is 6.54% and -4.24% minimum. These values fall within the modern range for Azraq $\delta^{18}O$ in precipitation, although dampened due to sample resolution (Britton et al 2011). The $^{87}Sr/^{86}Sr$ (0.70822 to 0.70766%) also fall within parameters established for the geology. Finally, $\delta^{13}C$ (-4.76 to -10.79%), after making a +1.5% adjustment to allow for the pre-industrial absence of the fossil fuel effect (Cerling & Harris 1999), convincingly equates to varying contributions of C_3 and C_4 plants typical of gazelle feeding ethology.

		$M_{\it 3}$				M 2								
Kharai	Kharanah IV Area A. context details		Wear stage	Side	GH	DMA (san	CO₃ nple nur	Sr nber)	Wear stage	Side	GH	DMA (san	CO 3	<i>Sr</i> nber
AR36	100		adult	right	22.2	2	2	2						
AR35	1100	FN871174	9	left	20.7	4	4	4						
AS36	100	FN871150	10	left	10.5	5	X	x						
AT35	34	FN871032	X	left	X	8	X	X						
AP35	034 or 035		10	left	10	17	X	X						
AS42	115		9	right	10	20	X	x						
AQ36	80	FN871628	X	left	20.5	21	X	X						
AT36	34		10	right	11.6	22	X	X						
AT35	100		9	right	20.4	26	26	26						
AR36	34		adult	left	23.2	30	30	X						
AR36	34		young adult	right	13.6	31	31	X						
AV71	176	545075/1	9	left	9	78	X	X						
AS42	99	FN871101	X	left	15.6	35	35	35	5	X	11.2	X	X	38
AW73	158	545200/1	7	left	24.2	79	79	79	5	left	15.8	99	X	X
AS35	34	FN871094							4	left	23.3	49	49	49
AQ35	80								5	left	19.2	51	51	51
AR35	100								x	left	x	59	X	x
AY72	43	545193/1-2							х	X	26.4	x	74	74
AX74	88								5	left	17.7	76	X	76
AU73	178	545035/27							5	left	17.5	x	X	89
22 tee	22 teeth from 20 individuals			4 M ₃ s		14	7	5		8 M ₂ s		5	3	6

Table 1. Analysed specimens with contextual and zooarchaeological information and details of analyses undertaken

1 2 The results for each gazelle tooth are displayed in Figure 7. Sp38 (M₂ with grey border) and Sp35 (M₃ with 3 white border) are from the same individual (top row Fig. 7) and, considered together, provide insight into how 4 enamel precipitation timing, and therefore generated isotopic curves, relate sequentially between tooth-types. 5 Consequently, all specimens (Fig. 7 M₃s in rows 2 & 3 and M₂s, with grey borders, in rows 4 & 5) can be 6 discussed in relation to each other. 7 8 In addition to tooth specimens with missing data sets, it is clear that not all time capsules span a whole year of 9 life; this is to be expected as M_2 s have <1 year formation period and toothwear has truncated M_3 records. 10 Nevertheless, M_3 specimens 2, 4, 30 and 79 show clear seasonal variation in δ^{18} O with seasonally linked 11 variation in δ^{13} C. The 87 Sr/ 86 Sr sequences span the same length of time although without seasonal linkage. 12 Although spanning a more limited period, a seasonal link between $\delta^{I8}O$ and $\delta^{13}C$ can be observed in M₂ 13 specimens 51, 74 and 49. 14 15 **DMA** 16 Turning to the DMA results (Table 2) fourteen M₃s (black) and five M₂s (open circles) are analysed using the 17 simplest DMA, 2-feature pit to striation discrimination, and displayed in Figure 8 with pit percentages for each 18 on the x-axis and striation percentages on the y-axis. In this research we confine ourselves to this analysis as we 19 do not have a modern comparator on which to base more nuanced DMA, and as recorders using this semi-20 quantitative recording method tends to miscount but not misidentify features (Grine et al 2002; Henton 2010 21 p328). 22 23 The results show the gazelle died having eaten a range of diets. The highest striation percentage was 79.5% and the lowest was 27.6%. Eight M₃s had more striations (>50%) than pits, and six had less; of the M₂s, three had 24 25 more and two had less. The bar chart, inset on the left of Figure 8, clearly shows a relatively even distribution of 26 results, only slightly higher than the 50% mark, with no evidence of any single or bimodal clustering. Whilst 27 some adjustment should be made for the known increase in M_2 s striation numbers compared to M_3 s in the same 28 tooth row, this might be offset by gazelle feeding behaviour; as ground feeders in more arid environments, 29 gazelle species ingest grit and have higher pit numbers than many ungulates (Rivals et al 2011; Schulz et al 30 2013; Solounias & Semprebon 2002). 31

Specimen	Dental Microwear results				Isotope results									
M ₃ white M ₂ grey	Total feature Number	Pit Number	Striation Number	a. Di	stance of s b. δ ¹⁸ O ₀	sampling ba	nds (1 to 6 c. δ ¹³ C		/enamel jur d. ⁸⁷ Sr/		Mean	Maximum Minimum		Range
2	70	43	27	a b c d	19.5 -3.74 -5.92 0.70808	19.8 -3.71 -5.57 0.70808	15.9 -2.50 -7.03 0.70812	12 -0.95 -8.42 0.70814	8.9 -0.14 -9.11 0.70813	5.3 -1.73 -9.20 0.70810	-2.13 -7.54 0.70811	-0.14 -5.57 0.70814	-3.74 -9.20 0.70808	3.60 3.64 0.00006
4	82	48	34	a b c d	18.4 -3.02 -4.72 0.70814	15 -0.41 -6.20 0.70816	11.4 3.32 -9.74 0.70815	8.1 5.27 -10.75 0.70815	6 4.92 -10.79 0.70809	3 3.85 -10.50 0.70814	2.32 -8.78 0.70814	5.27 -4.72 0.70816	-3.02 -10.79 0.70809	8.28 6.07 0.00007
5	38	13	25											
8	127	26	101											
17	90	50	40											
20	46	31	15											
21	79	49	30											
22	118	51	67											
26	138	56	82	a b c d	19.2 -0.86 -5.50 0.70811	12.8 0.16 -7.09	0.70822	7.4	4.9 0.70818	1.4	-0.35 -6.29 0.70817	0.16 -5.50 0.70822	-0.86 -7.09 0.70811	1.02 1.60 0.00011
30	176	46	90	a b c d	18.3 -3.99 -5.65	14.9 -4.24 -5.85	11.6 -4.11 -6.49	5.2 -0.02 -8.77	3.4 1.24 -9.85	1.5 0.41 -9.64	-1.78 -7.71	1.24 -5.65	-4.24 -9.85	5.48 4.20
31	89	44	45	a b c d	13.2 -0.50 -8.82	10.3 0.96 -10.12	8.2 1.61 -9.76	5.5 1.82 -9.72	3.1 1.08 -9.50	1.8 -1.48 -8.19	0.58 -9.35	1.82 -8.19	-1.48 -10.12	3.30 1.94
35 (the M2 is 38)	134	60	74	a b c d		12.3 -0.72 -5.97 0.70815	10.3 1.71 -8.21 0.70816	8.2 3.64 -9.68 0.70814	6.2 4.54 -9.31 0.70811	4.1 5.44 -8.3 0.70805	2.92 -8.29 0.70812	5.44 -5.97 0.70816	-0.72 -9.68 0.70805	6.16 3.71 0.00011
38 (the M3 is 35)				a b c d				9.4	6.5 0.70817	3.5 0.70814	0.70818	0.70822	0.70814	0.00008
49	87	63	24	a b c d	19.2 3.7 -8.08 0.70814	17.2 1.91 -7.69 0.70812	15 1.07 -7.41 0.70814	12.2 -1.16 -6.62 0.70815	9.3 -2.56 -6.23 0.70816	6.1 -4.24 -5.3 0.70815	-0.21 -6.89 0.70814	3.70 -5.30 0.70816	-4.24 -8.08 0.70812	7.94 2.78 0.00003
51	72	44	28	a b c d		16.3 -0.39 -7.66 0.70766	13 -2.51 -6.72	10.4 -3.45 -5.67 0.70777	7.3 -3.27 -5.08	5.2 -2.93 -4.76 0.70796	-2.51 -5.98 0.70779	-0.39 -4.76 0.70796	-3.45 -7.66 0.70766	3.06 2.90 0.00031
59	204	93	111											
74				a b c d		20.5 6.54 -7.87 0.70811	16.7 4.96 -7.79	13.1 3.55 -8.19 0.70810	9.9 2.17 -8.07	6.5 -0.21 -7.49 0.70811	3.40 -7.88 0.70811	6.54 -7.49 0.70811	-0.21 -8.19 0.70810	6.75 0.70 0.00001
76	96	36	60	a b c d		0.70812		9.1		3.2 0.70812	0.70811	0.70812	0.70809	0.00003
78	93	50	43	a b c d										
79	148	59	89	a b c d	20.9 -3.98 -5.56 0.70821	17.7 -4.03 -6.02 0.70820	14.2 -0.81 -9.14 0.70821	11.5 1.68 -10.48 0.70819	9.5 3.54 -10.62 0.70818	6.1 4.88 -10.30 0.70812	1.05 -9.31 0.70818	4.88 -6.02 0.70821	-4.03 -10.62 0.70812	8.91 4.60 0.00009
89				a b c d		0.70810		0.70812	7.2 0.70812	5.4	0.70812	0.70812	0.70810	0.00002
99	66	29	37	a b c d										

Table 2. Results of analyses for dental microwear and isotopes of oxygen, carbon and strontium in all specimens (M_2 s are shaded grey)

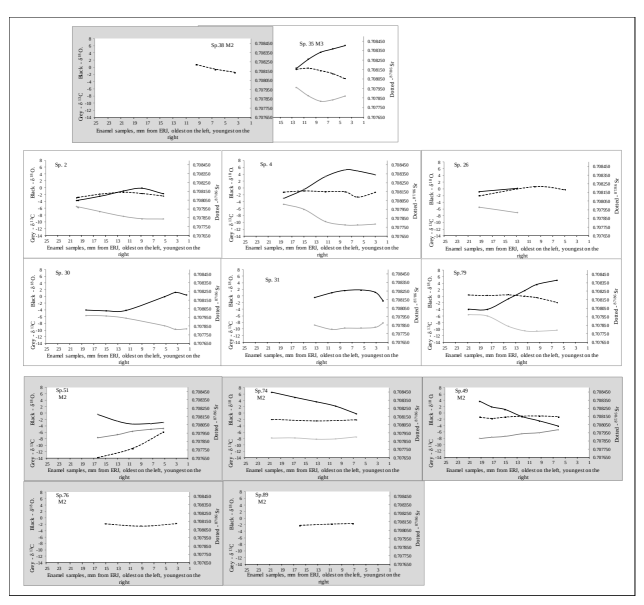


Fig 7. Plots of all isotope sequences for individual gazelle specimens. Top row: M_2 and M_3 from the same specimen. Rows 2 & 3: all other M_3 s. Rows 4 & 5: all other M_2 s (grey borders)

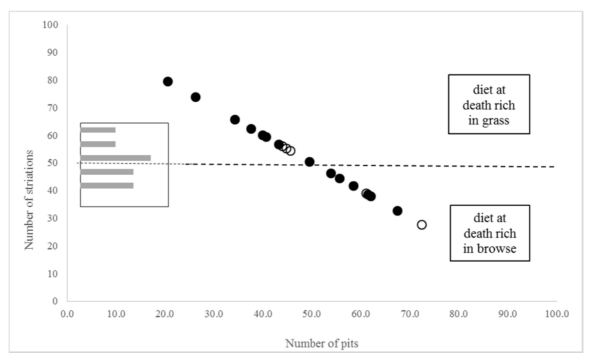


Fig 8. Chart of DMA results for all specimens, plotting percentage of pits against percentage of striations. M₃s are black circles and M₂s are open circles. The inset horizontal bar chart shows distribution in 10% ranges either side of 50%

DISCUSSION

One sub-adult year

To discuss seasonal mobility of Epipalaeolithic gazelle herds, specimen sequences are re-grouped by isotopic dataset (Fig.~9, $\delta^{18}O$ top row, ${}^{87}Sr/{}^{86}Sr$ middle row, $\delta^{13}C$ bottom row). The M_2 charts are to the left of M_3 charts suggestive of their developmental sequence. The two tooth-types (M_2 and M_3) together provide data that span an annual cycle, as can be seen from the fully sinusoidal $\delta^{18}O$ curves.

Starting with the δ^{18} O curves, it is tentatively argued that the specimens cluster, albeit loosely, to reflect animals living year-round in one climate regime in one location. Firstly, there are sharp seasonal differences between summer peaks and winter troughs mirroring those for local precipitation (*Fig.* 6) which markedly contrast cold, wet winters and hot, arid summers; inter-specimen differences reflect inter-annual weather differences but each maintain strong seasonal differences. Had herds avoiding the hot, arid summers of the steppe by moving into cooler Jordanian Highland the highly seasonal summer signature would be lost and the resultant peak reduced. the M3 specimen 'two' might fall into this category, but, secondly, differences proceed smoothly along all curve, more in keeping with seasonal change than with the halting process of migration across climate zones.

The data also show, with greater certainty that these gazelle were born in the same season, as the curves show enamel formation proceeding at approximately the same time in all teeth (highest values all towards the final section of the curve).

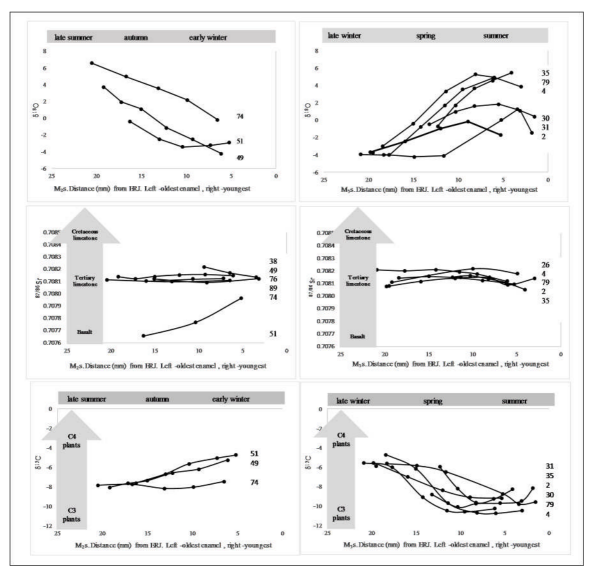


Fig 9. Charts showing all specimens on one chart for each isotope dataset, all M₂s on the left and M₃s on the right

 Turning to ${}^{87}\text{Sr}/{}^{86}\text{Sr}$ curves, it is apparent that, with one exception later revisited, these animals lived on the Tertiary limestone series and did not travel long distances either westwards into the Cretaceous limestone hills or north-east over the basalt lava flows. This evidence strongly corroborates the more tentative $\delta^{18}\text{O}$ interpretation. Strontium results alone do not allow us to determine how far animals moved over the limestone steppe however we argue that the likelihood is that the ${}^{87}\text{Sr}/{}^{86}\text{Sr}$ results, strongly clustered throughout the annual cycle, represent the local Tertiary limestone location. Firstly, zooarchaeological analysis of body-part representation suggests hunting was local to the site (Martin et al 2010 p123, Fig.7) and therefore herds were nearby at some point in the annual cycle; and secondly, where no advantage would be gained from extensive movement within a similarly resourced landscape, gazelle movement would be limited to minimal dispersal in order to conserve energy and resources.

Finally, the consistent link between δ^{13} C and δ^{18} O curves (*Fig.7*) show unstressed C₃ plants dominating food intake through late winter, into spring, and on into early summer, followed by increasing amounts of arid-

stressed C₃ plants, possibly including some C₄ plants in high summer and autumn. The modelled
 palaeovegetation proposes a longer growing season for annual grasses than today, explaining why C₃ species

remain available through the earlier summer. The ⁸⁷Sr/⁸⁶Sr data interpretation establishes that dietary changes

were not due to animal seasonal movement to more mesic vegetation zones, so it is argued that variation is due

5 to seasonal forage availability or to seasonal feeding preference; it is unnecessary to disentangle which as

feeding ethology and food seasonality are interrelated. In addition, for gazelle herds local to Kharaneh, gazelle

ethology allows us to assume a spring birth season when nutritious grasses flourished.

The weeks before death

If the above interpretations are correct, and these sampled specimens are representative of wider herd behaviour, it appears that gazelle herds in the vicinity of Kharaneh gave birth in spring and did not undertake long distance seasonal movements or migrations. We now turn to the evidence from DMA to assess hunting seasonality.

DMA evidence (*Fig.* 8) points to some of the sampled herd being hunted (local to Kharaneh) between late winter and early summer when grasses flourished and had not yet died back, and when feeding preference for lactating females in particular meant grasses were essential. Others were hunted when grasses had died off and the most palatable, and only available, food was browse which persists into arid periods (tree leaves, soft growth on woody shrubs, perennials and annuals such as chenopods). Yet others died on a mixed diet, possibly in early summer when sparse grasses had to be supplemented with browse, or in late winter when the first grass shoots were appearing and supplement the browse diet. The important point is that dental microwear evidence shows no indication of clustering in either one or two seasons, which might be expected if hunting had been seasonal or if gazelle presence had been seasonal; rather, DMA clearly points to hunting at Kharaneh being practiced in more than one season.

We have shown that gazelle remained on the limestone steppe year-round and were hunted in multiple seasons. The microwear gives a stronger signature for spring hunting than fusion data which noted a peak in winter culling, and helps define the broader spring/summer category identified by the cementum study. Each zooarchaeological method for determining cull seasonality has limitations, only allowing assessment of particular seasons rather than the whole annual cycle. The current study lends support, however, to other site evidence (Maher et al 2012a) indicating multi-seasonal and prolonged occupation of Kharaneh.

CONCLUSION

Discussion of hunter-gatherer mobility and seasonality in the Levantine Epipalaeolithic often relies on assumptions about underlying animal and plant resource availability. Some researchers draw on traditional models of radiating and circulating mobility around resource patches (Bar-Yosef & Belfer-Cohen 1989; Lieberman 1993), others look to recently observed pastoralist (Bedouin) seasonal movements in interpreting prehistoric hunter-gatherer mobility (Henry 1989). Maher and colleagues (2012b) question the validity of applying uniform mobility models across the Levant, where high degrees of ecological heterogeneity define the region. Indeed, the large-scale, multi-phase, repeatedly revisited site of Kharaneh (and nearby Wadi Jilat 6,

1 Garrard & Byrd 2013) attests to settlement organization quite different to the small campsites characterising the 2 Early/Middle Epipalaeolithic elsewhere. 3 4 This study makes an important contribution to the debate on settlement dynamics in the Epipalaeolithic Azraq 5 basin. Isotopic evidence suggests these gazelle herds were sustained year-round in the eastern Jordan limestone 6 steppe without recourse to long-distance seasonal migrations. An alternative model to that predicting large 7 migrating herds crossing the steppe might be one supporting localised seasonal aggregation and dispersal 8 consistent with seasonal variation in social groupings following vegetation availability and herd reproductive 9 cycles, similar to that seen amongst goitered gazelle in Saudi Arabia (Cunningham & Wronski 2011). This 10 scenario offers a different picture of wildlife availability in the Early/Middle Epipalaeolithic. Hunter-gatherers 11 could potentially have inhabited the steppes year-round, possibly aggregating and dispersing with herds. 12 13 Wadi Kharaneh may well have been an attractive winter location for hunting or mass-trapping large mixed 14 herds. But spring gazelle hunting probably targeted smaller herd groupings requiring different hunting 15 techniques such as stalking. We do not assume an ecologically deterministic relationship between hunters and 16 prey, the research presented here merely shows possibilities of how hunting practices may have varied 17 seasonally following wildlife behaviour. This study cautions against using recent and historical animal 18 migration patterns uncritically, especially where landscapes have been heavily impacted by humans and 19 livestock. 20 21 Finally, we return briefly to Specimen 51, the M₂ outlier with a strontium sequence indicating movement from 22 the basalt towards the Tertiary limestone series during the year (Fig. 9). This specimen (with no special 23 depositional conditions) serves as a reminder of the complexity of attempting reconstruction of past animal 24 mobility, and that regional (and indeed temporal) variation is to be expected. Future research, following the 25 methods used here, aims to include other Azraq Basin prehistoric sites, in order to explore spatial and temporal 26 variation of the animal behaviour that underpinned human use of the steppe in prehistory. 27 28 **ACKNOWLEDGEMENTS** 29 The research presented here was undertaken as part of a Leverhulme Trust funded multi-disciplinary project 30 (RPG-2013-223) and we thank the Trust for their support. We would also like to thank Ass. Prof. Lisa Maher 31 (Co-Director of EFAP) for permission to study materials from Kharaneh IV, and the Jordan Department of 32 Antiquities for allowing export of specimens. We are grateful for support provided by the CBRL, especially the 33 Director of the British Institute in Amman, Dr. Carol Palmer. Many thanks also to Sandra Bond, Tom Gregory 34 and Morag McBride at UCL's Institute of Archaeology and Christina Manning at Royal Holloway College UL. 35 36 **REFERENCES**

39 *206*, 187-221. 40

37

38

Akhani, H., Trimborn, P. Ziegler, H. 1997. Photosynthetic pathways in Chenopodiaceae from Africa, Asia and

Europe with their ecological, phytogeographical and taxonomical importance. Plant systematics and evolution

- 1 Ayliffe, L., Chivas, A. 1990. Oxygen isotope composition of the bone phosphate of Australian kangaroos.
- 2 Geochimica et Cosmochimica Acta, 54: 2603-2609.

- 4 Balasse, M. 2002. Reconstructing dietary and environmental history from enamel isotopic analysis: time
- 5 resolution of intra-tooth sequential sampling. *International Journal of Osteoarchaeology 12*, 155-165.

6

- 7 Balasse, M., Smith, A., Ambrose, S., Leigh, S. 2003. Determining sheep birth seasonality by analysis of tooth
- 8 enamel oxygen isotope ratios: the Late Stone Age site of Kasteelberg (South Africa). Journal of Archaeological
- 9 *Science 30*, 205-215.

10

- 11 Bar-Oz, G., Zeder, M., Hole, F. 2011. Role of mass-kill hunting strategies in the extirpation of Persian gazelle
- 12 (*Gazella subgutturosa*) in the northern Levant, *PNAS 108*, 7345–7350.

13

- 14 Bar-Yosef, O., Belfer-Cohen, A. 1989. The Origins of Sedentism and Farming Communities in the Levant.
- 15 Journal of World Prehistory 3, 447-498.

16

17 Batanouny, K. 2001. Plants in the deserts of the Middle East. Berlin: Springer.

18

- 19 Belfer-Cohen, A., Goring-Morris, A. 2013. The Upper Palaeolithic and Earlier Epi-Palaeolithic of Western
- Asia. In (Eds. Renfrew, A., Bahn, P.). The Cambridge World Prehistory, vol. 3, 1381-1407. Cambridge
- 21 University Press, Cambridge.

22

Bender, F. 1974. *Geology of Jordan*. Berlin: Borntraeger.

24

- 25 Bender, M. 1971. Variations in the ¹³C/¹²C ratios of plants in relation to the pathways of photosynthetic carbon
- dioxide fixation. *Phytochemistry 10*, 1239-1245.

27

- Bentley, A. 2006. Strontium isotopes from the earth to the archaeological skeleton: a review. Journal of
- 29 Archaeological Method and Theory 13, 135-187.

30

- 31 Blaise, E., Balasse, M. 2011. Seasonality and season of birth of modern and late Neolithic sheep from south-
- 32 eastern France using tooth enamel d18O analysis. Journal of Archaeological Science 38, 3085-3093.

33

- 34 Blank, D., Ruckstuhl, K., Yang, W. 2012. Influence of population density on group sizes in goitered gazelle
- 35 (Gazella subgutturosa Guld. 1780), European Journal of Wildlife Research 58, 981-989.

36

- Bocherens, H., Mashkour, M., Billou, D., Pellé, E., Mariotti, A. 2001. A new approach for studying prehistoric
- 38 herd management in arid areas: intra-tooth isotopic analyses of archaeological caprine from Iran. Earth and
- 39 Planetary Sciences 322, 67-74.

- 1 Britton, K., Grimes, V., Niven, L., Steele, T., McPherron, S., Soressi, M., Kelly, T., Jaubert, J., Hublin, J-J.,
- 2 Richards, M. 2011. Strontium isotope evidence for migration in later Pleistocene Rangifer: implications for
- 3 Neanderthal hunting strategies at the Middle Palaeolithic site of Jonzac, France. Journal of Human Evolution,
- 4 *61*, 176-185.

- 6 Bryant, J. Froelich, P. 1995. A model of oxygen isotope fractionation in body water of large mammals.
- 7 Geochimica et Cosmochimica Acta 59. 4523-4537.

8

- 9 Byrd, B., Garrard, A., Brandy, P. 2016. Modelling foraging ranges and spatial organization of Late Pleistocene
- 10 hunter-gathers in the southern Levant a least-cost GIS approach. *Quaternary International 396*, 62-78.

11

- 12 Cerling, T., Harris J. 1999. Carbon isotope fractionation between diet and bioapatite in ungulate mammals and
- implications for ecological and paleoecological studies. *Oecologia 120*, 347–363.

14

- 15 Colledge, S. 2001. Plant exploitation on Epipalaeolithic and early Neolithic sites in the Levant. Oxford, BAR
- 16 S986.

17

- 18 Cunningham, P., Wronski, T. 2011. Seasonal changes in group size and composition of Arabian Sand gazelle
- 19 Gazella subgutturosa marica Thomas 1987, during a period of drought in central western Saudi Arabia. Current
- 20 Zoology 57, 36-42.

21

Dansgaard, W. 1964. Stable isotopes in precipitation. *Tellus 16*, 436-468.

23

24 Davis, S. 1980. A note on the dental and skeletal ontogeny of Gazella. *Israel Journal of Zoology* 29, 129–134.

25

- de Niro, M., Epstein, S. 1978. Influence of diet on the distribution of carbon isotopes in animals. Geochimica et
- 27 *Cosmochimica Acta 42*, 495–506.

28

- Ehleringer, J., Cerling, T., Helliker, B. 1997. C₄ photosynthesis, atmospheric CO₂, and climate. *Oecologia 112*,
- 30 285-299.

31

- 32 Enzel, Y., Amit, R., Dayan, U., Crouvi, O., Kahana, R., Ziv, B., Sharon, D. 2008. The climatic and
- 33 physiographic controls of the eastern Mediterranean over the late Pleistocene climates in the southern Levant
- and its neighbouring deserts. Global and Planetary Change 60, 165-192.

35

Faure, G., Powell, J. 1972. Strontium Isotope Geology. New York: Springer.

- Feranec, R. 2007. Stable carbon isotope values reveal evidence of resource partitioning among ungulates from
- 39 modern C3-dominated ecosystems in North America. Palaeogeography, Palaeoclimatology, Palaeoecology
- 40 252, 575-585.

- 2 Fricke, H., Clyde, W., O'Neil J. 1998. Intra-tooth variations in δ^{18} O (PO₄) of mammalian tooth enamel as a
- 3 record of seasonal variations in continental climate variables. Geochimica et Cosmochimica Acta 62, 1839-
- 4 1850.

- 6 Fricke, H., O'Neil, J. 1996. Inter- and intra- tooth variations in the oxygen isotope composition of mammalian
- 7 tooth enamel phosphate: implications for palaeoclimatological and palaeobiological research. *Palaeogeography*,
- 8 Palaeoclimatology, Palaeoecology 126, 91-99.

9

- 10 Frison, G. 2004. Survival by hunting: prehistoric human predators and animal prey. University of California
- 11 Press.

12

- 13 Frumkin, A., Ford, D., Schwarcz, H. 1999. Continental oxygen isotopic record of the last 170,000 years in
- Jerusalem. *Quaternary Research* 51, 317–327.

15

- Garrard, A., Byrd, B. 2013. Beyond the Fertile Crescent: Late Palaeolithic and Neolithic communities of the
- 17 Jordanian Steppe. The Azraq Basin Project, Volume 1. Levant Supplementary Series 13, Oxford: Oxbow
- 18 Books.

19

- Garrard, A., Byrd, B., Harvey, P., Hivernel, F. 1985. Prehistoric environment and settlement in the Azraq
- Basin. A report on the 1982 survey season. *Levant* 17: 1-28.

22

- Gordon, K. 1982. A study of chimpanzee molars: implications for dental microwear analysis. *American Journal*
- of Physical Anthropology 59, 195-215.

25

- Goring-Morris, N. 1995. Ch. 10: Complex hunter-gatherers at the end of the Palaeolithic (20,000-10,000 BP),
- 27 142-167, in (Ed.) Levy, T. The Archaeology of Society in the Holy Land, Leicester University Press.

28

- Goring-Morris, N., Hovers, E., Belfer-Cohen, A. 2009. The dynamics of Pleistocene and Early Holocene
- 30 settlement patterns and human adaptations in the Levant: an overview. In (Eds.) Shea, J., Lieberman, D.
- 31 Transitions in Prehistory, 185-252. Oxbow Books.

32

- Graustein, W. 1989. ⁸⁷Sr/⁸⁶Sr ratios measure the sources and flow of strontium in terrestrial ecosystems. In
- 34 (Eds.). Rundel, P., Ehleringer, J., Nagy, K. Stable isotopes in ecological research. Springer-Verlag, New York,
- 35 491–512.

36

- Grine, F., Ungar, P., Teaford, M. 2002. Error rates in dental microwear quantification using scanning electron
- 38 microscopy. *Scanning* 24, 144-153.

- 1 Heaton, T. 1999. Spatial, species, and temporal variations in the 13C/12C ratios of C3 Plants: implications for
- 2 Palaeodiet studies. *Journal of Archaeological Science* 26, 637-649.

4 Henry, D. 1989. From Foraging to Agriculture. University of Pennsylvania Press.

5

- 6 Henton, E. 2010. (Unpublished PhD thesis). Herd management and the social role of herding at Neolithic
- 7 Catalhöyük: an investigation using oxygen isotope and dental microwear evidence in sheep. Institute of
- 8 Archaeology: University College, London.

9

- Henton, E., 2012. The combined use of oxygen isotopes and microwear in sheep teeth to elucidate seasonal
- 11 management of domestic herds: the case study of Çatalhöyük, central Anatolia. Journal of Archaeological
- 12 Science 30, 3264-3276.

13

- Hillman, G. 1996. Late Pleistocene changes in wild plant-foods available to hunter-gatherers of the northern
- 15 Fertile Crescent: possible preludes to cereal cultivation. In (Ed.) Harris, D. The origins and spread of
- *agriculture and pastoralism in Eurasia*. Washington D.C.: Smithsonian Institution Press, 159-203.

17

- Hoppe, K., Stuska, S., Amundson, R. 2005. The implications for paleodietary and paleoclimatic reconstructions
- of intrapopulation variability in the oxygen and carbon isotopes of teeth from modern feral horses. *Quaternary*
- 20 Research 64, 138-146.

21

- Hunt, C., Garrard, A., 2013. Section B: The Late Palaeolithic geological context. In: Garrard, A., Byrd, B.,
- 23 2013. Beyond the Fertile Crescent: Late Palaeolithic and Neolithic communities of the Jordanian steppe. The
- 24 Azraq Basin Project Volume 1. Levant Supplementary Series 13, Oxbow Books, Oxford, pp. 53-135.

25

- Iacumin, P., Longinelli, A. 2002. Relationship between $\delta^{18}O$ values for skeletal apatite from reindeer and foxes
- 27 and yearly mean δ^{18} O values of environmental water. Earth and Planetary Science Letters 201, 213-219.

28

29 IAEA/WMO. 2014. Global network of isotopes in precipitation. *The GNIP Database*. http://www.iaea.org/water

30

- Jones, J. 2012. Using gazelle dental cementum studies to explore seasonality and mobility patterns of the Early-
- 32 Middle Epipalaeolithic Azraq Basin, Jordan. *Quaternary International* 252, 195-201.

33

- Jones, M., Maher, L., Macdonald, D., Ryan, C., Rambeau, C., Black, S., Richter, T. 2016. The environmental
- setting of Epipalaeolithic aggregation site Kharaneh IV. Quaternary International 396, 95-104.

36

- 37 Jones, M., Richter, T. 2011. Paleoclimatic and archeological implications of Pleistocene and Holocene
- 38 environments in Azraq, Jordan. Quaternary Research 76, 363-372.

- 1 Jordan National Geographic Centre 1984. National Atlas of Jordan. Part I Climate and Agroclimatology.
- 2 Amman: Jordan National Geographic Centre.

- 4 Julien, M-A., Bocherens, H., Burke, A., Drucker, D., Patou-Mathis, M., Krotova, O., & Péan S. 2012. Were
- 5 European steppe bison migratory? ¹⁸O, ¹³C and Sr intra-tooth isotopic variations applied to a palaeoethological
- 6 reconstruction. *Quaternary International 271*, 106–119.

7

8 Kingswood, S., Blank, D. 1996. Gazella subgutturosa. Mammalian Species 518, 1-10.

9

- 10 Kohn, M., Schoeninger, M., Valley, J. 1996. Herbivore tooth oxygen isotope compositions: effects of diet and
- physiology. Geochimican et Cosmochimica Acta 60, 3889–3896.

12

- Kohn, M., Schoeninger, M., Valley, J. 1998. Variability in oxygen isotope compositions of herbivore teeth:
- reflections of seasonality or developmental physiology? *Chemical Geology 152*, 97-112.

15

16 Legge, A., Rowley-Conwy, P. 1987. Gazelle killing in Stone-age Syria. Scientific American 255, 88-95.

17

- Legge, A., Rowley-Conwy, P. 2000. The exploitation of animals. In (Eds.) Moore, A. Hillman, G., Legge, A.
- 19 *Village on the Euphrates–from foraging to farming at Abu Hureyra.* Oxford: Oxford University Press, 455-471.

20

- 21 Lieberman, D. 1993. The rise and fall of seasonal mobility among hunter-gatherers: the case of the southern
- 22 <u>Levant. Current Anthropology 34: 599-631</u>

23

- Lubinski, P., 2013. What is adequate evidence for mass procurement of ungulates in zooarchaeology?
- 25 Quaternary International 297, 167-175.

26

- Maher, L., Macdonald, D. 2013. Assessing typo-technological variability in Epipalaeolithic assemblages:
- Preliminary results from two case studies from the Southern Levant. In (Eds.): Borrell, F., Molist, M., Ibanez, J.
- 29 The state of stone: terminologies, continuities and contexts in Near Eastern lithics. Studies in Early Near
- 30 Eastern production, subsistence and environment 14. Ex oriente, Berlin, 29-44.

31

- 32 Maher, L., MacDonald, D., Allentuck, A., Martin, L., Spyrou, A., Jones, M. 2016. Occupying wide open
- spaces? Late Pleistocene hunter-gatherer activities in the Eastern Levant. *Quaternary International* 396, 79-94.

34

- Maher, L., Richter, T., Macdonald, D., Jones, M., Martin, L., Stock, J. 2012a. Twenty thousand-year-old huts at
- a hunter-gatherer settlement in Eastern Jordan, *PLoS ONE*, 7, 2, e31447.

37

- 38 Maher, L., Richter, T., Jones, M., Stock, J. 2011. The Epipalaeolithic Foragers in Azraq Project: prehistoric
- landscape change in the Azraq Basin, eastern Jordan. CBRL Bulletin 6:21-27.

- 1 Maher, L., Richter, T., Stock, J. 2012b. The Pre-Natufian Epipalaeolithic: long-term behavioural trends in the
- 2 Levant, Evolutionary Anthropology 21, 69-81.

- 4 Mainland, I. 1998. Dental microwear and diet in domestic sheep (Ovis aries) and goats (Capra hircus):
- 5 distinguishing grazing and fodder-fed ovicaprids using a quantitative analytical approach. Journal of
- 6 Archaeological Science 25, 1259-1271.

7

- 8 Martin, L. 2000. Gazelle (*Gazella* spp.) behavioural ecology: predicting animal behaviour for prehistoric
- 9 environments in south-west Asia. *Journal of Zoology, London* 250, 13-30.

10

- Martin, L., Edwards, Y., Garrard, A. 2010. Hunting practices at an Eastern Jordanian Epipalaeolithic
- aggregation site: the case of Kharaneh IV. *Levant 42*, 107-135.

13

- Merceron, G., de Bonis, L., Viriot, L., Blondel, C. 2005. Dental microwear of fossil bovids from northern
- 15 Greece: paleoenvironmental conditions in the eastern Mediterranean during the Messinian. *Palaeogeography*,
- 16 Palaeoclimatology, Palaeoecology 217, 173-185.

17

- 18 Merceron, G., Schulz, E., Kordos, L., Kaiser, T. 2007. Paleoenvironment of *Dryopithecus brancoi* at
- 19 Rudabánya, Hungary: evidence from dental meso- and micro-wear analyses of large vegetarian mammals.
- 20 Journal of Human Evolution 53, 331-349.

21

- Montgomery, J., Evans, J., Horstwood, M. 2010. Evidence for long-term averaging of strontium in bovine
- enamel using TIMS and LA-MC-ICP-MS strontium isotope intra-molar profiles. *Environmental Archaeology*
- **24** *15*, 32-42.

25

- Moore, A., Hillman, G., Legge, A. 2000. Village on the Euphrates. From foraging to farming at Abu Hureyra.
- 27 Oxford: Oxford University Press.

28

- Munro, N., Bar-Oz, G., Stutz, A. 2009. Aging mountain gazelle (Gazella gazelle): refining methods of tooth
- 30 eruption and wear and bone fusion. *Journal of Archaeological Science 36*, 752-763.

31 32

32 O'Leary, M. 1988. Carbon isotopes in photosynthesis. *Bioscience* 38, 328–336.

33

- Pearson, J., Grove, M. 2013. Counting sheep: sample size and statistical inference in stable isotope analysis and
- palaeodietary reconstruction. World Archaeology 45, 373-387.

36

- Poage, M., Chamberlain, C. 2001. Empirical relationships between elevation and the stable isotope composition
- 38 of precipitation and surface waters: considerations for studies of palaeoelevation change. American Journal of
- 39 Science 301, 1-15.

- 1 Price, T., Meiggs, D., Weber, M-J., Pike-Tay, A. 2015. The migration of Late Pleistocene reindeer: isotopic
- 2 evidence from northern Europe, *Archaeological and Anthropological Sciences*, 1-15.

- 4 Ramsey, M., Rosen, A. 2016. Wedded to wetlands: exploring Late Pleistocene plant-use in the Eastern Levant.
- 5 Quaternary International 396, 5-19.

6

- 7 Richter, T., Maher, L., Garrard, A., Edinborough, K., Jones, M., Stock, J. 2013. Epipalaeolithic settlement
- 8 dynamics in southwest Asia: New radiocarbon evidence from the Azraq Basin. Journal of Quaternary Science
- 9 28, 467-479.

10

- 11 Rivals, F., Prignano, L., Semprebon, G., Lozano, S. 2015. A tool for determining duration of mortality events in
- archaeological assemblages using extant ungulate microwear. Scientific Reports 5:17330 DOI:
- 13 *10.1038/srep17330*, 1-7.
- 14 Rivals, F., Solounias, N., Schaller, G. 2011. Diet of Mongolian gazelles and Tibetan antelopes from steppe
- habitats using premaxillary shape, tooth mesowear and microwear analyses. *Mammalian Biology* 76, 358-364.

16

- 17 Schulz, E., Fraas, S., Kaiser, T., Cunningham, P., Ismail, K., Wronski, T. 2013. Food preferences and tooth
- wear in the sand gazelle (*Gazella marica*). *Mammalian Biology* 78, 55-62.

19

- Shewan, L. 2004. Natufian settlement systems and adaptive strategies: the issue of sedentism and the potential
- 21 of strontium isotope analysis. In (Ed). Delage, C. The last hunter-gatherer societies in the Near East. BAR
- 22 (International Series) 1320: Oxford.

23

- Shomer-Ilan, A., Nissenbaum, A., Waisal, Y. 1981. Photosynthetic pathways and the ecological distribution of
- 25 the *Chenopodiaceae* in Israel. *Oecologia 48*, 244-248.

26

- Sillen, A., Hall, G., Richardson, S., Armstrong, R. 1998. 87Sr/86Sr ratios in modern and fossil food-webs of the
- 28 Sterkfontein Valley: implications for early hominid habitat preference. Geochimica et Cosmochimica Acta 62,
- **29** 2463-2473.

30

- 31 Solounias, N., Hayek, L-A. 1993. New methods of tooth microwear analysis and application to dietary
- determination of two extinct antelopes. *Journal of Zoology London* 229, 421-445.

33

- 34 Solounias, N., Moelleken, S. 1992. Tooth microwear analysis of *Eotragus sansaniensis* (Mammalia:
- Ruminantia), one of the oldest known bovids. *Journal of Vertebrate Paleontology 12*, 113-121.

36

- 37 Solounias, N., Semprebon, G. 2002. Advances in the reconstruction of ungulate ecomorphology with application
- 38 to early fossil equids. *American Museum Novitates 3366*, 1-49.

- 1 Spyrou, A. 2014. Gazelle carcass utilisation during the Early/Middle Epipalaeolithic of the southern Levant:
- 2 zooarchaeological and ethnographic implications of animal nutrient storage among human foragers. Assemblage
- 3 *PFAZ*, 1-18.

- 5 Teaford M. & Walker A. 1984. Quantitative differences in dental microwear between primate species with
- 6 different diets and a comment on the presumed diet of Sivapithecus. American Journal of Physical
- 7 Anthropology 64, 191-200.

8

- 9 Tieszen, L. 1991. Natural variations in the carbon isotope values of plants: implications for archaeology,
- ecology and paleoecology. *Journal of Archaeological Science* 18, 227–248.

11

- Tornero, C., Bălășescu, A., Ughetto-Monfrin, J., Voinea, V., Balasse, M. 2013. Seasonality and season of birth
- in early Eneolithic sheep from Cheia (Romania): methodological advances and implications for animal
- economy. *Journal of Archaeological Science 40*, 4039-4055.

15

- 16 Ungar, P. 2002. Microwear image analysis software, version 4.02 (04.10.02) Buffalo Edition (Retrieved
- 17 10.01.05).

18

- Vogel, J., Fuls, A., Danin, A. 1986. Geographical and environmental distribution of C₃ and C₄ grasses in the
- Sinai, Negev, and Judean deserts. *Oecologia* 70, 258-265.

21

- Wilkie, T., Mainland, I., Albarella, U., Dobney, K., Rowley-Conwy, P. 2007. A dental microwear study of pig
- diet and management in Iron Age, Romano-British, Anglo-Scandinavian, and Medieval contexts in England. In
- 24 (Eds.) Albarella, U., Dobney, K., Ervynck, A., Rowley-Conwy, P. Pigs and humans: 10,000 years of
- *interaction.* Oxford: Oxford University Press, 241-254.

26

- Zazzo, A., Mariotti, A., Lécuyer, C., Heintz, E. 2002. Intra-tooth isotope variations in late Miocene bovid
- 28 enamel from Afghanistan: paleobiological, taphonomic, and climatic implications. *Palaeogeography*,
- 29 Palaeoclimatology, Palaeoecology 186, 145-161.

30

31 Zohary, M. 1973. Geobotanical foundations of the Middle East, Vols 1 & 2. G. Fischer: Stuttgart.

33

32

34

35

36 37

38

SUPPLEMENTARY MATERIAL

Supp 1. - 87Sr/86Sr in modern plants from three locations

This supplement provides a little background information on the strontium isotope baseline that is referred to in this paper and used to contribute to the identification of key geological locations. The supplement forms a preliminary report on a small part of a more comprehensive isotopic baseline using modern plant collections, which is currently in publication preparation. The ⁸⁷Sr/⁸⁶Sr results reported here are those collected from key geological endmembers within and around the Azraq Basin: the Cretaceous limestone series of the Highlands to the west, the Basalt lava flow to the northeast and the Early Tertiary limestone series near Kharaneh IV.

The leaves and soft shoots of moderately rooted plants were collected in one season (April) and used to measure labile strontium. Each sampling aliquot included material from three individual plants and duplicates were run to test intra-species variation at one location. Plants were exported to UCL Institute of Archaeology with the permission of the Department of Antiquities of Jordan. Samples were first washed in milli-Q water then, in the Earth Sciences Department at Royal Holloway University of London, labile strontium was extracted from homogenised material followed standard protocols, and analyses were made in a VG354 thermal ionisation mass spectrometer. Typical external reproducibility for 87 Sr/ 86 Sr is ± 0.000014 (2sd).

Results are displayed below in Supplementary 1, Figure 1. It is immediately clear that the plant labile ⁸⁷Sr/⁸⁶Sr values cluster into three groups that discriminate between the three locations and convincingly reflect underlying geologies (Bentley 2006; Hartman & Richards 2014). The averaged ⁸⁷Sr/⁸⁶Sr readings for the Cretaceous Highland location is 0.708941‰, the Early Tertiary limestone steppe 0.708087‰, and the basalt steppe is 0.707720‰.

Taxa * duplicate analyses from one plant	Location UTM (north hemisphere)	87Sr/86Sr average of 3 plants	Bedrock geology
Phlomis fruticosa Salvia heirosolymitana Anthemis sp.	Upper Wadi Yabis 36: 7.597 E x 35.881 N	0.708479 0.708453 0.708543	Cretaceous limestone series (c2)
Hordeum glaucum Achillea fragrantissima Malva parviflora	Kharaneh IV 37. 37: 2.587 E x 35.128 N	0.708115 0.708069 0.708078	Early Tertiary limestone series (tt1)
Achillea fragrantissima* Achillea fragrantissima*	Shubayqa 1 37: 3.334 E x 35.868 N	0.707731 0.707737	Pliocene/Oligocene basalt (B4)

Hordeum glaucum*	0.707775	
Hordeum glaucum*	0.707780	
Malva parviflora*	0.707642	
Malva parviflora*	0.707656	

Supp. 1, Table 1. Labile 87Sr/86Sr in plants collected from 3 locations on different geologies



Supp. 1, Fig. 1. Chart showing results of modern plant ⁸⁷Sr/⁸⁶Sr analyses at 3 geological locations in the Azraq

Basin and the Jordanian Highlands