


Late-Holocene faunal and landscape change in the Bahamas

The Holocene
2014, Vol. 24(2) 220–230
© The Author(s) 2014
Reprints and permissions:
sagepub.co.uk/journalsPermissions.nav
DOI: 10.1177/0959683613516819
hol.sagepub.com


David W Steadman,¹ Nancy A Albury,² Perry Maillis,³ Jim I Mead,⁴
John Slapcinsky,¹ Kenneth L Krysko,¹ Hayley M Singleton¹
and Janet Franklin⁵

Abstract

We report an intertidal, bone-rich peat deposit on the windward (Atlantic Ocean) coast of Abaco, The Bahamas. The age of the Gilpin Point peat (c. 950–900 cal. yr BP) is based on five overlapping radiocarbon dates (one each from single pieces of wood of buttonwood *Conocarpus erectus* and sabal palm *Sabal palmetto*, and single bones of the Cuban crocodile *Crocodylus rhombifer*, Albury's tortoise *Chelonoidis alburyorum*, and green turtle *Chelonia mydas*). The short time interval represented by the charcoal-rich peat suggests rapid sedimentation following initial anthropogenic fires on Abaco. The site's diverse snail assemblage is dominated by terrestrial and freshwater species. The peat is exposed today only during exceptionally low tides, suggesting a lower sea level at the time of deposition as well as a degrading shoreline during the past millennium. Fossils from Gilpin Point represent a late-Holocene vertebrate community at the time of first human presence; only 10 of the 17 identified species of amphibians, reptiles, birds, and mammals still live on Abaco. Numerous unhealed bite marks on the inside of the thick carapaces of the green turtle attest to consumption by Cuban crocodiles, which probably scavenged turtles butchered by humans. This concept, along with the dense concentration of bones in the peat, and charring on some bones of the green turtle and Abaco tortoise, suggests a cultural origin of the bone deposit at Gilpin Point, where the only Amerindian artifact recovered thus far is a shell bead.

Keywords

Bahamas, Holocene, landscape change, peat, snails, vertebrates

Received 30 September 2013; revised manuscript accepted 15 November 2013

Introduction

The late-Quaternary record of vertebrates on West Indian islands originates mainly from two very different sources of sediment. The first (and often richest) source occurs in karst features (dry or flooded limestone caves, rockshelters, and sinkholes) that accumulate bones most often through non-cultural means such as natural trap activity or predator roosts (e.g. Pregill, 1981; Pregill et al., 1994; Steadman et al., 2007). The second source is from open cultural (archaeological) sites where bones were deposited by prehistoric peoples (e.g. Jones O'Day, 2002; Steadman et al., 1984). In this article, we describe an unusual sedimentary setting for the deposition of ancient bones on a West Indian island, namely, a coastal peat deposit discovered on Great Abaco Island in the northern Bahamas. This peat has yielded an intriguing set of molluscan and vertebrate fossils that was deposited rapidly when humans first arrived in the northern Bahamas.

Site setting

The Bahamas consists of oceanic islands that lie just off the eastern coast of Florida (Figure 1). The archipelago features 23 major islands and many smaller ones lying on shallow carbonate banks separated by deep water. All exposed bedrock in this tectonically stable archipelago are Quaternary eolianitic subtidal limestone (Hearty and Kaufman, 2000; Mylroie, 2008). The Bahamian Archipelago stretches 980 km from c. 27°N and 79°W in the northwest to c. 21°N and 71°W in the southeast and consists politically of the independent Commonwealth of the Bahamas (or 'The Bahamas') and the Turks and Caicos Islands, a British Protectorate.

Our study focuses on Great Abaco Island (hereafter 'Abaco'), the second largest Bahamian island (1214 km²). Gilpin Point is an area of irregular limestone outcrops and calcareous beach sand on the windward (eastern, Atlantic Ocean) coast of southern Abaco (26.10457°N, 77.177666°W, Datum WGS84; Figure 2). The Gilpin Point site was discovered by Sabrina Bethel and landowner Perry Maillis on 8 and 10 May 2009, during a very low spring tide associated with a full moon on 9 May 2009. The dark, peaty sediment at Gilpin Point is inundated today by the ocean under normal circumstances, as well as covered by calcareous sand. Access to the sediment typically is possible only during very low tides (spring tides) associated with certain new and full moons during non-summer months (October–May). Even then, only the most exceptional spring tides are low enough to expose the sediment subaerially. PM also collected bones on 6 June 2009; 21, 22, 27, and 30 November 2009; 5 December 2009; sporadically from 28 September to 2 December 2012; 30 March 2013; and 10 May 2013. During our group's field work on 8 and 13 November 2012

¹University of Florida, USA

²The National Museum of the Bahamas, The Bahamas

³Gilpin Point, The Bahamas

⁴East Tennessee State University, USA

⁵Arizona State University, USA

Corresponding author:

David W Steadman, Florida Museum of Natural History, University of Florida, Gainesville, FL 32611, USA.

Email: dws@flmnh.ufl.edu

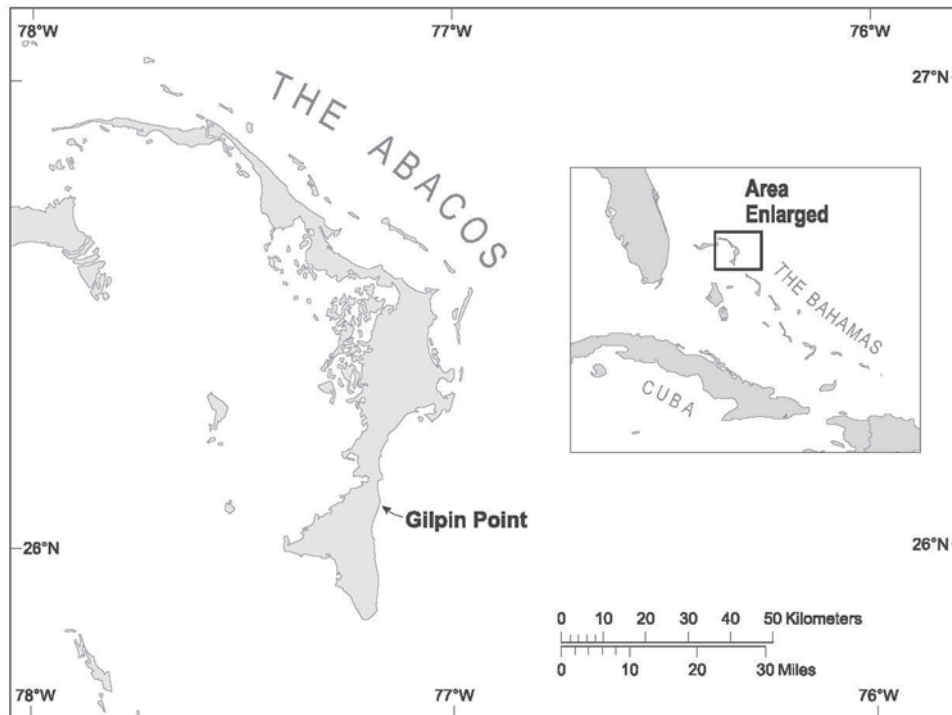


Figure 1. The Bahamian islands, showing the location of Abaco and other islands mentioned in the text.

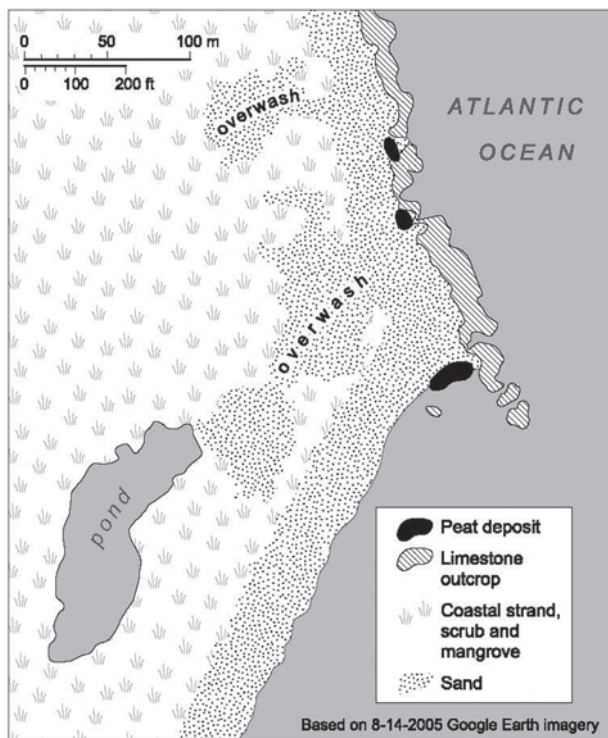


Figure 2. Gilpin Point, Abaco, The Bahamas, showing the location of the bone-bearing peat deposit.

(new moon on 13 November), the deposit remained under *c.* 10 cm of water even during the lowest spring tide. During another round of group field work on 1 December 2012 (full moon on 28 November), the deposit was exposed subaerially for 30 min during the lowest spring tide (Figure 3).

Aside from tidal cycles, another factor that limits access to the peat deposit is sand transport. For example, during our visit to Gilpin Point on 10 June 2013 (new moon on 8 June), the site was covered by sand at least 1 m deep. Prevailing easterly winds

off the Atlantic Ocean tend to build up sand during the warm months, as well as inhibit extremely low spring tides. During the cooler months, when Abaco receives north winds from periodic frontal systems, sand is more often removed from this section of shoreline, thereby exposing the peat layer during extremely low spring tides. The peat can be exposed and then completely covered by sand over a single tidal cycle.

Methods

Because the peat deposit faces the open ocean on a high-energy coast, it would be prohibitively complicated and expensive to build a break-wall to improve access to the site. Therefore, our collection methods at Gilpin Point consisted of removing blocks of the peat (working quickly because of the narrow time interval of the lowest tidal level) with a square shovel (spade; Figure 4). These blocks of sediment, each overlain and underlain by calcareous sand, measured from 14 to 20 cm wide, 10 to 16 cm long, and 10 to 22 cm deep (*c.* 1.5–6 L per sample). For the three largest species of vertebrates recovered (*Crocodylus rhombifer*, *Chelonoidis alburyorum*, *Chelonia mydas*), *c.* 60% of the fossils were collected in situ because of being partially exposed at the top of the peat. Fossils of the other, smaller species were collected by screen-washing *c.* 0.2 m³ (200 L) of sediment. We processed *c.* 50% of this sediment in the field through nested sieves of 12.5, 6.4, and 1.6 mm mesh; the remainder of the sediment was processed in the laboratory through five nested sieves of 12.5, 5.6, 3.35, 2.0, and 1.0 mm mesh. Snails and small bones occurred throughout the vertical sequence of sediment.

The fossils from Gilpin Point are housed in the scientific collections of the National Museum of the Bahamas in Marsh Harbour, Abaco, under the care of NAA. Some of the fossils are currently on loan to DWS at the Florida Museum of Natural History, University of Florida (UF). Identifications of the fossils are based on direct comparisons with modern specimens of mollusks and vertebrates in the UF collections. In 'Results' section, we provide the common, scientific, and family name for each species at its first mention.



Figure 3. Peat deposit exposed, 1 December 2012, Gilpin Point, Abaco, The Bahamas. DWS and PM pictured. Photo by JF.



Figure 4. DWS collecting blocks of peat with a square spade during the lowest spring tide, 1 December 2012, Gilpin Point, Abaco, The Bahamas. Photo by JF.

Results

Sediment description

The sediment at Gilpin Point is poorly stratified, variably silty, and sandy peat (Table 1). The organic matter consists of charcoal (pieces ranging in size from microscopic to 14 mm diameter), unburned plant material, and bones. The peat deposit is 20–22 cm thick. We found no vertical trends in any characteristics of the peat. The peat deposit is much less organic and has considerably fewer mollusks and bones in the two small outcrops to the north of the main collection area, which is the largest and southernmost

black area in Figure 2. All five sediment samples are from the main collection area; the two that are the least organic (samples 2 and 4) are from the northern margin of the main collection area. We did not quantify the charcoal other than to point out that it dominates the organic matter combusted to generate the loss on ignition (LOI) data (Table 1). The charcoal occurred throughout the deposit; we found no rich concentrations that would suggest a cultural feature such as a hearth. The deposit contained no high-energy overwash sediments of marine origin.

Chronology

Our age estimate for the peat deposit at Gilpin Point is based on five radiocarbon (^{14}C) dates from individual specimens of plants and animals identified to species (Table 2). The four accelerator-mass spectrometer (AMS) ^{14}C dates consist of one on wood of the buttonwood *Conocarpus erectus* (Combretaceae) and three on purified collagen from individual bones of the Cuban crocodile *Crocodylus rhombifer* (Crocodylidae), the extinct Albury's tortoise *Chelonoidis alburyorum* (Testudinidae), and green turtle *Chelonia mydas* (Cheloniidae). The single conventional ^{14}C date is from a large sample of wood of the sabal palm *Sabal palmetto* (Arecaceae). The two ^{14}C dates on wood overlap (at 2σ) at 920–900, 860–830, 810–800 cal. yr BP, whereas the three ^{14}C dates on bone overlap (at 2σ) at 950–920 cal. yr BP. Both the buttonwood and palm were found in an upright growth position (Figure 5), suggesting that they may have continued to live after deposition of the bones. This mechanism would explain the slightly younger age determinations for the wood samples than the bone samples.

With a ^{14}C age of 970–920 cal. yr BP, Beta-338510 is the youngest direct age determination for the Cuban crocodile in the Bahamas and therefore helps to pinpoint the time of its extirpation on Abaco. The earliest ^{14}C age previously determined directly on a Cuban crocodile bone in the Bahamas was 2880–2750 cal. yr BP (Beta-225508) from Sawmill Sink blue hole, Abaco (Steadman et al., 2007). (A blue hole is a flooded cave or sinkhole.) Nevertheless, crocodile bite marks exist on an Albury's tortoise specimen from Sawmill Sink that has a ^{14}C age of 970–920 cal. yr BP (Beta-298219; Hastings et al., in press), which is identical to our age determination for the crocodile from Gilpin Point. With a ^{14}C age of 960–910 and 840–840 cal. yr BP (Beta-338511), the Albury's tortoise from Gilpin Point overlaps with the two youngest previous age determinations for this species, namely, 970–920 cal. yr BP (Beta-298219; Sawmill Sink) and 920–780 cal. yr BP (Beta-298220; Lost Reel Cave, Abaco; Hastings et al., in press).

The 'marine reservoir effect' reflects the fact that marine organisms and ocean waters are depleted in ^{14}C (i.e. contain 'dead' carbon) and therefore may yield ^{14}C ages that are older than those based on contemporary terrestrial organic materials (Goodwin, 1993). The precise range of the surface marine reservoir correction for the Bahamas is not known, so we used an approximation of 350 years (from global estimates in Stuiver and Braziunas, 1993) to correct the AMS ^{14}C date for the green turtle.

Stable isotopes

The carbon stable isotope data (^{13}C values) of the ^{14}C -dated samples (Table 2) are reasonable for each species, given what is known about their biology. Like most woody plants, the buttonwood and sabal palm follow the C_3 (Calvin) photosynthetic pathway; their determined ^{13}C values (–28.0‰ and –28.4‰, respectively) are as expected for C_3 plants (average –28.5‰, with a typical range of –37‰ to –20‰; Kohn, 2010; O'Leary, 1981, 1988).

The Cuban crocodile was a terrestrial carnivore in the Bahamas (Morgan and Albury, 2013; Steadman et al., 2007); the ^{13}C value for crocodile from Gilpin Point (–19.8‰) falls within the values reported elsewhere for modern and prehistoric terrestrial

Table 1. Description of sediment samples from Gilpin Point. Fraction organic matter based on loss on ignition for 2 h at 550°C.

Sediment sample no.	Sediment type	Color		Fraction organic matter
		Dry	Wet	
1	Peaty, slightly silty, angular fine calcareous sand	5YR 7/1; light gray	5YR 3/1; very dark gray	0.25
2	Slightly peaty, slightly sandy silt	5YR 8/1; white	7.5YR 6/2; pinkish gray	0.12
3	Peaty, silty, angular fine calcareous sand	5YR 5/1; gray	5YR 3/1; very dark gray	0.22
4	Peaty, silty, angular fine to medium calcareous sand	5YR 6/1; gray	5YR 4/1; dark gray	0.11
5	Very peaty, slightly silty, angular fine to medium calcareous sand	5YR 3/1; very dark gray	7.5YR 2/0; black	0.30

Table 2. Radiocarbon (^{14}C) chronology for the peat deposit at Gilpin Point, Abaco, Bahamas. All determinations (done at Beta Analytic, Inc., Miami, FL), except Beta-345519, are accelerator-mass spectrometer (AMS) dates. Calibration of radiocarbon ages to calendar dates follows the IntCal9 database (Heaton et al., 2009; Reimer et al., 2009). For further details of laboratory and calibration methods, see www.radiocarbon.com. The three ^{14}C dates on bone are on ultrapurified collagen. For Beta-338512, the ^{14}C age in brackets is uncorrected for the marine reservoir effect; the ^{14}C age without brackets considers an approximate marine surface reservoir correction of 350 years, following Figures 5A and 15A in Stuiver and Braziunas (1993).

Material dated	Sample weight (as submitted; g)	Sample number	$\delta^{13}\text{C}$ (‰)	Conventional age	^{14}C age (cal. yr BP, 2σ)
Wood (<i>Sabal palmetto</i>)	50.8	Beta-345519	-28.0	990 \pm 30 BP	960–900, 860–830, 810–800
Wood (<i>Conocarpus erectus</i>)	3.9	Beta-338518	-28.4	900 \pm 30 BP	920–740
Bone (postorbital of <i>Crocodylus rhombifer</i>)	5.0	Beta-338510	-19.8	1020 \pm 30 BP	970–920
Bone (left first costal of <i>Chelonoidis alburysorum</i>)	7.8	Beta-338511	-21.6	1010 \pm 30 BP	960–910, 840–840
Bone (left first costal of <i>Chelonia mydas</i>)	15.6	Beta-338512	-9.6	1340 \pm 30 BP	[1300–1260, 1200–1190] 950–910, 850–840

carnivores (c. -21% to -17% ; Bocherens and Drucker, 2003; Coltrain et al., 2004) and the values for Holocene Cuban crocodile fossils from other sites on Abaco (-20.9% to -16.4% , $N = 8$; Hastings et al., in press). The Abaco tortoise was a terrestrial herbivore believed to have consumed mainly C_3 plants (Franz and Franz, 2009; Steadman et al., 2007); the ^{13}C value for tortoise from Gilpin Point (-21.6%) falls within the values reported elsewhere for modern and prehistoric terrestrial herbivores (c. -24% to -19% ; Bocherens and Drucker, 2003; Coltrain et al., 2004), including Holocene tortoise fossils from elsewhere on Abaco (-22.9% to -21.1% , $N = 5$; Hastings et al., in press).

In the Greater Caribbean Region, the green turtle is a marine herbivore as an adult, although juveniles can be omnivorous or carnivorous (Bjorndal, 1997). Adult green turtles (females only) come ashore only to nest on sandy beaches. The ^{13}C value for the dated green turtle bone from Gilpin Point (-9.6%) falls within the values reported for modern specimens from Long Island, Bahamas (-12.2% to -6.4% ; Vander Zanden et al., 2013).

Plants

The peat at Gilpin Point contains abundant wood, often in nearly vertical growth position, of buttonwood (*C. erectus*). The tops of dead buttonwood branches were also exposed in growth position on the sandy beach near the present shoreline c. 25 m north of where the peat was excavated (Figure 5), suggesting that the peat deposit extends at least this far northward. Buttonwood has lower tolerance to soil salinity than other mangroves (Medina, 1999; Rada et al., 1989). In the Bahamas, buttonwood typically is found inland from the three other species of mangroves (black mangrove *Avicennia germinans*, Acanthaceae; white mangrove *Laguncularia racemosa*, Combretaceae; red mangrove *Rhizophora mangle*, Rhizophoraceae) and is associated with slightly brackish or freshwater wetlands (Areces-Mallea et al., 1999).



Figure 5. Black trunks of *Conocarpus erectus* in growth position in water on left side, originating in peat deposit, 13 November 2012, Gilpin Point, Abaco, The Bahamas. Photo by KLK.

The stump of the ^{14}C -dated sabal palm (*S. palmetto*) also was found in growth position. In the Bahamas, sabal palm tolerates poorly drained soils and is associated with ephemeral freshwater ponds and areas with subterranean freshwater lenses near the surface (Areces-Mallea et al., 1999; Sullivan-Sealey et al., 2002). Sabal palm is prone to mortality when coastal flooding from sea-level rise exposes it to increasing salinity (DeSantis et al., 2007; Perry and Williams, 1996). Such flooding may have killed the individual that was ^{14}C -dated, suggesting that this inundation took place no later than c. 800 cal. yr BP.

Mollusks

The sediment at Gilpin Point contains abundant remains of small gastropods as well as much rarer small bivalves. The molluscan assemblage was analyzed by JDS and JIM in detail from a single

Table 3. Snails identified from a single sediment sample at Gilpin Point, Abaco, The Bahamas. B: bivalve; all others are gastropods. Macrohabitat categories – E: estuarine; F: freshwater; M: marine; T: terrestrial.

Taxon	Count	Macrohabitat	Microhabitat
Assimineidae	31	E, M	Marine/brackish ponds and mangrove areas
<i>Brachidontes exustus</i> (B)	1	M	Protected creeks, exposed shores, seagrass beds, under rocks
<i>Ctenocardia guppyi</i> (B)	2	M	Sand bottoms, seagrass beds
<i>Cenchritys muricatus</i>	1	M	Exposed rocky coasts on supratidal rock surfaces
<i>Cerion</i> sp.	1	T	Leaf litter and on trees near the coast
<i>Cerithidea costata</i>	53	E	Mangroves and pond edges, exposed on mud and sand
<i>Cerithium lutosum</i>	26	E, M	Marine/brackish ponds, coves and mangrove areas on algae, coral, rock, sand, and seagrass beds
<i>Chione elevata</i> (B)	1	M	Muddy sand, seagrass beds
<i>Crenella divaricata</i> (B)	1	M	Silty sand in high intertidal
<i>Drepanotrema cimex</i>	2	F	Eutrophic ponds or slow-moving water on vegetation
<i>Eoacmaea pustulata</i>	1	M	Under rocks or on seagrass
<i>Ervilia concentrica</i>	2	M	Intertidal or shallow sand
<i>Gastrocopta</i> cf. <i>pentodon</i>	14	T	Open grassy areas under leaf litter and stones
<i>Gemma gemma</i> (B)	3	E, M	Sandy bottoms, brackish ponds and creeks
<i>Hawaiiia minuscula</i>	13	T	Leaf litter in open forests and grasses
<i>Helicina rawsoni</i>	4	T	Forests on trees and in leaf litter
<i>Hemitrochus varians</i>	1	T	Open forest on trees and in leaf litter
<i>Littoridinops</i> cf. <i>tenuipes</i>	366	E, F	Fresh and brackish ponds
<i>Echinolittorina</i> cf. <i>angustior</i>	3	M	Upper intertidal on exposed rocky coasts
<i>Littoraria angulifera</i>	1	M	Supratidal on mangroves
<i>Melampus bidentatus</i>	5	E, M	Edge of brackish ponds and creeks at high tide mark
<i>Melampus bullaoides</i>	1	E, M	Under rocks at high tide line in protected creeks
Oleacinidae	6	T	Unknown, presumed to be in leaf litter or under rocks in forests
<i>Olivella</i> sp.	3	M	Buried in open sand or sand pockets
<i>Physella</i> sp. 1	98	F	Eutrophic ponds or slow-moving water on vegetation
<i>Physella</i> sp. 2	159	F	Eutrophic ponds or slow-moving water on vegetation
<i>Planorbidae</i> sp.	1	F	Eutrophic ponds or slow-moving water on vegetation
<i>Polinices lacteus</i>	1	M	Sand or sand pockets
<i>Polygyra plana</i>	22	T	Open grassy areas, open forest
<i>Strobellops hubbardi</i>	35	T	Leaf litter in moist forest
<i>Succinea</i> cf. <i>ochracina</i>	5	T	Dry, open grassy areas
<i>Timoclea pygmaea</i> (B)	1	M	Sand bottoms, seagrass beds
<i>Transennella cubaniana</i> (B)	3	M	Sand bottoms
<i>Varicella gracillima bahamensis</i>	1	T	Woodlands under leaf litter and rocks
<i>Vertigo</i> cf. <i>ovata</i>	2	T	Wet woodlands and grasslands
<i>Zonitoides arboreus</i>	11	T	Forests and forest edges under wood and in leaf litter

3-L sediment sample taken from the area of densest bone concentration. Of 36 taxa recognized, 12 are terrestrial, 4 are freshwater, 1 is freshwater/estuarine, 2 are estuarine, 4 are estuarine/marine, and 13 are marine (Table 3). The majority of the most commonly occurring species are snails that require terrestrial or freshwater habitats. We interpret this trend in habitat preference to mean that the main sources of snails in the peat deposit originated inland (washing in with the charcoal) or in situ (in the freshwater or slightly brackish swamp).

The species of mollusks in Table 3 are minute and do not include the large, edible, marine species (such as conchs, whelks, and top shells) that often are abundant in Amerindian middens in the Bahamas and southern coastal Florida (Jones O'Day, 2002; Keegan et al., 2008; Marquardt, 2010a, 2010b). We note, however, that a single specimen of Queen conch (*Strombus gigas*, Strombidae) and two of the West Indian top shell (*Cittarium pica*, Tegulidae) were recovered in situ alongside bones of the green turtle, only c. 2 m south of the snail sediment sample. These three specimens, each recovered in a broken condition typical of Amerindian middens (see Jones O'Day and Keegan, 2001), constitute the only mollusks from the Gilpin Point site that are likely to have been consumed or used as tools by prehistoric peoples.

Vertebrates

We identified 20 species of vertebrates from the fossils at Gilpin Point (Table 4). The five identifiable fish bones represent three species, the bonefish *Albula vulpes* (Albulidae; 2 vertebrae), bar jack *Caranx ruber* (Carangidae; palatine), and midnight parrotfish *Scarus coelestinus* (Scaridae; atlas, hypural). Each of these edible estuarine or marine species occurs regularly in West Indian archaeological contexts (Newsom and Wing, 2004).

Of the 17 species of amphibians, reptiles, birds, and mammals found in the Gilpin Point material, we first will document the seven that no longer occur on Abaco. The Cuban crocodile *Crocodylus rhombifer* is represented by c. 60 cranial and post-cranial bones (e.g. Figure 6) that depict two individuals (an adult and a sub-adult). Albury's tortoise *Chelonoidis alburyorum* is represented by c. 60 post-cranial bones that are dominated by pieces of the carapace and plastron of a single adult individual with apparently healed bite marks from the Cuban crocodile on the outside of its carapace (Figure 6). Five post-cranial elements are referred to the rock iguana *Cyclura* cf. *carinata* (Iguanidae), another rather large, edible species (Carlson and Keegan, 2004; Keegan and DeNiro, 1988).

The Gilpin Point material of Bermuda petrel *Pterodroma cahow* (Procellariidae) consists of a nearly complete coracoid and

Table 4. Species of vertebrates identified from late-Holocene fossils at Gilpin Point, Abaco, Bahamas. 'Status on Abaco' categories – EA: extirpated on Abaco, but still exists elsewhere; EG: extinct globally; SE: still exists on Abaco. 'Habitat preference' categories: E: estuarine; F: freshwater; M: marine; T: terrestrial. *Chelonia mydas* and *Pterodroma cahow* are regarded as both marine and terrestrial because they most must come ashore to nest. Numbers in brackets represent the minimum number of individuals represented by the fossils of the five most common species. The species names for *Leiocephalus*, *Cyclura*, *Chilabothrus*, and *Cubophis* are based on the modern species that inhabit Abaco or nearby islands; the fossils from Gilpin Point for these three genera are diagnostic to genus but not to species.

Scientific name	Common name	Status on Abaco	Number of fossils	Habitat preference
Fish				
<i>Albula vulpes</i>	Bonefish	SE	2	E, M
<i>Caranx ruber</i>	Bar jack	SE	1	E, M
<i>Scarus coelestinus</i>	Midnight parrotfish	SE	2	M
Amphibians				
<i>Osteopilus septentrionalis</i>	Cuban tree frog	SE	1	T
Reptiles				
<i>Crocodylus rhombifer</i>	Cuban crocodile	EA	~60 [2]	E, F, T
<i>Chelonoidis alburyorum</i>	Albury's tortoise	EG	~60 [1]	T
<i>Chelonia mydas</i>	Green turtle	SE	~80 [2]	M, T
Gekkonidae sp.	Unknown small gecko	SE	1	T
<i>Leiocephalus cf. carinatus</i>	Bahamian curly tailed lizard	SE	1	T
<i>Anolis cf. distichus</i>	Bahamian bark anole	SE	1	T
<i>Anolis cf. sagrei</i>	Cuban anole	SE	3	T
<i>Cyclura cf. carinatus</i>	Bahamian rock iguana	EA	5	T
<i>Chilabothrus cf. exsul</i>	Bahamian boa	SE	2	T
<i>Cubophis cf. vudii</i>	Bahamian racer	SE	12 [1]	T
Birds				
<i>Pterodroma cahow</i>	Bermuda Petrel	EA	3	M, T
<i>Eudocimus albus</i>	White ibis (juvenile)	EA	2	E, F
<i>Nyctanassa violacea</i>	Yellow-crowned night-Heron	SE	1	E
Rallidae sp.	Unknown small rail	SE	1	E, F
<i>Corvus nasicus</i>	Cuban crow	EA	8 [2]	T
Mammals				
<i>Geocapromys ingrahami</i>	Bahamian hutia	EA	1	T

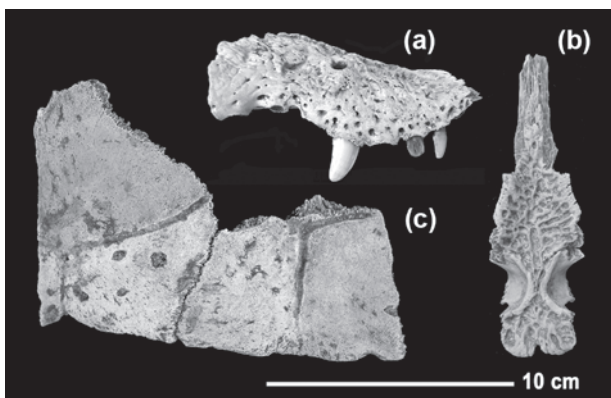


Figure 6. (a and b) Specimens of Cuban crocodile (*Crocodylus rhombifer*) and (c) Albury's tortoise (*Chelonoidis alburyorum*) from Gilpin Point, Abaco, The Bahamas. (a) Lateral view of right maxilla (NMB.AB62.12), (b) dorsal view of frontal and parietal (NMB.AB62.11), and (c) dorsal view of nuchal and left peripheral I (NMB.AB62.1). Photo by NAA.

two fragmentary humeri. These specimens agree with skeletal elements of *P. cahow* rather than those of the confamilial Audubon's Shearwater *Puffinus lherminieri* (extant in the Bahamas) in these characters: larger overall size; medial surface of humeral end of coracoid singly concave (doubly concave in *P. lherminieri*); in medial or lateral aspect, procoracoid less flared dorsally; and humeral shaft less compressed (more circular in cross-section). The length of the coracoid (23.06 mm) agrees with that of fossils of *P. cahow* from Bermuda (22.5–23.7 mm, $N = 15$; Olson and Hilgartner, 1982).

The two pedal phalanges of white ibis *Eudocimus albus* (Threskiornithidae) from Gilpin Point represent nestlings or very young fledglings of a species that no longer breeds in the Bahamas. Eight fossils (sternum, radius, two carpometacarpi, four pedal phalanges) of the Cuban crow *Corvus nasicus* (Corvidae) occur at Gilpin Point. They represent the first record of *C. nasicus* from the Little Bahama Bank and the northernmost occurrence of species today confined to Cuba and the Caicos Islands. Finally, among the extirpated species, the Bahamian hutia *Geocapromys ingrahami* (Capromyidae) is recorded from Gilpin Point by a single molar. Common as a late-Quaternary fossil on many Bahamian islands (Morgan, 1989), *G. ingrahami* now survives only on several very small islands (see 'Discussion' section).

In all, 10 species of terrestrial vertebrates found in the Gilpin Point material certainly or probably still exist on Abaco. A single humerus represents the Cuban tree frog *Osteopilus septentrionalis* (Hylidae). The green turtle *C. mydas* is the most commonly occurring species of vertebrate in the deposit, represented by c. 80 bones of the carapace, plastron, and limbs from two very large individuals. Because unhealed bite marks from a crocodile are found on the inside (rather than outside) of the carapace of one individual (Figure 7), we interpret the bite marks to represent scavenging of a green turtle that people had butchered. Being so large, adult green turtles likely were butchered near the beach prehistorically and then shared among multiple households so that the bones of a single individual might be deposited in more than one midden (Newsom and Wing, 2004). With further exploration, perhaps more remains of fish and shellfish will be found at Gilpin Point.

A single caudal vertebra has the morphology of a gecko (Gekkonidae) but is not diagnostic beyond family level. A single genus of geckos (*Sphaerodactylus*) occurs on Abaco today (Schwartz and Henderson, 1991). A complete tibia is referred to the curly

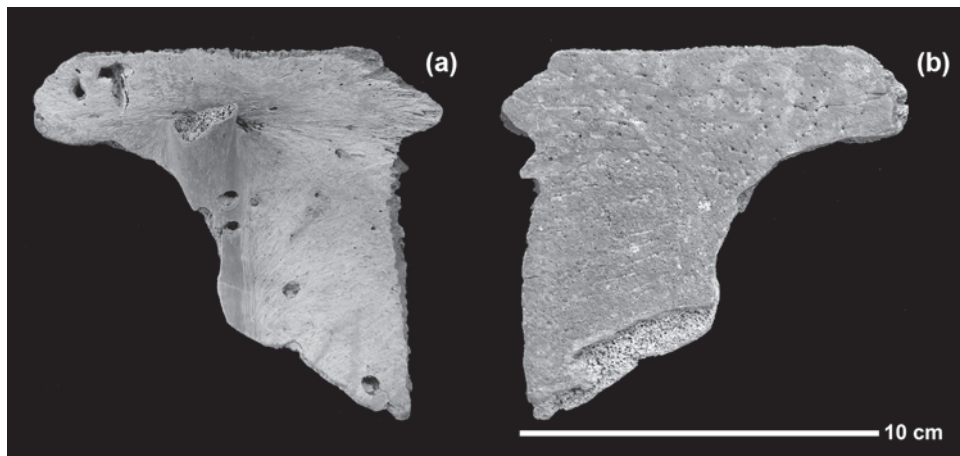


Figure 7. Specimen of green turtle (*Chelonia mydas*; NMB.AB62.7) from Gilpin Point, Abaco, The Bahamas, in dorsal (a) and ventral (b) aspects. Photo by NAA.

tailed lizard *Leiocephalus carinatus* (Leiocephalidae). This fossil tibia is 19.83 mm long, compared with 17.33 mm for the tibia of an adult modern specimen of *L. carinatus* from Cuba with a snout-vent length (SVL) of 98 mm and total length (TL) of 226 mm. Thus, the prehistoric curly tailed lizard from Gilpin Point had a SVL of *c.* 112 mm and a TL of *c.* 259 mm.

Four specimens are identified as anoles (*Anolis*, Polychrotidae), which are diverse and widespread in the West Indies, including the Bahamas (Henderson and Powell, 2009). One dentary and two maxillae of *Anolis* cf. *sagrei* are recorded from Gilpin Point by these characters: central cusp low, wide, and not excessively higher than marginal cusps; dentary robust and wide. This species exists in Abaco today. Another dentary is identified as *Anolis* cf. *distichus* because the diagnostic sharply pointed central cusp is distinctly higher than the well-pointed marginal cusps (see Pregill, 1982). Buckner et al. (2012) questioned whether or not *A. distichus* was historically introduced to Abaco; the Gilpin Point dentary is evidence that *A. distichus* is indeed indigenous to the island.

A single caudal vertebra and an edentulous fragment of dentary from a boa *Chilabothrus* cf. *exsul* (Boidae) are noticeably larger than that in the trope (dwarf boa) *Tropidophis* (Tropidophiidae). A mid-trunk vertebra is needed for more precise identification. The most commonly recovered small reptile from Gilpin Point is the snake (Bahamian racer) *Cubophis* cf. *vudii* (Dipsadidae; 12 specimens), identified by mid-trunk vertebrae with long, thin neural spines, elliptical prezygapophyses, and a thin, straight hemal keel with slight swelling and overhang at the condyle.

The yellow-crowned night-heron *Nyctanassa violacea* (Ardeidae) is a common resident species in the Bahamas, both today and as a fossil (Steadman et al., 2007; White, 1998). Our identification of a small rail (Rallidae sp.) is based on a fragmentary tibiotarsus of a size that may represent either the sora *Porzana carolina* or Virginia rail *Rallus limicola*; the specimen lacks diagnostic species-level characters. Both species presently occur in the Bahamas as non-breeding birds in freshwater and estuarine habitats.

Discussion

Landscape change

The setting and contents of the coastal peat deposit at Gilpin Point demonstrate how dynamic the Bahamian physical landscape can be over geologically short time frames. At both local and regional (inter-island) scales, these low islands can be affected dramatically by changes in sea level. The estimated sea level in the eastern Atlantic at 950 cal. yr BP (when humans arrived on Abaco)

was ~20 cm lower than at present but was rising in response to the 'Medieval Warm Period' (Curtis et al., 1996; Keigwin, 1996; Kemp et al., 2011; Walker, 2013). Within a century or so, the shorelines degraded, inundating any coastal habitation sites that had been located on beaches seaward from the buttonwood and sabal palm wetlands at Gilpin Point. This wetland was also affected by the encroaching saltwater in at least two ways. First, the increased salinity and direct contact with the ocean would have made the wetland uninhabitable by buttonwood and sabal palm. Second, the peat deposit would have begun to get covered by beach sand.

We found no high-energy overwash sediment within the peat deposit, that is, no coarse sandy to pebbly to cobbly storm-surge facies such as those found in sediment cores in saline lakes on San Salvador (Park, 2012). The short interval of time represented by the peat is compatible with rapid sedimentation from inland areas following initial anthropogenic fires on Abaco. The time of deposition of peat at Gilpin Point corresponds with when people first arrived on Abaco (Steadman et al., 2007). Prehistoric deforestation, accompanied by fire, may have increased the rate of sediment deposition in Abaco's wetlands, which has been reported independently on Abaco from the sediments of Emerald Pond blue hole (Slayton, 2010). The charcoal at Gilpin Point also corresponds in time with charcoal reported in sediments from a blue hole on Andros (Kjellmark, 1996). Similar increases in charcoal-rich sediment influx, believed to be associated with human activity, have been reported on other Caribbean islands and even the Atlantic coast of North America (e.g. Burney et al., 1994; Hilgartner and Brush, 2006; Horn et al., 2000; Lane et al., 2009). At Gilpin Point, the rapid deposition of peat (in less than a century, from *c.* 950 to 900 cal. yr BP) was followed by breaching of the former coastline (at least 200 m east of the modern one) and deposition of beach sand over the peat.

Aside from a lower sea level, deposition of the peat was enhanced by the limestone outcrop that exists just seaward from the peat deposit. This formerly continuous outcrop, which once would have served as a dam for the charcoal-rich sediment washing in from inland during deposition of the Gilpin Point site, has been broken and breached in several places in the past two decades during storms (e.g. Figures 5 and 8; PM, personal observation).

Faunal change

With a late-Holocene age of ~950–900 cal. yr BP, the fossils from Gilpin Point represent a vertebrate community at the time of initial Amerindian (Lucayan Taíno) presence; 7 of its 17 species



Figure 8. Section of limestone ridge broken by wave action, 13 November 2012, Gilpin Point, Abaco, The Bahamas. Note as well the black trunks of *Conocarpus erectus*. Photo by KLK.

(41%) of amphibians, reptiles, birds, and mammals no longer live on Abaco. Among well-dated assemblages of West Indian vertebrates similar in age to those of Gilpin Point, much lower percentages of extirpated species are found on islands where humans arrived in the middle Holocene. For example, at the Trouing Jean Paul site in Hispaniola (^{14}C -dated from ~1600 to 600 cal. yr BP), only 1 of 23 species (4%) of birds identified by Steadman and Takano (2013) no longer occurs on the island. Amerindians had arrived in Hispaniola four or five millennia earlier and already had extirpated the most vulnerable species of vertebrates (Steadman et al., 2005). A similar situation took place on some Pacific islands, where prehistoric human colonization led to huge losses of species within a century or two (Steadman et al., 2002). The species that survived the initial shock of human presence were likely to persist into modern times.

Four of the seven extirpated species (Cuban crocodile, Albury's tortoise, white ibis, and Bahamian hutia) have been reported previously from other Holocene sites on Abaco (Franz and Franz, 2009; Franz et al., 1995; Morgan and Albury, 2013; Steadman et al., 2007). The three other extirpated species (Bahamian rock iguana, Bermuda petrel, Cuban crow) are new records for the island.

The morphology, systematics, and paleoecology of the Cuban crocodile and Albury's tortoise on Abaco have been reviewed recently (Franz and Franz, 2009; Morgan and Albury, 2013; Steadman et al., 2007). Until their extermination shortly after human arrival, these two species were the largest terrestrial carnivore and herbivore on Abaco, respectively. The AMS ^{14}C date on Cuban crocodile from Gilpin Point is the youngest direct age determination on the species and provides additional evidence that the species existed on Abaco when Amerindians colonized the island. The Bahamian rock iguana is unknown on Abaco in modern times. We suspect that it was a food item for Amerindians on Abaco, as has been well documented on Grand Turk Island (Carlson and Keegan, 2004). Late-Quaternary fossils of rock iguana are also reported from Banana Hole, New Providence Island (Pregill, 1982).

The Bermuda petrel breeds today only on Bermuda, where it is common as a fossil (Olson and Meylan, 2009). This seabird was believed to have been extinct for several centuries until its rediscovery in 1951 and subsequent increase through intensive conservation efforts (Howell, 2012: 176–178; Wingate, 1972). An endangered species, the Bermuda petrel, does not occur in the Bahamas today, although six prehistoric bones were reported from Crooked Island (Olson and Hilgartner, 1982).

The white ibis has been recorded uncommonly in small numbers on several Bahamian islands (but not Abaco) in modern

times but is not known to breed in the island group (White, 1998: 234). Both white ibis fossil specimens (pedal phalanges) from Gilpin Point represent nestlings or very young fledglings. Fossils of nestling and fledgling white ibises also occur in the late-Holocene sediments of Sawmill Sink on Abaco (Steadman et al., 2007), further substantiating that this large aquatic bird once bred on the island.

Nowadays in the Bahamian archipelago, the Cuban crow lives only in the Caicos Islands, far to the south of Abaco (Buden, 1987). Its main geographic range is on Cuba and Isle of Pines. The Cuban crow is known from late-Quaternary fossils on New Providence Island, Little Exuma Island, Crooked Island, and Eleuthera Island (Olson and Hilgartner, 1982; Steadman, unpublished data); the fossils from Gilpin Point represent its northernmost occurrence and its first record from any island on the Little Bahama Bank. They also demonstrate that the Cuban crow was still widespread in the Bahamas during the late Holocene.

The Bahamian hutia also was formerly widespread in the Bahamas but survives on tiny East Plana Cay (28 km east of Acklins Island) with introduced populations on certain islands in the Exuma Chain (Borroto-Páez et al., 2012; Morgan, 1989; Olson and Pregill, 1982). As with the Cuban crow (above), the range contraction of the Bahamian hutia was mostly a late-Holocene phenomenon.

Among extant species, the green turtle is a widespread marine herbivore that is well represented at archaeological sites throughout the West Indies (Newsom and Wing, 2004: 63). Adult female green turtles are vulnerable to predation by people when they come ashore to nest on sandy beaches. Our identification of *A. cf. distichus* is evidence that this anole was not introduced to Abaco in modern times but is indigenous to the island (contra Buckner et al., 2012). The Bahamian boa and Bahamian racer occur on Abaco today (Buckner et al., 2012; Krysko et al., 2013). Both of these snakes also were recovered from the late Pleistocene owl roost deposit in Sawmill Sink, Abaco, with the boa *C. exsul* reported as *Epicrates striatus* and the racer *C. vudii* reported as *Alsophis* sp. (Steadman et al., 2007; our updated nomenclature follows Hedges et al., 2009, and Reynolds et al., 2013).

Cultural origin of the bone deposit

Numerous unhealed bite marks on the thick carapaces of the green turtle are evidence of consumption by Cuban crocodiles. The fact that the bite marks are well developed on the insides of the carapaces but not evident on the outer surfaces, suggests that the crocodiles scavenged butchered green turtles rather than preying on living turtles. Humans are the only terrestrial predators in the Bahamas capable of splitting open the shell of an adult green turtle to expose the meat and organs inside the carapace and plastron. Human involvement in deposition of bones at Gilpin Point is supported further by the dense, midden-like concentration of large bones (crocodile, green turtle, and tortoise) in the peat, and the fact that some bones of both the green turtle and Abaco tortoise are charred (burned). Berman et al. (2013) suggested that Lucayans likely cooked/roasted sea turtles on their back in a hearth, which is compatible with the charring on the outside of the carapaces.

Other species in the assemblage, such as the three fish, the rock iguana, petrel, and crow, are also edible. The few large marine shells (queen conch, West Indian top shell) are broken in a way characteristic of prehistoric human processing. Nevertheless, the only unequivocal Amerindian artifact recovered thus far is a single shell bead (Figure 9). This round, well-polished specimen is 3.4 mm wide, 1.4 mm deep, with a drilled hole 1.5 mm wide. Similar shell beads have been reported from the Coralie Site on Grand Turk Island (Keegan, 1997: Figure 8.2). We note as well that the charcoal occurs throughout the sediments rather than in local concentrations suggestive of hearths.

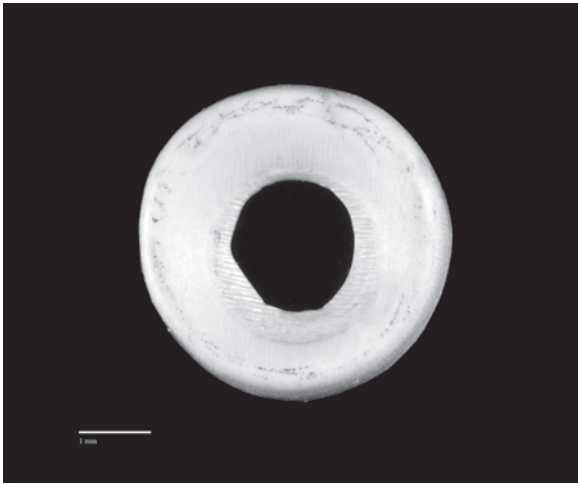


Figure 9. Ground and polished shell bead from Gilpin Point, Abaco, The Bahamas. Photo by HMS.

In spite of using fine-mesh sieves (1.6 mm in the field, 1.0 mm in the laboratory), fossils of the three largest species of vertebrates (crocodile, tortoise, green turtle) far outnumber those of the 16 much smaller species at Gilpin Point (*c.* 200 vs 46 specimens). While prehistoric West Indian peoples typically preferred larger species when available (Carlson and Keegan, 2004), they also often deposited high numbers of small species in their middens (Newsom and Wing, 2004). Among the 16 smaller species in the deposit, the scarcity of fish in particular suggests that we have not yet sampled the densest concentrations of bones in the site because fish typically dominate the bone assemblages from prehistoric archaeological sites in the region (e.g. Jones O'Day, 2002; Wing, 2001).

The fossil sample from Gilpin Point probably is based on the prey remains left behind by humans as well as the Cuban crocodile. We found, nevertheless, very little non-bone evidence of prehistoric human activity at the site, such as pottery or shell midden (often dominated by conch shells *Strombus* spp.), both of which occur routinely and often in abundance in subsurface archaeological sites in the Bahamas and coastal Florida (Jones O'Day, 2002; Keegan, 2007; Keegan et al., 2008; Marquardt, 2010a, 2010b). Our failure to find any pottery or rich shell midden at Gilpin Point might be due to inadequate sampling; if the site extends inland beneath the beach ridge (which seems likely), then the peaty sediment that we have observed would represent less than 1% of the entire site. It could be that *c.* 950–900 years ago the Lucayans lived (at least seasonally) on a barrier beach located on the seaward side of the current beach (and the prehistoric peat deposit). Wave breaks at high tide may have contributed some relatively gentle overwash sand to the peat deposit (and brought small marine mollusks with them), but as noted, evidence of high-energy overwash sediment is lacking. The snails recovered in the deposit are consistent with sheet-wash transport of local terrestrial leaf litter and associated sediments into the peat layer that was forming in the wetland.

Conclusion

The Gilpin Point site is a late-Holocene (*c.* 950–900 cal. yr BP) intertidal, snail-rich, bone-rich peat deposit on the Atlantic coast of Abaco, Bahamas. The site is significant from several standpoints. The fossil plants and mollusks suggest that the site was a freshwater or somewhat brackish coastal wetland at the time the peat was deposited. The short time interval represented by the peat is compatible with rapid sedimentation following initial

human-set fires on Abaco. The peat is exposed today only during exceptionally low tides, suggesting a lower sea level at the time of deposition as well as a degrading shoreline during the past millennium.

As a sample of a late Holocene vertebrate community at the time of first Amerindian presence, the fossils from Gilpin Point demonstrate that Abaco has undergone major biotic changes during the past *c.* 950 years of human occupation. Only 10 of the 17 identified species of non-fish vertebrates still live on Abaco. The loss of seven species of reptiles, birds, and mammals included Abaco's largest terrestrial carnivore (Cuban crocodile) and its three largest terrestrial herbivores (Albury's tortoise, rock iguana, and hutia). Unhealed crocodile bite marks on the inside of the carapaces of green turtles suggest scavenging of turtles that had been butchered by Lucayans.

The dense concentration of large bones in the peat, the charring on some bones of the green turtle and Albury's tortoise, and the discovery of a shell bead, all point to a cultural origin of the bone deposit at Gilpin Point. A challenge now is to search the beaches of Abaco's windward side to begin to learn whether the Gilpin Point site is truly unique or merely represents a more common situation that heretofore has been overlooked. During the lowest spring tides of the coming winter, we look forward to this search.

Acknowledgements

We thank the National Museum of the Bahamas/Antiquities, Monuments, and Museums Corporation (Michael Pateman, Keith Tinker), Abaco Friends of the Environment (Michael Albury, Olivia Patterson, Kristin Williams), and the Bahamas National Trust (Eric Carey, Lynn Gape, David Knowles) for furnishing permits, logistical support, and many other courtesies. Sabrina Bethel, Richard Franz, and Gary Morgan helped with preliminary collections. William Kenny (UF Department of Geology) kindly provided the loss on ignition data for our sediment samples. For comments that improved the manuscript, we thank Karen Bjorndal, Alan Bolton, Sally Horn, and Michelle LeFebvre.

Funding

Financial support was provided by the US National Science Foundation (grant BCS-1118369 to DWS, and BCS-1118340 to JF). This work represents the findings of the authors, and does not necessarily reflect the opinion of the sponsors.

References

- Areces-Mallea AE, Weakley AS, Li X et al. (1999) *A Guide to Caribbean Vegetation Types: Preliminary Classification System and Descriptions*. Washington, DC: The Nature Conservancy.
- Berman MJ, Gnivecki PL and Pateman MP (2013) The Bahama archipelago. In: Keegan WF, Hofman CL and Rodríguez Ramos R (eds) *The Oxford Handbook of Caribbean Archaeology*. Oxford: Oxford University Press, pp. 264–280.
- Bjorndal KA (1997) Foraging ecology and nutrition of sea turtles. In: Lutz PL and Musick JA (eds) *The Biology of Sea Turtles*. Boca Raton, FL: CRC Press, pp. 259–273.
- Bocherens H and Drucker D (2003) Trophic level isotopic enrichment of carbon and nitrogen in bone collagen: Case studies from recent and ancient terrestrial ecosystems. *International Journal of Osteoarchaeology* 13: 46–53.
- Borrito-Páez R, Mancina CA, Woods CA et al. (2012) Updated checklist of endemic terrestrial mammals of the West Indies. In: Borrito-Páez R, Woods CA and Sergile FE (eds) *Terrestrial Mammals of the West Indies*. Micanopy, FL: Wacahoota Press, pp. 389–415.
- Buckner SD, Franz R and Reynolds RG (2012) Bahama Islands and Turks & Caicos Islands (pp. 93–110). In: Powell R and Henderson RW (eds) *Island lists of West Indian amphibians*

- and reptiles. *Bulletin of the Florida Museum of Natural History* 51: 85–166.
- Buden DW (1987) *The Birds of the Southern Bahamas* (B.O.U. Check-list no. 8). London: British Ornithologists' Union.
- Burney DA, Burney LP and MacPhee RDE (1994) Holocene charcoal stratigraphy from Laguna Tortuguero, Puerto Rico, and the timing of human arrival on the island. *Journal of Archaeological Science* 21: 273–281.
- Carlson LA and Keegan WF (2004) Resource depletion in the prehistoric northern West Indies. In: Fitzpatrick SM (ed.) *Voyages of Discovery: The Archaeology of Islands*. Westport, CT: Praeger, pp. 85–101.
- Coltrain JB, Harris JM, Cerling TE et al. (2004) Rancho La Brea stable isotope biogeochemistry and its implications for the palaeoecology of late Pleistocene, coastal southern California. *Palaeogeography, Palaeoclimatology, Palaeoecology* 205: 199–219.
- Curtis JH, Hodell DA and Brenner M (1996) Climate variability on the Yucatan Peninsula (Mexico) during the past 3500 years, and its implications for Maya cultural evolution. *Quaternary Research* 46: 37–47.
- DeSantis LRG, Bhotika S, Williams K et al. (2007) Sea-level rise and drought interactions accelerate forest decline on the Gulf Coast of Florida, USA. *Global Change Biology* 13: 2349–2360.
- Franz R and Franz SE (2009) A new fossil land tortoise in the genus *Chelonoidis* (Testudines: Testudinidae) from the northern Bahamas, with an osteological assessment of other Neotropical tortoises. *Bulletin of the Florida Museum of Natural History* 49: 1–44.
- Franz R, Morgan GS, Albury N et al. (1995) Fossil skeleton of a Cuban crocodile (*Crocodylus rhombifer*) from a blue hole on Abaco, Bahamas. *Caribbean Journal of Science* 31: 149–152.
- Goodwin ID (1993) Holocene deglaciation, sea-level change, and the emergence of the Windmill Islands, Budd Coast, Antarctica. *Quaternary Research* 40: 70–80.
- Hastings AK, Krigbaum J, Steadman DW et al. (in press) Reptile-dominated terrestrial food web in the Bahamas prior to human occupation. *Journal of Herpetology*.
- Hearty PJ and Kaufman DS (2000) Whole-rock aminostratigraphy and Quaternary sea-level history of the Bahamas. *Quaternary Research* 54: 163–173.
- Heaton TJ, Blackwell PG and Buck CE (2009) A Bayesian approach to the estimation of radiocarbon calibration curves: The INTCAL09 methodology. *Radiocarbon* 51: 1151–1164.
- Hedges SB, Couloux A and Vidal N (2009) Molecular phylogeny, classification, and biogeography of West Indian racer snakes of the Tribe Alsophiini (Squamata, Dipsadidae, Xenodontinae). *Zootaxa* 2067: 1–28.
- Henderson RW and Powell R (2009) *Natural History of West Indian Reptiles and Amphibians*. Gainesville, FL: University Press of Florida.
- Hilgartner WB and Brush GS (2006) Prehistoric habitat stability and post-settlement habitat change in a Chesapeake Bay freshwater tidal wetland, USA. *The Holocene* 16: 479–494.
- Horn SP, Orvis KH, Kennedy LM et al. (2000) Prehistoric fires in the highlands of the Dominican Republic: Evidence from charcoal in soils and sediments. *Caribbean Journal of Science* 36: 10–18.
- Howell SNG (2012) *Petrels, Albatrosses & Storm-Petrels of North America*. Princeton, NJ: Princeton University Press.
- Jones O'Day S (2002) Late prehistoric Lucayan occupation and subsistence on Middle Caicos Island, northern West Indies. *Caribbean Journal of Science* 38: 1–10.
- Jones O'Day S and Keegan WF (2001) Expedient shell tools from the northern West Indies. *Latin American Antiquity* 12: 274–290.
- Keegan WF (1997) *Bahamian Archaeology*. Nassau: Media Publishing.
- Keegan WF (2007) *Taino Indian Myth and Practice: The Arrival of the Stranger King*. Gainesville, FL: University Press of Florida.
- Keegan WF and DeNiro MJ (1988) Stable carbon- and nitrogen-isotope ratios of bone collagen used to study coral-reef and terrestrial components of Bahamian diet. *American Antiquity* 53: 320–336.
- Keegan WF, Fitzpatrick SM, Sullivan-Sealey K et al. (2008) The role of small islands in marine subsistence strategies: Case studies from the Caribbean. *Human Ecology* 36: 635–654.
- Keigwin LD (1996) The Little Ice Age and Medieval Warm Period in the Sargasso Sea. *Science* 274: 1504–1508.
- Kemp AC, Horton BP, Donnelly JP et al. (2011) Climate related sea-level variations over the past two millennia. *Proceedings of the National Academy of Sciences* 108: 11017–11022.
- Kjellmark E (1996) Late Holocene climate change and human disturbance on Andros Island, Bahamas. *Journal of Paleolimnology* 15: 133–145.
- Kohn MJ (2010) Carbon isotope compositions of terrestrial C₃ plants as indicators of paleoecology and paleoclimate. *Proceedings of the National Academy of Sciences* 107: 19691–19695.
- Krysko KL, Steadman DW, Mead JI et al. (2013) New island records for reptiles on the Little Bahama Bank, Commonwealth of the Bahamas. *IRCF Reptiles & Amphibians* 20: 152–154.
- Lane CS, Horn SP, Mora CI et al. (2009) Late-Holocene paleoenvironmental change at mid-elevation on the Caribbean slope of the Cordillera Central, Dominican Republic. *Quaternary Science Reviews* 28: 2239–2260.
- Marquardt WH (2010a) Mounds, middens, and rapid climate change during the Archaic-Woodland transition in the southeastern United States. *Anthropological Papers of the American Museum of Natural History* 93: 253–271.
- Marquardt WH (2010b) Shell mounds in the Southeast: Middens, monuments, temple mounds, rings, or works? *American Antiquity* 75: 551–570.
- Medina E (1999) Mangrove physiology: The challenge of salt, heat, and light stress under recurrent flooding. In: Yáñez-Arancibia A and Lara-Domínguez AL (eds) *Ecosistemas de Manglar em América Tropical*. Instituto de Ecología AC Xalapa, México. Silver Spring, MD: UICN/ORMA (Costa Rica), NOAA/NMFS, pp. 109–126.
- Morgan GS (1989) Fossil Chiroptera and Rodentia from the Bahamas, and the historical biogeography of Bahamian mammals. In: Woods CA (ed.) *Biogeography of the West Indies: Past, Present and Future*. Gainesville, FL: Sandhill Crane Press, pp. 685–740.
- Morgan GS and Albury NA (2013) The Cuban crocodile (*Crocodylus rhombifer*) from Late Quaternary fossil deposits in the Bahamas and Cayman Islands. *Bulletin of the Florida Museum of Natural History* 52: 161–236.
- Mylroie JE (2008) Late Quaternary sea-level position: Evidence from Bahamian carbonate deposition and dissolution cycles. *Quaternary International* 183: 61–75.
- Newsom LA and Wing ES (2004) *On Land and Sea: Native American Uses of Biological Resources in the West Indies*. Tuscaloosa, AL: University of Alabama Press.
- O'Leary MH (1981) Carbon isotopic fractionation in plants. *Phytochemistry* 20: 553–567.
- O'Leary MH (1988) Carbon isotopes in photosynthesis. *BioScience* 38: 328–336.
- Olson SL and Hilgartner WB (1982) Fossil and subfossil birds from the Bahamas. *Smithsonian Contributions to Paleobiology* 48: 22–56.

- Olson SL and Meylan PA (2009) A second specimen of the Pleistocene Bermuda tortoise, *Hesperotestudo bermudae* Meylan and Sterrer. *Chelonian Conservation and Biology* 8: 211–212.
- Olson SL and Pregill GK (1982) Introduction to the paleontology of Bahaman vertebrates. *Smithsonian Contributions to Paleobiology* 48: 1–7.
- Park LE (2012) Comparing two long-term hurricane frequency and intensity records from San Salvador, Bahamas. *Journal of Coastal Research* 28: 891–902.
- Perry L and Williams K (1996) Effects of salinity and flooding on seedlings of cabbage palm (*Sabal palmetto*). *Oecologia* 105: 428–434.
- Pregill GK (1981) *Late Pleistocene herpetofaunas from Puerto Rico* (Miscellaneous Publication no. 71). Lawrence, KS: University of Kansas Museum of Natural History.
- Pregill GK (1982) Fossil amphibians and reptiles from New Providence Island, Bahamas. *Smithsonian Contributions to Paleobiology* 48: 8–21.
- Pregill GK, Steadman DW and Watters DR (1994) *Late Quaternary Vertebrate Faunas of the Lesser Antilles: Historical Components of Caribbean Biogeography* (Bulletin of the Carnegie Museum of Natural History no. 30). Pittsburgh, PA: Carnegie Museum of Natural History.
- Rada F, Goldstein G, Orozco A et al. (1989) Osmotic and turgor relations of three mangrove ecosystem species. *Functional Plant Biology* 16: 477–486.
- Reimer PJ, Baillie MGL, Bard E et al. (2009) INTCAL09 and MARINE09 radiocarbon age calibration curves, 0–50,000 years cal BP. *Radiocarbon* 51: 1111–1150.
- Reynolds RG, Niemiller ML, Hedges SB et al. (2013) Molecular phylogeny and historical biogeography of West Indian boid snakes (*Chilabothrus*). *Molecular Phylogenetics and Evolution* 68: 461–470.
- Schwartz A and Henderson RW (1991) *Amphibians and Reptiles of the West Indies: Descriptions, Distributions, and Natural History*. Gainesville, FL: University Press of Florida.
- Slayton IA (2010) *A vegetation history from Emerald Pond, Great Abaco Island, the Bahamas, based on pollen analysis*. M.S. Thesis, University of Tennessee.
- Steadman DW and Takano OM (2013) Late Holocene birds from Haiti: Refining the chronology of vertebrate extinction in the West Indies. *The Holocene* 23: 934–942.
- Steadman DW, Pregill GK and Burley DV (2002) Rapid prehistoric extinction of birds and iguanas in Polynesia. *Proceedings of the National Academy of Sciences* 99: 3673–3677.
- Steadman DW, Franz R, Morgan GS et al. (2007) Exceptionally well-preserved late Quaternary plant and vertebrate fossils from a blue hole on Abaco, Bahamas. *Proceedings of the National Academy of Sciences* 104: 19897–19902.
- Steadman DW, Martin PS, MacPhee RDE et al. (2005) Asynchronous extinction of late Quaternary sloths on continents and islands. *Proceedings of the National Academy of Sciences* 102: 11763–11768.
- Steadman DW, Watters DR, Reitz EJ et al. (1984) Vertebrates from archaeological sites on Montserrat, West Indies. *Annals of Carnegie Museum* 53: 1–29.
- Stuiver M and Braziunas (1993) Modeling atmospheric ¹⁴C influences and ¹⁴C ages of marine samples to 10,000 BC. *Radiocarbon* 35: 137–189.
- Sullivan-Sealey K, Brunnick B, Harzen S et al. (2002) *An Ecoregional Plan for the Bahamian Archipelago*. Jupiter, FL: Taras Oceanographic Foundation.
- Vander Zanden HB, Arthur KE, Bolten AB et al. (2013) Trophic ecology of a green turtle breeding population. *Marine Ecology Progress Series* 476: 237–249.
- Walker KJ (2013) The Pineland Site complex: Environmental contexts. In: Marquardt WH and Walker KJ (eds) *The Archaeology of Pineland: A Coastal Southwest Florida Site Complex, A.D. 50–1710*. Gainesville, FL: Institute of Archaeology and Paleoenvironmental Studies, University of Florida, pp. 23–52.
- White A (1998) *A Birder's Guide to the Bahama Islands* (Including Turks and Caicos). Colorado Springs, CO: American Birding Association, Inc.
- Wingate DB (1972) First successful hand-rearing of an abandoned Bermuda Petrel chick. *Ibis* 114: 97–101.
- Wing ES (2001) Native American use of animals in the Caribbean. In: Woods CA and Sergile FE (eds) *Biogeography of the West Indies: Patterns and Perspectives*. Boca Raton, FL: CRC Press, pp. 481–518.