Cave paleozoology in the Judean Desert: assembling records of Holocene wild mammal communities

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ABSTRACT: Long temporal records of Holocene wild mammal communities are essential to examine the role of human impacts and climatic fluctuations in the configuration of modern ecosystems. We show that such records can be assembled through extensive radiocarbon dating of faunal remains obtained from biogenic cave deposits. We dated 110 mammalian remains from 19 different cave sites in the Judean Desert. We use the dates in combination with archaeological survey data and bone collagen/apatite δ¹³C values to study faunal succession in the context of Holocene climate change and human settlement history in the region. Our results suggest a change in the late Holocene, expressed in fewer observations of Arabian leopard (Panthera pardus nimr) and gazelle (Gazella spp.), and an increase of Syrian striped hyena (Hyaena hyaena syriaca), fox (Vulpes spp.), Nubian ibex (Capra nubiana) and rock hyrax (Procavia capensis); suids (Sus scrofa) appear for the first time. According to the data distribution, however, the probability of finding a bone diminishes exponentially with time, which implies that the Judean Desert cave paleozoological record is temporally biased. The weight of evidence ultimately favors an explanation of the observed patterns as the consequence of a combined anthropogenic and climatic impact on local food webs.

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Introduction

Paleozoological data can inform on species abundance and distribution in the past (Faith and Lyman, 2019), with broad relevance to topics ranging from climate change to extinctions and conservation biology (Boivin et al., 2016; Tyler and Schneider, 2018). Such data, however, are difficult to obtain for the later Holocene, when larger wild mammals almost disappear from the paleozoological record and are replaced by domesticated animals (Diamond, 2002; Tsahar et al., 2009). As a result, tracking the dynamics of later Holocene wild mammal community structure and its relation to anthropogenic and climatic impacts remains a challenge for paleozoologists, especially those working in the earliest centers of human societal complexity, which were also cradles of domestication (Arbuckle, 2018). The problem posed by the deterioration of the ‘wildlife signal’ in the paleozoological record with the transition to agriculture can be tackled in specific regions where there exists, on the one hand, high-resolution data on human settlement history, and, on the other, natural archives that preserve millennium-scale record of contemporary wildlife (e.g. Byers and Broughton, 2004; Stewart et al., 2021). These regions can be used as windows to the multiplex interactions between agricultural societies and wildlife communities in the Holocene (Fig. 1).

Such a rare juxtaposition of Holocene human and wildlife records can be found in the southern Judean Desert, west of the Dead Sea (Fig. 2). There, the punctuated settlement history at the major oasis in the region, Ein Gedi, is known from years of survey and excavations and is supplemented by historical sources (Hadas, 2005; Davidovich, 2013; Ussishkin, 2014). This archaeological settlement record is matched by well-preserved biogenic assemblages of faunal remains found in the hundreds of karstic caves dotting the high-relief topography of the region (Horwitz et al., 2002; Frumkin, 2015; Lazagabaster et al., 2021a,b). Together, the archaeological and paleozoological data provide an opportunity to study Holocene wild mammal communities from a binocular perspective, against the background of environmental studies that have been conducted in the Dead Sea (e.g. Neumann et al., 2010; Stewart et al., 2021). We use radiocarbon-dated remains of larger wild mammals from the Judean Desert caves in tandem with Holocene settlement dynamics and climatic data to observe diachronic change in the medium to large (>1 kg) mammal community. Ultimately, we investigate whether observed change can be ascribed to biotic interactions (including with humans), environmental change or a mixture of both. To the published paleoenvironmental assessments of the area (e.g. Bar-Matthews et al., 1999; Enzel et al., 2003; Vaks et al., 2003;
Litt et al., 2012), we add new bone collagen/apatite δ¹³C results from our paleobiological collection. Disentangling the role of Anthropocene environmental and human influence on a large mammal community is expected to advance our understanding of transformative interactions between humans, other faunas and the environment in the later Holocene, which are difficult to track using other methods.

Paleozoological record

The paleozoological record of the southern Judean Desert caves has been mostly formed by animal feeding and denning activities (Kerbis-Peterhans and Horwitz, 1990; Horwitz et al., 2002; Lazagabaster et al., 2021b). The importance of caves for animals is amplified in this hyper-arid region by the absence of other shelters, such as trees and thickets. In the near-absence of fluvial and eolian sedimentation, cave deposits are typically shallow (<1 m) and almost entirely biogenic, comprising decayed dung, nesting materials and bones from in situ mortality and carnivore accumulation (Frumkin, 2015; Frumkin et al., 2017). The dry, loose and shallow sediments have been constantly turbated and reworked by animals, and, in the last century or so, by cave looters. The result of this deposition history, combined with the pristine condition of dry-preserved organics (Weiner, 2010, p. 10), is that there is no stratigraphic or visual way to estimate the date of faunal remains from the caves without direct radiocarbon chronometry. This archaeologically hopeless situation of sediment mixing implies that the cave paleozoological collection represents a time-random sample of the fauna that has roaming the region mostly during the Holocene and the Late Pleistocene (Lazagabaster et al., 2021a,b).

In contrast to the central importance of the caves to animals in the Judean Desert, these karstic spaces have seen intermittent human use (Davidovich et al., 2015). Archaeological remains in the Judean Desert cliff caves are from three main episodes: late fifth millennium BCE, late 4th millennium BCE and 2nd century CE (Porat et al., 2007; Davidovich, 2013). Human occupation during these episodes was possibly related to temporary refuge and spanned weeks or months and has had little if any input to the faunal accumulation. For instance, we have not found evidence of butchery on any of the studied Holocene wild mammal remains. Visits to certain caves are
documented for other periods as well, yet again connected with ephemeral activities (Davidovich et al., 2015). Human contribution to cave wild mammal assemblage formation on a thousand-year scale is therefore considered negligible.

**Settlement record**

Many of the faunal remains found in the caves, however, are of domesticated animals (Horwitz et al., 2002). These represent hunted or scavenged household animals from the focal point of human settlement in the region: the Ein Gedi Oasis. This oasis, the major habitable locality along the southern Dead Sea coast (Hirschfeld, 2007; Stem, 2007), has attracted human settlement throughout the late Holocene (see settlement history in the Results). The intensity of distinct settlement episodes in the Ein Gedi oasis is an important variable that can potentially relate to wild animal community structure in the region through time (Lazagabaster et al., 2021a). As the major perennial water source in the region, human settlement and activity at the oasis could limit access to that critical resource. Furthermore, human settlement could have influenced the composition of wild mammal communities by either bottom-up or top-down effects, such as overgrazing, tree-cutting or ungulate hunting, on the one hand, and large predator killing, on the other (Dietl and Flessa, 2017; Thomas, 2020). Scavenging opportunities and protection from predators near settlements would probably join other subtle niche construction by humans that would have affected local biodiversity (Boivin et al., 2016; Zeder, 2016).

**Climate**

Another independent variable affecting mammal community structure is climate. Numerous paleoenvironmental studies in the region have revealed the complexity of Quaternary climate (e.g. Litt et al., 2012; Törnqvist et al., 2015; Kiro et al., 2016; Vaks et al., 2018; Lazagabaster et al., 2021b). From these we learn that during the late Pleistocene the terminal Lake Lisan stand was much higher than that of the Dead Sea, its Holocene successor, probably indicating a lower evaporation rate and potentially higher precipitation in the northern headwaters of the River Jordan that provides most of the water input to the lake. The early Holocene, until the fourth millennium BCE, may have been a wetter period than the late Holocene (Rambau, 2010). Although the difference is not on par with the Pleistocene/Holocene sharp climatic changes (Törnqvist et al., 2015; Lazagabaster et al., 2021b), small climatic fluctuations can be important in marginal regions due to the increase in the flow of spring waters or slight increases in humidity that can determine the survival of resilient flora. Because of a rain shadow effect, however, the Judean Desert would have been ultimately dry under the Mediterranean storm track regime of the Holocene.

**Preliminary observations and research questions**

Our team has surveyed caves in the southern Judean Desert and has recovered a substantial number of identified large mammal remains. We assumed that the probability of dating a bone is directly related to that taxon’s frequency in the life assemblage from the region at a specific period. Using this ‘dates as data’ (Rick, 1987; Brierley et al., 2018) approach, we tracked the succession of large carnivore remains from the paleozoological cave survey in the Holocene Judean Desert, which suggested a late Holocene decline in the frequency of leopards (*Panthera pardus nimr*) and a concomitant increase in striped hyenas (*Hyaena hyaena syriaca*) (Lazagabaster et al., 2021a). This turnover is temporally correlated with settlement intensification in the region, which suggests that humans could have had an impact on Holocene wild mammal communities. However, the role of aridification leading to a reduction in prey population – and, ultimately, leopard populations – could not be tested based solely on carnivore data.

In this study, we substantially broaden the scope of our previous investigation by adding new radiocarbon dates and stable isotope readings of herbivores to the carnivore dataset. Using these extended data, the Holocene large mammal succession in the study region can be evaluated also from the primary consumer perspective. This is very important, as it can resolve the open question of reduced prey availability as a possible cause of the carnivoran turnover from leopards to hyenas and shed light on the carrying capacity of the marginal desert environment throughout the Holocene. More broadly, the new data will be used to discuss large mammal biotic interactions and response to human- and climate-induced landscape change in the region. Specifically, by tracing change in the distribution of radiocarbon dates of primary consumers from the Judean Desert cave sites we would like to see if there is change through time in the composition of the herbivore community in the study region. In case we observe succession, the faunal changes can then be discussed in relation to observed changes in human settlement intensity or the local climatic record.

We state four hypotheses closely related to each other concerning predator–prey dynamics (Hypothesis 1), climate change (Hypothesis 2), intraguild competition (Hypothesis 3) and human impact (Hypothesis 4). Hypothesis 1 suggests that with changes in the carnivore community (e.g. the decrease in leopard population) we will observe changes in the frequency of prey (e.g. an increase in the frequency of leopards’ main prey species, the Nubian ibex and the rock hyrax) (Wallach et al., 2017; Winnie and Creel, 2017). An alternative hypothesis (Hypothesis 2) to explain potential changes in large mammal herbivores is climate change: we expect that the increasing trend in heat and aridity in the region towards the late Holocene drier period would adversely affect large herbivore demography (Broughton et al., 2008); this may, in turn, influence predator density. Hypothesis 3 states that with changes in the frequency in one or various carnivore species, other carnivorans will show an increase or decrease in numbers (e.g. increase in the frequency of foxes because of the depressed leopard frequencies) (Prugh et al., 2009; Ritchie and Johnson, 2009; Ripple et al., 2013). Alternatively (Hypothesis 4), changes in the density of certain carnivoran taxa may be related to human impact (e.g. in the case of striped hyenas, due to new foraging opportunities around human settlements).

**Material and methods**

**Archaeological survey**

In the last two decades, R.P. and U.D. have led cave and site surveys and excavations in the region. Their results were integrated with the works of other scholars to a fine-grained synthesis of regional settlement dynamics during the Holocene (see Lazagabaster et al., 2021a: supplementary information for references). In 2019–2020, additional survey and excavation work in the Ein Gedi Oasis was led by R.P. and U.D., aiming to clarify the settlement history during the 5th and the 1st millennia BCE. Their work included polygon surveys, 16 random probes and four small-scale excavations in parts of
the oasis flagged by the new surveys as containing potentially important clues to the spatial and temporal extent of settlement based on pottery techno-typology. The archaeological data collected by the team were compared with previous published surveys using ArcGIS and used to categorize local human settlement intensity into low (<0.5 ha), medium (0.5–1.0 ha) and high (>1.0 ha) levels based on the area of convex hulls drawn around settlements and find spots (see Lazagabaster et al., 2021a: Table 1).

**Paleozoological cave survey**

A survey carried out during 2019–2020 recovered 36 faunal assemblages from 43 caves in the southern Judean Desert (Fig. 2). We surveyed caves that could be accessed by experienced persons without rappelling gear, as such caves are approachable by terrestrial mammals and therefore reflect hunting and scavenging in the immediate region. Potentially identifiable bone and tooth fragments of macro-mammals and hyena coprolites were collected from the surface and from crevices and sediment pockets within the surveyed caves by the field teams led by M.U. and R.P. To our survey data we added faunal collections recovered from the Cave of Skulls, excavated by Amir Ganor, Eitan Klein, R.P., M.U. and U.D., on behalf of the Israel Antiquities Authority and the Hebrew University in 2016; and a single carnivore bone (EG-039) collected in an excavation conducted by Gideon Hadas in Ein Gedi.

**Archaeozoological analysis**

Animal remains from the survey were identified using the comparative collection of the Laboratory of Archaeozoology at the University of Haifa, with reference to the collections at the Steinhardt Museum of Natural History at Tel Aviv University and to the National Collections at the Hebrew University in Jerusalem. The resulting database comprises 2123 identified specimens. A total of 110 bones from 19 sites were sent for radiocarbon dating and stable isotope analysis; 64 belong to primary consumers and are reported here for the first time.

We prioritized dating and isotopic analysis of the large (>21 kg) carnivores *P. pardus nimir* and *H. hyaena*, acknowledging their role as keystone taxa and their relative rarity in the mammal community. Although the carnivore guild of the Judean Desert also contains wolves and jackals, we could not differentiate the small local variants of the Arabian wolf (*Canis lupus arabs*) and the golden jackal (*Canis aureus*) from domestic dogs (*Canis lupus familiaris*), and their remains were not dated. We also dated Blanford’s fox (*Vulpes cana*) and *Vulpes sp.* (either Rüppell’s fox *V. rueppellii* or red fox *V. vulpes*) specimens. From among the herbivores, we dated only bones that could be identified with certainty as belonging to a wild taxon: gazelles (*Gazella gazella, G. dorcas*), rock hyrax (*Procavia capensis*), suids (*Sus scrofa*) and Nubian ibex (*Capra nubiana*). Ibex specimens were selected only when they could be distinguished with confidence from those of domestic goats (*Capra hircus*), as in occipital bones and horn cores. Information on the ecological background (habitat and feeding behaviors) of the local populations of these taxa is summarized in Table 1.

In all cases, effort was made to maximize the number of caves from which the sample of each taxon was obtained (Table 2); this was done both to obtain a spatially balanced sample of the study region, and to reduce potential interdependence between specimens. Specimens from the same taxon but from different caves were considered as belonging to two different individuals; specimens identified to the same taxon but originating in the same cave were considered different individuals if their dates were different by at least two radiocarbon year error intervals.

**Radiocarbon dating and δ13C**

A total of 110 bones representing at least 89 individuals from 19 sites are included in this study (Appendix S1; Table 2). Primary consumers (*n* = 64) include suids (*n* = 14), Nubian ibex (*n* = 10), gazelle (*n* = 15) and rock hyrax (*n* = 25). Secondary consumers (*n* = 46) include striped hyena (*n* = 20), leopard (*n* = 12) and fox (*n* = 14). While these sample sizes for some taxa are relatively small, they are probably sufficient to

| Taxon              | Habitat                                      | Feeding                                      |
|--------------------|----------------------------------------------|----------------------------------------------|
| *Gazella cf. Gazella dorcas* | Desert, ˂150 mm per annum                  | Leaves, flowers and pods of *Acacia* trees, twigs and leaves of other bushes; grazing on annuals in season       |
| *Capra nubiana*                              | Cliffs in desert regions                    | Grasses and herbs during winter, browsing in the summer       |
| *Procavia capensis*                          | Rocky terrain in different environments. In desert regions they are more numerous near water sources | Herbs and grass in the winter and spring, leaves in the summer. Can subsist of plants with high concentration of secondary metabolites |
| *Panthera pardus nimir*                      | Desert oases, wadis and cliffs              | Mainly hyrax and ibex                                      |
| *Sus scrofa*                                  | Dense thickets, forests, riverine habitats  | Omnivorous: small vertebrates, insects, carrion, nuts, acorns, berries, rhizomes, seeds |
| *Hyaena hyaena*                               | Generalist                                  | Carrion, bone, hunting rare                             |
| *Vulpes cana*                                 | Rocky slopes, canyons and cliffs in arid and hot regions | Insectivores, frugivores                               |

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Table 1. Description of the habitat and feeding ecology of the mammal taxa mentioned in this work. Most information is summarized from Mendelssohn and Yom-Tov (1999).
compare central tendencies in the dates between taxa. The selected bones were dated in the Oxford Radiocarbon Accelerator Unit (ORAU; n = 57) and at the Center for Applied Isotope Studies (CAIS), University of Georgia (n = 55). When possible, organic carbon from collagen was used for radiocarbon dating; otherwise, carbon from bioapatite was dated.

Although bioapatite is susceptible to contamination by soil carbonates, in arid environments with dry sediments this method is considered reliable (Zasso and Saliege, 2011). Nevertheless, we dated two specimens using both collagen and bioapatite to check for the possible effect of contamination on bioapatite carbon results. The dates obtained were comparable, suggesting no significant contamination of soil carbonates (see Lazagabaster et al., 2021a).

The $^{13}$C/$^{12}$C ($\delta^{13}$C) ratios were measured separately using a stable isotope ratio mass spectrometer and expressed as $\delta^{13}$C with respect to PDB, with an error of <0.1‰. The $\delta^{13}$C values are used to discuss possible shifts in animal diet and environment through time in terrestrial ecosystems. Due to differences between isotope spacing in different tissue and mineral fractions (collagen and bone apatite), an average $\Delta^{13}$C$_{\text{coll-bon}}$ spacing of 5‰ correction fraction was applied to the bioapatite samples ($n = 10$) (Lee-Thorp et al., 1989; France and Owsley, 2015; Codron et al., 2018). Nitrogen isotopes were not used because of contamination, to which we were alerted when specimens with high nitrogen content were found not to have preserved collagen. The source of contamination is probably animal urine.

In the presentation of the results and in the analyses, the samples are temporally grouped following the formal subdivision of the Holocene epoch into Early, Middle and Late according to Walker et al. (2019): the Early–Middle Holocene Boundary is situated at 8200 a BP and the Middle–Late Holocene Boundary at 4200 a BP.

The laboratory protocols of the radiocarbon dating facilities are available online (CAIS: https://cais.uga.edu/facilities/radiocarbon-ams-facility/; ORAU: https://c14.arch.ox.ac.uk/methods.html).

Data analysis

The radiocarbon ages were calibrated using the Intcal20 curve. Our primary means for interpreting changes in large mammal communities is bootstrap ($n = 1000$) confidence intervals (95%) calculated for the median cal a BP dates of each taxon, presented along with the jitter of calibrated dates and probability density plots. This gives a simple and robust measure for the peak frequency of a taxon in the sample, with minimal assumptions. A comparison between median dates employed a Kruskal–Wallis test and a post-hoc Dunn test with Bonferroni correction. Finally, to account for potential preservation biases on the overall sample, we fit various probability distributions to the data and choose the best fit based on Akaike’s information criterion (AIC) values. Fitting distributions to data is a very common task in statistics and consists in choosing a probability distribution modeling the random variable, as well as finding parameter estimates for that distribution (Motulsky and Christopoulos, 2004). Data analysis was facilitated by libraries ‘rcarbor’, ‘tidyverse’, ‘FSA’ and ‘fitdistrplus’ in R, using R Studio.

Results

Historical human settlement intensity in Ein Gedi

Three distinct phases of human settlement can be observed at the oasis of Ein Gedi (Fig. 3). The first was during an early stage of the Late Chalcolithic period, probably around the third quarter of the 5th millennium BCE and is best known by the isolated building complex above the main spring known as the Ein Gedi Shrine (Ussishkin, 1980). Several nodes of activity from the same period observed along the Ein Gedi spring slope indicate the existence of a sparsely populated hamlet (Shai et al., 2007). The second phase of settlement in the Ein Gedi oasis occurred from the 7th century BCE to the 6th century CE, with short (decadal scale) occupation gaps. During this span, from the later Iron Age to the Byzantine era (Hirschfeld, 2007; Stern, 2007). Settlement remains from these periods are abundant, and agricultural terraces and water systems turned the oasis into an intensively managed landscape severely impacted by human activities, which peaked in the Roman–Byzantine era (Hadas, 2012). The area occupied by

![Figure 3](https://example.com/figure3.png)
Roman period remains could have been as large as 27.7 ha, reduced to 5.9 ha during the Byzantine period (Lazagabaster et al., 2021a). Following a long abandonment, the oasis was briefly resettled during the Mamluk period in the 14th–15th centuries CE, when a small hamlet was built above the Byzantine ruins (Hirschfeld, 2007).

Distribution of radiocarbon dates and taxonomic succession

The earliest concentration of dates corresponds to early Holocene gazelles, with a range of 10 560–7829 cal a BP (95% confidence interval (CI); median = 9578 cal a BP). The leopard dates also peak early, between 6082 and 5174 cal a BP (95% CI; median 6082 cal a BP). A similarly early 10 560–7829 cal a BP (95% CI) range brackets the peak frequency of early Holocene gazelles (median 9578 cal a BP). The dates of the other taxa, however, appear to climaxed significantly later: the ibex (5164–949 cal a BP; median 1773 cal a BP), the hyrax (3064–854 cal a BP; median 1226 cal a BP) and the group of Late Holocene gazelles (2320–591 cal a BP; median 1838 cal a BP). Foxes (4036–1031 cal a BP; median 2283 cal a BP), suids (2717–425 cal a BP; median 890 cal a BP) and hyenas (11 109–145 cal a BP; median 668 cal a BP) share even later peak frequencies (Figs. 3 and 4). These observations on the dissimilar distribution of dates between taxa are supported by a Kruskal–Wallis test ($\chi^2 = 47.66$, d.f. = 6, $p < 0.0001$; see Table 3 for Kolgomorov–Smirnov multiple-comparison tests).

We also consider the possibility that the temporal patterns emerging from the radiocarbon dates are due to inherent preservation biases of the paleozoological record. We fit three common probability distributions to the data, including normal, logarithmic and exponential (Fig. 5). The data are better explained by an exponential distribution with an exponential parameter of 0.329 (Table 4). This means that the probability of finding a bone decreases exponentially with time. If this is true, then we cannot simply assume that more dates of a particular taxon in the paleozoological record reflect more overall abundance in the environment. Rather, the discussion of dates needs weighting each sample relative to its age. This matter is discussed in the following sections.

Table 3. The results of statistical comparisons of median cal a BP dates between taxa using non-parametric Kruskal–Wallis and multiple pairwise Kolgomorov–Smirnov tests with False Discovery Rate correction. Significant statistical differences are highlighted in bold.

| Taxon comparisons | D-stat | p-value |
|-------------------|--------|---------|
| Gazelle1 Fox      | 0.38   | 0.5091  |
| Gazelle2 Fox      | 1.00   | <0.0001 |
| Hyena Fox         | 0.56   | 0.0115  |
| Hyrax Fox         | 0.27   | 0.6482  |
| Ibex Gazelle1     | 0.30   | 0.8930  |
| Leopard Fox       | 0.77   | 0.0022  |
| Suid Fox          | 0.43   | 0.2410  |
| Gazelle2 Gazelle1 | 1.00   | 0.0011  |
| Hyena Gazelle1    | 0.50   | 0.1431  |
| Hyrax Gazelle1    | 0.28   | 0.8390  |
| Ibex Gazelle1     | 0.30   | 0.8930  |
| Leopard Gazelle1  | 0.92   | 0.0020  |
| Suid Gazelle1     | 0.43   | 0.3593  |
| Hyena Gazelle2    | 1.00   | <0.0001 |
| Hyrax Gazelle2    | 0.92   | 0.0008  |
| Ibex Gazelle2     | 0.90   | 0.0022  |
| Leopard Gazelle2  | 0.92   | 0.0028  |
| Suid Gazelle2     | 1.00   | 0.0001  |
| Hyrax Hyena       | 0.40   | 0.1066  |
| Ibex Hyena        | 0.50   | 0.1090  |
| Leopard Hyena     | 0.95   | <0.0001 |
| Suid Hyena        | 0.21   | 0.9124  |
| Ibex Hyrax        | 0.22   | 0.9124  |
| Leopard Hyrax     | 0.76   | 0.0008  |
| Suid Hyrax        | 0.27   | 0.6534  |
| Leopard Ibex      | 0.73   | 0.0115  |
| Suid Ibex         | 0.37   | 0.4283  |
| Suid Leopard      | 0.92   | 0.0002  |
Carbon stable isotopes

The δ¹³C results are presented in Table 5 and Fig. 6, and in Appendix S1 for the three major primary consumer taxa that existed in the Judean Desert throughout the Holocene: gazelle, ibex and hyrax.

Carbon isotope ratios of bone collagen are predominantly routed from dietary protein and represent the weighted average of foods consumed over a period of several years (Froehle et al., 2010). Herbivore tissue δ¹³C therefore reflects the carbon composition of the plants that they consumed (Jim et al., 2004; Richards, 2020). Three photosynthetic pathways influence the δ¹³C values of plants: C₃, C₄ and Crassulacean Acid Metabolism (CAM). Most plants that use the C₃ photosynthetic pathway (trees, most forbs and shrubs) have δ¹³C values that range between −35 and −20‰ with a mean of ca. −27‰; C₄ plants include dry adapted grasses, and some forbs and shrubs, and exhibit higher values than C₃, ranging between −15 and −7‰ with a mean of ca. −12‰; CAM plants (cacti, succulents and epiphytes) have δ¹³C values that range between −22 and −10‰ (Kohn, 2010; Sage et al., 2012; Munroe et al., 2021).

Previous research in modern local C₃ vegetation of Tell es-Safi/Gath (west coast of Israel) shows δ¹³C values ranging from −29 to −25‰ (Arnold et al., 2016).

Our samples show mean δ¹³C values in herbivores of −17.6‰ in the Early Holocene, −17‰ in the Middle Holocene and −18.2‰ in the Late Holocene. These results suggest herbivores such as gazelles were eating vegetation from a mixed C₃/C₄ environment, which is consistent with grazing typical of this antelope in the Levant (see Hartman, 2012) during the early period, with more strong presence of C₄ vegetation for two of the hyrax samples, as reflected by their carbon isotope values, which range from −14.3 to −18.5‰. Instead, towards the recent Holocene, a slight increment on C₃ diet appears to occur for hyraxes as δ¹³C values are more depleted (ranging from −18 to −20.6‰). Ibex samples from the last two periods show a larger component of C₃ in their diet, consistent with caprine mixed feeding behavior (Hartman, 2012). Unfortunately, with only one sample of ibex from the earlier Holocene, we cannot monitor major changes in their diet.

Discussion

Dates equal data? Understanding the distribution of radiocarbon dates in the paleozoological record

The cave biogenic assemblages of the Judean Desert are natural archives of the wildlife living in the region throughout history. The formation of the different cave deposits and skeletal assemblages reported in this study can vary greatly depending on the size and spatial arrangement of the caves, the different inhabitants of the caves (including humans), the predators that accumulated bones, the impact of anthropogenic activities (including modern alteration by looters, hikers, archaeologists, local Bedouins) and abiotic processes (e.g. water, temperature, soil pH, mineral diagenesis). The study of these caves is challenging as the bone assemblages often present hardly any stratigraphy, with anachronic remains mixed and dispersed throughout (Lazagabaster et al., 2021a,b).

Thus, the potential to overcome these biases and understand how the deposits were formed is limited. Radiocarbon dating gets us as close as we could to a true paleozoological record. Our underlying assumption is simple, yet useful: among other factors the possibility of finding a bone of a certain taxon in the paleozoological record is related to the abundance of that taxon in the surroundings. Other factors might include ecological guild, life history and innate behaviors. Thus, when...
there are apparent gaps in the record, it does not necessarily imply that a given taxon disappears from the region, but rather that its population was probably lower than in other times. The paleontological motto ‘absence of evidence is not evidence of absence’ readily applies in this context. This is exemplified by the absence of leopard remains younger than 3500 cal aBP in the record (Lazagabaster et al., 2021a). We know from historical sources, molecular analyses and modern sightings that leopards had been present historically in the Judean Desert until the last one was killed in a road accident 15 years ago (Perez et al., 2006). Therefore, the absence of leopards in the paleozoological record probably reflects a reduction in their population instead of its complete disappearance from the region.

Table 5. Summary statistics of carbon stable isotope data of primary consumers. EH, Early Holocene; LH, Late Holocene; max, maximum, MH; mid-Holocene; min, minimum; SD, standard deviation.

| Taxon | Period | LH min | LH max | LH mean | LH SD | MH min | MH max | MH mean | MH SD | EH min | EH max | EH mean | EH SD |
|-------|--------|--------|--------|---------|-------|--------|--------|---------|-------|--------|--------|---------|-------|
| Gazelle | LH     | −18.5  | −15.3  | −16.8   | 1.1   | −18.1  | −16.1  | −17.0   | 1.0   | −18.0  | −15.4  | −17.4   | 1.3   |
| Hyrax  | LH     | −18.8  | −14.5  | −18.6   | 1.4   | −18.5  | −14.3  | −17.0   | 1.9   | −18.8  | −14.3  | −17.7   | NA    |
| Ibex   | LH     | −18.3  | −15.9  | −18.5   | NA    | −17.4  | −17.0  | −17.2   | 0.8   | −18.1  | −17.1  | −18.8   | 0.3   |
| Suid   | LH     | −20.9  | −24.5  | −20.7   | 1.2   | −18.3  | −20.7  | −18.1   | NA    | −20.9  | −27.1  | −20.7   | NA    |

**Figure 6.** Changes through time in bone collagen/bioapatite $\delta^{13}$C of primary consumers in the Holocene Judean Desert suggest stability in plant composition, and hyrax population growth in the oasis [Color figure can be viewed at wileyonlinelibrary.com]
patterns emerge for the different taxa analyzed. The data suggest significant change in the fauna of the Holocene Judean Desert. The early and mid-Holocene saw the peak in frequency of both leopards and gazelles. The late Holocene marks a drastic change. Gazelle and leopard numbers drop; ibex and hyrax become the dominant herbivores; smaller foxes, and then synanthropic hyena, step to the frontline of the carnivore guild. The late Holocene also marks the first appearance of suids in the regional record; these may be the ancestors of the small isolated wild boar population of Mount Sodom on the coast of the Dead Sea. Preliminary morphological work (Marom et al., 2019) suggests that the extant Dead Sea boar population is morphologically indistinguishable from wild suids in the Mediterranean regions of the country. Thus, we suggest that the Dead Sea boars could have been introduced during the Late Holocene and subsequently became feral; this hypothesis is awaiting further investigation.

When all the data are pooled together, however, it is evident that the distribution of radiocarbon dates is skewed, with dates concentrated towards recent times. We have fit various common probability distributions and we show that the data approach an exponential distribution. This means that the probability of finding a bone in the Judean Desert Holocene record diminishes exponentially with time. This effect may be due to preservation or taphonomic factors. As time passes, natural degradation of the bone matrix can result in bone destruction (Keenan and Engel, 2017; Kendall et al., 2018); the low sedimentation in the caves would explain why there is almost no fossil diagenesis and why pre-Holocene remains are usually not preserved. This explanation, however, has a caveat: the state of bones in the caves remains physically pristine, even with specimens beyond the limit of radiocarbon dating maintaining full structural integrity. Moreover, the abundance and preservation of skeletal remains in caves is greatly influenced by spatial variation in preservation conditions and the mineralogical composition. Karkanas et al. (2000) proposed stepwise mineral diagenetic processes rather than uniform gradual transformations to explain why some materials could be affected differently. An alternative explanation could be the increase in the presence of hyenas in the Late Holocene. Many of the most frugivorous caves are active or inactive hyena dens and hyenas are known to be efficient bone accumulators (Kerbis-Petersans and Horwitz, 1990; Orbach and Yeshurun, 2021). These are not the only bone accumulators in the desert though; certainly porcupines, foxes and other small carnivores are cave dwellers and can contribute to the assemblages with their remains and that of their prey (Horwitz et al., 2002; Lazagabaster et al., 2021a). Hyraxes also seek refuge in caves and crevices and contribute to the in situ mortality (Yom-Tov, 1993). Finally, large ibex and gazelle remains are brought to the caves by large carnivores, including leopards, wolves and hyenas. The importance of caves as feeding and resting places to bone accumulators is expected to have increased as trees disappeared from the region due to late Holocene dryness and human exploitation.

Nevertheless, the exponential decrease in the likelihood of finding a bone with time demands that we interpret the data with caution. If more ‘weight’ is given to bones found in older time periods than those found in recent time periods, then the Late Holocene peak in abundance flattens, except for hyenas and suids. The populations of ibex, gazelle, hyrax, fox and leopard would have been higher during the early and mid-Holocene, with numbers reduced in the Late Holocene. If we assume that the high abundance of prey (ibex, gazelle, hyrax, suid) in the Late Holocene is a taphonomic artifact – either due to negative diagenetic processes or to increased hyena activity – then the absence of leopard remains in the paleozoological record is even more telling and would tend to reinforce our previous suggestion that leopard abundance in the Judean Desert drastically declined after ~3500 cal a BP (Lazagabaster et al., 2021a). The Late Holocene would have consequently seen a generalized downgrading of the ecosystem probably due to the combined effect of top-down cascading effects, aridification and human impacts.

**Faunal carbon stable isotope ecology in relation to Holocene environmental fluctuations**

Although the magnitude of regional Holocene climatic oscillations was smaller in comparison with those that occurred during the Pleistocene (Bar-Matthews et al., 1999; Liszer et al., 2010; Torfstein et al., 2013, 2015; Frumkin and Comay, 2021), the Judean Desert has seen environmental fluctuations throughout the Holocene that could have affected the composition of mammalian communities. Today, the Judean Desert shows arid conditions with high temperatures and mean annual rainfall below 150 mm. The region’s environment is determined by Mediterranean seasonal climate regimes, characterized by wet, cool winters and dry, hot summers (Soto-Berlov et al., 2015). Rainfall patterns are influenced by synoptic-scale circulation patterns forced by an upper-level Mediterranean trough (Armon et al., 2019). Active subtropical jets and active Red Sea troughs, in addition to heavy precipitation in the highlands, can also contribute to sporadic rainstorms that produce intense floods in the wadis that discharge into the Dead Sea. The Dead Sea water level, however, is mainly controlled by Mediterranean cyclones and the amount of water discharged into the Dead Sea, especially from the Lower Jordan River (Armon et al., 2019). The vegetation of the Judean Desert consists mostly of Saharo-Arabian flora, with some Irano-Turanian and Sudanian components, the latter especially in the areas surrounding the Dead Sea (Miebach et al., 2019). Arid-adapted grasses using the C₄ photosynthetic pathway represent 17% of the vegetation in the Judean uplands, but this number increases to 50% in the Dead Sea valley, including around the area of Ein Gedi. The C₃ plants are mostly associated with rocky terrain, crevices and wadi slopes, while C₃ vegetation dominates in open plains and loess soils (Vogel et al., 1986). There are few trees and bushes, and these are mostly concentrated near water springs, such as in Ein Gedi.

Regional paleoclimatic records derived from the analysis of cave speleothems (e.g. Soreq Cave) show that the modern seasonal regime of wet winters and dry summers in the southern Levant has been in place since the beginning of the Holocene (Orland et al., 2012). However, the reduction in precipitation in the western hills during periods of Mediterranean drought could have influenced the local availability of water and the amount of vegetation cover. For example, it has been suggested that a drop in annual rainfall of about ~200 mm on average occurred between 4600 and 4000 cal a BP (Bar-Matthews and Ayalon, 2004). Pollen remains recovered from a sediment core drilled near Ein Gedi suggest that three major Holocene climatic time intervals characterize the region (Litt et al., 2012): a relatively dry and warm period in the Early-Pottery Neolithic and Pottery Neolithic (~10–6.5 ka cal a BP; though this interpretation seems contradictory with the high Dead Sea lake stand, see Migowski et al., 2006); a relatively wet and cool interval between the Chalcolithic and the Late Bronze Age (~6.3–3.3 ka cal a BP); and a rapid change to drier and warmer conditions at ~3200 cal a BP. The Late Holocene drying trend coincides with a relatively sharp drop in the Dead Sea level at ~3500 cal a BP (Frumkin et al., 2001; Enzel et al., 2003; Bookman et al., 2004;
Holocene, with fewer and relatively smaller oscillations (Frumkin et al., 2001; Bar-Matthews and Ayalon, 2004; Morin et al., 2019). The absence of travertine deposition during at least 2000 years in Morowa Cave, situated near the Ein Gedi spring, confirms that precipitation during the Late Holocene was not enough to support speleothem deposition (Lisker et al., 2007). Plant remains found in the Ein Gedi spring, confirms that precipitation during the Late Holocene period (Lisker et al., 2007). Plant remains found in the ‘Caves of the Spear’ from around the time of the Bar-Kokhba revolt (~135 CE) indicate the presence of relict vegetation that was more abundant and diverse in the area than it is today (Simchoni and Kislev, 2009). Thus, it is evident that the aridification trend has persisted until today and is likely to continue in the future.

The vegetation cover of the rocky slopes of the Judean Desert produces pasture for local herbivores, mainly for the Nubian ibex and the rock hyrax (Vogel et al., 1986; Alba, 2016). The carbon stable isotope data on the bones of these herbivorous mammals do not show significant changes in diet throughout the Holocene. However, there is a trend towards more negative δ13C values with time (Fig. 6). This trend may indicate a slight reduction in C4 grasslands and a change to feeding more often in dense vegetation spots with more abundant C3 plants near water springs. The change towards diets with a higher C3 plant component is evident in the hyrax during the Late Holocene. Thus, it is possible that these values reflect a general reduction in the Judean Desert’s vegetation outside of the local aquifers and springs that irrigate the different oases. Furthermore, the diminishment of leopards (the main predator) could have produced a change in the ecology of hyraxes, which would have ventured more safely next to the water springs. In general, the δ13C values of C3 plants are more negative in those growing in wetter conditions.

By contrast, water availability does not appear to influence C4 plant δ13C values, as they are generally less sensitive to environmental variation (Zhao et al., 2010). However, some desert plants obtain most of their water from secondary sources, namely water channeled by local topographic features rather than direct rainfall, as in the case of oasis vegetation, resulting in more negative δ13C values than plants from exposed ridge desert microhabitats (Hartman and Danin, 2010).

According to the Israeli Nature and Parks Authority, the Ein Gedi Natural Reserve has a diverse flora, including some of the densest concentrations of tropical plants growing in Israel and the northernmost point at which some of these tropical species can be found. Although some of these plants are less abundant today due to the use of part of the reserve’s water, the Ein Gedi Oasis is still a biodiversity hotspot of fauna and flora in the desert. Thus, it is likely that with the increase in aridity towards the Late Holocene and the diminishment of vegetation, the whole faunal community of the Judean Desert turned to these humid and vegetated refugia for food and water.

Table 6. Description of the animal, human and environmental changes in the Holocene of the Judean Desert described in this study.

| Parameter                      | Early and mid-Holocene | Late Holocene | Summary                            |
|--------------------------------|------------------------|---------------|------------------------------------|
| Human settlement               | Hamlet, ritual, ephemeral Oasis, grassland | Large village, intensive agriculture Oasis, no grassland | Intensification Aridification |
| Environment                    |                        | More gazelle, more C₄ grazing Leopards | Increase + introduction & feralization Mesopredators & scavengers |
| Mammal community structure     | Herbivores             | More ibex and hyrax, introduction of pigs, larger numbers | Fox and hyena |
|                               | Grazing                |                            |                                    |
| Carnivores                     |                        |                            |                                    |

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Trophic cascades and ecological collapse: towards an integrative ecological view of Holocene human–environment interactions

The Dead Sea area has been the subject of extensive paleoclimatic and paleoenvironmental studies, including regarding Dead Sea level fluctuations (Enzel et al., 2003; Bookman et al., 2004; Migowski et al., 2006; Kagan et al., 2015), cave speleothems (Bar-Matthews et al., 1999; Vaks et al., 2006; Lisker et al., 2007; Vaks et al., 2018), dating of cave wood detritus (Frumkin et al., 1991), pollen archives (Leroy, 2010; Neumann et al., 2010; Litt et al., 2012; Langgut et al., 2014; Miebach et al., 2017; Chen and Lieb, 2018) and sedimentology (Dayan and Morin, 2006; Kiro et al., 2016). However, no record of Holocene faunal changes in the Judean Desert had previously been published. The paleoecological survey and radiocarbon dates of cave skeletal assemblages is the first attempt to reconstruct changes in wild faunal communities in the Judean Desert (Lazagabaster et al., 2021a,b). This is a step forward in the integration of multiple avenues of data with the overarching goal of providing a holistic view of the evolution of the ecosystem in the area through time and examining historical human–environment interactions. Furthermore, paleoenvironmental reconstructions of the Judean Desert derived from Dead Sea pollen records and cave speleothems should be taken with caution, because they reflect regional, rather than local, environmental conditions (see discussion in Lazagabaster et al., 2021a,b). The cave paleozoological assemblages, by contrast, are local archives of the fauna that lived near the caves.

We suggest that the later part of the Holocene, and especially the first millennium BCE and the early first millennium CE, has seen changes in the environment, human settlement intensity and large mammal community structure (Table 6). With regard to human settlement intensity, our fieldwork has confirmed and quantified previous observations on the extent of human use of the major oasis of Ein Gedi throughout the Holocene, as a proxy for the general impact of humans on the southern Judean Desert (Lazagabaster et al., 2021a). Surveys and excavation in the oasis have not discovered a substantial mid-Holocene settlement but have enlarged the known size of the first millennium BCE and CE settlements.

Circling back to the hypotheses of our study, is it possible to ascribe a cause for the observed changes to predator–prey dynamics, climatic factors, intraguild competition or anthropogenic impacts? The answer is, as usual, not clear-cut. Interpreting the results at face value, the increase in hyrax and ibex numbers, which goes against the tide of adverse climate and more intensive human settlement, is suggested to have been caused by relief of predation stress following a reduction of the leopard population, confirming Hypothesis 1. Hypothesis 2, which stated that the number of larger ungulates would decrease in the late Holocene drier period due to the increasing trend in heat and aridity, can be rejected, as...
both ibex, hyrax, and suids seem to proliferate. Despite the possible loss of earlier Holocene habitats and the decline in the population of gazelles, the overall primary consumer community has not been impoverished. Foxes would also have benefited from the disappearance of the top predator in the local ecosystem (Prugh et al., 2009), so there is reason to believe that, to some extent, carnivoran intraguild competition has also played an important factor in shaping the Holocene large mammal communities of the Judean Desert (Hypothesis 3).

However, we also show the possibility that the paleozoological record of the Judean Desert is temporally biased (Fig. 5), which implies that the apparent abundance of ibex and hyrax populations in the Late Holocene could be an artifact of the record. Regardless of the different interpretations of the data, what is noticeable from the radiocarbon dates is that the diminishment of leopard populations occurs before the peak in dated remains of its major prey, ibex, gazelle and hyrax. The herbivore succession should have supported the leapord population throughout the Late Holocene and thus our previous hypothesis that the decrease in leopards and the increase in hyenas was caused by human activities (e.g. hunting and waste) is strongly supported (Hypothesis 4; Zagazbaster et al., 2021a). These conclusions are, of necessity, still preliminary, and await the results of an ongoing genetic study on the Holocene demographic trends of these key carnivore and herbivore taxa. The new research presented here, however, takes another step in the effort to unravel the intricate interactions between animals, humans and the environment in the difficult depositional conditions of the post-domestication Holocene of the Judean Desert.

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Data Availability Statement

The data and R code that support the findings of this study are openly available in Open Science Framework at https://osf.io/x6w24/.

Supporting information

Additional supporting information can be found in the online version of this article.

Appendix S1. Specimen list and radiocarbon and carbon isotope data.

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CAVE PALEOZOOLOGY IN THE JUDEAN DESERT 661
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