Contents lists available at ScienceDirect





CrossMark

# **Earth-Science Reviews**

journal homepage: www.elsevier.com/locate/earscirev

# Late Pleistocene and Holocene mammal extinctions on continental Africa

## J. Tyler Faith \*

Archaeology Program, School of Social Science, University Of Queensland, Brisbane, QLD 4072, Australia

## ARTICLE INFO

Article history: Received 23 April 2013 Accepted 14 October 2013 Available online 24 October 2013

Keywords: Quaternary Megafauna Overkill Climate change Paleoecology

## ABSTRACT

Understanding the cause of late Quaternary mammal extinctions is the subject of intense debate spanning the fields of archeology and paleontology. In the global context, the losses on continental Africa have received little attention and are poorly understood. This study aims to inspire new discussion of African extinctions through a review of the extinct species and the chronology and possible causes of those extinctions. There are at least 24 large mammal (>5 kg) species known to have disappeared from continental Africa during the late Pleistocene or Holocene, indicating a much greater taxonomic breadth than previously recognized. Among the better sampled taxa, these losses are restricted to the terminal Pleistocene and early Holocene, between 13,000 and 6000 yrs ago. The African extinctions preferentially affected species that are grazers or prefer grasslands. Where good terrestrial paleoenvironmental records are present, extinctions are associated with changes in the availability, productivity, or structure of grassland habitats, suggesting that environmental changes played a decisive role in the losses. In the broader evolutionary context, these extinctions represent recent examples of selective taxonomic winnowing characterized by the loss of grassland specialists and the establishment of large mammal communities composed of more ecologically flexible taxa over the last million years. There is little reason to believe that humans played an important role in African extinctions.

© 2013 Elsevier B.V. All rights reserved.

#### Contents

1.	Introduct	10	)6
2.	Africa and	extinctions debate	)6
	2.1. Af	n the global context	)6
	2.2 Ex	for research in Africa	)7
3	The extin	ammals 10	18
5.	31 Or	Artiodactula 10	18
	3.1	Aenveeros sn	18
	3	Antidorcos australis	18
	3.1	Antidorcas bondi	18
	3.1	Ros nrimigenius	19
	3.1	Damaliscus hypsodon	9
	3.1	Damalisaus niro	19
	3.1	Gazella atlantica	)9
	3.1	Gazella tineitana	19
	3.1	Himotragis Jeuconhaeus	9
	3	Meralogran isance miscus	19
	3	Rusingruv atonocranion 11	1
	3	Swncerus antianus	1
	3	Unnamed carrine	1
	3	Camelus 11	2
	3	Megazernides algeriaus	2
	3	Matridichamu	2
	2	Kolnochaerus	2
	J.	Roipochocrus	-

<sup>\*</sup> Tel.: +61 7 336 53314; fax: +61 7 336 51544. *E-mail address:* j.faith@uq.edu.au.

<sup>0012-8252/\$ -</sup> see front matter © 2013 Elsevier B.V. All rights reserved. http://dx.doi.org/10.1016/j.earscirev.2013.10.009

3.2.1.       Equus algericus		. 112
3.2.2.       Equus capensis		. 112
3.2.3.       Equus mauritanicus         3.2.4.       Equus melkiensis         3.2.5.       Stephanorhinus         3.3.       Order Proboscidea		. 112
3.2.4.       Equus melkiensis		. 113
3.2.5.         Stephanorhinus		. 113
3.3. Order Proboscidea		. 113
		. 113
3.3.1. Elephas iolensis		. 113
3.4. Order Tubulidentata		. 113
3.4.1. Orycteropus crassidens		. 113
3.5. Problematic species		. 113
3.5.1. Hipparion lybicum	• •	. 113
3.5.2. Giraffa gracilis	• •	. 113
3.5.3. Parmularius	• •	. 114
3.5.4. Equus quagga	• •	. 114
3.5.5. Gazella rufina (=Eudorcas rufina)	• •	. 114
4. Chronology and explanations	• •	. 114
4.1. Southern Africa	• •	. 114
4.2. East Africa	• •	. 115
4.3. North Atrica	• •	. 115
5. Extinctions in ecological and evolutionary context	• •	. 115
6. Conclusion	• •	. 116
Acknowledgments	• •	. 11/
References	• •	. 11/

## 1. Introduction

The last ~100,000 yrs witnessed massive extinctions of large mammals across the continents. Understanding what caused these extinctions is one of the most contentious problems in Quaternary science, with debate focusing primarily on the extent to which anthropogenic or climatic drivers are to blame (MacPhee, 1999; Barnosky et al., 2004; Koch and Barnosky, 2006; Wroe et al., 2013). Compared to the losses in North America, Australia, and Eurasia, the late Quaternary extinctions (LQE) on continental Africa have received little attention and are considered to be poorly understood (Martin and Steadman, 1999; Barnosky et al., 2004; Koch and Barnosky, 2006).

Klein (1984b) provided the most comprehensive review of African LQE to date, with particular attention to the South African record. However, within the context of the global megafaunal extinction debate, it is Maglio and Cooke's (1978) volume on the evolution of African mammals that provides the ultimate primary reference for most accounts of African species thought to have disappeared during the late Pleistocene and Holocene (e.g., Martin, 1984; Smith et al., 2003; Barnosky et al., 2004; Lyons et al., 2004; Koch and Barnosky, 2006). The species lists generated from this volume have yet to be updated or vetted by contemporary standards, despite substantial gains in African paleontology over the last several decades (Werdelin and Sanders, 2010). The result is that species now known to have disappeared during the middle Pleistocene routinely appear on lists of taxa thought to have disappeared within the last ~100,000 yrs (e.g., the bovid Parmularius and the equid Hipparion) (e.g., Barnosky et al., 2004; Lyons et al., 2004; Koch and Barnosky, 2006), whereas more recent additions do not appear at all (e.g., Bagtache et al., 1984; Pickford and Thomas, 1984; Brink, 1999; Faith et al., 2011, 2012).

A comprehensive account of the mammals involved is needed if Quaternary scientists are to begin to understand the causes of African LQE. The goal of this paper is to provide such a synthesis for the late Pleistocene (126,000 to 12,000 yrs ago) and Holocene (<12,000 yrs ago) of continental Africa, in addition to an assessment of the extinction chronology and possible explanations.

## 2. Africa and the extinctions debate

#### 2.1. Africa in the global context

Despite being poorly understood, the African LQE play an important role in the global extinction debate. The African losses are often characterized as less severe than extinctions elsewhere (Martin, 1984; Barnosky et al., 2004; Koch and Barnosky, 2006). Martin (1984: 380) remarks that the African mammals "escaped being greatly altered by late Pleistocene extinctions" while Koch and Barnosky (2006: 221) consider Africa "a fortunate anomaly." This is true in the sense that Africa supports an exceptionally diverse large mammal community today (Kingdon, 1982). For example, there are 38 genera of extant African megafauna (>44 kg), compared to fewer than 20 on other continents (Eurasia: 17; North America: 13; South America: 10; Australia: 2) (Koch and Barnosky, 2006).

The apparently anomalous survival of Africa's large mammals is thought by some to reflect long-term co-evolution of human predators and their prey (Martin, 1984; Lyons et al., 2004). The earliest representatives of our genus (Homo habilis) appeared in Africa ~2.3 million yrs ago (Kimbel et al., 1997; Antón, 2012), with fossil and genetic evidence pointing to an African origin of Homo sapiens between 200,000 and 100,000 yrs ago (White et al., 2003; McDougall et al., 2005; Gonder et al., 2007; Campbell and Tishkoff, 2010; Brown et al., 2012). African hominins incorporated animal tissues into their diet perhaps as early as 3.4 million yrs ago (McPherron et al., 2010; but see Domínguez-Rodrigo et al., 2010) and there is ample evidence for the exploitation of terrestrial prey by ~2.6 to 1.5 million yrs ago (Bunn, 1981; Potts and Shipman, 1981; Bunn and Kroll, 1986; Domínguez-Rodrigo et al., 2005; Pobiner et al., 2008; Domínguez-Rodrigo et al., 2009; Ferraro et al., 2013). Although there is debate about when early hominins became proficient hunters (Marean and Assefa, 1999; Klein, 2001; Domínguez-Rodrigo, 2002; Faith, 2008; Ferraro et al., 2013), the longterm development of hunting behavior is argued to have translated to the evolution of defense mechanisms in African prey, rendering them relatively immune to the impacts of prehistoric hunters (Martin, 1984).

The co-evolution argument is central to Martin's (1967, 1984, 2005) overkill hypothesis, which proposes that human hunters caused

massive extinctions as they dispersed throughout the continents during the last ~70,000 yrs. According to Martin, the greater severity of extinctions in Australia and the Americas was the result of naïve prey that lacked the behavioral defense mechanisms of their African counterparts (see Wroe et al., 2004 for a critique). The presumed vulnerability of non-African faunas was such that invading human hunters were able to wipe them out in a geological instant, meaning that little archeological evidence for overkill is to be expected. Martin also argued that the survival of Africa's megafauna is evidence that late Quaternary climate change is incapable of driving mass extinction of large mammals. No matter what their potential, albeit contentious (e.g., Grayson and Meltzer, 2003; Wroe et al., 2004, 2013), applicability to extinctions elsewhere, these arguments offer little insight into the losses that occurred in Africa.

## 2.2. Extinction research in Africa

The study of Pleistocene African mammals was fueled by the discovery of hominin fossils from sites in Southern and East Africa, including the Sterkfontein Caves and Olduvai Gorge (Fig. 1), following World War II. These discoveries stimulated interest in the associated fauna in order to establish the biochronological, paleoenvironmental, and evolutionary context of the hominin fossils (Maglio and Cooke, 1978). This tradition continues today, with much of the recent progress in Cenozoic African paleontology driven by paleoanthropological research (Sanders and Werdelin, 2010). The extinct mammals discussed here, although corresponding to a much more recent phase of human prehistory, are no exception. The majority of sites yielding extinct late Pleistocene and Holocene mammals are the focus of archeological investigations into the behavioral, ecological, and environmental context of early *H. sapiens* (Fig. 1). Most of these are caves or rockshelters that were inhabited by Middle Stone Age (MSA: 280,000 to 40,000 yrs ago) or Later Stone Age (LSA: <40,000 yrs ago) humans, with the associated faunal remains typically representing discarded food refuse. While evidence that humans hunted extinct megafauna is scant in the Americas or Australia (Grayson and Meltzer, 2002; Barnosky et al., 2004; Koch and Barnosky, 2006; Wroe and Field, 2006), there is abundant evidence that MSA and LSA foragers routinely preyed upon extinct mammals in Africa (for two exceptional cases see Marean, 1997; Milo, 1998).

Klein (1972, 1974b, 1980, 1984b) was the first to address African LQE in the context of the global extinction debate. His critically important work focused on the southern African record, where a long history of archeological research provided the most detailed late Quaternary faunal sequence for the continent. At the time, this record documented the extinction of a handful of open grassland ungulates near the Pleistocene–Holocene transition, between 12,000 and 9500 yrs ago. The association of these extinctions with a major episode of faunal turnover, characterized by the replacement of ungulate grazers by ungulate browsers, suggested that environmental changes were at least partly responsible. Because the extinct taxa survived previous glacialinterglacial transitions, however, Klein reasoned that human impacts



Fig. 1. Location of sites referred to in the text: (1) Jebel Irhoud, (2) El Harhoura 1 & 2, (3) Mugharet el 'Aliya, (4) Tighenif, (5) Beausejour Farm, (6) Les Phacochères, (7) Tamar Hat, (8) Jebel Thaya, (9) Hammam el Meskhoutine, (10) Haua Fteah, (11) Natodameri, (12) Shungura Formation, (13) Rusinga & Mfangano islands, (14) Lainyamok, (15) Lukenya Hill, (16) Gol Kopjes, (17) Olduvai Gorge, (18) Kisese II, (19) Redcliff Cave, (20) Chelmer Farm, (21) Kruger Cave, (22) Sterkfontein & Swartkrans, (23) Border Cave, (24) Wonderwerk Cave, (25) Modder River, (26) Florisbad & Vlakkraal, (27) Koffiefontein, (28) Colwinton, (29) Elands Bay Cave, (30) Elandsfontein, (31) Swartklip, (32) Die Kelders Cave, (33) Boomplaas Cave, (34) Nelson Bay Cave.

108

were also a necessary contributor. He argued that the Pleistocene–Holocene transition was unique in that it was the first glacial–interglacial transition when large mammal communities were subject to predation by highly effective hunters armed with LSA technology. According to Klein, MSA human populations, although anatomically modern, lacked the cognitive and technological prowess of their LSA successors and had little impact on animal populations.

Subsequent research on African LQE has favored environmental extinction hypotheses (Brink, 1987; Marean and Gifford-Gonzalez, 1991; Brink and Lee-Thorp, 1992; Marean, 1992; Brink, 1994, 1999; Faith, 2011c; Faith et al., 2011, 2012, in press). Anthropogenic hypotheses have fallen out of favor, in part because archeological research over the last several decades has shown the behavioral contrast between MSA and LSA humans proposed by Klein (1980, 1984b) to be untenable (e.g., Marean and Assefa, 1999; McBrearty and Brooks, 2000; Henshilwood and Marean, 2003; Faith, 2008). A better understanding of how environmental change or human impacts may have contributed to the extinctions first requires a detailed account of the species involved.

## 3. The extinct mammals

There are at least 24 large mammal (>5 kg) species known to have disappeared from continental Africa during the late Pleistocene (126,000 to 12,000 yrs ago) or Holocene (<12,000 yrs ago), all of which are globally extinct (Table 1, Fig. 1). These include species from six genera that are globally extinct (Megalotragus, Rusingoryx, Megaceroides, Metridiochoerus, Kolpochoerus, Stephanorhinus), two that are represented in Africa today only by domesticates (Bos and Camelus), and one that survives elsewhere (Elephas). An extinct caprine antelope from South Africa (Brink, 1999) likely accounts for a seventh globally extinct genus (see below). Although the precise extinction chronology for many of these taxa is poorly known, all of the taxa listed in Table 1 have been recovered from late Pleistocene or Holocene contexts as established by radiometric dates or associated archeological industries. Fossil occurrences associated with late Pleistocene or Holocene radiometric dates are reported in Table 2. Absolute ages associated with extinct mammals are derived primarily from radiocarbon (<sup>14</sup>C), Uranium

#### Table 1

Extinct	late Pleistocene	and Holocer	e African	mammals.	Taxa in	bold	represent	genera
that are	globally extinct	t.						

Order	Family	Taxon
Artiodactyla	Bovidae	Aepyceros sp. Antidorcas australis Antidorcas bondi Bos primigenius <sup>a</sup> Damaliscus hypsodon Damaliscus niro Gazella atlantica Gazella tingitana Hippotragus leucophaeus <b>Megalotragus priscus</b> <b>Rusingoryx atopocranion</b> Syncerus antiquus Unnamed caprine antelope
	Camelidae	Camelus sp. <sup>a</sup>
	Cervidae	Megaceroides algericus
	Suidae	Metridiochoerus sp.
Perissodactyla	Equidae	Kolpochoerus sp. Equus algericus Equus capensis Equus mauritanicus Fauus melkiensis
	Rhinocerotidae	Stephanorhinus
Proboscidea	Elephantidae	Elephas iolensis <sup>b</sup>
Tubulidentata	Orycteropodidae	Orycteropus crassidens

<sup>a</sup> Genus represented in Africa by domesticates only.

<sup>b</sup> Genus no longer represented in Africa, but survives elsewhere.

series (U-series or U/Th), Electron Spin Resonance (ESR), or luminescence (TL/OSL) techniques. Radiocarbon ages are calibrated ( $2\sigma$ ) using Calib (Stuiver and Reimer, 1993) and the INTCAL09 calibration curve (Reimer et al., 2009). Age ranges discussed here (in calendar years), often derived from multiple sites and dating techniques (Table 2), are rounded to the nearest 1000 yrs and given the suffix "ka." A summary of the extinct taxa is provided below.

## 3.1. Order Artiodactyla

#### 3.1.1. Aepyceros sp.

The evolutionary history of impalas (Aepyceros spp.) was once characterized as a single, gradually changing lineage leading to the modern species (Aepyceros melampus) (e.g., Gentry, 1978). However, recent research has documented the presence of several extinct impalas in the Pliocene (Geraads et al., 2009; Gentry, 2011; Geraads, 2012) and Pleistocene (Brink et al., 2012; Faith et al., in press). These include a large-bodied and markedly hypsodont impala recovered from late Pleistocene (100 to 33 ka) deposits along the margins of Lake Victoria in Kenya (Faith et al., in press). Due to the lack of a suitable type specimen, this extinct impala remains to be formally named. Its exceptionally deep mandibles and hypsodont teeth suggest a greater emphasis on grazing in open and dry environments compared to A. melampus. The adaptive implications of elevated hypsodonty is consistent with the associated faunal communities, which indicate the dominance of open and seasonally arid grasslands (Tryon et al., 2010, 2012), although provisional isotopic evidence indicates that the extinct impala consumed some dicots (Garret et al., 2010).

#### 3.1.2. Antidorcas australis

First recovered from middle Pleistocene deposits at Swartklip along the southern coast of South Africa, the southern springbok (Antidorcas australis) was originally described as a subspecies of the modern springbok (as A. marsupialis australis) (Hendey and Hendey, 1968). Vrba (1973) subsequently elevated it to specific status, noting that its presence at Swartkrans suggests a wide temporal and geographic distribution worthy of a valid species. She later expressed doubt concerning her attribution of the Swartkrans fossils to A. australis (Vrba, 1976), suggesting that the material may instead represent a pre-marsupialis variant. Although the taxonomic status of the Swartkrans material remains uncertain (de Ruiter, 2003; Gentry, 2010a), Hendey (1974) supported the taxonomic validity of A. australis when it became clear that it was found together with Antidorcas recki, the presumed ancestor of modern springbok, in the early-to-middle Pleistocene deposits at Elandsfontein (see also Klein et al., 2007). With definitive records occurring only along the southern and western coasts of South Africa, the southern springbok is documented in the late Pleistocene (~60 to 50 ka) at Die Kelders Cave (Feathers and Bush, 2000; Klein and Cruz-Uribe, 2000; Schwarcz and Rink, 2000) and perhaps also in the very latest Pleistocene deposits at Nelson Bay Cave  $(11,950 \pm 150^{14}$ C yrs BP = 13,430 to 14,140 cal yrs BP) as A. cf. australis (Klein, 1980, 1983). Where it is found, A. australis is associated with open-habitat species including equids and alcelaphine antelopes. Like extant springbok, the southern springbok was probably a mixed feeder, capable of browsing dicots when grasses became unpalatable or scarce (Klein, 1980).

#### 3.1.3. Antidorcas bondi

Cooke and Wells (1951) described *Gazella bondi* from Quaternary deposits at Chelmer Farm in Zimbabwe and Vlakkraal in South Africa. In her analysis of antelope remains from Swartkrans, Vrba (1973) later placed the species within *Antidorcas* on the basis of its cranial morphology. Bond's springbok is very well documented in the late Pleistocene to early Holocene (89 to 8 ka) of southern Africa (Klein, 1977; Cruz-Uribe, 1983; Klein, 1984a; Brown and Verhagen, 1985; Klein et al., 1991; Brink and Lee-Thorp, 1992; Plug and Engela, 1992; Plug, 1993, 1997; de Ruiter et al., 2008). Its exceptionally hypsodont teeth,

together with isotopic evidence, suggest that this small springbok was a specialized grazer (Vrba, 1973; Brink and Lee-Thorp, 1992). Gentry (2010a) notes that in near-mature individuals, the extreme hypsodonty results in incomplete ossification along the inferior border of the mandible's horizontal ramus. In the stratified sequence from Border Cave in South Africa, its abundance tracks that of other open grassland species (Klein, 1977).

## 3.1.4. Bos primigenius

The aurochs (Bos primigenius) is the extinct progenitor of domestic cattle. Although it is typically considered a Eurasian taxon (Martinez-Navarro et al., 2007), aurochs fossils are well known from numerous late Pleistocene and early Holocene (115 to 6 ka) sites across North Africa (Churcher, 1972; Churcher and Smith, 1972; Churcher, 1974, 1999; Gautier, 1976; Wendorf and Schild, 1976; Klein and Scott, 1986; Marks et al., 1987; Michel, 1992; Peters, 1992; Vermeersch, 2000; Wengler et al., 2002; Aouraghe, 2004; Nespoulet et al., 2008; Michel et al., 2009; Monchot and Aouraghe, 2009; Bougariane et al., 2010; Dibble et al., 2012). North African aurochs typically co-occurs with several species of gazelle (Gazella spp.) and equids (Equus spp.). Genetic evidence suggests that North African aurochs underwent indigenous African domestication near the onset of the Holocene (Bradley et al., 1996; Hanotte et al., 2002). Although this has prompted some to treat the disappearance of aurochs as a pseudo-extinction (Turvey, 2009), African aurochs persisted well into the Holocene (Table 2), meaning that the domestication process did not involve all populations of the species. Those that were not domesticated ultimately disappeared ~6 ka.

## 3.1.5. Damaliscus hypsodon

Fossils of a small alcelaphine (wildebeest and allies) now known as Damaliscus hypsodon were initially reported by Marean (1990) from late Pleistocene (>11,950  $\pm$  460<sup>-14</sup>C yrs BP = 12,776 to 15,268 cal yrs BP) archeological contexts at Lukenya Hill in southcentral Kenya. It was subsequently reported from the late Pleistocene at Kisese II Rockshelter (>10,720  $\pm$  132  $^{14}$ C yrs BP = 12,375 to 12,946 cal yrs BP) and Gol Kopjes (undated) in northern Tanzania (Marean and Gifford-Gonzalez, 1991) and from late Pleistocene sites (100 to 33 ka) in Kenya's Lake Victoria Basin (Tryon et al., 2010, 2012). This small alcelaphine remained unnamed until recently, when middle Pleistocene specimens from Lainyamok (Kenya) that were previously attributed to Damaliscus cf. dorcas (Potts and Deino, 1995) were formally described as *D. hypsodon* (Faith et al., 2012). The dietary and locomotor adaptations of *D. hypsodon*, together with its association with oryx (Oryx beisa) and Grevy's zebra (Equus grevyi) (Faith et al., 2013), suggest that it grazed in open and arid grasslands. As suggested by its name, and like A. bondi, its teeth are characterized by greater hypsodonty than any extant African bovid, overlapping with some species of Equus.

## 3.1.6. Damaliscus niro

Hopwood (1936) designated a horn core collected by L.S.B. Leakey from Bed IV Olduvai Gorge as the type specimen of *Hippotragus niro*, a species later placed within *Damaliscus* by Gentry (1965). There are numerous records of *D. niro* from the early-to-middle Pleistocene of East and southern Africa (Cooke, 1974; Gentry and Gentry, 1978; Brink, 1987; Vrba, 1997; Suwa et al., 2003; Geraads et al., 2004; Thackeray and Brink, 2004). A late Pleistocene record of *D. niro* was recently reported by de Ruiter et al. (2008) from Plovers Lake in South Africa. The Plovers Lake specimen came from deposits that were disturbed by mining activity. However, the vast majority of *in situ* fossils from this site are from a deposit bracketed by flowstones and dated by U-Series and ESR to between 89 and 63 ka. There are some fossil remains from beneath the primary fossil deposit, but these are also dated to the late Pleistocene (92 ka). Thus, the Plovers Lake *D. niro* most likely dates to between 89 and 63 ka, or perhaps slightly older. Isotopic evidence from middle Pleistocene specimens in South Africa indicates a diet dominated by  $C_4$  grasses (Codron et al., 2008).

### 3.1.7. Gazella atlantica

The Atlantic gazelle (*Gazella atlantica*) was described by Bourguignat (1870) from late Quaternary cave deposits at Jebel Thaya in Algeria. This North African species is common in late Pleistocene (100 to 37 ka) archeological and paleontological contexts across Algeria and Morocco (Aouraghe, 2004; Raynal et al., 2008; Michel et al., 2009; Bougariane et al., 2010; Geraads, 2012; Steele, 2012). Klein (1984b) suggests that *G. atlantica* may have persisted into the middle Holocene, although this remains to be substantiated (Turvey, 2009). In the stratified sequence from El Harhoura 2 in Morocco, the occurrences of *G. atlantica* coincide with those of wildebeest (*Connochaetes taurinus*) and hartebeest (*Alcelaphus buselaphus*) (Michel et al., 2009), suggesting a preference for grasslands.

#### 3.1.8. Gazella tingitana

In his survey of North African Pleistocene gazelles, Arambourg (1957) described *Gazella tingitana* from Aterian levels at Mugharet el 'Aliya in Morocco, now dated to between 85 and 37 ka (Wrinn and Rink, 2003). Amani and Geraads (1993) subsequently documented this poorly known gazelle from middle-to-late Pleistocene (190 to 90 ka) deposits at Jebel Irhoud (Grün and Stringer, 1991; Smith et al., 2007). There is little paleoecological information available for this taxon, although like other gazelles it may have been a mixed feeder that preferred open habitats.

#### 3.1.9. Hippotragus leucophaeus

The first large African mammal to become extinct in historic times, the blue antelope (Hippotragus leucophaeus) was endemic to southernmost South Africa and disappeared ~1800 AD (Klein, 1974a; Kerley et al., 2009; Skead, 2011). Although its historic range was limited to some 4300 km<sup>2</sup> to the east of Cape Town (Kerley et al., 2009), late Pleistocene fossil evidence and rock art suggests a much broader distribution across southern Africa (Klein, 1974a; Loubser et al., 1990; Faith and Thompson, 2013). The blue antelope is commonly found in late Pleistocene assemblages from the southern and western Cape of South Africa, and it is typically more numerous in assemblages dominated by open grassland species, including equids and alcelaphine bovids (Klein, 1983). Evidence from mortality profiles indicates that blue antelope preferentially calved in the western margin of its range and migrated east-west across the southern coast, presumably to track seasonal rains (Faith and Thompson, 2013). A combination of long-term and more recent mechanisms have been invoked to account for its extinction, including climate-driven vegetation change and disruption of migration routes (Faith and Thompson, 2013), competition with domestic livestock within the past 2000 yrs (Klein, 1974a), overhunting by European colonists (Klein, 1974a; IUCN SSC Antelope Specialist Group, 2008b; but see Kerley et al., 2009), and habitat loss and fragmentation due to agricultural expansion during the colonial era (Faith and Thompson, 2013).

#### 3.1.10. Megalotragus priscus

The giant wildebeest (*Megalotragus priscus*) was described by Broom (1909a) as *Bubalis priscus* on the basis of a cranial fragment from late Quaternary deposits along the Modder River in central South Africa. The taxon has many synonyms (e.g., *Bubalis helmei, Pelorocerus elegans, Connochaetes grandis*), although most authorities attribute all giant alcelaphine remains from the late Pleistocene of southern Africa to *M. priscus* (Gentry, 1978; Gentry and Gentry, 1978; Klein, 1980; Brink et al., 1995; Brink, 2005; Gentry, 2010a). The molars of *Megalotragus* are hypsodont (Gentry and Gentry, 1978) and isotopic data indicate a diet of C<sub>4</sub> grasses (Lee-Thorp and Beaumont, 1995; Codron et al., 2008). Together with its massive body size, which exceeds extant alcelaphine antelopes, this suggests a diet involving the unselective

## Table 2

Late Pleistocene and Holocene radiometric dates associated with extinct African mammals (excluding blue antelope). ALG = Algeria, EGY = Egypt, KEN = Kenya, LYB = Lybia, MOR = Morocco, SA = South Africa, SUD = Sudan, TAN = Tanzania, TUN = Tunisia, ZIM = Zimbabwe.

Taxon	Site	Date	Reference
Aepyceros sp.	Rusinga Island KEN	$>$ 28,670 $\pm$ 600 $^{14}$ C yrs BP to 100 ka	Faith et al. (in press)
Antidorcas australis	Die Kelders Cave SA	50.7 $\pm$ 4.7 to 79.7 $\pm$ 15.6 ka (TL/OSL)	Feathers and Bush (2000) and Klein and Cruz-Uribe (2000)
	Nelson Bay Cave SA	$11,950 \pm 150$ to $18,660 \pm 100^{14}$ C yrs BP	Klein (1983) and Deacon (1984)
Antidorcas bondi	Border Cave SA	$37,700 \pm 590^{14}$ C yrs BP to $60 \pm 3$ ka (ESR)	Klein (1977) and d'Errico et al. (2012)
	Equus Cave SA	$9830 \pm 120$ to 27,230 $\pm 340^{-14}$ C yrs BP	Klein et al. (1991) and Lee-Thorp and Beaumont (1995)
	Heuningneskrans SA	$19,980 \pm 260$ <sup>14</sup> C yr BP	Klein (1984a)
	Rruger Cave SA	$7570 \pm 60^{-1}$ C yrs BP	Brown and Vernagen (1985) de Ruiter et al. (2008)
	PIOVEIS Lake SA Rose Cottage Cave SA	$02.9 \pm 1.5 \text{ to } 86.7 \pm 1.0 \text{ Kd} \text{ (U)}$ $9250 \pm 70^{-14} \text{C yr BP}$	Wedley (1991) and Plug and Engels (1992)
	Redcliff Cave ZIM	$25650 \pm 100$ Cyrbi 25650 + 1800 <sup>14</sup> C yrs BP	Cruz-Uribe (1983)
Bos primigenius	Contrebandiers Cave MOR	96 + 4  to  115 + 3  (OSL)	Dibble et al. (2012)
I B	El-Kilh E71P1 EGY	$16,950 \pm 300$ to $17,800 \pm 300^{14}$ C yrs BP	Wendorf and Schild (1976)
	El-Kilh E71P5 EGY	$11,560 \pm 80^{14}$ C yrs BP	Wendorf and Schild (1976)
	Fayum E29H1 EGY	$8070 \pm 115$ <sup>14</sup> C yrs BP	Wendorf and Schild (1976)
	Isna E71K1 EGY	$18,020 \pm 300^{-14}$ C yrs BP	Wendorf and Schild (1976)
	Isna E71K3 EGY	$17,590 \pm 300^{-14}$ C yrs BP	Wendorf and Schild (1976)
	Isna E/IK9 EGY	$9330 \pm 160$ to 16,830 $\pm 290^{14}$ C yrs BP	Wendorf and Schild (1976)
	ISRA E / IK 14 EGY	$12,690 \pm 240$ °C yrs BP 12,660 + 280 to 12,220 + 170 $^{14}$ C yrs BP	Vermoorsch (2000)
	Makhadma 4 FCV	$12,000 \pm 200$ to $13,330 \pm 170^{-14}$ C yis BP 12 320 + 70 to 12 940 + 130 <sup>-14</sup> C yrs BP	Vermeersch (2000)
	Shuwikhat 1 FGY	$12,520 \pm 70.00$ $12,540 \pm 150$ C yrs BP	Vermeersch (2000)
	Haua Fteah LYB	7300 + 30 to $47.000 + 1500$ <sup>14</sup> C vrs BP	Klein and Scott (1986)
	El Harhoura 1 MOR	$5400 \pm 290^{14}$ C yr BP to >66.5 $\pm$ 5.2 ka (U)	Monchot and Aouraghe (2009) and Raynal and Occhietti (2012)
	El Harhoura 2 MOR	$<\!51.6\pm3.6$ to 102.6 $\pm$ 5.7 (OSL)	Michel et al. (2009) and Jacobs et al. (2012)
	El Mnasra MOR	$71\pm5.3$ to $108.8\pm6.6$ (OSL)	Nespoulet et al. (2008) and Jacobs et al. (2012)
	Rhafas Cave MOR	70 (TL) to $107 \pm 12$ ka (OSL)	Michel (1992) and Mercier et al. (2007)
	Wadi Assaka MOR	$>30,200 \pm 2900^{-14}$ yrs BP	Wengler et al. (2002)
	Khashm el Girba KG15 SUD	$10,230 \pm 270^{14}$ C yr BP	Marks et al. (1987)
Damaliscus nypsoaon	Lukenya Hill GvJm19 KEN	$13,705 \pm 430^{+1}$ C yrs BP 12,720 + 420 to 17,700 + 760 <sup>14</sup> C yrs PD	Marcan (1992)
	Lukenya Hill Gyjni22 KEN	$13,730 \pm 430$ to $17,700 \pm 700$ C yis BP 19 330 $\pm$ 945 to 20 780 $\pm$ 1050 <sup>14</sup> C yrs BP	Marean (1992)
	Lukenya Hill GvJm40 KEN	$11,950 \pm 460$ to $21,535 \pm 980$ <sup>14</sup> C vrs BP	Marean (1992)
	Mfangano Island KEN	$>30,530 \pm 390^{14}$ C yrs BP	Tryon et al. (2012)
	Rusinga Island KEN	$>$ 28,670 $\pm$ 600 $^{14}$ C yrs BP to 100 ka	Tryon et al. (2010, 2012)
	Kisese II TAN	$>10,720 \pm 132$ <sup>14</sup> C yrs BP	Marean and Gifford-Gonzalez (1991)
Damaliscus niro	Plovers Lake SA	$62.9\pm1.3$ to $88.7\pm1.6$ ka (U)	de Ruiter et al. (2008)
Gazella atlantica	El Harhoura 1 MOR	$>66.5 \pm 5.2$ ka (U)	Monchot and Aouraghe (2009) and Raynal and Occhietti (2012)
	El Harhoura 2 MOR	$<51.6 \pm 3.6$ to $99.9 \pm 5.8$ ka (OSL)	Michel et al. (2009) and Jacobs et al. (2012)
Cazella tingitana	Mugharet el 'Aliya MOR	$37 \pm 4$ to $85 \pm 9$ ka (ESR)	Steele (2012) and Wrinn and Rink (2003) Steele (2012) and Wrinn and Rink (2002)
Megalotragus priscus	Boomplaas Cave SA	$37 \pm 4000 \pm 9 \text{ Ka} (ESK)$ 21 100 + 180 <sup>14</sup> C yr BP	Klein (1983) and Deacon (1984)
meguioriugus priseus	Equus Cave SA	$9830 + 120 \text{ to } 27.230 + 340^{14} \text{C yrs BP}$	Klein et al. (1991) and Lee-Thorp and Beaumont (1995)
	Nelson Bay Cave SA	$16,700 \pm 240^{-14}$ C yrs BP	Klein (1983) and Deacon (1984)
	Plovers Lake SA	$62.9 \pm 1.3$ to $88.7 \pm 1.6$ ka (U)	de Ruiter et al. (2008)
	Rose Cottage Cave SA	12,690 $\pm$ 120 to 13,360 $\pm$ 150 $^{14}$ C yrs BP	Wadley (1991) and Plug and Engela (1992)
	Sibudu Cave SA	$61.7 \pm 2$ to $64.7 \pm 2.3$ ka (OSL)	Clark (2011)
	Wonderwerk Cave SA	$7430 \pm 60^{14}$ C yrs BP	Thackeray et al. (1981)
Megalotragus	Rusinga Island KEN	$> 28,670 \pm 600$ ° C yrs BP to 100 ka	Iryon et al. $(2010)$ and Faith et al. $(2011)$
κασιησοι γχ ατοροει απιοπ	Rusinga Island KEN	$>30,330 \pm 390$ $>28,670 \pm 600^{14}$ C yrs BP to 100 ka	Faith et al. $(2012)$ Faith et al. $(2011)$ and Tryon et al. $(2012)$
Syncerus antiauus	Amekni ALG	5500 + 250 to $8670 + 150$ <sup>14</sup> C vrs BP	Gautier (1987) and Gautier and Muzzolini (1991)
	Medjez II ALG	$8550 \pm 150$ <sup>14</sup> C yrs BP	Gautier and Muzzolini (1991)
	Meniet ALG	$5400 \pm 150 {}^{14}\text{C}\text{yrs}\text{BP}$	Gautier (1987) and Gautier and Muzzolini (1991)
	Lukenya Hill GvJm22 KEN	13,730 $\pm$ 430 to 17,700 $\pm$ 760 $^{14}$ C yrs BP	Marean (1992)
	Mfangano Island KEN	$>30,530 \pm 390$ to 41,846 $\pm 458$ <sup>14</sup> C yrs BP	Tryon et al. (2012)
	Rusinga Island KEN	$>28,670 \pm 600$ <sup>14</sup> C yrs BP to 100 ka	Tryon et al. (2010, 2012) Man de tan d'Assessible (2000) and Basessible d'Oschistri (2012)
	El Harhoura I MOR	$>$ bb.5 $\pm$ 5.2 Ka (U) 51.6 $\pm$ 3.6 to 106.7 $\pm$ 6.6 kg (OSL)	Monchot and Aouragne (2009) and Raynal and Occhietti (2012)
	El Hallioula 2 MOR Mugharet el 'Aliva MOR	$31.0 \pm 3.0$ to $100.7 \pm 0.0$ Ka (OSL) $37 \pm 4$ to $85 \pm 9$ ka (FSR)	Steele (2012) and Wrinn and Rink (2003)
	Boomplaas Cave SA	$12.060 \pm 105 \text{ to } 21.110 \pm 180^{14} \text{C vrs BP}$	Klein (1983) and Deacon (1984)
	Die Kelders Cave SA	$50.7 \pm 4.7$ to $79.7 \pm 15.6$ ka (TL/OSL)	Feathers and Bush (2000) and Klein and Cruz-Uribe (2000)
	Nelson Bay Cave SA	$11,950 \pm 150$ to $18,660 \pm 100^{14}$ C yrs BP	Klein (1983) and Deacon (1984)
	Klasies River Mouth SA	$43.4 \pm 3.0$ to $106.8 \pm 12.6$ ka (OSL)	Klein (1976) and Feathers (2002)
	Sibudu Cave SA	57.6 $\pm$ 2.1 to 59.6 $\pm$ 2.3 ka (OSL)	Clark (2011)
Unnamed caprine	Boomplaas Cave SA	$9100 \pm 135$ <sup>14</sup> C yr BP to $66 \pm 7$ ka	Faith (2012) and Vogel (2001)
Comoluo	Colwinton Shelter SA	$62/0 \pm 40$ <sup>12</sup> C yr BP	Brink (1999) Courtier (1992) and Stalies (1992)
Cameius	BIF TATTAWI 14 EGY	$1.4 \pm 8.2/- 1.1$ to $129.2 \pm 7.7$ ka (USL)	Gautier (1993) and Stokes (1993)
	oued el Alzrit TLIN	$> 30,200 \pm 2300$ C yis Dr 19 900 + 750 to > 34 000 $^{14}$ C yrs PD	Run et al. (2002) Run et al. (1988)
Megaceroides algericus	Tamar Hat AIG	$17.040 + 400^{14}$ C vr BP	Merzoug and Sari (2008)
Metridiochoerus	Redcliff Cave ZIM	$25,650 \pm 1800^{-14}$ C vrs BP	Cruz-Uribe (1983)
Kolpochoerus	Rusinga Island KEN	$>$ 28,670 $\pm$ 600 $^{14}$ C yrs BP to 100 ka	Tryon et al. (2010); this study
Equus algericus	El Harhoura 1 MOR	5400 $\pm$ 290 $^{14}\text{C}$ yrs BP to >66.5 $\pm$ 5.2 (U)	Monchot and Aouraghe (2009) and Raynal and Occhietti (2012)
Equus capensis	Apollo 11 NAM	12,510 $\pm$ 90 to 14,550 $\pm$ 90 $^{14}$ C yrs BP	Thackeray (1979)
	Equus Cave SA	9830 $\pm$ 120 to 27,230 $\pm$ 340 $^{14}$ C yrs BP	Klein et al. (1991) and Lee-Thorp and Beaumont (1995)

Table 2 (continued)

Taxon	Site	Date	Reference
	Boomplaas Cave SA	12,060 $\pm$ 105 to 17,830 $\pm$ 180 $^{14}\mathrm{C}\mathrm{yr}\mathrm{BP}$	Klein (1983) and Faith (2012)
	Border Cave SA	$47 \pm 2$ ka (ESR)	Klein (1977) and d'Errico et al. (2012)
	Bushman Rock Shelter SA	>47,000 <sup>14</sup> C yr BP	Badenhorst and Plug (2012)
	Byneskranskop 1 SA	$12,730 \pm 185 {}^{14}$ C yr BP	Schweitzer and Wilson (1982)
	Die Kelders Cave 1 SA	50.7 $\pm$ 4.7 to 79.7 $\pm$ 15.6 (TL/OSL) ka	Feathers and Bush (2000) and Klein and Cruz-Uribe (2000)
	Elands Bay Cave SA	9600 $\pm$ 90 to 12,450 $\pm$ 280 $^{14}$ C yrs BP	Klein and Cruz-Uribe (1987)
	Rose Cottage Cave SA	12,690 $\pm$ 120 to 13,360 $\pm$ 150 $^{14}$ C yrs BP	Wadley (1991) and Plug and Engela (1992)
	Wonderwerk Cave SA	$10,000 \pm 70^{14}$ C yrs BP	Thackeray et al. (1981)
	Redcliff Cave ZIM	$25,650 \pm 1800$ <sup>14</sup> C yrs BP	Cruz-Uribe (1983)
Equus mauritanicus	Contrebandiers Cave MOR	$96 \pm 4$ to $115 \pm 3$ (OSL)	Dibble et al. (2012)
	El Harhoura 1 MOR	5400 $\pm$ 290 $^{14}$ C yrs BP to >66.5 $\pm$ 5.2 ka (U)	Monchot and Aouraghe (2009) and Raynal and Occhietti (2012)
	Rhafas Cave MOR	5400 $\pm$ 100 $^{14}$ C yrs BP to <107 $\pm$ 12 (OSL)	Michel (1992) and Mercier et al. (2007)
	Oued el Akarit TUN	19,900 $\pm$ 750 to >34,000 <sup>14</sup> C yrs BP	Brun et al. (1988)
Equus melkiensis	Mugharet el 'Aliya MOR	$37 \pm 4$ to $85 \pm 9$ ka (ESR)	Eisenmann (2006) and Wrinn and Rink (2003)
Stephanorhinus	Contrebandiers Cave MOR	$107 \pm 9$ (OSL)	Dibble et al. (2012)
	Haua Fteah LYB	16,070 $\pm$ 100 to 47,000 $\pm$ 1500 $^{14}$ C yrs BP	Klein and Scott (1986)
	El Harhoura 2 MOR	${<}52.6\pm3.3$ to $61.9\pm3.5$ (OSL)	Michel et al. (2009) and Jacobs et al. (2012)
	El Mnasra MOR	$108.8\pm6.6$ ka (OSL)	Nespoulet et al. (2008) and Jacobs et al. (2012)
Elephas iolensis	Natodameri SUD	35 ka (U)	Coppens et al. (1978)
Orycteropus crassidens	Rusinga Island KEN	$>$ 28,670 $\pm$ 600 $^{14}$ C yrs BP to 100 ka	Tryon et al. (2010, 2012)

bulk grazing of low quality forage (Bell, 1971; Jarman, 1974; Marean, 1992). Relative to extant alcelaphines, the metapodials of Megalotragus are long compared to its upper limbs (Gentry and Gentry, 1978), suggesting a preference for open habitats (Scott, 1985; Plummer and Bishop, 1994). Megalotragus priscus is documented in numerous late Pleistocene archeological and paleontological localities (89 to 12 ka) across southern Africa (Klein, 1980; Cruz-Uribe, 1983; Klein, 1983, 1984a; Klein et al., 1991; Plug and Engela, 1992; Brink et al., 1995; Robbins et al., 2000; Clark and Plug, 2008; de Ruiter et al., 2008), with a single early Holocene record dating to 7430  $\pm$  60  $^{14}$ C yrs BP (8160 to 8392 cal yrs BP) at Wonderwerk Cave (Thackeray et al., 1981; Thackeray, 1983). In addition to the southern African occurrences, Megalotragus remains have also been recovered from late Pleistocene deposits (100 to 33 ka) in Kenya's Lake Victoria Basin (Tryon et al., 2010; Faith et al., 2011; Tryon et al., 2012). It is unclear whether these represent a late form of Megalotragus kattwinkeli, known from earlier in the East African Pleistocene, M. priscus, or a new species.

## 3.1.11. Rusingoryx atopocranion

Pickford and Thomas (1984) described a new genus of alcelaphine bovid on the basis of a partial cranium recovered from the Wasiriya Beds of Rusinga Island, Kenya. Rusingoryx atopocranion was later subsumed into Megalotragus by Harris (1991), a decision supported by Vrba's (1997) phylogenetic analysis of fossil alcelaphines. However, renewed fieldwork on Rusinga Island (Tryon et al., 2010) recovered a large sample of Rusingoryx remains that provided morphological and phylogenetic evidence supporting its distinction from Megalotragus, which is also present in the fossil collection (Faith et al., 2011). The Rusinga specimens are now dated to the late Pleistocene (100 to 33 ka) and additional remains have been recovered from roughly contemporaneous deposits (>35 ka) on nearby Mfangano Island (Tryon et al., 2012). Dental mesowear analysis, isotopic analysis, and its extreme hypsodonty indicate that *R. atopocranion* was a grazer, while its association with Grevy's zebra (E. grevyi) and oryx (O. beisa) suggests a preference for dry grassland or scrub habitats (Faith et al., 2011).

## 3.1.12. Syncerus antiquus

The giant buffalo (*Syncerus antiquus*) holds the distinction of being the largest African bovid and the first fossil bovid described on the continent. Believing it to be closely related to the Asian water buffalo (*Bubalus bubalis*), Duvernoy (1851) founded the taxon as *Bubalus antiquus* on the basis of a cranial fragment recovered from Quaternary deposits in Algeria. *Bubalus* was subsequently used for several species of giant buffalo, including *Bubalus bainii* from South Africa (Seeley, 1891) and *Bubalus nilssoni* from Kenya (Lönnberg, 1933). Bate (1951) recognized that these buffaloes were conspecific and not related to Bubalus, placing them within Homoioceras (as Homoioceras antiquus), which was established for a specimen from Sudan (Homoioceras singae) (Bate, 1949). Although *H. antiquus* is still in use by some (Brink, 2005; Codron et al., 2008), the type specimen of *H. singae* appears to be a primitive form of extant African buffalo (Syncerus caffer) (Gentry, 1978; Gentry and Gentry, 1978), rendering Homoioceras unavailable as the genus for antiquus. Gentry and Gentry (1978) assigned antiquus to Pelorovis, as do many today (Klein, 1994; Klein et al., 2007; Tryon et al., 2010), although morphological and systematic evidence is leading to a growing consensus that *antiquus* should be placed within *Syncerus* (Gautier and Muzzolini, 1991; Geraads, 1992; Klein, 1994; Peters et al., 1994; Hadjouis, 2002; Martinez-Navarro et al., 2007; Gentry, 2010a). Syncerus antiquus had the broadest geographic distribution of any of the extinct species considered here, occurring in the late Pleistocene (107 to 13 ka) of southern Africa (Klein, 1980, 1983; Cruz-Uribe, 1983; Clark and Plug, 2008; Rector and Reed, 2010; Faith, 2012, 2013b), East Africa (Marean and Gifford-Gonzalez, 1991; Marean, 1992; Tryon et al., 2010, 2012), Central Africa (Peters, 1990), and North Africa (Gautier and Muzzolini, 1991; Aouraghe, 2004; Monchot and Aouraghe, 2009; Steele, 2012). As indicated by fossil remains and rock art, the giant buffalo persisted into the middle Holocene (~6 ka) in North Africa (Gautier and Muzzolini, 1991). Isotopic and mesowear evidence indicate that *S. antiquus* was a grazer (Codron et al., 2008; Stynder, 2009), and its massive body size suggests that it consumed large quantities of low quality forage. The horns of larger individuals extended upwards of 3 m from tip to tip (versus up to 1.4 in S. caffer), limiting S. antiquus to open habitats (Klein, 1994).

#### 3.1.13. Unnamed caprine

Brink (1999) reported an extinct caprine antelope from the late Pleistocene to early Holocene (66 to 7 ka) at several localities in mountainous regions of South Africa. Morphologically similar fossils were subsequently found in middle Pleistocene (830 to 578 ka) deposits at Gladysvale Cave (Lacruz et al., 2002). This extinct caprine's teeth are markedly hypsodont and its post-crania match those of other caprines adapted to rugged habitats. Due to the lack of a suitably complete type specimen, the taxon remains to be formally described. The species may belong within the extinct genus *Makapania* (Brink, 1999; Lacruz et al., 2002), which is typically thought to belong to the tribe ovibovini (muskoxen and allies) (Gentry, 1970, 2010a), although Brink's (1999) analysis suggests numerous caprine affinities. In the stratified sequence from Boomplaas Cave, its disappearance between 9100  $\pm$  135 and 6400  $\pm$  75 <sup>14</sup>C yrs BP (9866 to 10,095 and 7171 to 7433 cal yrs BP) is associated with the replacement of grazing ungulates by browsers (Faith, 2012, 2013b).

## 3.1.14. Camelus

An extinct camel that perhaps belongs within *Camelus thomasi* is documented at a handful of North African late Pleistocene sites ranging in age from >24 ka to 130 ka (Gautier, 1966, 1993; Wendorf, 1977; Brun et al., 1988; Wengler et al., 2002). *Camelus thomasi* was described by Pomel (1893) from the early/middle Pleistocene at Tighenif in Algeria. Late Pleistocene *Camelus* is often assigned to the same taxon (Gautier, 1966, 1993; Wendorf, 1977), although Harris et al. (2010a) caution that these identifications are suspect as they were made without reference to the Tighenif material. Due to a rather sparse record, it is possible that late Pleistocene *Camelus* represents a different species or that multiple species are represented. The relationship of late Pleistocene *Camelus* to the domestic dromedary (*Camelus dromedarius*) is uncertain (Harris et al., 2010a). Late Pleistocene specimens are 20–30% larger than *C. dromedarius* (Gautier, 1966).

## 3.1.15. Megaceroides algericus

Lydekker (1890) founded Cervus algericus on the basis of a maxilla recovered from late Quaternary deposits at Hammam el Meskhoutine in Algeria. The taxonomic position of this and other megalocerine deer is contentious (Lister et al., 2005), and various authorities place algericus within Megaloceros (Hamilton, 1978; Gentry, 2010b) or Megaceroides (Joleaud, 1914; Hadjouis, 1990; Azzaroli and Mazza, 1993; Pfeiffer, 2002; Abbazzi, 2004; Geraads, 2010a). Megaceroides is used here, as the large sample of antlers from Les Phacochères in Algeria shows a number of features that distinguish it from Eurasian Megaloceros, including the lack of a posterior tine (Hadjouis, 1990). Megaceroides algericus is known from various late Quaternary sites across Algeria and Morocco (Arambourg, 1938; Saxon et al., 1974; Hadjouis, 1990; Merzoug and Sari, 2008). In the stratified archeological sequence from Tamar Hat in Algeria, its last appearance is dated to 17,040  $\pm$ 400 <sup>14</sup>C yrs BP (19,411 to 21,306 cal yrs BP) (Saxon et al., 1974; Merzoug and Sari, 2008). Possible artistic renditions of the species raise the possibility that it may have persisted into the Holocene (Arkell, 1962; Saxon et al., 1974), although these have been questioned (Turvey, 2009). The brachydont teeth and shallow horizontal mandibular ramus of *M. algericus* suggest it was a browser (Abbazzi, 2004).

## 3.1.16. Metridiochoerus

An extinct giant warthog (*Metridiochoerus* sp.) is reported from the middle-to-late Pleistocene of southern Africa (Klein, 1980; Cruz-Uribe, 1983; Klein, 1984b). The *Metridiochoerus* lineage thrived earlier in the Pleistocene (van Hoepen and van Hoepen, 1932; Harris and White, 1979; Potts, 1998; Suwa et al., 2003; Geraads et al., 2004; Brink, 2005; Bishop, 2010), and Klein (1980, 1984b) suggests that it persists into the late Pleistocene at four southern African localities (Florisbad, Koffiefontein, Vlakkraal, and Redcliff Cave). However, the Florisbad specimens may belong to extant *Phacochoerus* (Brink, 1987) and are probably middle Pleistocene in age (279 to 121 ka) (Grün et al., 1996). Those from Koffiefontein and Vlakkraal were assigned to the late Pleistocene on the basis of their association with MSA artifacts, although the

oldest MSA is now dated to ~280 ka (Tryon and McBrearty, 2006; Morgan and Renne, 2008). This leaves the *Metridiochoerus* specimens from Redcliff Cave in Zimbabwe, which are associated with a radiocarbon date of  $25,650 \pm 1800$  <sup>14</sup>C yr BP (26,257 to 34,125 cal yrs BP), as the sole late Pleistocene record of the taxon (Klein, 1980; Cruz-Uribe, 1983). The teeth of the Redcliff specimens are extremely hypsodont (Klein, 1980) and isotopic data from earlier Pleistocene *Metridiochoerus* indicate that the giant warthog was a grazer with limited dependence on water (Harris and Cerling, 2002).

#### 3.1.17. Kolpochoerus

The suid genus Kolpochoerus has a long history that extends into the earliest Pliocene (Brunet and White, 2001). The youngest previously documented records of Kolpochoerus (Kolpochoerus majus) were middle Pleistocene (~500 ka) in age (Geraads et al., 2004; McBrearty and Jablonski, 2005; Faith et al., 2012). Recent collections from the late Pleistocene (100 to 33 ka) Wasiriya Beds on Rusinga Island in Kenya (Tryon et al., 2010, 2012) recovered several dental specimens belonging to Kolpochoerus, the most complete of which is illustrated here in Fig. 2. The level of compression of the pillars and cementum most closely matches K. majus, although a secure identification will require the recovery of more complete specimens. The Rusinga Island Kolpochoerus is associated with other ungulates interpreted as reflecting a dry grassland environment (Tryon et al., 2010; Faith et al., 2011; Tryon et al., 2012). Early Pleistocene Kolpochoerus from Lake Turkana was a grazer, but more dependent on water than Metridiochoerus (Harris and Cerling, 2002).

## 3.2. Order Perissodactyla

#### 3.2.1. Equus algericus

Bagtache et al. (1984) described a caballine horse (*Equus algericus*) from late Quaternary deposits of Les Phacochères (AKA Allobroges) in Algeria. *Equus algericus* has since been reported from the Neolithic and Aterian deposits at El Harhoura 1 in Morocco (Aouraghe and Debénath, 1999; Aouraghe, 2004; Monchot and Aouraghe, 2009). Its occurrences at this site range in age from the middle Holocene (5400  $\pm$  290<sup>14</sup>C yrs BP = 6683 to 6805 cal yrs BP) to >66.5  $\pm$  5.2 ka (Raynal and Occhietti, 2012). Little is known about its paleoecology, although like most large species of *Equus* it was probably a bulk grazer that preferred open habitats.

## 3.2.2. Equus capensis

The Cape zebra (*Equus capensis*) is the largest equid known from the African Quaternary, standing 150 cm tall at the withers and with a body mass exceeding 400 kg (Eisenmann, 2000, 2003). *Equus capensis* was founded by Broom (1909b) on the basis of a right mandible recovered from near Cape Town in South Africa. More than a dozen large equids were later described from South African Pleistocene sites in the early 1900s, all of which are now synonymized within *E. capensis* (Churcher and Richardson, 1978). *Equus capensis* is documented at many late



Fig. 2. Distal fragment of Kolpochoerus right mandibular M<sub>3</sub> from Rusinga Island (KMN-RU 49738) in (A) lingual, (B) occlusal, and (C) labial views.

Pleistocene sites (90 to 11 ka) across southern Africa, where it disappeared near the Pleistocene-Holocene transition (Klein, 1977; Thackeray, 1979; Klein, 1980; Thackeray et al., 1981; Schweitzer and Wilson, 1982; Cruz-Uribe, 1983; Klein, 1983; Klein and Cruz-Uribe, 1987; Klein et al., 1991; Plug and Engela, 1992; Klein and Cruz-Uribe, 2000; Robbins et al., 2000; Badenhorst and Plug, 2012; Faith, 2012, 2013b). Churcher (Churcher et al., 1999; Churcher, 2006) reports several E. capensis specimens spanning the middle Pleistocene and Holocene of Egypt. However, their taxonomic status is equivocal. Believing E. capensis to be the direct ancestor of E. grevyi (Churcher and Richardson, 1978; Churcher, 2006), Churcher treats the distinction between the two species loosely, using E. capensis for fossil remains and E. grevyi for modern animals. Ancient DNA evidence now shows that E. capensis is more closely related to extant plains zebra (Equus quagga – formerly Equus burchelli) than to E. grevyi (Orlando et al., 2009). Thus, the Egyptian specimens may belong to extant Grevy's zebra or to an extinct North African taxon, perhaps Equus mauritanicus. Isotopic data show that E. capensis was a grazer (Lee-Thorp and Beaumont, 1995; Codron et al., 2008) and Klein (1980) observes that it is more numerous in fossil localities from what are today more arid regions of southern Africa.

#### 3.2.3. Equus mauritanicus

The extinct zebra Equus mauritanicus is a poorly defined North African taxon created by Pomel (1897) from late Quaternary material in Algeria. Churcher and Richardson (1978) regarded it as a subspecies of the extant plains zebra (E. quagga – formerly E. burchelli). However, Eisenmann (1980) and Eisenmann and Baylac (2000) examined several skulls of E. mauritanicus and found them to be morphologically distinct from plains zebra, whereas its massive teeth overlap in size with southern African E. capensis (Eisenmann, 2003). Bernor et al. (2010) agree that E. mauritanicus is not E. quagga, but they caution that the taxon needs a more clear definition and further study. Although its taxonomic status is uncertain, there are no extant African zebras that it could reasonably belong to and E. mauritanicus is used here for the extinct taxon that is routinely reported from late Pleistocene to Holocene (115 to 6 ka) localities in North Africa (Higgs, 1967; Brun et al., 1988; Michel, 1992; Aouraghe and Debénath, 1999; Monchot and Aouraghe, 2009; Dibble et al., 2012; Steele, 2012).

## 3.2.4. Equus melkiensis

Bagtache et al. (1984) described an extinct equid (*Equus melkiensis*) probably related to the wild ass (*Equus africanus*) from Les Phacochères in Algeria, also the type site of *E. algericus*. Eisenmann (2006) reports *E. melkiensis* from several presumably late Pleistocene sites in Algeria and Morocco, although the record from Mugharet el 'Aliya is the only one that can be securely assigned to the late Pleistocene (85 to 37 ka) (Wrinn and Rink, 2003). Like many of the extinct North African taxa, there is little known about its paleoecology. Extant wild asses inhabit arid environments and are able to subsist on very poor quality forage (Bauer et al., 1994).

#### 3.2.5. Stephanorhinus

The extinct rhinoceros *Stephanorhinus* is a Eurasian taxon that migrated into North Africa near the end of the middle Pleistocene (Amani and Geraads, 1993; Geraads, 2012). The phylogeny and systematics of Pleistocene African rhinoceroses are problematic and have not kept pace with work in Eurasia (Fortelius et al., 1993; Heissig, 1999; Billia, 2008; Geraads, 2010b). As a result, *Dicerorhinus* is sometimes used for *Stephanorhinus*, although the former now has a more restricted definition (Fortelius et al., 1993). At least one, but perhaps two species of *Stephanorhinus* disappeared from North Africa during the late Pleistocene. The narrow-nosed rhinoceros (*Stephanorhinus hemitoechus*) is reported from a handful of late Pleistocene (109 to 53 ka) localities in North Africa (Nespoulet et al., 2008; Raynal et al., 2008; Michel et al., 2009; Jacobs et al., 2012). *Stephanorhinus kirchbergensis* is also reported

from the late Pleistocene (>40 to 19 ka) at the Huah Fteah in Libya (Klein and Scott, 1986). The two species have often been confused in the past (Billia, 2008) and Geraads (2010b) suggests that only *S. hemitoechus* is present in the North African late Pleistocene.

## 3.3. Order Proboscidea

## 3.3.1. Elephas iolensis

The terminal representative of one of the more long-lived and widespread African elephant lineages, *Elephas iolensis* was founded by Pomel (1895) on the basis of a lower molar from Beausejour Farm in Algeria. The species is known from middle/late Pleistocene sites across Africa (Maglio, 1973; Coppens et al., 1978; Abbate et al., 2010; Sanders et al., 2010). A specimen of *E. iolensis* from Natodameri in South Sudan has been directly dated by U–Th to 35 ka and provides the latest record of the taxon (Coppens et al., 1978). The molars of *E. iolensis* are more hypsodont than any other African elephant (Maglio, 1973; Coppens et al., 1978), suggesting a specialized grazing adaptation.

## 3.4. Order Tubulidentata

#### 3.4.1. Orycteropus crassidens

MacInness (1956) described a large extinct aardvark (*Orycteropus crassidens*) from the late Pleistocene Wasiriya Beds (100 to 33 ka) on Rusinga Island in Kenya (Tryon et al., 2012). The taxon is also known from the early-to-middle Pleistocene at nearby Kanjera (Behrensmeyer et al., 1995; Lehman, 2009). Pickford (1975) regarded *O. crassidens* as a subspecies of extant *Orycteropus afer*, although Lehman (2009) summarizes morphological and phylogenetic evidence supporting the validity of the species.

#### 3.5. Problematic species

There are a handful of taxa absent from Table 1 that are often cited as having become extinct during the late Pleistocene or Holocene (Martin, 1984; MacPhee and Flemming, 1999; Barnosky et al., 2004; Lyons et al., 2004; Koch and Barnosky, 2006; Turvey, 2009), but whose taxonomic status or last appearances require revision.

## 3.5.1. Hipparion lybicum

A three-toed (hipparionine) horse referred to as *Hipparion lybicum* often appears on lists of extinct late Pleistocene taxa (Barnosky et al., 2004; Lyons et al., 2004; Koch and Barnosky, 2006). This can be traced to Churcher and Richardson (1978), who synonymized over a dozen taxa from across Africa into *H. lybicum* and suggested that it persisted into the late Pleistocene at a number of sites. *Hipparion lybicum* is no longer recognized as a valid taxon (Bernor et al., 2010) and none of the supposed late Pleistocene occurrences can be confidently placed in this time frame. Current evidence indicates that hipparionine horses disappeared from Africa early in the middle Pleistocene (~500 ka) (Geraads et al., 2004; Bernor et al., 2010; Geraads, 2010a). All Pliocene and Pleistocene three-toed horses from Africa are now assigned to the genus *Eurygnathohippus* (Bernor et al., 2010).

## 3.5.2. Giraffa gracilis

Arambourg (1947) described a lightly built species of giraffe (*Giraffa gracilis*) from the Shungura Formation in Ethiopia, although the taxon is now regarded as a junior synonym of *Giraffa stillei* (Harris et al., 2010b). Churcher (1978) describes the age range of this taxon as late Pliocene to late Pleistocene. However, Churcher does not mention any specific late Pleistocene occurrences and none of his tables or figures provide age ranges indicating the presence of *G. gracilis* or *G. stillei* beyond the middle Pleistocene. In the decades since then, there are still no records of an extinct late Pleistocene giraffe (Harris et al., 2010b).

## 3.5.3. Parmularius

*Parmularius* is an extinct genus of alcelaphine antelope most closely related to *Damaliscus*, which includes extant blesbok (*Damaliscus dorcas*) and topi (*Damaliscus lunatus*). Its misplaced status as an extinct late Pleistocene taxon can be traced to Maglio's (1978) list of Cenozoic African mammals, which indicates an age range for *Parmularius* from the early to the late Pleistocene. This conflicts with Gentry's (1978) assessment of *Parmularius* in the same volume and with current evidence indicating the extinction of the genus near the early-to-middle Pleistocene boundary (Vrba, 1997; Geraads et al., 2004; Gentry, 2010a).

#### 3.5.4. Equus quagga

The extinct quagga (*Equus quagga quagga*), noted for the lack of stripes on its rear half, was historically widespread on the South African plains south of the Orange River. It disappeared in the wild in the second half of the 19th century and the last individual died in the Amsterdam Zoo in 1883. From the 1980s until recently there was substantial debate over its taxonomic status and relationship to the more widespread plains zebra (formerly *E. burchelli*) (Bennett, 1980; Thackeray, 1988; Klein and Cruz-Uribe, 1995; Thackeray, 1997; Klein and Cruz-Uribe, 1999; Eisenmann and Brink, 2000; Groves and Bell, 2004). Genetic evidence has resolved this debate by showing that quagga and plains zebra belong to the same species (Leonard et al., 2005), a position also adopted by the IUCN (Hack and Lorenzen, 2008) and pale-ontological authorities (Bernor et al., 2010). The correct name for extant plains zebra is now *E. quagga*, which has priority over *E. burchelli*.

## 3.5.5. Gazella rufina (=Eudorcas rufina)

The red gazelle (*Eudorcas rufina*) was thought to have disappeared from Algeria near the end of the 19th century. The only known specimens (n = 3) were purchased from markets in northern Algeria and the taxon has never been documented in the wild. One of these specimens has since been identified as belonging to extant red-fronted gazelle (*Eudorcas rufifrons*), prompting the IUCN to change its conservation status from extinct to data deficient in 2008 (IUCN SSC Antelope Specialist Group, 2008a).

## 4. Chronology and explanations

At the continental scale, the last appearance dates (LADs) of the extinct African taxa range in age from >63 ka (*D. niro*) to the middle Holocene (6 ka) (*B. primigenius, S. antiquus, E. mauritanicus, E. melkiensis*). There is good reason to believe that this chronology is structured to a large extent by sampling. Fig. 3 plots the number of dated sites against the youngest associated radiometric age for each extinct taxon, excluding the historically extinct blue antelope (data from Table 2). There is a



Fig. 3. The number of dated sites against the youngest associated radiometric date for each extinct taxon, excluding blue antelope (data from Table 2). Dashed line corresponds to the Pleistocene–Holocene transition.

significant negative relationship between the number of dated sites in which a taxon occurs and its youngest radiometric date (Spearman's rho: -0.670, p < 0.001). All of the taxa (n = 6) that have been documented in 7 or more dated sites are associated with fairly recent LADs (~13 to 6 ka), whereas those (n = 17) that occur in fewer dated sites, either because they are rare in the fossil record or because their occurrences are undated, are characterized by a broader range of LADs (63 to 6 ka). As has been argued in the context of North American extinctions (Faith and Surovell, 2009), this suggests that additional sampling of the fossil record, or dating of samples that have not been dated, will lead to the establishment of substantially younger LADs for some taxa. Although the better-sampled taxa suggest that the extinctions may have been restricted to the terminal Pleistocene and early Holocene, the regional perspective indicates that losses occurred at different times in different places.

#### 4.1. Southern Africa

An abundance of stratified late Quaternary archeological and paleontological sites across southern Africa provides the best regional chronology of large mammal extinctions on continental Africa (Table 2). In agreement with previous characterizations (Klein, 1980, 1984b), the last appearances of *E. capensis, S. antiquus*, and likely *A. australis* broadly correspond with the Pleistocene–Holocene transition. However, there is now good evidence from stratified and well-dated cave deposits that *M. priscus, A. bondi*, and the extinct caprine antelope persisted for several thousand years into the Holocene (Thackeray et al., 1981; Thackeray, 1983; Brown and Verhagen, 1985; Plug and Engela, 1992; Brink, 1999; Faith, 2012). The last appearances of *Metridiochoerus* and *D. niro* occur prior to the Last Glacial Maximum, although both are so rare in the late Pleistocene record (1 locality each) that they may have persisted for much longer without being detected (Faith and Surovell, 2009).

The extinct southern African species share a number of ecological characterizations. All show paleoecological associations or morphological adaptations indicating a preference for open grasslands. Many are characterized by extreme hypsodonty and, with the possible exception of *A. australis*, all were likely grazers. *Equus capensis*, *S. antiquus*, and



**Fig. 4.** Last appearances of extinct ungulates at Nelson Bay Cave (data from Klein, 1983) and Boomplaas Cave (data from Klein, 1983; Faith, 2012, 2013b) relative to the abundance of grazing ungulates across stratigraphic units. Shaded area indicates stratigraphic units dating to the Last Glacial Maximum (LGM: 24–18 ka) and Lateglacial (18–12 ka). The stratigraphy and chronology of both sites follow Deacon (1984), with radiocarbon age estimates calibrated using the INTCAL09 calibration curve (Reimer et al., 2009).

*Metridiochoerus* were among the largest of their respective lineages, suggesting a requirement for greater quantities of grassy forage and more expansive home ranges. In light of these similarities, the extinctions have been attributed primarily to a decline in the availability or productivity of grassland habitats since the end of the Last Glacial Maximum (Klein, 1980, 1984b; Brink, 1987, 1999; Brink and Lee-Thorp, 1992; Faith, 2011c, 2012). Evidence for this is derived primarily from large mammal faunal assemblages, which indicate progressive declines in grazer abundances and diversity over the last 18,000 yrs. This is illustrated here in Fig. 4, which shows how ungulate extinctions at Nelson Bay Cave (Klein, 1983) and Boomplaas Cave (Faith, 2012, 2013b) correspond to a long-term decline of grassland ungulates. Potential mechanisms behind this decline could include vegetation change due to altered rainfall regimes, increased seasonality, or elevated atmospheric CO<sub>2</sub> concentrations (Brink, 1987; Brink and Lee-Thorp, 1992; Brink, 2005; Faith, 2011c, 2013a, 2013b; see also Faith, 2011a).

Klein (1980, 1984b) argued that environmental change alone is an inadequate explanation for the extinctions. Noting that the extinct taxa endured glacial-interglacial transitions since at least the beginning of the middle Pleistocene, he suggests that the late Pleistocene and Holocene losses occurred because of intensified predation by human hunters armed with LSA technology. Their survival during previous episodes of pronounced climate change is attributed to the presumably ineffective technology and limited hunting ability of MSA humans. However, a growing body of evidence indicates that this behavioral contrast is probably inaccurate and there is no evidence for intensified LSA predation pressure on large mammals (Marean and Assefa, 1999; McBrearty and Brooks, 2000; Faith, 2008; Dusseldorp, 2010; Wadley, 2010; Faith, 2011b). More importantly, Klein's argument overlooks the substantial extinctions that did occur prior to the late Pleistocene.

The southern African fossil record documents the extinction of at least 22 large mammals from ~1.0 million yrs ago to the end of the middle Pleistocene, including the dirk-toothed cat Megantereon whitei, the giant baboon Theropithecus oswaldi, the hippopotamid Hippopotamus gorgops, the suids Kolpochoerus and three species of Metridiochoerus, two species of three-toed horse Eurygnathohippus, an unnamed wild ass, an elephant (Loxodonta atlantica), a sivathere (Sivatherium maurusium), and at least ten bovids, including Aepyceros helmoedi, A. recki, Damaliscus aff. lunatus, ?Damaliscus sp. nov., Gazella sp., Hippotragus gigas, Megalotragus eucornutus, ?Parmularius sp. nov., Rabaticeras arambourgi, and the enigmatic "spiral horn" antelope (Cooke, 1974; Brink, 1987, 1994, 2005; Klein et al., 2007; Bishop, 2010; Brink et al., 2012). Some of these taxa may have evolved into other forms (e.g. L. atlantica, A. recki, M. eucornutus, R. arambourgi), but most disappeared without leaving any surviving representatives. Like the species that disappeared during the late Pleistocene and Holocene, many of these taxa are known or thought to have been open habitat grazers (Codron et al., 2008; Stynder, 2009). With this long term history in mind, there is little reason to argue that extinctions during the late Pleistocene and Holocene were the unique result of LSA human impacts. Rather, they represent part of a long-term extinction pattern, perhaps the result of elevated climate variability (e.g., Potts, 1998; Faith et al., 2012) or increased niche specialization (Codron et al., 2008).

## 4.2. East Africa

The East African LQE are only beginning to come to light, with the relevant taxonomic and chronological data still being established (Marean and Gifford-Gonzalez, 1991; Faith et al., 2011, 2012; Tryon et al., 2012; Faith et al., in press). Emerging evidence indicates some parallels with southern Africa. In the stratified archeological sequence from Lukenya Hill in south-central Kenya, *D. hypsodon* and *S. antiquus* disappear between the end of the Last Glacial Maximum and the onset of the Holocene (Marean and Gifford-Gonzalez, 1991; Marean, 1992). Other extinct East African taxa include *Megalotragus* sp., *R. atopocranion*, the

unnamed impala (*Aepyceros* sp.), *Kolpochoerus* sp., and *Orycteropus crassidens*, but they are known only from open air deposits broadly dated to the late Pleistocene (33 to 100 ka) (Faith et al., 2011; Tryon et al., 2012; Faith et al., in press). The extinct ungulates were all associated with faunal communities indicative of arid grassland habitats and, with the exception of the extinct impala, all were likely grazers (Marean and Gifford-Gonzalez, 1991; Marean, 1992; Garret et al., 2010; Tryon et al., 2010; Faith et al., 2011; Tryon et al., 2012; Faith et al., 2013). The disappearance of *D. hypsodon* and *S. antiquus* is attributed to a loss of arid grasslands at the onset of the Holocene together with increased competition from mesic-adapted grazers (Marean and Gifford-Gonzalez, 1991; Marean, 1992; Faith et al., 2011, 2012, 2013), likely the result of a marked increase in precipitation (Bonnefille et al., 1990; Bonnefille and Chalie, 2000; Kiage and Liu, 2006; Wolff et al., 2011).

Evidence from Kenya indicates that although the magnitude of extinctions over the last ~100,000 yrs is not particularly large in terms of the number of taxa lost, the extinctions radically reconfigured the large mammal community by eliminating the dominant taxa (Faith et al., 2012). *Damaliscus hypsodon* is the most abundant large mammal recovered from the late Pleistocene deposits at Lukenya Hill (Marean, 1992). In the Lake Victoria Basin, the dominant species include *R. atopocranion* followed by *D. hypsodon* (Faith et al., 2011; Tryon et al., 2012). At both localities, extinct species account for more than 50% of the large mammals (>5 kg). This represents an important departure from the southern African record, where extinct taxa typically represent a smaller component of late Pleistocene faunas.

#### 4.3. North Africa

North Africa documents the largest number of extinct taxa (Table 2), although its losses have received the least attention. The extinctions chronology is weak, as many of the relevant paleontological and archeological sites were excavated between 1850 and 1950 with little stratigraphic control. Even the more recently excavated late Pleistocene sites lack absolute dates and precise chronologies, leaving the late Pleistocene record poorly resolved (Geraads, 2012). It is clear that the North African extinctions extended into the Holocene, as evidenced by the late survival of B. primigenius, S. antiquus, E. mauritanicus, and E. melkiensis (Table 2). The middle Holocene extinction of S. antiquus corresponds with an abrupt climatic shift toward extreme aridity (Lézine, 1989; Lézine et al., 1990; DeMenocal et al., 2000; Foley et al., 2003). Its extinction during this arid phase may be related to competition with pastoralists and livestock for increasingly scarce water (Gautier and Muzzolini, 1991), a factor linked to the decline of water-dependent African ungulates that inhabit arid to semi-arid habitats today (Moehlman et al., 1998; Williams, 2002; Moehlman et al., 2008a, 2008b; Faith et al., 2013).

The disappearance of two species of *Gazella*, three species of *Equus*, and other large-bodied hypsodont species (*E. iolensis*, *Camelus*, *B. primigenius*, *S. antiquus*) in North Africa indicates that grazers or open habitat species were predominately affected. These ecological parallels imply that environmental mechanisms likely played a role. However, pending the establishment of a refined extinction chronology and terrestrial paleoenvironmental records, the North African extinctions will remain poorly understood. With the recent surge of paleoanthropological interest in the region (Hublin and McPherron, 2012), substantial progress can be expected in the near future.

## 5. Extinctions in ecological and evolutionary context

Ecomorphological evidence, paleoenvironmental associations, or taxonomic analogy indicate that many of the extinct taxa preferred open grassland habitats (20 of 24; Table 3). This is particularly clear within the bovids and equids (Table 1). The bovid tribes Antilopini and Alcelaphini, both of which include species with hypsodont teeth and cursorial postcranial adaptations, are considered the archetypal

#### Table 3

Inferred dietary habits of extinct species. Those in bold likely preferred grassland habitats, as indicated by ecomorphological adaptations, paleoenvironmental associations, or taxonomic analogy (see Section 3).

Taxon	Diet
Aepyceros sp.	Mixed feeder
Antidorcas australis	Mixed feeder
Antidorcas bondi	Grazer
Bos primigenius	Grazer
Damaliscus hypsodon	Grazer
Damaliscus niro	Grazer
Gazella atlantica	Mixed feeder
Gazella tingitana	Mixed feeder
Hippotragus leucophaeus	Grazer
Megalotragus priscus	Grazer
Rusingoryx atopocranion	Grazer
Syncerus antiquus	Grazer
Unnamed caprine antelope	Grazer
Camelus sp.	Browser
Megaceroides algericus	Browser
Metridiochoerus sp.	Grazer
Kolpochoerus sp.	Grazer
Equus algericus	Grazer
Equus capensis	Grazer
Equus mauritanicus	Grazer
Equus melkiensis	Grazer
Stephanorhinus	?
Elephas iolensis	Grazer
Orycteropus crassidens	Insectivore

open grassland lineages (Vrba, 1980; Greenacre and Vrba, 1984; Bobe and Behrensmeyer, 2004; Bobe, 2006). Species belonging to these tribes account for 22 of 82 (27%) extant African bovids recognized by the IUCN (2012), yet they account for the majority (8 of 13 = 62%) of the extinct bovids (Fisher's exact test: p = 0.022). The four extant species of *Equus* (*E. quagga, E. zebra, E. grevyi*, and *E. africanus*) are bulk grazers that also prefer open habitats, spanning a continuum from extremely arid scrublands (*E. africanus*) to more mesic grasslands (*E. quagga*). The late Pleistocene and Holocene extinctions halved the number of African equids (Table 1). With this long-term pattern in mind, it is perhaps not surprising that the conservation outlook for two of the extant species is grim, with *E. grevyi* and *E. africanus* classified as endangered and critically endangered, respectively (Moehlman et al., 2008a, 2008b).

Diet is also a significant correlate of extinction. Table 3 reports the inferred diets of the extinct African large mammals. These are based on a combination of stable carbon isotopes, dental mesowear analysis, ecomorphology, or analogy with extant relatives (references provided in species descriptions). Focusing on the ungulates, which include all extinct taxa but *O. crassidens*, it is evident that grazers are substantially over-represented. Reed (1998) summarized the dietary habits of 67



Fig. 5. Dietary habits of extant (from Reed, 1998) and extinct African ungulates. Number of species indicated in parentheses.

extant African ungulates, illustrated here in Fig. 5. Species identified as fresh-grass grazers or fruit/leaf eaters are included in the grazer or browser category, respectively, as these finer dietary classes cannot be readily identified in fossil taxa. A chi-square test shows the distribution of dietary classes among extinct taxa (Fig. 5) to differ significantly from extant ungulates ( $\chi^2 = 11.994$ , p = 0.007). Adjusted residuals, which are to be read as standard normal deviates, show that this is driven by the dominance of grazers (AR = 3.28, p = 0.001) and rarity of browsers (AR = -2.81, p = 0.005) among extinct species. As reviewed below, there is reason to believe that the preferential loss of grazers and open habitat taxa is rooted in long-term environmental and evolutionary processes.

Paleoenvironmental evidence documents an expansion of open and arid-to-semi-arid C<sub>4</sub> grasslands throughout Africa since the late Miocene (Cerling and Hay, 1988; Cerling, 1992; Cerling et al., 1997; Bobe and Behrensmeyer, 2004; DeMenocal, 2004; Bobe, 2006; Levin et al., 2011). Among large herbivores, this is associated with the diversification of grassland lineages (e.g., the bovid tribes Alcelaphini and Antilopini) and morphological adaptations to grazing in open habitats, including an increase in hypsodonty, enlarged body size, and the development of cursoriality (reviewed in Potts and Behrensmeyer, 1992; Bobe, 2006). Fossil evidence from East Africa and southern Africa documents a reversal of this trend over the last million years, however, with the loss of many large-bodied or hypsodont forms (Potts, 1998; Brink, 2005; Codron et al., 2008; Faith et al., 2012). For example, ~15 species of Alcelaphine bovid disappeared in the last million years (Vrba, 1997; Gentry, 2010a; and Table 1), leaving only 6 survivors (A. buselaphus, Beatragus hunteri, D. dorcas, D. lunatus, Connochaetes gnou, C. taurinus). Consistent with the long-term trend, many of the extinct grazers reviewed here are exceptionally hypsodont (Aepyceros sp., D. hypsodon, R. atopocranion, A. bondi, E. capensis, Metridiochoerus, E. iolensis) or among the largest of their respective lineages (M. priscus, S. antiquus, Metridiochoerus, E. capensis, Camelus).

The disappearance of large-bodied hypsodont grazers over the last million years corresponds with the establishment of high-amplitude 100 kyr glacial cycles and increased climate variability (Potts, 1998; DeMenocal, 2004). Terrestrial records spanning the Pleistocene-Holocene transition indicate that extinctions are associated with substantial changes in the structure, distribution, or productivity of grassland habitats (Klein, 1980; Avery, 1982; Klein, 1983; Marean and Gifford-Gonzalez, 1991; Marean, 1992; Faith, 2011c, 2012) and the same was likely true of previous glacial-interglacial transitions. In light of this evidence, it is probable that the extinctions within the hyper-diverse grazing niche were the result of high-amplitude late Quaternary glacial-interglacial cycles that altered the structure of grassland habitats and eliminated the feeding niches of more specialized taxa during interglacials. The result of this process can be described as selective taxonomic winnowing, characterized by the gradual loss of specialists and the establishment of a large mammal community composed of more versatile species capable of persisting across a broader range of environmental conditions (see also Potts, 1998; Faith et al., 2012).

## 6. Conclusion

The most recent comprehensive examination of LQE on continental Africa identified 11 species and 6 genera that disappeared during the late Pleistocene and Holocene (Klein, 1984b). In the last few decades this figure has expanded to include at least 24 species and 9–10 extinct genera (Table 1), revealing a much greater taxonomic breadth than previously considered (Klein, 1984b; Barnosky et al., 2004; Koch and Barnosky, 2006). This represents the loss of 14% of Africa's large mammals (>5 kg) and 25% of its megafauna (>44 kg) (data from Smith et al., 2003; *Antidorcas* excluded from megafauna). In East Africa alone, the number of securely dated extinct late Pleistocene mammals has risen from two to seven in only the last several years (Tryon et al., 2010; Faith et al., 2011, 2012; Tryon et al., 2012; Faith et al., in press).

These recent increases, together with the geographic bias in sampling (Fig. 1), suggest that much more remains to be understood.

It is now clear that the magnitude of large mammal extinctions on continental Africa over the last ~100,000 yrs is substantial. In terms of the number of extinct taxa that are securely dated to this time frame, the African extinctions exceed those in Europe and Asia (Grayson, 2007; Louys et al., 2007), match those from Australia (Wroe and Field, 2006; Field et al., 2008; Faith and O'Connell, 2011; Wroe et al., 2013), and are surpassed only by the losses in the Americas (Grayson, 1991, 2007; Cione et al., 2009; Faith and Surovell, 2009). Nevertheless, Africa remains the "fortunate anomaly" (Koch and Barnosky, 2006:221) in the sense that the majority of its large mammals survived the extinctions, in contrast to the situation in the Americas or Australia.

The African extinctions can be broadly characterized as involving the loss of species that are grazers or prefer open habitats. Where good extinction chronologies and paleoenvironmental records are available, these losses are readily accounted for by environmental change. In the broader temporal context, these losses represent the more recent examples of a long-term environmentally-driven extinction process over the last million years. Taken together, there is little reason to believe that humans played a major role in African extinctions.

## Acknowledgments

This synthesis is the outcome of years of reflection and discussion, and I would like to acknowledge the people who have contributed directly or indirectly to the ideas presented here: Kay Behrensmeyer, René Bobe, James Brink, Alison Brooks, Don Grayson, Curtis Marean, Dan Peppe, Rick Potts, Christian Tryon, Bernard Wood, and Mindy Zeder. I thank the two anonymous referees and Don Grayson for their helpful comments on previous versions of this manuscript. The Lake Victoria Prehistory Project has contributed substantially to our knowledge of East African late Pleistocene faunas, and I thank my collaborators, especially Christian Tryon and Dan Peppe, for making this work enjoyable and productive. Darryl de Ruiter provided helpful discussions on the Plovers Lake fauna. This research was supported by a University of Queensland Postdoctoral Research Fellowship.

#### References

- Abbate, E., Albianelli, A., Awad, A., Billi, P., Bruni, P., Delfino, M., Ferretti, M.P., Filippi, O., Gallai, G., Ghinassi, M., Laurtzen, S.-E., Lo Vetro, D., Martinez-Navarro, B., Martini, F., Napoleone, G., Bedri, O., Papini, M., Rook, L., Sagri, M., 2010. Pleistocene environments and human presence in the middle Atbara valley (Khashm El Girba, Eastern Sudan). Palaeogeogr. Palaeoclimatol. Palaeoecol. 292, 12–34.
- Abbazzi, L, 2004. Remarks on the validity of the generic name *Praemegaceros* Portis 1920, and an overview on *Praemegaceros* species in Italy. Rend. Lincei 15, 115–132.
- Amani, F., Geraads, D., 1993. Le gisement moustérien du Djebel Irhoud, Maroc: précisions sur la faune et la biochronologie, et description d'un nouveau reste humain. C. R. Acad. Sci. Paris 316, 847–852.
- Antón, S.C., 2012. Early *Homo*: who, when, and where. Curr. Anthropol. 53, S278–S298.Aouraghe, H., 2004. Les populations de mammifères atériens d'El Harhoura 1 (Témara, Maroc). Bull. Archaeol. Marocaine 20, 83–104.
- Aouraghe, H., Debénath, A., 1999. Les équidés du Pléistocène supérieur de la grotte Zouhrah à El Harhoura, Maroc. Quaternaire 10, 283–292.

Arambourg, C., 1938. Mammifères fossiles du Maroc. Mem. Soc. Sci. Nat. Maroc 46, 1-65.

Arambourg, C., 1947. Contribution a l'étude géologique et paléontologique du Bassin du Lac Rodolphe et de la basse vallée de l'Omo. Mission Scientifique de l'Omo, 1932–1933. Géologie-AnthropologieMuséum National D'Histoire Naturelle, Paris 231–562.

- Arambourg, C., 1957. Observation sur les gazelles fossiles du Pléistocène supérieur de l'Afrique du Nord. Bull. Soc. Hist. Nat. Afr. Nord. 48, 49–77.
- Arkell, A.J., 1962. The petroglyphs of Wadi Zirmei in North-Eastern Tibesti. Actes du IV Congres Panafricain de Prehistoire et de l'Etude du Quaternaire, pp. 391–394.
- Avery, D.M., 1982. Micromammals as palaeoenvironmental indicators and an interpretation of the late Quaternary in the southern Cape Province, South Africa. Ann. S. Afr. Mus. 85, 183–274.
- Azzaroli, A., Mazza, P., 1993. Large early Pleistocene deer from Pietrafitta lignite mine, central Italy. Palaeontol. Ital. 80, 1–24.
- Badenhorst, S., Plug, I., 2012. The faunal remains from the Middle Stone Age levels of Bushman Rock Shelter in South Africa. S. Afr. Archaeol. Bull. 67, 16–31.
- Bagtache, B., Hadjouis, D., Eisenmann, V., 1984. Présence d'un Equus caballin (E. algericus n. sp.) et d'une autre espèce nouvelle d'Equus (E. melkiensis n. sp.) dans l'Atérien des Allobroges, Algérie. C. R. Acad. Sci. 14, 609–612.

- Barnosky, A.D., Koch, P.L., Feranec, R.S., Wing, S.L., Shabel, A.B., 2004. Assessing the causes of Late Pleistocene extinctions on the continents. Science 306, 70–75.
- Bate, D.M.A., 1949. A new African fossil long-horned buffalo. Ann. Mag. Nat. Hist. Lond. 12, 396–398.
- Bate, D.M.A., 1951. The Mammals of Singa and Abu Hugar: Fossil Mammals of Africa, No. 2. British Museum (Natural History), London 1–28.
- Bauer, I.E., McMorrow, J., Yalden, D.W., 1994. The historic ranges of three equid species in north-east Africa: a quantitative comparison of environmental tolerances. J. Biogeogr. 21, 169–182.
- Behrensmeyer, A.K., Potts, R., Plummer, T.W., Tauxe, L, Opdyke, N., Jorstad, T., 1995. The Pleistocene locality of Kanjera, Western Kenya: stratigraphy, chronology and paleoenvironments. J. Hum. Evol. 29, 247–274.
- Bell, R.H.V., 1971. A grazing ecosystem in the Serengeti. Sci. Am. 225 (1), 86-93.
- Bennett, D.K., 1980. Stripes do not a zebra make, part I: a cladistic analysis of *Equus*. Syst. Zool. 29, 272–287.
- Bernor, R.L., Armour-Chelu, M., Gilbert, W.H., Kaiser, T.M., Schulz, E., 2010. Equidae. In: Werdelin, L., Sanders, W.J. (Eds.), Cenozoic Mammals of Africa. University of California Press, Berkeley.
- Billia, E.M.E., 2008. Revision of the fossil material attributed to *Stephanorhinus kirchbergensis* (Jäger 1839) (Mammalia, Rhinocerotidae) preserved in the museum collections of the Russian Federation. Quat. Int. 179, 25–37.
- Bishop, L.C., 2010. Suoidea. In: Werdelin, L., Sanders, W.J. (Eds.), Cenozoic Mammals of Africa. University of California Press, Berkeley, pp. 821–842.
- Bobe, R., 2006. The evolution of arid ecosystems in eastern Africa. J. Arid Environ. 66, 564–584.
- Bobe, R., Behrensmeyer, A.K., 2004. The expansion of grassland ecosystems in Africa in relation to mammalian evolution and the origin of the genus *Homo*. Palaeogeogr. Palaeoclimatol. Palaeoecol. 207, 399–420.
- Bonnefille, R., Chalie, F., 2000. Pollen-inferred precipitation time-series from equatorial mountains, Africa, the last 40 kyr BP. Global Planet. Change 26, 25–50.
- Bonnefille, R., Roeland, J.C., Guiot, J., 1990. Temperature and rainfall estimates for the past 40,000 years in equatorial Africa. Nature 346, 347–349.
- Bougariane, B., Zouhri, S., Ouchaou, B., Oujaa, A., Boudad, L., 2010. Large mammals from the Upper Pleistocene at Tamaris I 'Grotte des gazelles' (Casablanca, Morocco): paleoecological and biochronological implications. Hist. Biol. 22, 295–302.
- Bourguignat, J.R., 1870. Histoire du Djebel-Thaya et des Ossements Fossiles Recueillis dans la Grande Caverne de la Mosquée. Challamel, Paris.
- Bradley, D.G., MacHugh, D.E., Cunningham, P., Loftus, R., 1996. Mitochondrial diversity and the origins of African and European cattle. Proc. Natl. Acad. Sci. U. S. A. 93, 5131–5135.
- Brink, J.S., 1987. The archaeozoology of Florisbad, Orange Free State. Mem. Natl. Mus. Bloemfontein 24, 1–151.
- Brink, J.S., 1994. An ass, Equus (Asinus) sp., from the late Quaternary mammalian assemblages of Florisbad and Vlakkraal, central southern Africa. S. Afr. J. Sci. 90, 497–500.
- Brink, J.S., 1999. Preliminary report on a caprine from the Cape mountains, South Africa. Archaeozoologia 10, 11–26.
- Brink, J.S., 2005. The Evolution of the Black Wildebeest, Connochaetes gnou, and Modern Large Mammal Faunas in Central South Africa. University of Stellenbosch, South Africa (Ph.D. Thesis).
- Brink, J.S., Lee-Thorp, J.A., 1992. The feeding niche of an extinct springbok, Antidorcas bondi (Antelopini, Bovidae), and its paleoenvironmental meaning. S. Afr. J. Sci. 88, 227–229.
- Brink, J.S., De Bruiyn, H., Rademeyer, L.G., van der Westhuizen, W.A., 1995. A new find of *Megalotragus priscus* (Alcelaphini, Bovidae) from the central Karoo, South Africa. Palaeontol. Afr. 32, 17–22.
- Brink, J.S., Herries, A.I.R., Moggi-Cecchi, J., Gowlett, J.A.L., Bousman, C.B., Hancox, J.P., Grün, R., Eisenmann, V., Adams, J.W., Rossouw, L., 2012. First hominine remains from a 1.0 million year old bone bed at Cornelia-Uitzoek, Free State Province, South Africa. J. Hum. Evol. 63, 527–535.
- Broom, R., 1909a. On a large extinct species of Bubalis. Ann. S. Afr. Mus. 7, 279–280.

Broom, R., 1909b. On evidence of a large horse recently extinct in South Africa. Ann. S. Afr.

- Mus. 7, 281–282.
  Brown, A., Verhagen, B., 1985. Two Antidorcas bondi individuals from the Late Stone Age site of Kruger Cave 35/83, Olifantshoek, Rustenburg District, South Africa. S. Afr. I. Sci. 81, 102.
- Brown, F.H., McDougall, I., Fleagle, J.G., 2012. Correlation of the KHS Tuff of the Kibish Formation to volcanic ash layers at other sites, and the age of early *Homo sapiens* (Omo I and Omo II). J. Hum. Evol. 63, 577–585.
- Brun, A., Guerin, C., Levy, A., Riser, J., Rognon, P., 1988. Steppic environments at the end of the Upper Pleistocene in southern Tunisia (Oued el Akarit). J. Afr. Earth Sci. 7, 969–980.
- Brunet, M., White, T.D., 2001. Deux nouvelles espèces de Suini (Mammalia, Suidae) du continent africain (Ethiopie; Tchad). C. R. Acad. Sci. 332, 51–57.
- Bunn, H.T., 1981. Archaeological evidence for meat-eating by Plio-Pleistocene hominids from Koobi Fora and Olduvai Gorge. Nature 291, 574–576.
- Bunn, H.T., Kroll, E.M., 1986. Systematic butchery by Plio-Pleistocene hominids at Olduvai Gorge, Tanzania. Curr. Anthropol. 27, 431–452.
- Campbell, M.C., Tishkoff, S.A., 2010. The evolution of human genetic and phenotypic variation in Africa. Curr. Biol. 20, R166–R173.
- Cerling, T.E., 1992. Development of grasslands and savannas in East Africa during the Neogene. Palaeogeogr. Palaeoclimatol. Palaeoecol. 97, 241–247.
- Cerling, T.E., Hay, R.L., 1988. An isotopic study of paleosol carbonates form Olduvai Gorge. Quatern. Res. 25, 63–78.
- Cerling, T.E., Harris, J.M., MacFadden, B.J., Leakey, M.G., Quade, J., Eisenmann, V., Ehleringer, J.R., 1997. Global vegetation change through the Miocene–Pliocene boundary. Nature 389, 153–158.

Churcher, C.S., 1972, Late Pleistocene vertebrates from archaeological sites in the Plain of Kom Ombo, Upper Egypt, R. Ont, Mus. Life Sci. Contrib. 82, 1–172.

Churcher, C.S., 1974. Relationships of the late Pleistocene vertebrate fauna from Kom

- Ombo, Upper Egypt, Ann. Geol. Surv. Egypt 4, 363–384. Churcher, C.S., 1978. Giraffidae. In: Maglio, V.J., Cooke, H.B.S. (Eds.), Evolution of African Mammals, Harvard University Press, Cambridge, pp. 509–535.
- Churcher CS 1999 Holocene faunas of the Dakleh Oasis In: Churcher CS Mills AI (Eds.), Reports From the Survey of the Dakleh Oasis, Western Desert of Egypt 1977–1987. Oxbow Books, Oxford, pp. 133–151.
- Churcher, C.S., 2006. Distribution and history of the Cape zebra (Equus capensis) in the Quaternary of Africa. Trans. R. Soc. S. Afr. 61, 89-95.

Churcher, C.S., Richardson, M.L., 1978. Equidae. In: Maglio, V.J., Cooke, H.B.S. (Eds.), Evolution of African Mammals. Harvard University Press, Cambridge, pp. 379–422.

- Churcher, C.S., Smith, P.E.L, 1972. Kom Ombo: preliminary report on the fauna of late
- Paleolithic sites in Upper Egypt. Science 177, 259–261. Churcher, C.S., Kleindienst, M.R., Schwarcz, H.P., 1999. Faunal remains from a Middle Pleistocene lacustrine marl in Dakleh Oasis, Egypt: paleoenvironmental reconstructions, Palaeogeogr, Palaeoclimatol, Palaeoecol, 154, 301-312,
- Cione, A.L., Tonni, E.P., Soibelzon, L., 2009. Did humans cause the late Pleistocene-early Holocene mammalian extinctions in South America in a context of shrinking open areas? In: Haynes, G. (Ed.), American Megafaunal Extinctions at the End of the Pleistocene. Springer, Dordrecth, pp. 125-144.
- Clark, J.L., 2011. The evolution of human culture during the later Pleistocene: using fauna to test models on the emergence and nature of "modern" human behavior. I. Anthropol. Archaeol. 30, 273–291.
- Clark, J.L., Plug, I., 2008. Animal exploitation strategies during the South African Middle Stone Age: Howiesons Poort and post-Howiesons Poort fauna from Sibudu Cave. I Hum Evol 54 886-898
- Codron, D., Brink, J.S., Rossouw, L., Clasuss, M., 2008. The evolution of ecological specialization in southern African ungulates: competition or physical environmental turnover. Oikos 117, 334-353.
- Cooke, H.B.S., 1974. The geology, archaeology and fossil mammals of the Cornelia Beds, Orange Free State. Mem. Natl. Mus. Bloemfontein 35, 1-109.
- Cooke, H.B.S., Wells, L.W., 1951. Fossil remains from Chelmer, near Bulawayo, Southern Rhodesia. S. Afr. J. Sci. 57, 205-209.
- Coppens, Y., Maglio, V.J., Madden, C.T., Beden, M., 1978. Proboscidea. In: Maglio, V.J., Cooke, H.B.S. (Eds.), Evolution of African Mammals. Harvard University Press, Cambridge, pp. 336-367.
- Cruz-Uribe, K., 1983. The mammalian fauna from Redcliff Cave, Zimbabwe. S. Afr. Archaeol. Bull. 38, 7-16.
- d'Errico, F., Backwell, L., Villa, P., Degano, I., Lucejko, J.J., Bamford, M.K., Higham, T.F.G., Colombini, M.P., Beaumont, P.B., 2012. Early evidence of San material culture represented by organic artifacts from Border Cave, South Africa. Proc. Natl. Acad. Sci. U. S. A. 109, 13214-13219.
- de Ruiter, D.J., 2003. Revised faunal lists for Members 1-3 of Swartkrans, South Africa. Ann. Transv. Mus. 40, 29-41.
- de Ruiter, D., Brophy, J.K., Lewis, P.J., Churchill, S.E., Berger, L.R., 2008. Faunal assemblage composition and paleoenvironment of Plovers Lake, a Middle Stone Age locality in Gauteng Province, South Africa. J. Hum. Evol. 2008, 1102-1117.
- Deacon, J., 1984. The Later Stone Age of Southernmost Africa. British Archaeological Reports International Series, Oxford.
- DeMenocal, P.B., 2004. African climate change and faunal evolution during the Pliocene-Pleistocene. Earth Planet. Sci. Lett. 220, 3-24.
- DeMenocal, P., Ortiz, J., Guilderson, T., Adkins, J., Sarnthein, M., Baker, L., Yarusinksy, M., 2000. Abrupt onset and termination of the African Humid Period: rapid climate responses to gradual insolation forcing. Quat. Sci. Rev. 19, 347-361.
- Dibble, H.L., Aldeias, V., Alvarez-Fernandez, E., Blackwell, B.A.B., Hallet-Desguez, E., Jacobs, Z., Goldberg, P., Lin, S.C., Morala, A., Meyer, M.C., Olszewski, D.I., Reed, K., Reed, D., Rezek, Z., Richter, D., Roberts, R.G., Sandgathe, D., Schurmans, U., Skinner, A.R., Steele, T.E., El Hajraoui, M.A., 2012. New excavations at the site of Contrebandiers Cave, Morocco. PaleoAnthropology 2012, 145-201.
- Domínguez-Rodrigo, M., 2002. Hunting and scavenging by early humans: the state of the debate. J. World Prehistory 16, 1-54.
- Domínguez-Rodrigo, M., Pickering, T.R., Semaw, S., Rogers, M.J., 2005. Cutmarked bone from Pliocene archaeological sites at Gona, Afar, Ethiopia: implications for the function of the world's oldest tools. J. Hum. Evol. 48, 109-121.
- Domínguez-Rodrigo, M., Mabulla, A., Bunn, H.T., Barba, R., Diez-Martín, F., Egeland, C.P., Espílez, E., Egeland, A., Yravedra, J., Sánchez, P., 2009. Unraveling hominin behavior at another anthropogenic site from Olduvai Gorge (Tanzania): new archaeological and taphonomic research at BK, Upper Bed II. J. Hum. Evol. 57, 260-283.
- Domínguez-Rodrigo, M., Pickering, T.R., Bunn, H.T., 2010. Configurational approach to identifying the earliest hominin butchers. Proc. Natl. Acad. Sci. U. S. A. 107. 20929-20934.
- Dusseldorp, G.L., 2010. Prey choice during the South African Middle Stone Age: avoiding dangerous prey or maximizing returns? Afr. Archaeol. Rev. 27, 107-133.
- Duvernoy, G.L., 1851. Note sur une espèce de buffle fossile, Bubalus (Arni) antiquus, découverte en Algérie, C. R. Hebd, Seances Acad, Sci, Paris 33, 595-597.
- Eisenmann, V., 1980. Les chevaux (Equus sensu lato) fossiles et actuels: crânes et dents jugales supérieures. Cahiers de Paléontologie.CNRS. Paris.
- Eisenmann, V., 2000. Equus capensis (Mammalia, Perissodactyla) from Elandsfontein. Palaeontol, Afr. 36, 91-96.

Eisenmann, V., 2003. Gigantic horses. In: Petculescu, A., Stiuca, E. (Eds.), Advances in Vertebrate Paleontology "Hen to Panta". Romanian Academy, Bucharest, pp. 31–40.

Eisenmann, V., 2006. Pliocene and Pleistocene equids: paleontology versus molecular biology. In: Kahlke, R.-D., Maul, L.C., Mazza, P. (Eds.), Late Neogene and Quaternary

Biodiversity and Evolution: Regional Developments and Interregional Correlations. Courier Forschungsinstitut, Senckenberg, pp. 71–89.

- Eisenmann, V., Baylac, M., 2000. Extant and fossil Equus (Mammalia, Perissodactyla) skulls: a morphometric definition of the subgenus Equus. Zool. Scr. 29, 89-100. Eisenmann, V., Brink, J.S., 2000. Koffiefontein quaggas and true Cape quaggas: the impor-
- tance of basic skull morphology. S. Afr. J. Sci. 96, 529-533.
- Faith, I.T., 2008. Eland, buffalo, and wild pigs: were Middle Stone Age humans ineffective hunters? J. Hum, Evol. 55, 24-36.
- Fith, J.T., 2011a. Late Pleistocene climate change, nutrient cycling, and the megafaunal extinctions in North America. Quat. Sci. Rev. 30, 1675–1680. Faith, J.T., 2011b. Ungulate biogeography, statistical methods, and the proficiency of
- Middle Stone Age hunters. J. Hum. Evol. 60, 315–317.
- Faith, J.T., 2011c. Ungulate community richness, grazer extinctions, and human subsistence behavior in southern Africa's Cape Floral Region. Palaeogeogr. Palaeoclimatol. Palaeoecol. 306. 219-227.
- Faith, J.T., 2012. Paleozoological insights into management options for a threatened mammal: southern Africa's Cape mountain zebra (Equus zebra zebra). Divers. Distrib. 18, 438-447
- Faith, J.T., 2013a. Ungulate diversity and precipitation history since the Last Glacial Maximum in the Western Cape, South Africa. Quat. Sci. Rev. 68, 191-199.
- Faith, J.T., 2013b. Taphonomic and paleoecological change in the large mammal sequence from Boomplaas Cave, Western Cape, South Africa. J. Hum. Evol. 65, 715-730.
- Faith, J.T., O'Connell, J.F., 2011. Revisiting the late Pleistocene mammal extinction record at Tight Entrance Cave, southwestern Australia. Quatern. Res. 76, 397-400.
- Faith, J.T., Surovell, T.A., 2009. Synchronous extinction of North America's Pleistocene mammals. Proc. Natl. Acad. Sci. U. S. A. 106, 20641-20645.
- Faith, J.T., Thompson, J.C., 2013. Fossil evidence for seasonal calving and migration of extinct blue antelope (Hippotragus leucophaeus) in southern Africa. J. Biogeogr. 40, 2108-2118.
- Faith, J.T., Choiniere, J.N., Tryon, C.A., Peppe, D.J., Fox, D.L., 2011. Taxonomic status and paleoecology of Rusingoryx atopocranion (Mammalia, Artiodactyla), an extinct Pleistocene bovid from Rusinga Island, Kenya. Quatern. Res. 75, 697-707.
- Faith, J.T., Potts, R., Plummer, T.W., Bishop, L.C., Marean, C.W., Tryon, C.A., 2012. New perspectives on middle Pleistocene change in the large mammal faunas of East Africa: Damaliscus hypsodon sp. nov. (Mammalia, Artiodactyla) from Lainyamok, Kenya. Palaeogeogr. Palaeoclimatol. Palaeoecol. 361-362, 84-93.
- Faith, J.T., Tryon, C.A., Peppe, D.J., Fox, D.L., 2013a. The fossil history of Grévy's zebra (Equus grevyi) in equatorial East Africa. J. Biogeogr. 40, 359-369.
- Faith, J.T., Tryon, C.A., Peppe, D.J., Beverly, E.J., Blegen, N., 2013. Biogeographic and evolutionary implications of an extinct late Pleistocene impala from the Lake Victoria Basin. J. Mamm. Evol. http://dx.doi.org/10.1007/s10914-013-9238-1 (in press).
- Feathers, J.K., 2002. Luminescence dating in less than ideal conditions: case studies from Klasies River Main Site and Duinefontein, South Africa. J. Archaeol. Sci. 29, 177-194.
- Feathers, J.K., Bush, D.A., 2000. Luminescence dating of Middle Stone Age deposits at Die Kelders. J. Hum. Evol. 38, 91-118.
- Ferraro, J.V., Plummer, T.W., Pobiner, B.L., Oliver, J.S., Bishop, L.C., Braun, D.R., Ditchfield, P.W., Seaman III, J.W., Binetti, K.M., Seaman Jr., J.W., Hertel, F., Potts, R., 2013. Earliest archaeological evidence of persistent hominin carnivory. PLos ONE 8, 62174
- Field, J., Fillios, M., Wroe, S., 2008. Chronological overlap between humans and megafauna in Sahul (Pleistocene Australia-New Guinea): a review of the evidence. Earth-Sci. Rev. 89.97-115.
- Foley, J.A., Coe, M.T., Scheffer, M., Wang, G., 2003. Regime shifts in the Sahara and Sahel: interactions between ecological and climatic systems in northern Africa. Ecosystems 6, 524-539.
- Fortelius, M., Mazza, P., Sala, B., 1993. Stephanorhinus (Mammalia: Rhinocerotidae) of the western European Pleistocene, with a revision of S. etruscus (Falconer, 1868). Palaeontogr. Ital. 80, 63-155.
- Garret, N., Fox, D., McNulty, K., Tryon, C., Peppe, D., 2010. Isotope paleoecology of the Pleistocene Wasiriya Beds of Rusinga Island, Kenya. J. Vertebr. Paleontol. 30, 94A (supplement).
- Gautier, A., 1966. Camelus thomasi from the northern Sudan and its bearing on the relationship C. thomasi:C. bactrianus. J. Paleontol. 40, 1368-1372.
- Gautier, A., 1976. Animal remains from localities near Dishna. In: Wendorf, F., Schild, R. (Eds.), Prehistory of the Nile Valley. Academic Press, New York, pp. 365-368.
- Gautier, A., 1987. Prehistoric men and cattle in North Africa: a dearth of data and a surfeit of models. In: Close, A.E. (Ed.), Prehistory of Arid North Africa: Essays in Honour of Fred Wendorf. Southern Methodist University Press, Dallas, pp. 163-187.
- Gautier, A., 1993. The Middle Palaeolithic archaeofaunas from Bir Tarfawi (Western Desert, Egypt). In: Wendorf, F., Schild, R., Close, A.E. (Eds.), Egypt During the Last Interglacial. Plenum Press, New York, pp. 121-143.
- Gautier, A., Muzzolini, A., 1991. The life and times of the giant buffalo alias Bubalus/ Homoioceras/Pelorovis antiquus in North Africa. Archaeozoologia 4, 39-92.
- Gentry, A.W., 1965. New evidence on the systematic position of Hippotragus niro Hopwood, 1936 (Mammalia). Ann. Mag. Nat. Hist. 8, 335-338.
- Gentry, A.W., 1970. Reclassification for Makapania broomi Wells and Cooke (Bovidae, Mammalia). Palaeontol. Afr. 13, 63-67.
- Gentry, A.W., 1978. Bovidae. In: Maglio, V.J., Cooke, H.B.S. (Eds.), Evolution of African Mammals. Harvard University Press, Cambridge, pp. 540-572.
- Gentry, A.W., 2010a. Bovidae. In: Werdelin, L., Sanders, W.J. (Eds.), Cenozoic Mammals of Africa. University of California Press, Berkeley, pp. 741–796.
- Gentry, A.W., 2010b. Cervidae. In: Werdelin, L., Sanders, W.J. (Eds.), Cenozoic Mammals of Africa. University of California Press, Berkeley, pp. 813–814.
- Gentry, A.W., 2011. Bovidae. In: Harrison, T. (Ed.), Paleontology and Geology of Laetoli, vol. 2. Springer, Dordrecth, pp. 363–465.

Gentry, A.W., Gentry, A., 1978. Fossil Bovidae of Olduvai Gorge, Tanzania. Bull. Br. Mus. Nat. Hist. Geol. Ser. 29, 289–446.

Geraads, D., 1992. Phylogenetic analysis of the tribe Bovini (Mammalia: Artiodactyla). Zool. J. Linn. Soc. 104, 193–207.

- Geraads, D., 2010a. Biogeographic relationships of Pliocene and Pleistocene Northwestern African mammals. Quat. Int. 212, 159–168.
- Geraads, D., 2010b. Rhinocerotidae. In: Werdelin, L., Sanders, W.J. (Eds.), Cenozoic Mammals of Africa. University of California Press, Berkeley, pp. 669–683.
- Geraads, D., 2012. The faunal context of human evolution in the late middle/late Pleistocene of northwestern Africa. In: Hublin, J.-J., McPherron, S.P. (Eds.), Modern Origins: A North African Perspective. Springer. Dordrecht.
- Geraads, D., Alemseged, Z., Reed, D., Wynn, J., Roman, D.C., 2004. The Pleistocene fauna (other than Primates) from Asbole, lower Awash Valley, Ethiopia, and its environmental and biochronological implications. Geobios 37, 697–718.
- Geraads, D., Melillo, S., Haile-Selassie, Y., 2009. Middle Pliocene Bovidae from the Hominid-bearing sites in the Waronso-Mille area, Afar region, Ethiopia. Palaeontol. Afr. 44, 57–68.
- Gonder, M.K., Mortensen, H.M., Reed, F.A., de Sousa, A., Tishkoff, S.A., 2007. Whole mtDNA genome sequence analysis of ancient African lineages. Mol. Biol. Evol. 24, 757–768.
- Grayson, D.K., 1991. Late Pleistocene mammalian extinctions in North America. J. World Prehistory 5, 193–231.
- Grayson, D.K., 2007. Deciphering North American Pleistocene extinctions. J. Anthropol. Res. 63, 185–213.
- Grayson, D.K., Meltzer, D.J., 2002. Clovis hunting and large mammal extinction: a critical review of the evidence. J. World Prehistory 16, 313–359.
- Grayson, D.K., Meltzer, D.J., 2003. A requiem for North American overkill. J. Archaeol. Sci. 30, 585–593.
- Greenacre, M.J., Vrba, E.S., 1984. Graphical display and interpretation of antelope census data in African wildlife areas using correspondence analysis. Ecology 65, 984–997.
- Groves, C.P., Bell, C.H., 2004. New investigations on the taxonomy of the zebras genus *Equus*, subgenus *Hippotigris*. Mamm. Biol. 69, 182–196.
- Grün, R., Stringer, C.B., 1991. Electron spin resonance dating and the evolution of modern humans. Archaeometry 33, 153–199.
- Grün, R., Brink, J.S., Spooner, N.A., Taylor, L., Stringer, C.B., Franciscus, R.G., Murray, A.S., 1996. Direct dating of the Florisbad hominin. Nature 382, 500–501.
- Hack, M.A., Lorenzen, E., 2008. Equus quagga, IUCN 2012. IUCN Red List of Threatened Species. Version 2012.2 (www.iucnredlist.org).
- Hadjouis, D., 1990. Megaceroides algericus (Lydekker, 1890), du gisement des Phacochères (Alger, Algérie). Etude critique de la position systématique de Megaceroides. Quaternaire 1, 247–258.
- Hadjouis, D., 2002. Un nouveau Bovini dans le faune du Pléistocène supérieur d'Algérie. Anthropologie 106, 377–386.
- Hamilton, W.R., 1978. Cervidae and Palaeomerycidae. In: Maglio, V.J., Cooke, H.B.S. (Eds.), Evolution of African Mammals. Harvard University Press, Cambridge, pp. 496–508. Hanotte, O., Bradley, D.G., Ochieng, J.W., Verjee, Y., Hill, E.W., Rege, J.E.O., 2002. African

pastoralism: genetic imprints of origins and migrations. Science 296, 336–339.

- Harris, J.M., 1991. Family Bovidae. In: Harris, J.M. (Ed.), Koobi Fora Research Project. The Fossil Ungulates: Geology, Fossil Artiodactyls, and Palaeoenvironments, 3. Clarendon Press, Oxford, pp. 139–320.
- Harris, J.M., Cerling, T.E., 2002. Dietary adaptations of extant and Neogene African suids. J. Zool. Soc. Lond. 256, 45–54.
- Harris, J.M., White, T.D., 1979. Evolution of the Plio-Pleistocene African suidae. Trans. Am. Philos. Soc. 69, 1–128.
- Harris, J.M., Geraads, D., Solounias, N., 2010a. Camelidae. In: Werdelin, L., Sanders, W.J. (Eds.), Cenozoic Mammals of Africa. University of California Press, Berkeley, pp. 815–820.
- Harris, J.M., Solounias, N., Geraads, D., 2010b. Giraffoidea. In: Werdelin, L., Sanders, W.J. (Eds.), Cenozoic Mammals of Africa. University of California Press, Berkeley, pp. 797–811.
- Heissig, K., 1999. Family Rhinocerotidae. In: Rossner, G., Heissig, K. (Eds.), The Miocene Land Mammals of Europe. Pfiel, Munich, pp. 175–188.
- Hendey, Q.B., 1974. The late Cenozoic carnivora of the south-western Cape Province. Ann. S. Afr. Mus. 63, 1–369.
- Hendey, Q.B., Hendey, H., 1968. New Quaternary fossil sites near Swartklip, Cape Province. Ann. S. Afr. Mus. 52, 43–73.
- Henshilwood, C.S., Marean, C.W., 2003. The origin of modern human behavior. Curr. Anthropol. 44, 627–651.
- Higgs, E.S., 1967. Environment and chronology: the evidence from mammalian fauna. In: McBurney, C.B.M. (Ed.), The Haua Fteah (Cyrenaica) and the Stone Age of the Southeast Mediterranean. Cambridge University Press, Cambridge, pp. 149–164.
- Hopwood, A.T., 1936. New and little-known fossil mammals from the Pleistocene of Kenya Colony and Tanganyika Territory. Ann. Mag. Nat. Hist. Ser 10 10 (17), 636–641.
- Hublin, J.-J., McPherron, S.P. (Eds.), 2012. Modern Origins: A North African Perspective. Springer, Dordrecht.
- IUCN SSC Antelope Specialist Group, 2008a. Eudorcas rufina. IUCN 2012. IUCN Red List of Threatened Species. Version 2012.2 (<www.iucnredlist.org>).
- IUCN SSC Antelope Specialist Group, 2008b. Hippotragus leucophaeus. IUCN 2012. IUCN Red List of Threatened Species, Version 2012.2. <a href="https://www.iucnredlist.org">www.iucnredlist.org</a>>.
- Jacobs, Z., Roberts, R.G., Nespoulet, R., El Hajraoui, M.A., Debénath, A., 2012. Single-grain OSL chronologies for Middle Paleolithic deposits at El Mnasra and El Harhoura 2, Morocco: implications for Late Pleistocene human–environment interactions along the Atlantic coast of northwest Africa. J. Hum. Evol. 62, 377–394.
- Jarman, P.J., 1974. The social organization of antelope in relation to their ecology. Behavior 58, 215–267.
- Joleaud, L, 1914. Sur le Cervus (Megaceroides) algericus Lydekker (1890). C. R. Soc. Biol. Paris 76, 737–739.

- Kerley, G.I.H., Sims-Castley, R., Boshoff, A.F., Cowling, R.M., 2009. Extinction of the blue antelope *Hippotragus leucophaeus*: modeling predicts non-viable global population size as the primary driver, Biodivers, Conserv. 18, 3235–3242.
- Kiage, L.M., Liu, K., 2006. Late Quaternary paleoenvironmental changes in East Africa: a review of multiproxy evidence from palynology, lake sediments, and associated records. Prog. Phys. Geogr. 30, 633–658.
- Kimbel, W.H., Johanson, D.C., Rak, Y., 1997. Systematic assessment of a maxilla of *Homo* from Hadar, Ethiopia. Am. J. Phys. Anthropol. 103, 235–262.
- Kingdon, J., 1982. East African Mammals, IIIC & IIID. University of Chicago Press, Chicago. Klein, R.G., 1972. The late Quaternary mammalian fauna of Nelson Bay Cave (Cape Province, South Africa): its implications for megafaunal extinctions and environmental and cultural change. Quatern. Res. 2, 135–142.
- Klein, R.G., 1974a. On the taxonomic status, distribution and ecology of the blue antelope, *Hippotragus leucophaeus* (Pallas, 1766), Ann. S. Afr. Mus. 65, 99–143.
- Klein, R.G., 1974b. A provisional statement on terminal Pleistocene mammalian extinctions in the Cape Biotic Zone (Southern Cape Province, South Africa). S. Afr. Archaeol. Soc. Goodwin Ser. 2, 39–45.
- Klein, R.G., 1976. The mammalian fauna of the Klasies River Mouth sites, Southern Cape Province, South Africa. S. Afr. Archaeol. Bull. 31, 75–98.
- Klein, R.G., 1977. The mammalian fauna from the Middle and Later Stone Age (Later Pleistocene) levels of Border Cave, Natal Province, South Africa. S. Afr. Archaeol. Bull. 34, 14–27.
- Klein, R.G., 1980. Environmental and ecological implications of large mammals from Upper Pleistocene and Holocene sites in southern Africa. Ann. S. Afr. Mus. 81, 223–283.
- Klein, R.G., 1983. Palaeoenvironmental implications of Quaternary large mammals in the fynbos region. In: Deacon, H.J., Hendey, Q.B., Lambrechts, J.J.N. (Eds.), Fynbos Palaeoecology: A Preliminary Synthesis, South African National Scientific Programmes Report No 75. Mills Litho, Cape Town, pp. 116–138.
- Klein, R.G., 1984a. Later Stone Age faunal samples from Heuningneskrans Shelter (Transvaal) and Leopard's Hill Cave (Zambia). S. Afr. Archaeol. Bull. 39, 109–116.
- Klein, R.G., 1984b. Mammalian extinctions and Stone Age people in Africa. In: Martin, P.S., Klein, R.G. (Eds.), Quaternary Extinctions: A Prehistoric Revolution. University of Arizona Press, Tucson, pp. 553–573.
- Klein, R.G., 1994. The long-horned African buffalo (*Pelorovis antiquus*) is an extinct species. J. Archaeol. Sci. 21, 725–733.
- Klein, R.G., 2001. Southern Africa and modern human origins. J. Anthropol. Res. 57, 1–16. Klein, R.G., Cruz-Uribe, K., 1987. Large mammal and tortoise bones from Eland's Bay Cave and nearby sites, Western Cape Province, South Africa. In: Parkington, J.E., Hall, M. (Eds.), Papers in the Prehistory of the Western Cape, South Africa. BAR International Series, 322, pp. 350–372 (Cambridge).
- Klein, R.G., Cruz-Uribe, K., 1995. The identification of Equus skulls to species, with particular reference to the craniometric and systematic affinities of the extinct South African quagga. In: Stewart, K.M., Seymour, K.L. (Eds.), Palaeoecology and Palaeoenvironments of Late Cenozoic Mammals: Tributes to the Career of C.S. Churcher. University of Toronto Press, Toronto, pp. 589–629.
- Klein, R.G., Cruz-Uribe, K., 1999. Craniometry of the genus Equus and the taxonomic affinities of the extinct South African quagga. S. Afr. J. Sci. 95, 81–86.
- Klein, R.G., Cruz-Uribe, K., 2000. Middle and Later Stone Age large mammal and tortoise remains from Die Kelders Cave 1, Western Cape Province, South Africa. J. Hum. Evol. 38, 169–195.
- Klein, R.G., Scott, K., 1986. Re-analysis of faunal assemblages from the Haua Fteah and other late Quaternary archaeological sites in Cyernaican Libya. J. Archaeol. Sci. 13, 515–542.
- Klein, R.G., Cruz-Uribe, K., Beaumont, P.B., 1991. Environmental, ecological, and paleoanthropological implications of the Late Pleistocene mammalian fauna from Equus Cave, Northern Cape Province, South Africa. Quatern. Res. 36, 94–119.
- Klein, R.G., Avery, G., Cruz-Uribe, K., Steele, T.E., 2007. The mammalian fauna associated with an archaic hominin skullcap and later Acheulean artifacts at Elandsfontein, Western Cape Providence, South Africa. J. Hum. Evol. 62, 164–186.
- Koch, P.L., Barnosky, A.D., 2006. Late Quaternary extinctions: state of the debate. Annu. Rev. Ecol. Evol. Syst. 37, 215–250.
- Lacruz, R.S., Brink, J.S., Hancox, P.J., Skinner, A.R., Herries, A., Schmid, P., Berger, L.R., 2002. Palaeontology and geological context of a middle Pleistocene faunal assemblage from Gladysvale Cave, South Africa. Palaeontol. Afr. 38, 99–114.
- Lee-Thorp, J.A., Beaumont, P.B., 1995. Vegetation and seasonality shifts during the late Quaternary deduced from <sup>13</sup>C/<sup>12</sup>C ratios of grazers at Equus Cave, South Africa. Quatern. Res. 43, 426–432.
- Lehman, T., 2009. Phylogeny and systematics of the Orycteropodidae (Mammalia, Tubulidentata). Zool. J. Linn. Soc. 155, 649–702.
- Leonard, J.A., Rohland, N., Glaberman, S., Fleischer, R.C., Caccone, A., Hofrieter, M., 2005. A rapid loss of stripes: the evolutionary history of the extinct quagga. Biol. Lett. 22, 291–295.
- Levin, N.E., Brown, F.H., Behrensmeyer, A.K., Bobe, R., Cerling, T.E., 2011. Paleosol carbonates from the Omo Group: isotopic records of local and regional environmental change in East Africa. Palaeogeogr. Palaeoclimatol. Palaeoecol. 307, 75–89.
- Lézine, A.-M., 1989. Late Quaternary vegetation and climate of the Sahel. Quatern. Res. 32, 317–334.
- Lézine, A.-M., Casanova, J., Hillaire-Marce, C., 1990. Across an early Holocene humid phase in western Sahara: pollen and isotope stratigraphy. Geology 18, 264–267.
- Lister, A.M., Edwards, C.J., Nock, D.A.W., Bunce, M., van Pijlen, I.A., Bradley, D.G., Thomas, M.G., Barnes, I., 2005. The phylogenetic position of the 'giant deer' *Megaloceros* giganteus. Nature 438, 850–853.
- Lönnberg, E., 1933. Description of a fossil buffalo from East Africa. Ark. Zool. 25A (17), 1–32 (Stockholm).

Loubser, J., Brink, J.S., Lourens, G., 1990. Paintings of the extinct blue antelope, *Hippotragus leucophaeus*, in the eastern Orange Free State, S. Afr. Archaeol. Bull. 45, 106–111.

Louys, J., Curnoe, D., Tong, H., 2007. Characteristics of Pleistocene megafauna extinctions in Southeast Asia. Palaeogeogr. Palaeoclimatol. Palaeoecol. 243, 152–173.

- Lydekker, R., 1890. On a cervine jaw from Algeria. Proc. Zool. Soc. Lond. 1890, 602–604. Lyons, S.K., Smith, F.A., Brown, J.H., 2004. Of mice, mastodons and men: human-mediated extinctions on four continents. Evol. Ecol. Res. 6, 339–358.
- MacInnes, D., 1956. Fossil tubulidentata from East Africa. Fossil Mamm. Afr. 10, 1–38.
  MacPhee, R.D.E. (Ed.), 1999. Extinctions in Near Time: Causes, Contexts, and Consequences. Kluwer/Plenum. New York.
- MacPhee, R.D.E., Flemming, C., 1999. Requiem æternam: the last five hundred years of mammalian species extinctions. In: MacPhee, R.D.E. (Ed.), Extinctions in Near Time: Causes, Contexts, and Consequences. Kluwer Academic/Plenum Publishers, New York, pp. 333–371.
- Maglio, V.J., 1973. Origin and evolution of the Elephantidae. Trans. Am. Philos. Soc. 63, 1–149.
- Maglio, V.J., 1978. Patterns of faunal evolution. In: Maglio, V.J., Cooke, H.B.S. (Eds.), Evolution of African Mammals. Harvard University Press, Cambridge, pp. 603–619.
- Maglio, V.J., Cooke, H.B.S. (Eds.), 1978. Evolution of African Mammals. Harvard University Press, Cambridge.
- Marean, C.W., 1990. Late Quaternary Paleoenvironments and Faunal Exploitation in East Africa. University of California, Berkeley (Ph.D. Thesis).
- Marean, C.W., 1992. Implications of late Quaternary mammalian fauna from Lukenya Hill (south-central Kenya) for paleoenvironmental change and faunal extinctions. Quatern. Res. 37, 239–255.
- Marean, C.W., 1997. Hunter–gatherer foraging strategies in tropical grasslands: model building and testing in the East African Middle and Later Stone Age. J. Anthropol. Archaeol. 16, 189–225.
- Marean, C.W., Assefa, Z., 1999. Zooarchaeological evidence for the faunal exploitation behavior of Neandertals and early modern humans. Evol. Anthropol. 8, 22–37.
- Marean, C.W., Gifford-Gonzalez, D., 1991. Late Quaternary extinct ungulates of East Africa and palaeoenvironmental implications. Nature 350, 418–420.
- Marks, A.E., Peters, J., Van Neer, W., 1987. Late Pleistocene and early Holocene occupations in the Upper Atbara River Valley, Sudan. In: Close, A.E. (Ed.), Prehistory of Arid North Africa: Essays in Honor of Fred Wendorf. Southern Methodist University Press, Dallas, pp. 137–162.
- Martin, P.S., 1967. Prehistoric overkill. In: Martin, P.S., Wright, H.E.J. (Eds.), Pleistocene Extinctions: The Search for a Cause. Yale University Press, New Haven, pp. 75–120.
- Martin, P.S., 1984. Prehistoric overkill: the global model. In: Martin, P.S., Klein, R.G. (Eds.), Quaternary Extinctions: A Prehistoric Revolution. University of Arizona Press, Tucson, pp. 354–403.
- Martin, P.S., 2005. Twilight of the Mammoths: Ice Age Extinctions and the Rewilding of North America. University of California Press, Berkeley.
- Martin, P.S., Steadman, D.W., 1999. Prehistoric extinctions on islands and continents. In: MacPhee, R.D.E. (Ed.), Extinctions in Near Time: Causes, Contexts, and Consequences. Klewer/Plenum, New York, pp. 17–52.
- Martinez-Navarro, B., Perez-Claros, J.A., Palombo, M.R., Rook, L., Palmqvist, P., 2007. The Olduvai buffalo *Pelorovis* and the origin of *Bos*. Quatern. Res. 68, 220–226.
- McBrearty, S., Brooks, A.S., 2000. The revolution that wasn't: a new interpretation of the origin of modern human behavior. J. Hum. Evol. 39, 453–563.
- McBrearty, S., Jablonski, N.G., 2005. First fossil chimpanzee. Nature 437, 105-108.
- McDougall, I., Brown, F.H., Fleagle, J., 2005. Stratigraphic placement and age of modern humans from Kibish, Ethiopia. Nature 433, 733–736.
- McPherron, S.P., Alemseged, Z., Marean, C.W., Wynn, J.G., Reed, D., Geraads, D., Bobe, R., Bearat, H.A., 2010. Evidence for stone-tool-assisted consumption of animal tissues before 3.39 million years ago at Dikika. Ethiopia 466, 857–860.
- Mercier, N., Wengler, L., Valladas, H., Joron, J.-L., Froget, L., Reyss, J.-L., 2007. The Rhafas Cave (Morocco): Chronology of the mousterian and aterian archaeological occupations and their implications for Quaternary geochronology based on luminescence (TL/OSL) age determinations. Quat. Geochronol. 2, 309–313.
- Merzoug, S., Sari, L., 2008. Re-examination of the Zone I material from Tamar Hat (Algeria): zooarchaeological and technofunctional analyses. Afr. Archaeol. Rev. 25, 57–73.
- Michel, P., 1992. Pour une meilleure connaissance du Quaternaire Continental Marocain: les vertébrés fossiles du Maroc Atlantique. Cent. Orient. Anthropologie 96, 643–656.
- Michel, P., Campmas, E., Stoetzel, E., Nespoulet, R., Abdeljalil El Hajraoui, A., Amani, F., 2009. La macrofaune du Pléistocène supérieur d'El Harhoura 2 (Témara, Maroc): données préliminaires. Anthropologie 113, 283–312.
- Milo, R.G., 1998. Evidence for hominid predation at Klasies River Mouth, South Africa, and its implications for the behavior of early modern humans. J. Archaeol. Sci. 25, 99–133.
- Moehlman, P.D., Kedebe, F., Yohannes, H., 1998. The African wild ass (*Equus africanus*): conservation status in the horn of Africa. Appl. Anim. Behav. Sci. 60, 115–124.
- Moehlman, P.D., Rubenstein, D.I., Kebede, F., 2008a. Equus grevyi. IUCN Red List of Threatened Species. Version 2012.2 (<www.iucnredlist.org>).
- Moehlman, P.D., Yohannes, H., Teclai, R., Kebede, F., 2008b. Equus africanus. IUCN Red List of Threatened Species. Version 2012.2 (<www.iucnredlist.org>).
- Monchot, H., Aouraghe, H., 2009. Deciphering the taphonomic history of an Upper Paleolithic faunal assemblage from Zouhrah Cave/El Harhoura 1, Morocco. Quaternaire 20, 239–253.
- Morgan, LE, Renne, P.R., 2008. Diachronous dawn of Africa's Middle Stone Age: New <sup>40</sup>Ar/<sup>39</sup>Ar ages from the Ethiopian Rift. Geology 36, 967–970.
- Nespoulet, R., El Hajraoui, M.A., Amani, F., Ben Ncer, A., Debénath, A., El Idrissi, A., Lacombe, J.-P., Michel, P., Oujaa, A., Stoetzel, E., 2008. Palaeolithic and Neolithic occupations in the Temara region (Rabat, Morocco): recent data on hominin contexts and behavior. Afr. Archaeol. Rev. 25, 21–39.

- Orlando, L., Metcalf, J.L., Alberdi, M.T., Telles-Antunes, M., Bonjean, D., Otte, M., Martin, F., Eisenmann, V., Mashkour, M., Morello, F., Prado, J.L., Salas-Gismondi, R., Shockey, B.J., Wrinn, P.J., Vasilev, S.K., Ovodov, N.D., Cherry, M.I., Hopwood, B., Male, D., Austin, J.J., Hänni, C., Cooper, A., 2009. Revising the recent evolutionary history of equids using ancient DNA. Proc. Natl. Acad. Sci. U. S. A. 106, 21754–21759.
- Peters, J., 1990. Late Pleistocene hunter-gatherers at Ishango (Eastern Zaire): the faunal evidence. Rev. Paléobiol. 9, 73–112.
- Peters, J., 1992. Late Quaternary mammalian remains from Central and Eastern Sudan and their palaeoenvironmental significance. Palaeoecol. Afr. Surrounding Islands Antarct. 23, 91–115.
- Peters, J., Gautier, A., Brink, J.S., Haenen, W., 1994. Late Quaternary extinction of ungulates in sub-Saharan Africa: a reductionist's approach. J. Archaeol. Sci. 21, 17–28.
- Pfeiffer, T., 2002. The first complete skeleton of *Megaloceros verticornis* (Dawkins, 1868) Cervidae, Mammalia, from Bilshausen (Lower Saxony, Germany): description and phylogenetic implications. Mitteilungen as dem Museum für Naturkunde in Berlin. Geowissenschaftliche Reihe, 5 289–308.
- Pickford, M., 1975. New fossil Orycteropodidae (Mammalia, Tubulidentata) from East Africa. Neth. J. Zool. 25, 57–88.
- Pickford, M., Thomas, H., 1984. An aberrant new bovid (Mammalia) in subrecent deposits from Rusinga Island, Kenya. Proc. K. Ned. Akad. Wet. B87, 441–452.
- Plug, I., 1993. The macrofaunal remains of wild animals form Abbot's Cave and Lame Sheep Shelter, Seacow Valley, Cape. Koedoe 36, 15–26.
- Plug, I., 1997. Late Pleistocene and Holocene hunter–gatherers in the eastern highlands of South African and Lesotho: a faunal interpretation. J. Archaeol. Sci. 24, 715–727.
- Plug, I., Engela, R., 1992. The macrofaunal remains from recent excavations at Rose Cottage Cave, Orange Free State. S. Afr. Archaeol. Bull. 47, 16–25.
- Plummer, T.W., Bishop, L.C., 1994. Hominid paleoecology at Olduvai Gorge, Tanzania, as indicated by antelope remains. J. Hum. Evol. 27, 47–75.
- Pobiner, B.L., Rogers, M.J., Monahan, C.M., Harris, J.W.K., 2008. New evidence for hominin carcass processing strategies at 1.5 Ma, Koobi Fora, Kenya. J. Hum. Evol. 55, 103–130.
- Pomel, A., 1893. Caméliens et cervidés. Carte Géologie de l'Algérie. Monographies de Paléontologie 1–52.
- Pomel, A., 1895. Les éléphants Quaternaires. Carte Géol. Alger. Monogr. Paléontol. 6, 1–68. Pomel, A., 1897. Les équidés. Carte Geol. Alger. Monogr. Paleontol. 12, 1–44.
- Potts, R., 1998. Variability selection in hominid evolution. Evol. Anthropol. 7, 81–96.
- Potts, R., Behrensmeyer, A.K., 1992. Late Cenozoic terrestrial ecosystems. In: Behrensmeyer, A.K., Damuth, J.D., DiMichele, W.A., Potts, R., Sues, H.-D., Wing, S.L. (Eds.), Terrestrial Ecosystems Through Time: Evolutionary Paleoecology of Terrestrial Plants and Animals. University of Chicago Press, Chicago, pp. 419–541.
- Potts, R., Deino, A., 1995. Mid-Pleistocene change in large mammal faunas of East Africa. Quatern. Res. 43, 106–113.
- Potts, R., Shipman, P., 1981. Cutmarks made by stone tools on bones from Olduvai Gorge, Tanzania. Nature 291, 577–580.
- Raynal, J.-P., Occhietti, S., 2012. Amino chronology and an earlier age for the Moroccan Aterian. In: Hublin, J.-J., McPherron, S.P. (Eds.), Modern Origins: A North African Perspective. Springer, Dordrecht, pp. 79–90.
- Raynal, J.-P., Amani, F., Geraads, D., Graoui, M., Magoga, L., Texier, J.-P., 2008. Felids Cave, a new Upper Pleistocene Palaeolithic site at Casablanca (Morocco). Anthropologie 112, 182–200.
- Rector, A.L., Reed, K.E., 2010. Middle and Late Pleistocene faunas of Pinnacle Point and their paleoecological implications. J. Hum. Evol. 59, 340–357.
- Reed, K.E., 1998. Using large mammal communities to examine ecological and taxonomic structure and predict vegetation in extant and extinct assemblages. Paleobiology 24, 384–408.
- Reimer, P.J., Baillie, M.G.L., Bard, E., Bayliss, A., Beck, J.W., Blackwell, P.G., Bronk Ramsey, C., Buck, C.E., Burr, G.S., Edwards, R.L., Friedrich, M., Grootes, P.M., Guilderson, T.P., Hajdas, I., Heaton, T.J., Hogg, A.G., Hughen, K.A., Kaiser, K.F., Kromer, B., McCormac, F.G., Manning, S.W., Reimer, R.W., Richards, D.A., Southon, J.R., Talamo, S., Turney, C.S.M., van der Plicht, J., Weyhenmeyer, C.E., 2009. IntCal09 and Marine09 radiocarbon age calibration curves, 0–50,000 years cal BP. Radiocarbon 51, 1111–1150.
- Robbins, L.H., Murphy, M.L., Brook, G.A., Ivester, A.H., Campbell, A.C., Klein, R.G., Milo, R.G., Stewart, K.M., Downey, W.S., Stevens, N.J., 2000. Archaeology, palaeoenvironment, and chronology of the Tsodilo Hills White Paintings Rock Shelter, Northwest Kalahari Desert, Botswana. J. Archaeol. Sci. 27, 1085–1113.
- Sanders, W.J., Werdelin, L., 2010. Introduction. In: Werdelin, L., Sanders, W.J. (Eds.), Cenozoic Mammals of Africa. University of California Press, Berkeley, pp. xv-xxi.
- Sanders, W.J., Gheerbrant, E., Harris, J.M., Saegusa, H., Delmer, C., 2010. Proboscidea. In: Werdelin, L., Sanders, W.J. (Eds.), Cenozoic Mammals of Africa. University of California Press, Berkeley, pp. 161–251.
- California Press, Berkeley, pp. 161–251.Saxon, E.C., Close, A., Cluzel, C., Morse, V., Shackleton, N.J., 1974. Results of recent investigations at Tamar Hat. Lybica 22, 49–91.
- Schwarcz, H.P., Rink, W.J., 2000. ESR dating of the Die Kelders 1 site, South Africa. J. Hum. Evol. 38, 121–128.
- Schweitzer, F.R., Wilson, M.L., 1982. Byneskranskop 1, a late Quaternary living site in the southern Cape Province, South Africa. Ann. S. Afr. Mus. 88, 1–203.
- Scott, K.M., 1985. Allometric trends and locomotor adaptations in the Bovidae. Bull. Am. Mus. Nat. Hist. 179, 197–288.
- Seeley, H.G., 1891. On Bubalus bainii (Seeley). Geol. Mag. Lond. 8 (3), 199-202.
- Skead, C.J., 2011. Historical incidence of the larger land mammals in the broader Western and Northern Cape. Centre for African Conservation Ecology.Nelson Mandela Metropolitan University, Port Elizabeth.
- Smith, F.A., Lyons, S.K., Jones, K.E., Kaufman, D.M., Dayan, T., Marquet, P.A., Brown, J.H., Haskell, J.P., 2003. Body mass of late Quaternary mammals. Ecology 84, 3402.
- Smith, T.M., Tafforeau, P., Reid, D.J., Grün, R., Eggins, S., Boutakiout, M., Hublin, J.-J., 2007. Earliest evidence of modern human life history in North African early Homo sapiens. Proc. Natl. Acad. Sci. U. S. A. 104, 6128–6133.

- Steele, T.E., 2012. Late Pleistocene human subsistence in Northern Africa: the state of our knowledge and placement in a continental context. In: Hublin, J.-J., McPherron, S.P. (Eds.), Modern Origins: A North African Perspective. Springer, Dordrecht, pp. 107–125.
- Stokes, S., 1993. Optical dating of sediment samples from Bir Tarfawi and Bir Sahara East: an initial report. In: Wendorf, F., Schild, R., Close, A.E. (Eds.), Egypt During the Last Interglacial. Plenum Press, New York, pp. 229–233.
- Stuiver, M., Reimer, P.J., 1993. Extended 14C database and revised CALIB radiocarbon calibration program. Radiocarbon 32, 215–230.
- Stynder, D.D., 2009. The diets of ungulates from the hominid fossil-bearing site of Elandsfontein, Western Cape, South Africa. Quatern. Res. 71, 62–70.
- Suwa, G., Nakaya, H., Asfaw, B., Saegusa, H., Amzaye, A., Kono, R.T., Beyene, Y., Katoh, S., 2003. Plio-Pleistocene terrestrial mammal assemblage from Konso, Southern Ethiopia. J. Vertebr. Paleontol. 23, 901–916.
- Thackeray, J.F., 1979. An analysis of faunal remains from archaeological sites in southern south west Africa (Namibia). S. Afr. Archaeol. Bull. 34, 18–33.
- Thackeray, J.F., 1983. On Darwin, extinctions and South African fauna. Discovery 16 (2), 2–11.
- Thackeray, J.F., 1988. Zebras from Wonderwerk Cave, northern Cape Province: attempts to distinguish Equus burchelli and E. quagga. S. Afr. J. Sci. 84, 99–101.
- Thackeray, J.F., 1997. Morphometric, paleoecological and taxonomic considerations of southern African zebras: attempts to distinguish the quagga. S. Afr. J. Sci. 93, 89–93.
- Thackeray, J.F., Brink, J.S., 2004. Damaliscus niro horns from Wonderwerk Cave and other Pleistocene sites: morphological and chronological considerations. Palaeontol. Afr. 40, 89–93.
- Thackeray, A.I., Thackeray, J.F., Beaumont, P.B., Vogel, J.C., 1981. Dated rock engravings from Wonderwerk Cave, South Africa. Science 214, 64–67.
- Tryon, C.A., McBrearty, S., 2006. Tephrostratigraphy of the Bedded Tuff Member (Kapthurin Formation, Kenya) and the nature of archaeological change in the later Middle Pleistocene. Quatern. Res. 65, 492–507.
- Tryon, C.A., Faith, J.T., Peppe, D.J., Fox, D.L., McNulty, K.P., Jenkins, K., Dunsworth, H., Harcourt-Smith, W., 2010. The Pleistocene archaeology and environments of the Wasiriya Beds, Rusinga Island, Kenya. J. Hum. Evol. 59, 657–671.
- Tryon, C.A., Peppe, D.J., Faith, J.T., Van Plantinga, A., Nightengale, S., Ogondo, J., 2012. Late Pleistocene artefacts and fauna from Rusinga and Mfangano islands, Lake Victoria, Kenya. Azania Archaeol. Res. Afr. 47, 14–38.
- Turvey, S.T., 2009. In the shadow of the megafauna: prehistoric mammal and bird extinctions across the Holocene. In: Turvey, S.T. (Ed.), Holocene Extinctions. Oxford University Press, Oxford, pp. 17–40.
- van Hoepen, E.C.N., van Hoepen, H.E., 1932. Vrystaatse wilde varke. Palaeontol. Res. Natl. Mus. Bloemfontein 2, 39–62.
- Vermeersch, P.M. (Ed.), 2000. Palaeolithic Living Sites in Upper and Middle Egypt. Leuven University Press, Leuven.

- Vogel, J.C., 2001. Radiometric dates for the Middle Stone Age in South Africa. In: Tobias, P.V., Raath, M.A., Maggi-Cecchi, J., Doyle, G.A. (Eds.), Humanity from African Naissance to Coming Millennia. Florence University Press. Florence. pp. 261–268.
- Vrba, E., 1973. Two species of Antidorcas Sundevall at Swartkrans (Mammalia: Bovidae). Ann. Transv. Mus. 28. 287–352.
- Vrba, E.S., 1976. The fossil Bovidae of Sterkfontein, Swartkrans and Kromdraai. Transvaal Museum Memoir No. 21.Transvaal Museum, Pretoria.
- Vrba, E.S., 1980. The significance of bovid remains as an indicator of environment and predation patterns. In: Behrensmeyer, A.K., Hill, A.P. (Eds.), Fossils in the Making. University of Chicago Press, Chicago, pp. 247–272.
- Vrba, E.S., 1997. New fossils of Alcelaphini and Caprinae (Bovidae, Mammalia) from Awash, Ethiopia, and phylogenetic analysis of Alcelaphini. Paleontol. Afr. 34, 127–198.
- Wadley, L, 1991. Rose Cottage Cave: background and preliminary report on the recent excavations. S. Afr. Archaeol. Bull. 46, 125–130.
- Wadley, L., 2010. Were snares and traps used in the Middle Stone Age and does it matter? A review and case study from Sibudu, South Africa. J. Hum. Evol. 58, 179–192.
- Wendorf, F., 1977. Late Pleistocene and recent climatic changes in the Egyptian Sahara. Geogr. J. 143, 211–234.
- Wendorf, F., Schild, R., 1976. Prehistory of the Nile Valley. Academic Press, New York.
- Wengler, L, Weisrock, A., Brochier, J.-E., Brugal, J.-P., Fontigne, M., Magnin, F., Mathieu, J., Mercier, N., Ouammou, A., Reyss, J.-L., Senegas, F., Valladas, H., Wahl, L., 2002. Enregistrement fluviatile et paléoenvironnements au Pléistocène supérieur sur la bordure méridionale atlantique de l'Anti-Atlas (Oued Assaka, SO Marocain). Quaternaire 13, 179–192.
- Werdelin, L., Sanders, W.J. (Eds.), 2010. Cenozoic Mammals of Africa. University of California Press, Berkeley.
- White, T.D., Asfaw, B., DeGusta, D., Gilbert, H., Richards, G.D., Suwa, G., Howell, F.C., 2003. Pleistocene Homo sapiens from the Middle Awash, Ethiopia. Nature 423, 742–747.
- Williams, S.D., 2002. Status and action plan for Grevy's zebra (*Equus grevyi*). In: Moehlman, P.D. (Ed.), Equids: Zebras, Asses, and Horses, Status Survey and Conservation Action Plan. IUCN, Gland, Switzerland, pp. 11–27.
- Wolff, C., Haug, G.H., Timmermann, A., Sinninghe Damste, J.S., Brauer, A., Sigman, D.M., Cane, M.A., Verschuren, D., 2011. Reduced interannual rainfall variability in East Africa during the last Ice Age. Science 333, 743–747.
- Wrinn, P.J., Rink, W.J., 2003. ESR dating of tooth enamel from Aterian levels at Mugharet el 'Aliya (Tangier, Morocco). J. Archaeol. Sci. 30, 123–133.
- Wroe, S., Field, J., 2006. A review of the evidence for a human role in the extinction of Australian megafauna and an alternative interpretation. Quat. Sci. Rev. 25, 2692–2703.
- Wroe, S., Field, J., Fullagar, R., Jermin, L.S., 2004. Megafaunal extinction in the late Quaternary and the global overkill hypothesis. Alcheringa 28, 291–331.
- Wroe, S., Field, J.H., Archer, M., Grayson, D.K., Price, G.J., Louys, J., Faith, J.T., Webb, G.E., Davidson, I., Mooney, S., 2013. Climate change frames debate over the extinction of megafauna in Sahul (Pleistocene Australia-New Guinea). Proc. Natl. Acad. Sci. U. S. A. 110, 8777–8781.