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RESEARCH ARTICLE

Paleodietary reconstruction of endemic rodents from the precolumbian Dominican Republic: Discriminating wild feeding behavior from diets linked to human niche construction activities

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Abstract

In the Greater Antilles, certain animal taxa that have long been theorized to have been managed by indigenous peoples prior to AD1492, the main candidates being a group of endemic caviomorph rodents known as hutias (Capromyinae). This isotopic study investigates the paleodiets of several species of endemic rodents from three late precolonial sites in the northern Dominican Republic: El Flaco (cal. AD 990–1452), El Carril (cal. AD 1030–1262), and La Entrada (cal. AD 840–900) to assess whether human influence over animal diets can be determined. We examined bone collagen carbon ($\delta^{13}\text{C}_{\text{co}}$) and nitrogen ($\delta^{15}\text{N}$) and tooth enamel carbon ($\delta^{13}\text{C}_{\text{en}}$) isotope values of three species of hutias, *Isolobodon portoricensis*, *Isolobodon montanus*, and *Plagiodontia aedium*, alongside edible rat (*Brotomys* sp.), and domestic guinea pig (*Cavia porcellus*). To estimate dietary source contributions, we employed a Bayesian dietary mixing model (FRUITS v.3.0) and ran three different permutations to assess the relative contributions of C_3 or C_4/CAM plants. The addition of an extra 79 wild C_3 and 40 wild C_4/CAM plant species' isotope values from published sources to an established isotopic foodweb for the Caribbean region enabled us to discriminate between wild and domestic C_3 and C_4/CAM plant food sources in two of these models. Our results provide evidence of the significant consumption of domestic C_4/CAM plants by some animals. This likely represents maize (*Zea mays*) consumption, which is known to have been ubiquitously cultivated by indigenous peoples in the region. This is particularly the case for *I. portoricensis*, as FRUITS modeling suggests that a few individuals consumed C_4/CAM plants well beyond their expected natural diets as determined from feeding studies of extant hutia species. This may indicate human influence over endemic rodent diets due to niche construction activities such as horticultural practices and may reflect either opportunistic feeding on human produce or the purposeful supplementation of hutia diets by humans.

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KEYWORDS

dietary mixing models, hutias, isotopic analysis, niche construction, precolumbian, zooarchaeology

1 | INTRODUCTION

Archaeological researchers have long suggested that indigenous peoples in the precolonial Caribbean may have been managing some animal species. These inferences are largely postulated due to the ubiquity and high frequencies of certain animals at archaeological sites, alongside indication of the translocation of these same species throughout the region by Amerindians (Allen, 1916; Flemming & MacPhee, 1999; Giovas, 2019; Giovas et al., 2016; LeFebvre, deFrance, et al., 2019; Newsom & Wing, 2004; Oswald et al., 2020; Wing, 2008, 2012). In the Greater Antillean islands of Hispaniola, Cuba, Jamaica, and Puerto Rico, certain species of the Echimyidae family (subfamily: Capromyinae) (Courcelle et al., 2019) of caviomorph rodents, known as hutias, are the primary candidates for having been managed. Hutias are a group of rodents that evolved after a single colonization event into the Greater Antilles islands during the Early Miocene, eventually diversifying into at least eight genera and 33 species over geological time to occupy a myriad of ecological niches throughout the islands (Courcelle et al., 2019; Fabre et al., 2014; MacPhee & Iturralde-Vinent, 1995; Woods et al., 2021). In Hispaniola and Puerto Rico, one species of this radiation, *Isolobodon portoricensis*, has been speculated as undergoing “incipient domestication,” based on historical mention of corrals witnessed by European chroniclers in 16th century Hispaniola, the abundance of hutia skeletal remains in many archaeological assemblages, and evidence of their translocation by humans between islands (de Las Casas, 1875; Newsom & Wing, 2004; Wing, 2008, 2012).

Although no physical traces of enclosures or other material evidence of captive management have been identified archaeologically, biomolecular studies have shed light on complex relations between humans and hutia. Studies have revealed evidence of zoonotic parasite transmission from hutias to humans suggesting a close cohabitation of space, (Wiscovitch-Russo et al., 2020); provided genetic evidence of the introduction of populations of hutias to new island environs (Oswald et al., 2020; Woods et al., 2021); and highlighted similarities in dietary stable isotope values with humans which may suggest some degree of commensalism, both in the Bahamas with *Geocapromys ingrahami* and in Hispaniola with *I. portoricensis* (LeFebvre, DuChemin, et al., 2019; Shev, Laffoon, & Hofman, 2021). These studies do not necessarily indicate that hutias were held in captivity but do suggest that there was sharing of foodways and possibly indicate close cohabitation between humans and hutias.

The current research builds on previously published results of stable isotope analysis of bone collagen from archaeological *I. portoricensis* remains and other endemic taxa from four precolonial sites in the modern-day Dominican Republic (Shev, Laffoon, & Hofman, 2021). Added to these collagen data are newly analyzed

enamel isotope values, which enabled us to employ Bayesian dietary mixing modeling (FRUITS v.3.0) on several individuals. Using at our disposal three isotope proxies for each individual animal and published isotopic foodweb values of plants from Hispaniola, allows us to more succinctly ascertain whether there were distinct dietary linkages between hutias and human and sheds light on a likely mutualistic relationship.

Dietary stable isotope analysis is a useful technique for distinguishing domestic, tame, or captive diets from that of an animal's natural diet (Guiry et al., 2021; LeFebvre, deFrance, et al., 2019; Makarewicz & Tuross, 2012; Sugiyama et al., 2015, 2020). In large swathes of the Americas, agricultural maize (*Zea mays*) was a staple crop. Maize utilizes a C₄ photosynthetic pathway resulting in plant tissues with higher carbon ($\delta^{13}\text{C}$) isotope values than all C₃ taxa that form most the world's plant species. Therefore, in many parts of the Americas, human dietary influence over animals can be observed in unexpectedly raised $\delta^{13}\text{C}$ values in animal bones and teeth, which may indicate opportunistic scavenging behaviors or the feeding of agricultural maize, or another C₄ plant, to animals (Lee-Thorp et al., 1989; LeFebvre, deFrance, et al., 2019; Makarewicz & Tuross, 2012; Sugiyama et al., 2015, 2020).

Human dietary influence as determined from relatively raised $\delta^{13}\text{C}$ values in animal tissues may directly correlate to animal management practices (Barton et al., 2009; Monagle et al., 2018; Sharpe et al., 2018; Zavodny et al., 2015). In the Americas, human–animal interactions took on many forms that did not necessarily lead to domestication, let alone constitute a relationship where humans had direct control over animal lifeways (Zeder, 2006, 2012, 2015). Due to these often-complicated forms of human–animal relations, we do not seek to establish from isotopic data whether hutias were being managed in captivity by indigenous peoples, let alone undergoing domestication. Human niche construction activities, such as altering plant communities by removing unwanted species and cultivating economically beneficial plants, are known to present adaptive pressures or benefits to animal species that inhabit an environment (Boivin et al., 2016; Odling-Smee et al., 1996; Smith, 2001, 2007, 2011; Zeder, 2015). In the tropical Americas, ecological and anthropological studies indicate that some species, particularly dietary generalists or synanthropes, can benefit from and be attracted to environmental changes brought on by human agricultural activities, which in turn may be hunted in close proximity to agricultural plots to which they are attracted to, a practice known as “garden hunting” (Arce-Peña et al., 2019; Linares, 1976; Loiselle & Blake, 1992; Ramírez-Barajas & Calmé, 2015; Smith, 2005). Therefore, interpreting dietary linkages between humans and animals as part of a teleological process towards domestication may not necessarily be a sound approach for studying past societies from much of the Neotropics. Rather, we seek to shed

light on possible dietary linkages between both species that may be the result of purposeful feeding by humans, or from animals opportunistically scavenging from human settlements and feeding on garden plots, which are both in essence mutualistic behaviors that likely lead to equifinality in dietary isotope values.

1.1 | Dietary isotope analysis of human–animal interactions in the precolumbian Caribbean

Stable isotope analyses of carbon and nitrogen establish general information about animal diets by comparing the ratio of heavier to lighter isotope values within a consumer's tissues to the values of consumed food sources. Carbon isotope values of hard and soft tissues are dictated by the particular photosynthetic processes of plant species at the base of food webs, which vary according to environmental constraints such as temperature, aridity, and light exposure (DeNiro & Epstein, 1978; Schoeninger & DeNiro, 1984). Most plant species have a C₃ metabolic pathway and have $\delta^{13}\text{C}$ values ranging between -20 and -37% . C₄ plants, which are more tolerant of tropical conditions, usually have values between -7 and -17% , while arid-adapted crassulic acid metabolism (CAM) plants demonstrate values that mostly overlap with C₄ species (Ehleringer, 1989; Kohn, 2010; O'Leary, 1988). Nitrogen ($\delta^{15}\text{N}$) values within bone collagen are primarily derived from consumed proteins. Nitrogen values see a stepwise increase with the higher trophic positioning of an animal, with baselines originating from the $\delta^{15}\text{N}$ values of plants at the bottom of a foodweb that is derived from absorbed nitrates in substrates. The trophic position of an organism can be established by examining its $\delta^{15}\text{N}$ values and can be used to differentiate the consumption of terrestrial and marine foods as marine environments generally have higher basal $\delta^{15}\text{N}$ values and more trophic levels in their food webs (Keegan & DeNiro, 1988; Minagawa & Wada, 1984; Schoeninger & DeNiro, 1984).

Numerous stable isotope studies have assessed human paleodiets of precolonial and early colonial period indigenous communities in the insular Caribbean (Chinique de Armas et al., 2015, 2016; Keegan & DeNiro, 1988; Krigbaum et al., 2013; Laffoon et al., 2013, 2019, 2020; Laffoon & Vos, 2011; Pestle, 2010, 2013; Stokes, 2008). Isotopic studies of animal paleomobility have been conducted to investigate the introduction or movement of animals into different Caribbean islands (Giovas, 2016; Giovas et al., 2016, 2019; Laffoon et al., 2015, 2019; LeFebvre, deFrance, et al., 2019), while only a few isotopic studies have looked specifically at animal diets in an attempt to highlight dietary linkages between the human and animals (Laffoon et al., 2019; LeFebvre, deFrance, et al., 2019; Shev et al., 2020; Shev, Laffoon, & Hofman, 2021).

Our previous study discussed the bone collagen values of hutia and other endemic animals, establishing that there was likely some degree of commensalism between humans, dogs (*Canis familiaris*), domestic guinea pig (*Cavia porcellus*), and some hutias (*I. portoricensis*) at four Late Ceramic Age (AD 500–1500) sites in the Dominican Republic (Shev et al., 2020; Shev, Laffoon, & Hofman, 2021). Nevertheless, the ability to determine the relative

portions of particularly food sources that an animal consumed is limited from the study of bone collagen carbon ($\delta^{15}\text{C}_{\text{co}}$) and nitrogen ($\delta^{15}\text{N}$) values alone. This is partly because bone collagen isotope values are determined mainly by dietary protein intake, compared with enamel (and bone bioapatite) $\delta^{13}\text{C}$ values that are the result of an averaging of values from all macronutrients (fats, proteins, and carbohydrates) (Ambrose & Norr, 1993; Schwarcz, 2002). There is also the issue of disentangling equifinality in isotopic values between humans and animals when only one or two isotopic proxies are examined (Fernandes et al., 2015). This may be due to different macronutrient routing mechanisms between species or due to consumption of different types of plants that have similar values. Another phenomenon is the “canopy effect” where plants of the same species residing in tropical forest canopies have higher $\delta^{13}\text{C}$ values than ground-dwelling plants. This can lead to organisms having different isotopic values although they ate the same species of plant or having similar isotopic values in their tissues even though they ate different food sources altogether (Blumenthal et al., 2016; Kohn, 2010; Quinn, 2019; Roberts et al., 2017).

1.2 | FRUITS

To provide a more accurate estimation of input percentages of various dietary sources, a combination of dietary isotope values from tooth enamel apatite ($\delta^{13}\text{C}_{\text{en}}$) and bone collagen ($\delta^{13}\text{C}_{\text{co}}$ and $\delta^{15}\text{N}$) provides greater fidelity in predicting dietary contributions when inputting these values into a Bayesian dietary mixing model. For our study, we have employed the software Food Reconstruction Using Isotopic Transferred Signals (FRUITS v.3.0) (Fernandes et al., 2014). Bayesian dietary mixing models analyze multiple isotopic values from a studied individual against the mean or median values and associated uncertainties of the presumed food source groups they consumed, providing probabilistic estimates of the relative portions of defined food sources that an organism likely ate (Fernandes et al., 2012, 2014; Hopkins & Ferguson, 2012).

Dietary mixing model studies have been used sparsely in precolumbian Caribbean archaeological investigations (Chinique de Armas et al., 2016; Pestle, 2010). They have been employed to assess paleodiets of indigenous peoples from Tutu, Virgin Islands, demonstrating that C₃ plants formed the majority of the diets of all inhabitants contrary to the evidence suggested from the zooarchaeological and paleobotanical data (Pestle & Laffoon, 2018). Bayesian dietary mixing models have also been employed to assess weaning ages from the study of juvenile and adult human bone collagen values from different population groups at several sites in Cuba (Chinique de Armas et al., 2017, 2022; Chinique de Armas & Pestle, 2018). Relevant to animal diets, FRUITS was employed to estimate domestic dog (*C. familiaris*) dietary inputs from El Flaco, Dominican Republic, and Morel and Cathedrale de Basse-Terre in Guadeloupe. These results suggest that dogs had broadly similar diets to humans, which were mainly composed of C₃ plant foods, followed in importance by terrestrial animals, C₄ plants, and then marine animals (Shev et al., 2020).

In this study, FRUITS modeling has been conducted on multiple species of endemic rodents and one domestic guinea pig (*C. porcellus*) from three precolumbian sites within Hispaniola: El Flaco, El Carril, and La Entrada (Figures 1 and 2). Examined endemic species include extinct edible rat (*Brotomys* sp.), and two species of hutia, the critically endangered Hispaniolan hutia (*Plagiodontia aedium*) and the extinct Puerto Rican hutia (*I. portoricensis*), the latter of which researchers have long speculated may have been managed by indigenous peoples. Applying Bayesian dietary mixing models to study the diets of these animals permits greater resolution in determining the probable proportions of good groups consumed; therefore, facilitating a more concise assessment of human–animal dietary entanglements than is possible from relying on one or two isotope values alone. To assess if there was a human influence on the diets of endemic animals because of niche construction activities, we made distinctions between wild and domestic C₃ plant food sources and wild and domestic (mainly maize) C₄ plant food sources in our FRUITS modeling.

1.3 | The feeding preferences of hutias

To examine human–hutia commensalism, an understanding of the feeding preferences and behavioral ecology of hutias is required to ensure an accurate application of isotopic foodwebs. Particularly with “wild” animals, the isotopic values of noncultivated plant foods that they may have consumed are needed, often educated by a priori information recorded in observation studies (Bond & Diamond, 2011; Phillips et al., 2014). As *I. portoricensis*, *I. montanus*, and edible rat (*Brotomys* sp.) likely went extinct shortly after the arrival of Europeans

(Turvey et al., 2007), dietary information can only be inferred from stable isotopic studies of skeletal remains and gleaned from feeding studies conducted on extant hutia species which shared morphological and functional similarities.

Cooke and Crowley (2018) studied tooth enamel isotopic values of carbon and oxygen ($\delta^{18}\text{O}$) for *I. portoricensis*, *P. aedium*, *Brotomys voratus*, and other species recovered from two Holocene cave sites on the Tiburon peninsular of southwest Haiti: Trou Jean Paul and Trouing Jérémie #5. *I. portoricensis* exhibited relatively low oxygen but intermediate carbon isotope values compared with other taxa, likely indicating a terrestrial lifestyle involving foraging in undergrowth, with isotopic values of food sources likely influenced by the “canopy effect” (Cooke & Crowley, 2018; van der Merwe & Medina, 1991). Comparatively low $\delta^{18}\text{O}$ may also indicate this species was more frugivorous than most other studied taxa. *Brotomys* sp. exhibited higher $\delta^{13}\text{C}$ values than *I. portoricensis*, but also had relatively low $\delta^{18}\text{O}$, indicating that this species may also have been frugivorous, but more arboreal and less subject to the relatively diminished carbon values of ground-dwelling plants due to the “canopy effect.” However, the isotopic values of these specimens may be more reflective of specific environmental constraints in southwest Haiti (Cooke & Crowley, 2018). Additionally, these two sites do not contain archaeological material, meaning that the dietary behavior of studied taxa was possibly not directly influenced by human activities, although fossil deposits at Trou Jean Paul may coincide with the Late Ceramic Age (AD 500–1500) occupation of Hispaniola by agroceramicist cultures ($\sim 1690 \pm 570$ cal BP) (Soto-Centeno et al., 2017), so feasibly environmental changes brought about by humans may have affected feeding behavior of these species. At both sites, Cooke and Crowley (2018)



FIGURE 1 Map showing locations of the three sites in northern Hispaniola, modern-day Dominican Republic: El Carril, El Flaco, and La Entrada. Listed are the number of individuals of each taxon from which bone collagen and tooth enamel was analyzed [Colour figure can be viewed at [wileyonlinelibrary.com](https://onlinelibrary.wiley.com/doi/10.1002/oa.3149)]

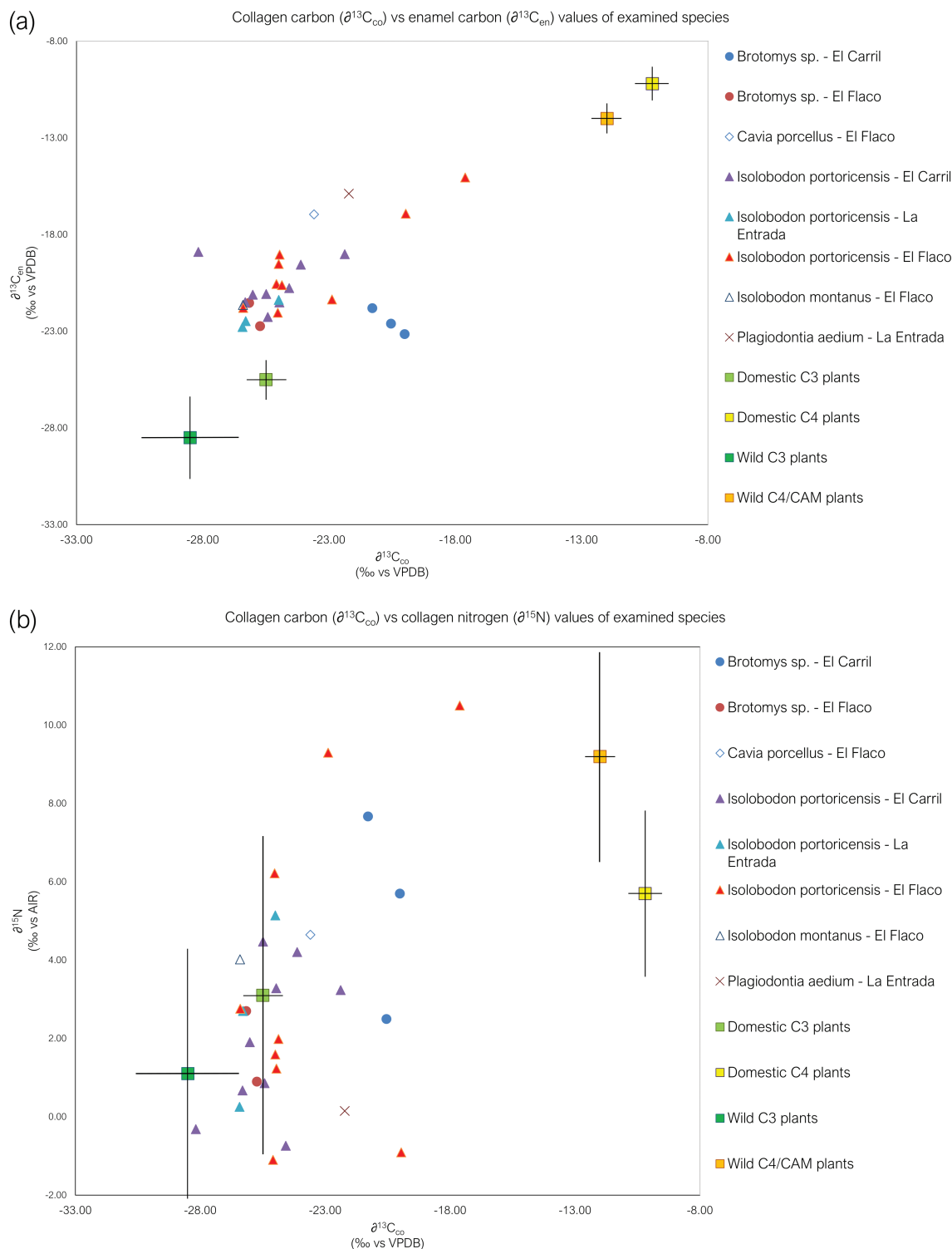


FIGURE 2 (a) Organic (collagen) and inorganic (enamel) carbon values of the studied fauna including trophic discrimination factors (TDFs). (b) Collagen carbon and nitrogen values of the studied fauna including TDFs. For both graphs, the mean isotope values are included for the four plant food source groups investigated. Bars indicate one standard deviation from the mean [Colour figure can be viewed at [wileyonlinelibrary.com](https://onlinelibrary.wiley.com)]

determined that all species likely consumed some C₄ foods, using a cutoff of $\delta^{13}C_{en} - 14.6\%$ based on a diet-bioapatite offset of $+9.4\%$ for *Rattus rattus* (Ambrose et al., 1997), accounting for the highest $\delta^{13}C$ value for C₃ forest ecosystems of $\delta^{13}C - 25.5\%$ (Kohn, 2010) and the Suess effect ($+1.5\%$) (Francey et al., 1999).

Feeding studies have been conducted on seven extant species of hutia, both in captive environments and observations in the wild. Observed in natural conditions, hutias will consume 173 different plant species from 63 families. Controlled feeding studies of captive hutias, in which different plants were introduced into their diets under

observation, determined that the studied hutias will readily consume 83 species of mostly cultivated forms but will also eat processed foods derived from animal meat. Most species demonstrate preferences for the consumption of tree barks, tender branches and petioles of trees and dicots, and all species were rarely observed eating monocots, especially grasses which can commonly be C_4 plants (Gramminae) (Boroto-Páez & Woods, 2012). All observed hutia species are primarily herbivorous but some have a tendency for omnivory, for example, Cuban hutias (*Capromys pilorides*) have been observed opportunistically catching insects and marine invertebrates in coastal habitats (Boroto-Páez & Woods, 2012; Eisenberg & Woods, 2012; Frias & Hernandez, 1985; Manójjina & Abreu, 1990). Wild *C. pilorides* favor plants from the Rosidae subclass, which includes red mangrove (*Rhizophora mangle*). Rosidae includes two families that have CAM species, Euphorbiaceae and Crussalaceae; however, no CAM species from these two families have been observed to be eaten. The majority of *C. pilorides* diets are composed of C_3 angiosperms; they will however consume wild pineapple (*Bromelia pinguin*), a CAM plant (Boroto-Páez & Woods, 2012; Keeley & Rundel, 2003). Species from the *Mesocapromys* genus, restricted to Cuba, also prefer plants from the Rosidae subclass, with one species *Mesocapromys melanurus* inhabiting mosaiced habitats consisting of wild and domestic plants, and favoring cultivated species within these. Jamaican hutia (*Geocapromys brownii*) has been observed eating some C_4 grasses, such as *Panicum maximum* and *Pharus glaber*, and sugar cane (*Saccharum officinarum*) which was introduced by Europeans into Caribbean; however, these foods accounted for only 5.2% of their observed diets. The sole extant hutia from Hispaniola, *P. aedium* has been observed eating one C_4 species, maize (*Z. mays*), although observation studies of this species in the wild are obscured by its rarity, nocturnality, and semi-arboreal behavior (Boroto-Páez & Woods, 2012; Eisenberg & Woods, 2012; Oliver et al., 1986). When observations have been made of *P. aedium* in the wild, they are noted to mainly consume the bark of twigs, upper branches, and trunks of wild cherry (*Prunus occidentalis*) and pine (*Pinus occidentalis*), but especially favoring the bark and fruits of wild avocado (*Persea anomala*) (Boroto-Páez & Woods, 2012; Woods & Ottenwalder, 1992). From these studies, it is that a wide variety of plants will readily be consumed by hutias; however, there is likely very few that are C_4 /CAM species. It can therefore be deduced that for most hutia species, their wild diets are predominantly composed of C_3 trees and other plants, which should be reflected within their dietary isotope values.

A multitude of domesticated plants were cultivated in the Caribbean region and have been identified within archaeological sites. Cultivated C_3 species include various tree fruits, manioc (*Manihot esculenta*), sweet potato (*Ipomoea batatas*), chili (*Capsicum* sp.), zamia (*Zamia* sp.), and legumes (Fabaceae) among others, whereas evidence of C_4 maize (*Z. mays*) consumption has been identified in starch grain and phytolith analysis, including at the site of El Flaco (Ciofalo et al., 2019, 2020; Mickleburgh & Pagan-Jimenez, 2012; Newsom & Wing, 2004; Pagán-Jiménez et al., 2020). Cultivated C_4 /CAM plants from the region that may have been targeted by hutia include maize, prickly pear (*Opuntia* sp.), agave (*Agave antillarum*), pineapple (*Ananas*

comosus), and amaranth (*Amaranthus* sp.) (Pestle, 2010); however, the paleoethnobotanical evidence suggests that maize was like the more ubiquitously cultivated crop in precolonial Hispaniola (Ciofalo et al., 2019; Figueredo, 2015; Mickleburgh & Pagan-Jimenez, 2012; Pagán-Jiménez et al., 2020; Pagán-Jiménez & Mickleburgh, 2022).

2 | MATERIALS AND METHODS

This study involves the examination of previously analyzed collagen samples from three precolonial sites in the Dominican Republic, El Flaco (cal. AD 990–1452), El Carril (cal. AD 1030–1262), and La Entrada (cal. AD 840–900), the site histories of which are discussed in previous publications (Hofman & Hoogland, 2015; Pagán-Jiménez et al., 2020; Shev, Laffoon, & Hofman, 2021). Accompanying these are 31 newly analyzed enamel samples from some of the same animals. Enamel from five species was analyzed; three hutia species *I. portoricensis* ($n = 24$), *Isolobodon montanus* ($n = 1$), and *P. aedium* ($n = 1$), edible rat (*Brotomys* sp.) ($n = 5$), and one example of domestic guinea pig (*C. porcellus*) that is one of only four instances of this animal recovered from precolonial contexts in Hispaniola (LeFebvre & deFrance, 2014).

Incisors were chosen over molariform teeth for enamel extraction for two reasons. Extracting adequate amounts of enamel from incisors is more easily accomplished than with molariform teeth of hutias which have thin enamel folds that disintegrate easily under a handheld rotary drill. Mandibular incisors are continually growing in all rodents, within rats at a rate of 0.4–0.6 mm per day, and within guinea pigs at 0.3 mm per day (Müller et al., 2015; Park et al., 2017). However, for many species of Capromyinae, molariform teeth are open-rooted and continuously regenerating, which is a useful adaptation for diets that often include tough material such as tree bark (Boroto-Páez & Woods, 2012; Hermanson & Woods, 2012). As the regeneration rates of molariform teeth in *I. portoricensis* are unknown, it was deemed that there was no extra benefit to sampling molariform teeth over incisors.

Collagen extraction protocols for previously analyzed samples are listed in that publication (see Shev, Laffoon, & Hofman, 2021). Enamel extraction follows a modified protocol from Bocherens et al. (2011). All enamel samples were taken from rodent incisors that were embedded in mandible bones that previously had collagen successfully extracted. Enamel powder was drilled from single incisors using a diamond-tipped Dremel handheld drill and inserted into polyethylene miniature test tubes, to which 1 ml of 2% to 3% NaOCl solution was added. Samples were vortexed and sat at room temperature for 20–24 h. Bleach was removed by centrifuging samples briefly and then removing solution with a sterile pipette, 1 ml of deionized water was added to each sample which was then centrifuged, and water was subsequently removed with a pipette. This step was repeated three to four times. To remove carbonates, 1 ml of 1 M acetic acid-CA acetate buffer (pH = 4.75) was added and samples were left at room temperature for 20–24 h. The cleaning step with deionized water was again repeated three to four times until no trace of acetic acid is left. Samples were then placed in an oven at 60°C overnight with caps open. Enamel samples were then weighed and transferred to small, labeled

glass vials (minimum sample weight 0.3 mg). Samples were analyzed at the Faculty of Earth Sciences, VU Amsterdam, using a Finnigan Delta-Plus Isotope Ratio Mass Spectrometer subsequent to reaction of the samples with H_3PO_4 [100%] and isolation of produced CO_2 on a Gasbench II universal automated interface. All $\delta^{13}\text{C}$, $\delta^{18}\text{O}$, and $\delta^{15}\text{N}$ values are stated in δ notation in parts per thousand (‰), carbon and oxygen relative to the international Pee Dee Belemnite (PDB) standard, and nitrogen relative to the international Ambient Inhalable Reservoir (AIR) standard.

2.1 | FRUITS protocol

FRUITS (v.3.0) was chosen due to its proficiency in integrating isotopic, elemental, and food macronutrient data, as well as uncertainties in source and consumer data, and its ability to discriminate animal-derived foods that are rich in protein and fat from plant sources which are generally carbohydrate rich but protein poor. A comprehensive overview of the application of FRUITS for the Caribbean region is listed in Pestle and Laffoon (2018).

There are important differences in dietary behavior, macronutrient routing and trophic fractionation calculations between rodents and humans that need to be considered. Before inputting all data into FRUITS, values were altered to reflect diet-tissue offsets, also known as trophic discrimination factors (TDFs), which are the average difference between the stable isotope values of foods consumed and the tissues of the consumer. As different animals undergo distinct processes of macronutrient routing, errors can occur using Bayesian dietary mixing models when assuming TDFs. Reliable TDFs must be obtained from controlled feeding studies of species that are taxonomically or functionally analogous (Bond & Diamond, 2011; Healy et al., 2017; Kurle et al., 2014; Rio et al., 2009). As three of the five species examined in this study are extinct, generic rodent feeding studies provided our TDFs. Measured isotope values were converted to account for fractionation with a diet-tissue discrimination factor of +9.9‰ for $\delta^{13}\text{C}_{\text{en}}$ following a previous application for rodents which was based on laboratory feeding studies (Ambrose & Norr, 1993; DeNiro & Epstein, 1978; Grimes et al., 2004). For $\delta^{15}\text{N}_{\text{co}}$, a TDF of +3.6‰ \pm 1.4‰ was applied based on numerous feeding studies conducted on omnivorous animals, including rodents (Ambrose, 2002; DeNiro & Epstein, 1981; Howland et al., 2003; Sponheimer et al., 2003). TDFs for $\delta^{13}\text{C}_{\text{co}}$ were calculated using the linear regression formula outlined by Pestle et al. (2015). On average, $\delta^{13}\text{C}_{\text{co}}$ TDFs were +4.9‰ \pm 1.2‰ and ranged from +3.8‰ to +6.7‰. All data were input into FRUITS accounting for an instrumental uncertainty of 0.1‰.

Some limitations of using dietary mixing models need to be considered when assessing model outcomes. The inclusion of too many sources may skew results (Cheung & Szpak, 2020; Fernandes, 2016; Phillips et al., 2014; Stock et al., 2018), so we have limited our models to a maximum of four source groups composed of only plant species. There is also potential for the missing of a source group having significant effects on model outcomes (Fernandes, 2016; Stock et al., 2018);

however, we rely on observational feeding studies that indicate that the Caribbean rodents we are studying are likely herbivorous, so we deemed that there was no need to include terrestrial or marine meats into our models. Another consideration is that the bioturnover rates between bone collagen and tooth enamel are vastly different. Collagen bioturnover has the slowest rate of any body tissue and likely represents an averaging of dietary sources over a long time (Gineyts et al., 2000), so when comparing collagen to tooth enamel values, particularly with rodent incisors that constantly remodel this needs to be taken into consideration. Therefore, using enamel and collagen isotope values together in a mixing model might mean an averaging of isotopic values from various time periods (Cheung & Szpak, 2020). As most species examined in this study are extinct, we have no information regarding their bone collagen bioturnover rates and tooth growth durations, so considering that we are not examining temporality or seasonal variations in diet we deem this as a minor limitation.

2.2 | Caribbean isotopic foodweb values

Isotopic values of established food sources of plants and animals consumed by indigenous peoples in the Caribbean were garnered from previous publications (Keegan & DeNiro, 1988; Pestle, 2010; Schwarcz et al., 1985; Stokes, 2008; von Fischer & Tieszen, 1995) but were altered to only contain plants found in Hispaniola and to accommodate presumed feeding behavior of the studied taxa (Table S1). According to captive feeding studies and observations in the wild, hutia species are generally herbivorous, but with some instances of opportunistic omnivory with certain observed hutia species such as *C. pilorides*. As it is likely that only some hutias are opportunistically carnivorous, we have worked under the assumption that any consumption of animal foods would be negligible; therefore, we have omitted terrestrial and marine animal food sources from our models. As hutia and edible rats were likely not subsisting solely or predominantly on cultivated plants, published isotopic values of wild plant species with natural ranges encompassing Hispaniola were added, including an extra 79 values of C_3 plants and 40 C_4/CAM values. These include some epiphytes, native C_4 amaranth species (*Amaranthus* spp.) and native grass species as possible wild C_4/CAM sources. When possible, published values of wild species were gleaned from studies conducted on Caribbean islands; however, due to a lack of prior research on isotopic baselines for the region, values of wide-ranging species were taken from studies in the tropical Neotropics, including Mexico, the Guianas, and Peru as long as these species had natural ranges encompassing Hispaniola (Table S1).

Three models were run: a two-source model examining the input of C_4/CAM and C_3 plants; a three-source model with C_3 plants and dividing C_4/CAM plants into domestic (maize and *Amaranthus amaranthus*) and wild species; and a four-source model dividing both C_4/CAM and C_3 plants into wild and domestic species. The mean and standard deviations of stable isotope values of food source groups are as listed in Table 1. According to Student's *t* tests, there is a significant difference between the mean carbon values of wild and domestic C_3

TABLE 1 Number of samples and mean isotopic values for the six food source groupings used in this study

Food group	No. of $\delta^{13}\text{C}$ samples	Mean $\delta^{13}\text{C}$ value (‰ vs. VPDB)	No. of $\delta^{15}\text{N}$ samples	Mean $\delta^{15}\text{N}$ value (‰ vs. N_{AIR})
C_3 plants	186	-26.8 ± 3.5	103	2.3 ± 4.4
C_4/CAM plants	68	-11.5 ± 1.4	35	7.5 ± 4.4
Domestic C_3	107	-25.5 ± 1.8	62	3.1 ± 2.1
Wild C_3	79	-28.5 ± 4.3	41	1.1 ± 6.2
Domestic C_4/CAM	28	-10.2 ± 1.5	17	5.7 ± 2.7
Wild C_4/CAM	40	-12.0 ± 1.3	18	9.2 ± 5.2

plants ($t(79) = -6.58425$, $p = 0.00001$) and between wild and domestic C_4 plants ($t(40) = 9.99412$, $p = 0.0001$). These significant differences allowed us to treat these as separate food source groups within FRUITS (Phillips et al., 2014). The reasons for the difference between wild and domestic plant species are unknown and may be accounted for by sampling biases; however, we also theorize it might be related to the “canopy effect.” Domestically grown plants may have been cultivated in environments that were subject to land clearance and likely had less canopy cover compared with forest-dwelling wild plants; so therefore, had higher carbon values.

3 | RESULTS

3.1 | Model outcomes

We achieved results from all three models with varying degrees of predictive accuracy and usefulness for our research aims of assessing dietary linkages between humans and animals. Listed below are the source contribution estimates generated by FRUITS, including an assessment of the utility from each of the two-, three-, and four-source models (Tables 2 and 3).

3.1.1 | Two-source model results

Across all examined species, our two-source model predicted that C_3 plants dominated the diets of all the examined animals, except for one *I. portoricensis* from El Flaco (FL2413) of which C_4/CAM plants comprised 69.5% of its diet (Table 3, Figure 3). The only domesticated animal in our study, guinea pig (*C. porcellus*), had a diet composed of 38.8% (SD = 13.7%) C_4/CAM plants with only three *I. portoricensis* and the Hispaniolan hutia (*P. aedium*) from the site of La Entrada exceeding this. On average across all sites, *I. portoricensis* diets were composed of 71.6% (SD = 9.2%) C_3 plants, which is similar to edible rat (*Brotomys* sp.) diets that on average was made up of 75.6% (SD = 9.9%) C_3 plants. A two-tailed Student's *t* test between these two species revealed no significant difference in their mean C_3 plant consumption ($t(26) = 0.72612$, $p = 0.47425$), suggesting similar diets. These two species both form the majority of terrestrial vertebrate remains recovered from the site of El Flaco (Shev, Ali, et al., 2021).

Although no direct information regarding the consumption of domesticated plants are provided by the two-source model, two noteworthy conclusions can be drawn: C_4/CAM plants comprised more of the diets of all examined species than what was expected from observational and feeding studies that have been conducted for hutia species; and *Brotomys* sp. and *I. portoricensis* broadly had overlapping isotopic niches.

3.1.2 | Three-source model results

The three-source model provided estimates as to the proportion of C_4/CAM wild or domestic species that comprised the diets of our examined taxa (Figure 4). On average across all sites, the two species that consumed the most domesticated C_4/CAM plants were guinea pig, and surprisingly the endangered Hispaniolan hutia. Further analysis of more *P. aedium* samples would help us ascertain whether the frequent consumption of C_4/CAM plants is natural for this species, as well as further study of archaeological guinea pig will aid us in establishing if this domestic species is a reliable proxy for a human-controlled diet. The one *I. montanus* consumed the lowest amount of domestic C_4/CAM at 12.2% (SD = 7.8%), followed by *Brotomys* sp. which averaged 13%. Three *I. portoricensis* consumed domesticated C_4/CAM plants in amounts that exceeded 20% of their total diets (CA15 = 20.2%, SD = 11.4%; FL1952 = 30.9%, SD = 14.6%; FL2413 = 20.3%, SD = 15.5%), but for FL2413, wild C_4/CAM rather than domestic C_4/CAM plants formed the majority (49.32%, SD = 14.1%) of its diet.

These results suggest a varied diet for *I. portoricensis* with FL1952 and FL2413 serving as examples of two hutias with considerably high proportions of C_4/CAM in their diets, but with different consumption rates of domestic or wild species. Importantly, 39.2% ($n = 11$) of *I. portoricensis* consumed more domestic than wild C_4/CAM plant varieties, suggesting that for a large part of the hutia population domestic C_4/CAM crops were a more important food source than were wild C_4/CAM plants. The guinea pig also consumed domestic C_4/CAM plants as 21.2% (SD = 12.9%) of their total diet but also consumed similar proportions of wild C_4/CAM plants (22.3%, SD = 7.8%).

From the two-source to the three-source model across all examined species the average consumption of C_3 plants was reduced from 70% to 66.4% of the diet. Alongside, this is a reduction in % C_3 standard deviation from 10.3% to 7% when comparing the two- to three-source models, indicating higher fidelity in dietary estimations for the

TABLE 2 Median estimated percentages of dietary contribution sources for each sample according to the two-, three-, and four-source models

	Two-source model				Three-source model				Four-source model									
	C ₃	SD	C ₄ /CAM	SD	C ₃	SD	C ₄ /CAM domestic	SD	C ₄ /CAM wild	SD	C ₃ domestic	SD	C ₄ /CAM domestic	SD	C ₄ /CAM wild	SD		
	<i>Brotomys</i> sp.																	
CA8	78.25%	12.26%	21.75%	12.26%	70.30%	5.88%	11.20%	9.13%	18.51%	8.03%	31.24%	23.53%	48.26%	19.02%	8.90%	6.38%	11.60%	7.90%
CA10	69.25%	9.43%	30.75%	9.43%	64.57%	7.03%	12.88%	10.81%	22.55%	9.17%	34.49%	20.77%	43.97%	21.57%	10.84%	7.44%	10.70%	7.26%
CA24	78.73%	10.77%	21.27%	10.77%	71.58%	7.13%	14.32%	9.47%	14.11%	8.88%	53.82%	24.64%	28.45%	21.92%	9.36%	6.87%	8.36%	6.18%
FL1774	77.83%	9.34%	22.17%	9.34%	77.04%	7.93%	11.92%	8.24%	11.03%	8.07%	27.83%	22.43%	51.04%	19.45%	10.92%	7.40%	10.21%	7.63%
FL4025	74.02%	7.54%	25.98%	7.54%	72.54%	7.73%	14.77%	8.51%	12.68%	9.11%	20.31%	15.59%	53.68%	10.18%	14.08%	9.61%	11.94%	8.46%
<i>Cavia porcellus</i>																		
FL1664	61.21%	13.67%	38.79%	13.67%	56.45%	6.09%	21.24%	12.85%	22.31%	13.05%	24.54%	21.50%	37.99%	19.97%	19.41%	12.00%	18.06%	13.18%
<i>Isolobodon montanus</i>																		
FL3504	81.61%	9.10%	18.39%	9.10%	75.13%	7.45%	12.23%	8.42%	12.63%	7.82%	17.39%	13.19%	57.56%	12.75%	12.84%	8.23%	12.21%	8.36%
<i>Isolobodon portoricensis</i>																		
CA13	82.52%	9.64%	17.48%	9.64%	76.19%	6.79%	12.25%	7.58%	11.56%	8.61%	11.03%	8.07%	69.24%	10.66%	11.06%	8.37%	8.66%	6.88%
CA15	57.94%	7.37%	42.06%	7.37%	61.21%	7.11%	20.15%	11.37%	18.64%	11.44%	29.90%	19.18%	31.18%	16.39%	20.87%	10.27%	18.05%	11.55%
CA17	82.38%	7.97%	17.62%	7.97%	72.40%	6.35%	15.03%	8.84%	12.57%	8.63%	42.61%	24.72%	38.79%	21.59%	8.79%	7.32%	9.81%	7.39%
CA21	78.92%	9.95%	21.08%	9.95%	74.64%	7.76%	14.19%	8.01%	11.17%	8.33%	24.07%	14.21%	50.73%	13.48%	13.16%	8.46%	12.04%	8.81%
CA23	73.25%	9.26%	26.75%	9.26%	70.43%	6.92%	16.78%	9.71%	12.79%	9.83%	14.66%	13.08%	53.87%	12.66%	16.42%	10.69%	15.05%	9.44%
CA27	85.22%	9.49%	14.78%	9.49%	74.07%	7.77%	13.31%	9.22%	12.62%	8.13%	24.29%	19.36%	55.27%	18.23%	8.72%	7.52%	11.72%	8.61%
CA30	65.31%	8.39%	34.69%	8.39%	65.70%	9.18%	18.15%	10.47%	16.14%	11.22%	29.15%	12.74%	39.94%	11.88%	14.64%	10.81%	16.27%	10.72%
CA31	79.13%	8.13%	20.87%	8.13%	70.01%	9.52%	13.89%	11.83%	16.10%	9.37%	35.36%	21.13%	42.38%	21.07%	11.40%	8.22%	10.85%	7.09%
CA32	75.41%	10.66%	24.59%	10.66%	68.16%	7.81%	14.30%	10.95%	17.55%	9.49%	29.92%	16.81%	44.20%	12.86%	12.80%	8.44%	13.08%	8.32%
EN1	76.20%	9.23%	23.80%	9.23%	74.64%	7.96%	11.45%	8.78%	13.92%	8.53%	39.42%	24.29%	40.67%	20.58%	7.83%	6.62%	12.08%	8.56%
EN6	82.93%	8.10%	17.07%	8.10%	76.52%	7.47%	11.79%	7.80%	11.69%	8.09%	40.28%	23.87%	44.10%	19.77%	8.00%	7.33%	7.62%	5.68%
EN7	74.22%	10.28%	25.78%	10.28%	69.44%	8.08%	14.04%	9.84%	16.53%	8.63%	45.56%	21.60%	30.40%	16.36%	9.07%	7.15%	14.97%	9.07%
FL1264	80.53%	10.82%	19.47%	10.82%	73.76%	7.73%	12.99%	9.71%	13.26%	8.93%	36.39%	27.99%	41.51%	23.57%	11.15%	8.62%	10.95%	8.62%
FL1421	65.67%	9.37%	34.33%	9.37%	65.23%	9.80%	15.39%	10.65%	19.38%	10.45%	12.79%	8.17%	52.35%	10.10%	12.70%	9.62%	22.17%	13.52%
FL1543	71.31%	7.83%	28.69%	7.83%	69.06%	7.23%	13.45%	10.13%	17.49%	9.04%	23.85%	14.37%	48.81%	9.98%	12.79%	10.12%	14.55%	9.23%
FL1952	57.51%	13.65%	42.49%	13.65%	50.05%	7.07%	30.85%	14.62%	19.09%	16.12%	8.17%	5.82%	39.67%	8.61%	25.53%	15.73%	26.63%	17.91%
FL1976	74.36%	9.74%	25.64%	9.74%	68.10%	7.44%	15.85%	10.71%	16.05%	10.04%	24.37%	12.22%	53.88%	13.62%	11.08%	7.59%	10.67%	8.47%
FL2009	72.74%	10.62%	27.26%	10.62%	68.91%	7.44%	15.28%	10.52%	15.81%	9.48%	23.19%	16.83%	49.86%	13.61%	12.79%	9.69%	14.16%	9.29%
FL2413	30.51%	7.28%	69.49%	7.28%	30.38%	4.47%	20.30%	15.49%	49.32%	14.07%	17.56%	7.85%	14.41%	7.92%	14.71%	9.24%	53.32%	10.64%
FL2542	70.62%	7.93%	29.38%	7.93%	69.68%	7.12%	15.02%	9.10%	15.30%	8.80%	13.61%	9.49%	53.49%	9.03%	14.43%	9.60%	18.48%	11.80%

TABLE 2 (Continued)

	Two-source model				Three-source model				Four-source model										
	C ₃	SD	C ₄ /CAM	SD	C ₃	SD	C ₄ /CAM	SD	C ₄ /CAM	SD	C ₃		C ₄ /CAM		C ₄ /CAM				
											domestic	wild	domestic	wild	domestic	SD	domestic	SD	
FL26	68.90%	9.62%	31.10%	9.62%	70.75%	8.33%	15.90%	8.50%	13.34%	9.50%	19.99%	14.28%	52.02%	10.45%	15.01%	9.53%	12.98%	8.54%	
FL4075	80.57%	8.52%	19.43%	8.52%	76.01%	5.96%	10.93%	7.66%	13.06%	6.85%	31.35%	22.92%	49.99%	18.50%	10.38%	6.81%	8.29%	5.95%	
FL438	61.69%	8.00%	38.31%	8.00%	61.75%	6.37%	19.26%	11.51%	18.99%	11.61%	31.44%	20.52%	35.11%	15.67%	17.60%	10.74%	15.86%	11.16%	
<i>Plagiodontia aedium</i>																			
EN5	59.89%	9.61%	40.11%	9.61%	61.53%	6.82%	22.74%	11.22%	15.73%	11.30%	10.78%	8.47%	51.00%	12.45%	19.33%	11.90%	18.89%	13.51%	

three-source model. Nevertheless, a Student's *t* test was run across all samples examining the difference in % C₃ between the two- and three-source models for *I. portoricensis*, confirming that there was no statistically significant difference ($t(22) = 1.2114$, $p = 0.116105$) in the outcomes regarding C₃ consumption between these two models.

3.1.3 | Four-source model results

In dividing C₃ plants into domestic and wild varieties, we achieved remarkably different results compared with the two- and three-source models (Figure 5). It is likely that the addition of an extra dietary source decreased the overall predictive accuracy of FRUITS (Cheung & Szpak, 2020; Fernandes, 2016; Phillips et al., 2014; Stock et al., 2018), with the standard deviations for both wild and domestic C₃ sources being considerably high, such was the case for one *I. portoricensis* (FL1264, % domestic C₃ = 36.4%, SD = 28%). Domestic C₃ consumption was also high on average of all *Brotomys* sp. individuals at 33.5% of the diet, although again a standard deviation of 21.4% suggests a wide margin of error. The three individuals with the highest domestic C₃ consumption were an edible rat (CA24, 53.8%, SD = 24.6%) and *I. portoricensis* (CA17, 42.6%, SD = 24.7%) from El Carril and *I. portoricensis* from La Entrada (EN7, 45.6%, SD = 21.6%), although the accuracy of these results again is questionable due to the high standard deviations produced in this model.

Although C₃ wild and domestic source standard deviation ranges were high, domestic, and wild C₄/CAM sources appeared to have performed better with smaller standard deviations, averaging SD = 9.3% for domestic C₄/CAM and 10.2% for wild C₄/CAM across all species. To assess differences in dietary predictions between the three- and four-source models, a *t* test was run on the % domestic C₄/CAM for *I. portoricensis* samples, confirming significant differences between the two models ($t(23) = 2.09459$, $p = 0.020999$). Counter to this, % wild C₄/CAM contribution predictions for *I. portoricensis* were not significantly different between the three- and four-source models ($t(23) = 0.4172$, $p = 0.339279$).

3.2 | Species and site comparison

To assess if the three models provided differences according to site, *I. portoricensis* from El Flaco, El Carril, and La Entrada, and *Brotomys* sp. from El Flaco and El Carril were compared (Table 4). According to the two-source model for both species, there was considerable overlap in mean % C₃ and % C₄/CAM values between El Flaco and El Carril; however, there was less consumption of C₄/CAM by *I. portoricensis* recovered from the earlier occupied site of La Entrada (22.2%, SD = 3.7). The three-source model predicted that hutias consumed more domestic C₄/CAM plants at El Flaco (16.8%, SD = 5.1) and El Carril (15.3%, SD = 2.4) compared with edible rat from the same sites (13.4% and 12.8%, respectively). The four-source model again showed that hutias ate more C₄/CAM plants than edible rat generally.

TABLE 3 Mean food group contributions for each species according to each source model

Two sources								
	C ₃	SD	C ₄ /CAM	SD				
<i>Brotomys</i> sp.	75.60%	9.90%	24.40%	9.90%				
<i>Cavia porcellus</i>	61.20%	13.70%	38.80%	13.70%				
<i>Isolobodon montanus</i>	81.60%	9.10%	18.40%	9.10%				
<i>Isolobodon portoricensis</i>	71.60%	9.20%	28.40%	9.20%				
<i>Plagiodontia aedium</i>	59.90%	9.60%	40.10%	9.60%				
All species	70.00%	10.30%	30.00%	10.30%				
Three sources								
	C ₃	SD	C ₄ wild	SD	C ₄ dom	SD		
<i>Brotomys</i> sp.	71.20%	7.10%	15.80%	9.20%	13.00%	8.70%		
<i>Cavia porcellus</i>	56.50%	6.10%	22.30%	12.90%	21.20%	13.10%		
<i>Isolobodon montanus</i>	75.10%	7.50%	12.60%	8.40%	12.20%	7.80%		
<i>Isolobodon portoricensis</i>	67.70%	7.50%	16.60%	10.10%	15.70%	9.80%		
<i>Plagiodontia aedium</i>	61.50%	6.80%	15.70%	11.20%	22.70%	11.30%		
All species	66.40%	7.00%	16.60%	10.40%	17.00%	10.10%		
Four sources								
	C ₃ dom	SD	C ₃ wild	SD	C ₄ dom	SD	C ₄ wild	SD
<i>Brotomys</i> sp.	33.50%	21.40%	45.10%	18.40%	10.80%	7.50%	10.60%	7.50%
<i>Cavia porcellus</i>	24.50%	21.50%	38.00%	20.00%	19.40%	12.00%	18.10%	13.20%
<i>Isolobodon montanus</i>	17.40%	13.20%	57.60%	12.80%	12.80%	8.20%	12.20%	8.40%
<i>Isolobodon portoricensis</i>	39.40%	24.30%	40.70%	20.60%	7.80%	6.60%	12.10%	8.60%
<i>Plagiodontia aedium</i>	10.80%	8.50%	51.00%	12.50%	19.30%	11.90%	18.90%	13.50%
All species	25.10%	17.80%	46.50%	16.80%	14.00%	9.30%	14.40%	10.20%

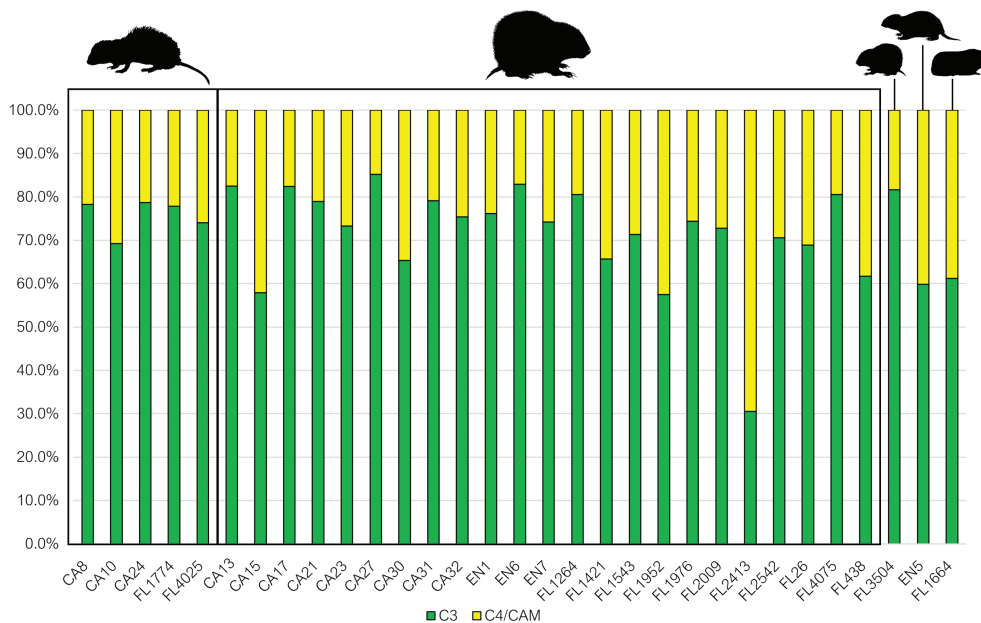
**FIGURE 3** Median dietary source contribution percentage estimates according to the two-source model. Taxa depicted from left to right are edible rat (*Brotomys* sp.), Puerto Rican hutia (*Isolobodon portoricensis*), montane hutia (*Isolobodon montanus*), Hispaniolan hutia (*Plagiodontia aedium*), and guinea pig (*Cavia porcellus*). Sample number prefixes indicate the site of origin: CA = El Carril, FL = El Flaco, EN = La Entrada [Colour figure can be viewed at wileyonlinelibrary.com]

FIGURE 4 Median dietary source contribution percentage estimates according to the three-source model. Taxa depicted from left to right are edible rat (*Brotomys* sp.), Puerto Rican hutia (*Isolobodon portoricensis*), montane hutia (*Isolobodon montanus*), Hispaniolan hutia (*Plagiodontia aedium*), and guinea pig (*Cavia porcellus*). Sample number prefixes indicate the site of origin: CA = El Carril, FL = El Flaco, EN = La Entrada [Colour figure can be viewed at wileyonlinelibrary.com]

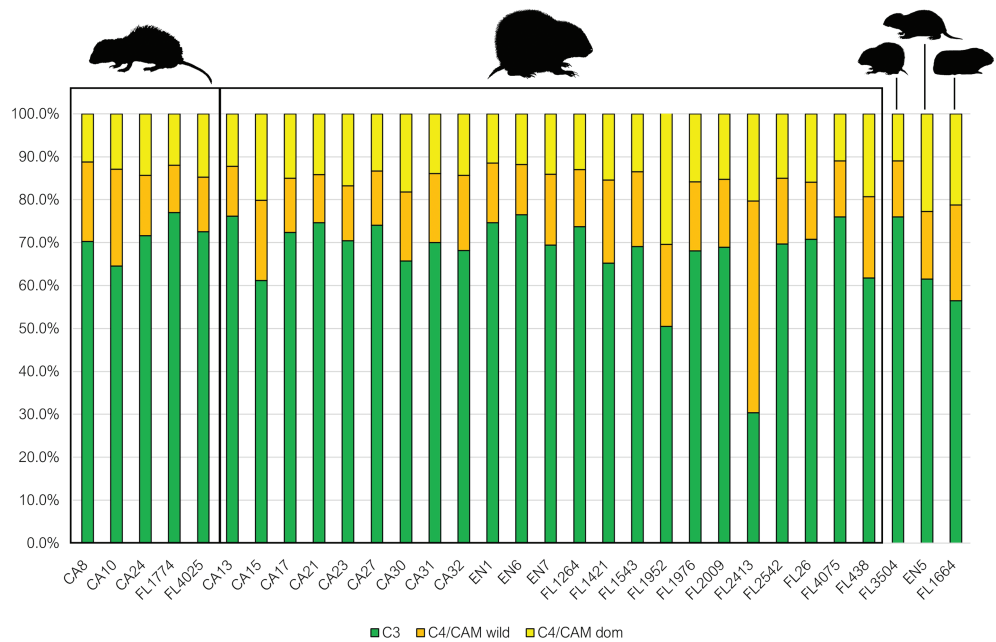
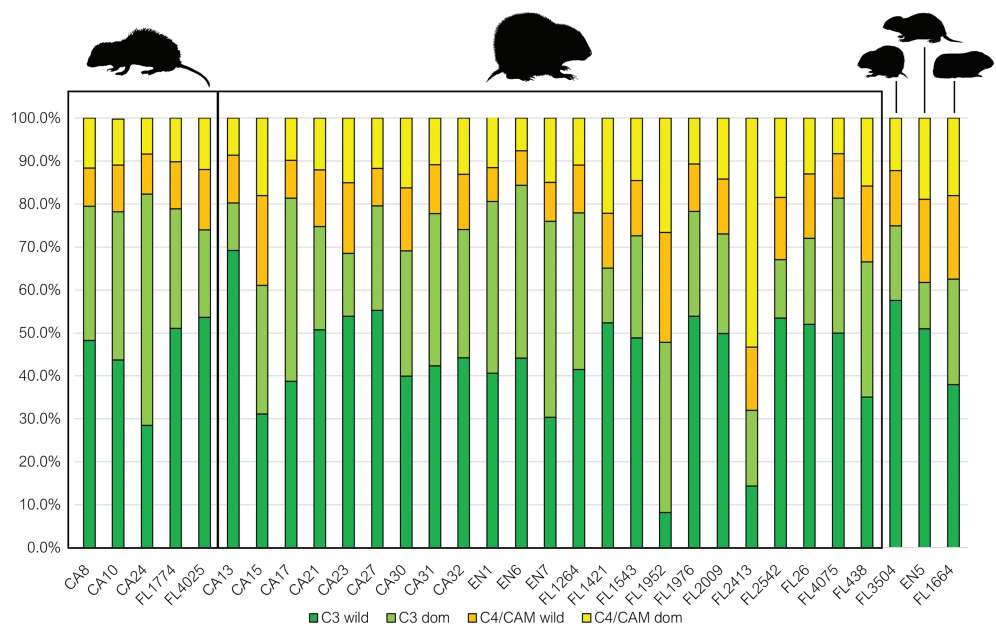


FIGURE 5 Median dietary source contribution percentage estimates according to the four-source model. Taxa are depicted from left to right: edible rat (*Brotomys* sp.), Puerto Rican hutia (*Isolobodon portoricensis*), montane hutia (*Isolobodon montanus*), Hispaniolan hutia (*Plagiodontia aedium*), and guinea pig (*Cavia porcellus*). Sample number prefixes illustrate the site of origin: CA = El Carril, FL = El Flaco, EN = La Entrada [Colour figure can be viewed at wileyonlinelibrary.com]



The results from the three- and four-source models generally indicate that *I. portoricensis* consumed more of both domestic and wild C_4 /CAM sources at El Flaco and El Carril than did edible rat. There also appears to be a difference in diet for *I. portoricensis* recovered from earlier dating La Entrada, with this species consuming less C_4 /CAM plants there than the later-occupied, inland sites.

3.3 | Comparisons the dietary source contributions of domestic guinea pig and wild hutia

We directly compared the dietary contribution estimates for one hutia (FL1952) that had the highest amount of C_4 /CAM consumption across all three models, and the one guinea pig recovered from El

Flaco (FL1664). This was done to assess whether this one hutia had similar dietary contributions to the only domesticated species sampled in our study (Figure 6).

The three-source model predicted overlapping contributions of all plant types between the two species, although the guinea pig likely consumed more C_3 plants and lesser quantities of domestic C_4 /CAM plants than the *I. portoricensis*. According to the four-source model, the hutia consumed more domestic than wild C_4 /CAM plants but compared with the guinea pig this hutia consumed greater quantities of wild C_3 than domestic C_3 plants. There is considerable overlap between both individuals suggesting similar diets. Nevertheless, this hutia likely consumed more domestic C_4 /CAM plants than the guinea pig according to both models, but less domestic C_3 plants than the guinea pig according to the four-source model; however, we consider

TABLE 4 Comparison of *Isolobodon portoricensis* and *Brotomys* sp. mean source consumption percentages across El Flaco, El Carril and La Entrada

	Two sources						Three sources						Four sources																																										
	C ₃		C ₄		SD		C ₃		C ₄		SD		C ₃ dom		C ₄ wild		SD		C ₃ wild		C ₄ dom		SD		C ₃ wild		C ₄ wild		SD																										
<i>Isolobodon portoricensis</i>																																																							
El Flaco	66.76%	13.28%	33.24%	13.28%	13.28%	63.97%	12.49%	16.84%	5.09%	19.19%	9.79%	22.06%	8.34%	44.65%	11.24%	14.38%	4.05%	18.91%	12.00%	75.56%	8.39%	24.44%	8.39%	4.48%	15.34%	2.39%	14.35%	2.61%	26.78%	9.18%	47.29%	10.65%	13.09%	3.63%	12.84%	2.91%																			
El Carril	77.78%	3.73%	22.22%	3.73%	73.53%	69.27%	8.47%	26.63%	8.47%	1.91%	24.08%	4.36%	24.59%	4.36%	68.82%	71.80%	2.65%	13.07%	1.35%	15.12%	2.14%	31.96%	6.86%	46.29%	4.91%	11.10%	1.20%	10.65%	1.11%	11.56%	3.02%	11.56%	3.02%	11.56%	3.02%	11.56%	3.02%																		
La Entrada	73.37%	8.47%	26.63%	8.47%	69.27%	69.27%	6.65%	14.87%	2.88%	15.86%	4.79%	30.20%	6.74%	43.44%	9.24%	11.92%	2.74%	14.44%	5.98%	73.37%	8.47%	26.63%	8.47%	6.65%	14.87%	2.88%	15.86%	4.79%	30.20%	6.74%	43.44%	9.24%	11.92%	2.74%	14.44%	5.98%	73.37%	8.47%	26.63%	8.47%	6.65%	14.87%	2.88%	15.86%	4.79%	30.20%	6.74%	43.44%	9.24%	11.92%	2.74%	14.44%	5.98%		
<i>Brotomys</i> sp.																																																							
El Flaco	75.93%	1.91%	24.08%	1.91%	74.79%	74.79%	2.25%	13.35%	1.42%	11.86%	0.83%	24.07%	3.76%	52.36%	1.32%	12.50%	1.58%	11.08%	0.87%	75.41%	4.36%	24.59%	4.36%	3.05%	12.80%	1.27%	18.39%	3.45%	39.85%	9.97%	40.23%	8.51%	9.70%	0.83%	10.22%	1.36%	75.67%	3.13%	24.33%	3.13%	71.80%	2.65%	13.07%	1.35%	15.12%	2.14%	31.96%	6.86%	46.29%	4.91%	11.10%	1.20%	10.65%	1.11%	
El Carril	75.41%	4.36%	24.59%	4.36%	68.82%	71.80%	2.65%	13.07%	1.35%	15.12%	2.14%	31.96%	6.86%	46.29%	4.91%	11.10%	1.20%	10.65%	1.11%	75.67%	3.13%	24.33%	3.13%	71.80%	2.65%	13.07%	1.35%	15.12%	2.14%	31.96%	6.86%	46.29%	4.91%	11.10%	1.20%	10.65%	1.11%	75.67%	3.13%	24.33%	3.13%	71.80%	2.65%	13.07%	1.35%	15.12%	2.14%	31.96%	6.86%	46.29%	4.91%	11.10%	1.20%	10.65%	1.11%
Total	75.67%	3.13%	24.33%	3.13%	71.80%	71.80%	2.65%	13.07%	1.35%	15.12%	2.14%	31.96%	6.86%	46.29%	4.91%	11.10%	1.20%	10.65%	1.11%	75.67%	3.13%	24.33%	3.13%	71.80%	2.65%	13.07%	1.35%	15.12%	2.14%	31.96%	6.86%	46.29%	4.91%	11.10%	1.20%	10.65%	1.11%	75.67%	3.13%	24.33%	3.13%	71.80%	2.65%	13.07%	1.35%	15.12%	2.14%	31.96%	6.86%	46.29%	4.91%	11.10%	1.20%	10.65%	1.11%

the results for the four-source model as less reliable than the three-source model, which we further elaborate upon below.

4 | DISCUSSION

All three models demonstrate some convergence regarding the premise that hutias and edible rats were consuming C₄/CAM plants, and for the three- and four-source models, that much of this was likely domesticated maize. This suggests that FRUITS is providing some accurate overview of the dietary behavior of the animals investigated, although the exact precision of the dietary estimates of each model is difficult to assess due to the extinct status of much of the fauna, and the use of independent dietary evidence gleaned from other sources. Discrepancies between models raise questions as to which model was the most accurate. The two-source model was theoretically the most precise due to it assessing only two dietary sources, although as our aim was to assess whether animals were consuming domestic crops this model shows limited utility for our study. In actuality, the three-source model may in fact be more accurate in determining C₃ percentages according to the observably lower standard deviations generated from this model when compared with the two-source model. In our opinion, the three-source model best achieved our goal of differentiating between domestic C₄/CAM plants (mostly agricultural maize) and wild C₄/CAM plants. Even so, the four-source model in our opinion is the most unreliable. Beyond the fact that increasing the amount of sources will likely decrease the predictive accuracy of FRUITS (Cheung & Szpak, 2020; Fernandes et al., 2014; Pestle & Laffoon, 2018; Stock et al., 2018), there are other reasons why the splitting of C₃ plants into wild and domestic varieties is problematic and led to wide-ranging predictions. First, the source plant data suggest a wider range of carbon values for C₃ plants ($\delta^{13}\text{C}$ –36.6‰ to –22.6‰, SD = 3.5) than for C₄ plants (–16.2‰ to –9.2‰ SD = 1.4), likely reducing predictive accuracy, as when splitting C₃ plants into two different source groups there was still high variability and overlap in isotopic values between these new groups. Second, dividing C₃ plants into domestic and wild species ignores the possibility that some domestic varieties that have been cultivated by humans may self-propagate in the wild without human involvement, complicating interpretations when trying to determine human influence over hutia dietary behavior. In effect, the consumption of wild or uncultivated but domesticated fruit tree species would lead to an equifinality in dietary isotope values. As domestic maize is not self-propagating, and is an introduced species that does not occur naturally in the wild, this problem does not arise in the three-source model that only splits C₄/CAM plants into domestic and wild groupings.

Our study demonstrates the potential for using FRUITS or other isotopic dietary mixing models to differentiate between domestic and wild plant dietary source contributions for synanthropic herbivores. There are however some limitations we encountered which cannot be overlooked. We had to gather the isotopic values of wild taxa from a variety of sources to bolster the collection of isotopic values from published sources that constituted our isotopic foodweb for

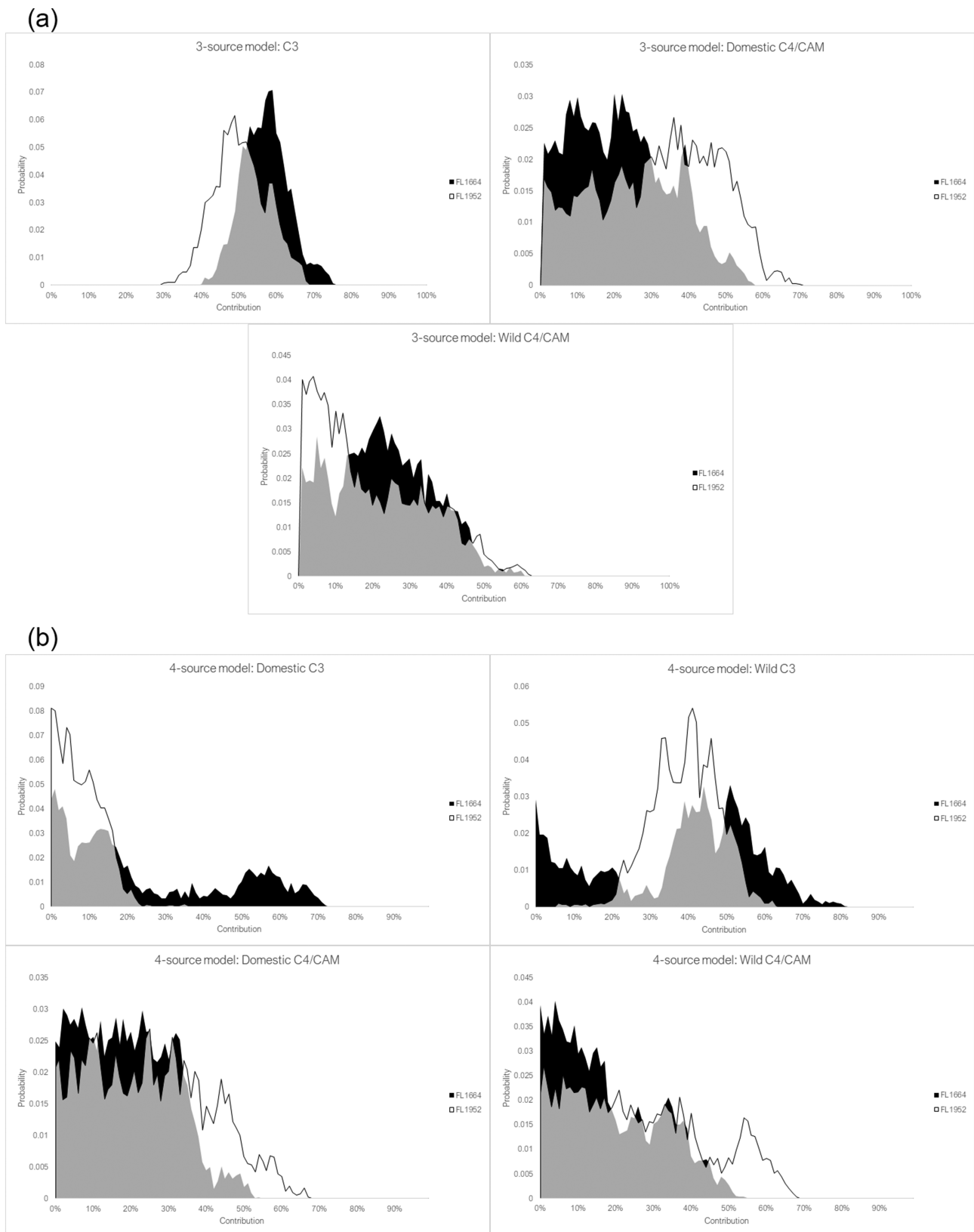


FIGURE 6 Contributions for *Isolobodon portoricensis* (FL1952) and guinea pig (FL1664) according to the (a) three-source model and (b) four-source model

Hispaniola. Higher accuracy would be achieved if observational feeding studies of studied species can be relied upon to precisely indicate what plants an animal will consume in the wild. As three of the five studied rodent species are now extinct, this information is unfortunately unavailable; therefore, we had to rely on feedings studies of other, extant hutia species to indicate their dietary preferences. Additionally, our application of FRUITS in trying to disseminate between “wild” and “domestic” food sources is perhaps best applied for herbivores only, as the inclusion of other sources such as terrestrial or marine proteins may generate too much overlap in values and confuse dietary contribution predictions.

According to all three models, there are notable differences in *I. portoricensis* diets from the sites of El Carril and El Flaco, and that of La Entrada, with hutias from the latter site likely consuming less C_4 /CAM plants. Although this might relate to the smaller sample size of La Entrada hutias, this may also be symptomatic of differences in food availability or the intensity of agricultural production between two different regions. La Entrada is situated along the northern coastline of Hispaniola and dates to the 9th century AD, with a faunal assemblage largely composed of marine foods such as fish, shellfish, and sea turtles (Chelonidae). El Flaco and El Carril are located within viewshed of one another and are nestled in the southern foothills of the Cordillera Septentrional overlooking the agriculturally productive Cibao Valley. The inhabitants of El Flaco and El Carril likely had stronger reliance on terrestrial animals according to zooarchaeological studies (Shev, Ali, et al., 2021) and were situated within an agriculturally productive region overlooking the fertile Cibao Valley from which sediment cores provide ample evidence for anthropogenic fire usage (Castilla-Beltrán et al., 2018, 2020; Hooghiemstra et al., 2018). It is possible that horticultural practices attracted hutias to the site and may indicate that this species was synanthropic with their predation by humans mostly reflecting garden hunting practices. This may explain the higher consumption of domestic C_4 /CAM plant in samples from these two sites. The inhabitants of La Entrada, because of its earlier date and coastal situation, may have not been as focused on agricultural production as they were on harvesting marine resources, which may be reflected in the lower percentage of C_4 /CAM plant consumption in *I. portoricensis* seen there.

The one analyzed Hispaniolan hutia (*P. aedium*) from La Entrada had the fourth highest contribution from C_4 /CAM plants. This same individual also likely ate mostly wild C_3 plants according to the four-source model, with equitable contributions of both wild and domestic C_4 /CAM plant sources. The high C_4 /CAM contribution is surprising, given that this species has not been observed consuming any C_4 /CAM plants apart from maize in captivity (Borroto-Páez & Woods, 2012; Eisenberg & Woods, 2012), and had the lowest mean $\delta^{13}C_{en}$ results for any hutia in the study by Cooke and Crowley (2018). We posit that the results for *P. aedium* from La Entrada may relate to an opportunistic consumption of marine foods, such as fish or shellfish, given the site's coastal location, and highlight the limitation of not including animal meats within the model. However, we acknowledge that this specimen is likely an outlier and that hutia diets are almost exclusively herbivorous. Further studies of the behavior

and isotope ecology of *P. aedium* are needed to precisely assess its dietary behavior.

When we compare our findings from previously conducted feeding studies on hutias, and the isotopic niche predictions of Hispaniolan rodents by Cooke and Crowley (2018), there is overall a greater predicted consumption of C_4 /CAM sources than expected for all examined endemic species. Nevertheless, this may be an artifact of FRUITS, as the inclusion of any source will mean that it will automatically generate some percentage estimation as part of the whole diet, even if that source was never actually consumed (Fernandes et al., 2014). Cooke and Crowley's (2018) prediction of edible rat (*Brotomys* sp.) being more frugivorous may be correct as we see high estimates of C_3 plant consumption with this species. Our three-source model, deemed by us to be the most reliable, indicates that *I. portoricensis* on average consumed more C_4 /CAM plants than edible rat, although with a high degree of variance between individual hutias of this species. These two species appear to have some considerable overlap in isotopic niches and were likely sympatric but may have not necessarily competed for food sources, with *I. portoricensis* perhaps being more ground-dwelling and *Brotomys* sp. more arboreal in nature (Cooke & Crowley, 2018) but with both species consuming plants with similar isotopic values. It is possible that both species were somewhat synanthropic and profited from human landscape changes such as slash-and-burn farming and the cultivation of garden plots and fruits trees.

The guinea pig evidently consumed significant quantities of both domestic and wild C_4 /CAM plants and can arguably be used as a proxy for what a domesticated rodent diet from precolonial contexts should look like. The FRUITS model outcomes suggest that there were no systematic feeding strategies affecting hutia diets; therefore, it is unlikely that the diets of all hutias were regimented by humans in the same way that the guinea pig likely was. Some individual hutias nevertheless were consuming C_4 /CAM plants beyond what would be expected of them from hutia feeding studies and in similar degrees to that of the guinea pig.

The *I. portoricensis* individuals that consumed high amounts of C_4 /CAM may have been consuming maize frequently enough to affect their isotopic values. It is a possibility that this may reflect some degree of dietary seasonality, with FL1952 perhaps exhibiting isotopic values that are reflective of its life cycle coinciding with maize growing and harvest seasons whereas other individuals did not. There are perhaps other scenarios in which an equifinality in isotopic values may have arisen due to unexpected dietary inputs for these extinct species. For example, it may be the case that some animals were consuming more wild plants of a variety that yields higher carbon values than the wild C_4 /CAM plants investigated in this study, contrary to what modern feeding studies would suggest about extant hutia dietary behavior. Notwithstanding these possibilities, we believe it is a more likely scenario that some individual hutias were being purposefully fed agricultural maize while most others were opportunistically feeding on maize crops. It is likely that horticultural practices such as slash-and-burn farming may have led to the creation of mosaic plant communities. These anthropogenic environments likely supported and

bolstered populations of hutias, and possibly edible rats, which profited from these anthropogenic environments and drew them into closer contact with humans.

An expansion of this study to include samples from other sites in Hispaniola, both coastally and inland situated, would be greatly beneficial for assessing whether there were differences in food production strategies at these different locales. We expect that there may be a broad trend of more domestic plant consumption by commensal species in locations where the food production was largely focused on horticulture, compared with coastal sites that may have been more reliant on the gathering of marine resources. This method allows greater insight into the dietary effects on animals resulting from human niche construction activities beyond what can only be inferred from the study of animals remains from archaeological sites at which hutia remains often comprise a significant amount of the faunal assemblage (Shev, Ali, et al., 2021).

The implementation of FRUITS to examine dietary linkages between humans and animals can be extended to other archaeological contexts and islands in the Caribbean. In the Lesser Antilles, there is some evidence to suggest that rice rats (*Oryzomys* spp.) may have also been introduced either intentionally or unintentionally to new island environments and likely served as an important food source in some islands that were relatively depauperate of other terrestrial fauna (Brace et al., 2015; Durocher et al., 2021). It is probable that these species served similar roles as hutias did in the Greater Antilles.

5 | CONCLUSIONS

The functioning of animal management strategies at the examined sites remains opaque, and there are a multitude of explanations for why some hutias had diets that were composed of considerable amounts of domestic C_4 /CAM plants. It is probable that most hutias were attracted to and opportunistically feeding from human garden plots, given that domestic plant consumption is well demonstrated in the FRUITS modeling, and more so for *I. portoricensis* than for edible rats. This in no way indicates that all hutias were being kept in captivity, although some individuals consumed domestic C_4 /CAM plants in such high proportions that are unlikely to be the result of an unfettered opportunistic consumption of maize from garden plots; therefore, indicating that these individuals had diets that were supplemented by humans. This may constitute a degree of animal management in the form of the purposeful feeding of some hutia to possibly attract others of the same species close to human settlements.

This study highlights the applicability of multiple stable isotope analyses and dietary mixing models (FRUITS) in the study of dietary linkages between humans and endemic animals in the Neotropics. This novel approach likely has more pertinency in situations where the examined species are herbivorous, allowing us to reduce to amount of potential food source groups; therefore, increasing the predictive accuracy of the dietary mixing models. Additionally, the prevalence of C_4 maize in the Americas as a ubiquitous food crop enabled

us to easily distinguish between “wild” and “domestic” food groups in archaeological settings. Therefore, this application of FRUITS on herbivorous rodent species, like hutias and edible rats, appears to accurately convey that there was some degree of human involvement variably affecting the diets of animals at the studied sites.

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CONFLICT OF INTEREST

The authors have no conflict of interest to declare especially regarding the use of funding, the acquisition and analysis of materials the results discussed in this paper.

DATA AVAILABILITY STATEMENT

The data that support the findings of this study are available in the Supporting Information of this article.

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REFERENCES

- Allen, J. A. (1916). An extinct octodont from the island of Porto Rico, West Indies. *Annals of the New York Academy of Sciences*, 27, 17–22. <https://doi.org/10.1111/j.1749-6632.1916.tb55182.x>
- Ambrose, S. H. (2002). Controlled diet and climate experiments on nitrogen isotope ratios of rats. In S. H. Ambrose & M. A. Katzenberg (Eds.), *Biogeochemical approaches to paleodietary analysis* (pp. 243–259). Springer US. https://doi.org/10.1007/0-306-47194-9_12
- Ambrose, S. H., Butler, B. M., Hanson, D. B., Hunter-Anderson, R. L., & Krueger, H. W. (1997). Stable isotopic analysis of human diet in the Marianas Archipelago, Western Pacific. *American Journal of Physical Anthropology*, 104, 343–361. [https://doi.org/10.1002/\(SICI\)1096-8644\(199711\)104:3<343::AID-AJPA5>3.0.CO;2-W](https://doi.org/10.1002/(SICI)1096-8644(199711)104:3<343::AID-AJPA5>3.0.CO;2-W)

- Ambrose, S. H., & Norr, L. (1993). Experimental evidence for the relationship of the carbon isotope ratios of whole diet and dietary protein to those of bone collagen and carbonate. In J. B. Lambert & G. Grupe (Eds.), *Prehistoric human bone: Archaeology at the molecular level* (pp. 1–37). Springer. https://doi.org/10.1007/978-3-662-02894-0_1
- Arce-Peña, N. P., Arroyo-Rodríguez, V., San-José, M., Jiménez-González, D., Franch-Pardo, I., Andresen, E., & Ávila-Cabadilla, L. D. (2019). Landscape predictors of rodent dynamics in fragmented rainforests. *Biodiversity and Conservation*, 28, 655–669. <https://doi.org/10.1007/s10531-018-1682-z>
- Barton, L., Newsome, S. D., Chen, F.-H., Wang, H., Guilderson, T. P., & Bettinger, R. L. (2009). Agricultural origins and the isotopic identity of domestication in northern China. *Proceedings of the National Academy of Sciences*, 106, 5523–5528. <https://doi.org/10.1073/pnas.0809960106>
- Blumenthal, S. A., Rothman, J. M., Chritz, K. L., & Cerling, T. E. (2016). Stable isotopic variation in tropical forest plants for applications in primatology. *American Journal of Primatology*, 78, 1041–1054. <https://doi.org/10.1002/ajp.22488>
- Bocherens, H., Sandrock, O., Kullmer, O., & Schrenk, F. (2011). Hominin palaeoecology in Late Pliocene Malawi: First insights from isotopes (^{13}C , ^{18}O) in mammal teeth. *South African Journal of Science*, 107(3), 1–6. <https://doi.org/10.4102/sajs.v107i3.4.331>
- Boivin, N. L., Zeder, M. A., Fuller, D. Q., Crowther, A., Larson, G., Erlandson, J. M., Denham, T., & Petraglia, M. D. (2016). Ecological consequences of human niche construction: Examining long-term anthropogenic shaping of global species distributions. *Proceedings of the National Academy of Sciences*, 113, 6388–6396. <https://doi.org/10.1073/pnas.1525200113>
- Bond, A. L., & Diamond, A. W. (2011). Recent Bayesian stable-isotope mixing models are highly sensitive to variation in discrimination factors. *Ecological Applications*, 21, 1017–1023. <https://doi.org/10.1890/09-2409.1>
- Borroto-Páez, R., & Woods, C. A. (2012). *Feeding habits of the capromyid rodents* (pp. 71–92). Florida Museum of Natural History and Wacahoota Press.
- Brace, S., Turvey, S. T., Weksler, M., Hoogland, M. L. P., & Barnes, I. (2015). Unexpected evolutionary diversity in a recently extinct Caribbean mammal radiation. *Proceedings of the Biological Sciences*, 282, 20142371. <https://doi.org/10.1098/rspb.2014.2371>
- Castilla-Beltrán, A., Hooghiemstra, H., Hoogland, M. L. P., Donders, T. H., Pagán-Jiménez, J. R., McMichael, C. N. H., Rolefes, S. M. F., Olijhoek, T., Herrera-Malatesta, E., Hung, J. U., & Hofman, C. L. (2020). Ecological responses to land use change in the face of European colonization of Haytí island. *Quaternary Science Reviews*, 241, 106407. <https://doi.org/10.1016/j.quascirev.2020.106407>
- Castilla-Beltrán, A., Hooghiemstra, H., Hoogland, M. L. P., Pagán-Jiménez, J., van Geel, B., Field, M. H., Prins, M., Donders, T., Herrera Malatesta, E., Ulloa Hung, J., McMichael, C. H., Gosling, W. D., & Hofman, C. L. (2018). Columbus' footprint in Hispaniola: A paleoenvironmental record of indigenous and colonial impacts on the landscape of the central Cibao Valley, northern Dominican Republic. *Anthropocene*, 22, 66–80. <https://doi.org/10.1016/j.ancene.2018.05.003>
- Cheung, C., & Szpak, P. (2020). Interpreting past human diets using stable isotope mixing models. *Journal of Archaeological Method and Theory*, 28, 1106–1142. <https://doi.org/10.1007/s10816-020-09492-5>
- Chinique de Armas, Y., Buhay, W. M., Rodríguez Suárez, R., Bestel, S., Smith, D., Mowat, S. D., & Roksandic, M. (2015). Starch analysis and isotopic evidence of consumption of cultigens among fisher-gatherers in Cuba: The archaeological site of Canimar Abajo, Matanzas. *Journal of Archaeological Science*, 58, 121–132. <https://doi.org/10.1016/j.jas.2015.03.003>
- Chinique de Armas, Y., Mavridou, A.-M., Garcell Domínguez, J., Hanson, K., & Laffoon, J. (2022). Tracking breastfeeding and weaning practices in ancient populations by combining carbon, nitrogen and oxygen stable isotopes from multiple non-adult tissues. *PLoS ONE*, 17, e0262435. <https://doi.org/10.1371/journal.pone.0262435>
- Chinique de Armas, Y., & Pestle, W. (2018). Assessing the association between subsistence strategies and the timing of weaning among indigenous archaeological populations of the Caribbean. *International Journal of Osteoarchaeology*, 28, 492–509. <https://doi.org/10.1002/oa.2695>
- Chinique de Armas, Y., Roksandic, M., Nikitović, D., Suárez, R. R., Smith, D., Kanik, N., Jordá, D. G., & Buhay, W. M. (2017). Isotopic reconstruction of the weaning process in the archaeological population of Canimar Abajo, Cuba: A Bayesian probability mixing model approach. *PLoS ONE*, 12, e0176065. <https://doi.org/10.1371/journal.pone.0176065>
- Chinique de Armas, Y., Roksandic, M., Suárez, R. R., Smith, D. G., & Buhay, W. M. (2016). Isotopic evidence of variations in subsistence strategies and food consumption patterns among “fisher-gatherer” populations of Western Cuba. In *Cuban archaeology in the Caribbean*. University Press of Florida. <https://doi.org/10.5744/florida/9781683400028.003.0009>
- Ciofalo, A. J., Sinelli, P. T., & Hofman, C. L. (2019). Late precolonial culinary practices: Starch analysis on griddles from the northern Caribbean. *Journal of Archaeological Method and Theory*, 26, 1632–1664. <https://doi.org/10.1007/s10816-019-09421-1>
- Ciofalo, A. J., Sinelli, P. T., & Hofman, C. L. (2020). Starchy shells: Residue analysis of precolonial northern Caribbean culinary practices. *Archaeometry*, 62, 362–380. <https://doi.org/10.1111/arc.12524>
- Cooke, S. B., & Crowley, B. E. (2018). Deciphering the isotopic niches of now-extinct Hispaniolan rodents. *Journal of Vertebrate Paleontology*, 38, e1510414. <https://doi.org/10.1080/02724634.2018.1510414>
- Courcelle, M., Tilak, M.-K., Leite, Y. L. R., Douzery, E. J. P., & Fabre, P.-H. (2019). Digging for the spiny rat and hutia phylogeny using a gene capture approach, with the description of a new mammal subfamily. *Molecular Phylogenetics and Evolution*, 136, 241–253. <https://doi.org/10.1016/j.ympev.2019.03.007>
- de Las Casas, B. (1875). *Historia de las Indias, Tomo I*. Imprenta de Miguel Ginesta.
- del Rio, C. M., Wolf, N., Carleton, S. A., & Gannes, L. Z. (2009). Isotopic ecology ten years after a call for more laboratory experiments. *Biological Reviews*, 84, 91–111. <https://doi.org/10.1111/j.1469-185X.2008.00064.x>
- DeNiro, M. J., & Epstein, S. (1978). Influence of diet on the distribution of carbon isotopes in animals. *Geochimica et Cosmochimica Acta*, 42, 495–506. [https://doi.org/10.1016/0016-7037\(78\)90199-0](https://doi.org/10.1016/0016-7037(78)90199-0)
- DeNiro, M. J., & Epstein, S. (1981). Influence of diet on the distribution of nitrogen isotopes in animals. *Geochimica et Cosmochimica Acta*, 45, 341–351. [https://doi.org/10.1016/0016-7037\(81\)90244-1](https://doi.org/10.1016/0016-7037(81)90244-1)
- Durocher, M., Nicolas, V., Perdikaris, S., Bonnissent, D., Robert, G., Debue, K., Evin, A., & Grouard, S. (2021). Archaeobiogeography of extinct rice rats (*Oryzomyini*) in the Lesser Antilles during the Ceramic Age (500 BCE–1500 CE). *Holocene (Sevenoaks)*, 31, 433–445. <https://doi.org/10.1177/0959683620972785>
- Ehleringer, J. R. (1989). Carbon isotope ratios and physiological processes in aridland plants. In *Stable isotopes in ecological research* (pp. 41–54). Springer. https://doi.org/10.1007/978-1-4612-3498-2_3
- Eisenberg, J. F., & Woods, C. A. (2012). Review of captive studies of Capromyidae with comments on their natural history. In *Terrestrial mammals of the West Indies: Contributions* (pp. 143–150). Florida Museum of Natural History and Wacahoota Press.
- Fabre, P.-H., Vilstrup, J. T., Raghavan, M., Der Sarkissian, C., Willerslev, E., Douzery, E. J. P., & Orlando, L. (2014). Rodents of the Caribbean: Origin and diversification of hutias unravelled by next-generation museomics. *Biology Letters*, 10, 20140266. <https://doi.org/10.1098/rsbl.2014.0266>

- Fernandes, R. (2016). A simple(R) model to predict the source of dietary carbon in individual consumers. *Archaeometry*, 58, 500–512. <https://doi.org/10.1111/arc.12193>
- Fernandes, R., Grootes, P., Nadeau, M.-J., & Nehlich, O. (2015). Quantitative diet reconstruction of a Neolithic population using a Bayesian mixing model (FRUITS): The case study of Ostorf (Germany). *American Journal of Physical Anthropology*, 158, 325–340. <https://doi.org/10.1002/ajpa.22788>
- Fernandes, R., Millard, A. R., Brabec, M., Nadeau, M.-J., & Grootes, P. (2014). Food Reconstruction Using Isotopic Transferred Signals (FRUITS): A Bayesian model for diet reconstruction. *PLoS ONE*, 9, e87436. <https://doi.org/10.1371/journal.pone.0087436>
- Fernandes, R., Nadeau, M.-J., & Grootes, P. M. (2012). Macronutrient-based model for dietary carbon routing in bone collagen and bioapatite. *Archaeological and Anthropological Sciences*, 4, 291–301. <https://doi.org/10.1007/s12520-012-0102-7>
- Figueredo, A. E. (2015). Manioc dethroned and maize triumphant: Interpretations on the ethnohistory and archaeology of the Bahamas with sundry notes on relations of production. *Journal of Caribbean Archaeology*, 15, 120–134.
- Flemming, C., & MacPhee, R. D. E. (1999). *Redetermination of holotype of Isolobodon portoricensis (Rodentia, Capromyidae): With notes on Recent mammalian extinctions in Puerto Rico*. American Museum Novitates; no. 3278. American Museum of Natural History.
- Francey, R. J., Allison, C. E., Etheridge, D. M., Trudinger, C. M., Enting, I. G., Leuenberger, M., Langenfelds, R. L., Michel, E., & Steele, L. P. (1999). A 1000-year high precision record of $\delta^{13}\text{C}$ in atmospheric CO_2 . *Tellus Series B: Chemical and Physical Meteorology*, 51, 170–193. <https://doi.org/10.3402/tellusb.v51i2.16269>
- Frias, A. I., & Hernandez, N. (1985). Observaciones sobre conducta alimentaria en jutía (Capromys). *Miscelánea Zoológica*, 26, 2–3.
- Gineyts, E., Cloos, P. A. C., Borel, O., Grimaud, L., Delmas, P. D., & Garner, P. (2000). Racemization and isomerization of type I collagen C-telopeptides in human bone and soft tissues: Assessment of tissue turnover. *Biochemical Journal*, 345(3), 481–485. <https://doi.org/10.1042/bj3450481>
- Giovas, C. M. (2016). Though she be but little: Resource resilience, Amerindian foraging, and long-term adaptive strategies in the Grenadines, West Indies. *The Journal of Island and Coastal Archaeology*, 11, 238–263. <https://doi.org/10.1080/15564894.2016.1193572>
- Giovas, C. M. (2019). The beasts at large—Perennial questions and new paradigms for Caribbean translocation research. Part I: Ethnozoogeography of mammals. *Environmental Archaeology*, 24, 182–198. <https://doi.org/10.1080/14614103.2017.1315208>
- Giovas, C. M., Kamenov, G. D., Fitzpatrick, S. M., & Krigbaum, J. (2016). Sr and Pb isotopic investigation of mammal introductions: Pre-Columbian zoogeographic records from the Lesser Antilles, West Indies. *Journal of Archaeological Science*, 69, 39–53. <https://doi.org/10.1016/j.jas.2016.03.006>
- Giovas, C. M., Kamenov, G. D., & Krigbaum, J. (2019). $^{87}\text{Sr}/^{86}\text{Sr}$ and ^{14}C evidence for peccary (Tayassuidae) introduction challenges accepted historical interpretation of the 1657 Ligon map of Barbados. *PLoS ONE*, 14, e0216458. <https://doi.org/10.1371/journal.pone.0216458>
- Grimes, S. T., Collinson, M. E., Hooker, J. J., Matthey, D. P., Grassineau, N. V., & Lowry, D. (2004). Distinguishing the diets of coexisting fossil theridomyid and glirid rodents using carbon isotopes. *Palaeogeography, Palaeoclimatology, Palaeoecology*, 208, 103–119. <https://doi.org/10.1016/j.palaeo.2004.02.031>
- Guiry, E., Orchard, T. J., Needs-Howarth, S., & Szpak, P. (2021). Isotopic evidence for garden hunting and resource depression in the late woodland of Northeastern North America. *American Antiquity*, 86, 90–110. <https://doi.org/10.1017/aaq.2020.86>
- Healy, K., Kelly, S. B. A., Guillerme, T., Inger, R., Bearhop, S., & Jackson, A. L. (2017). *Predicting trophic discrimination factor using Bayesian inference and phylogenetic, ecological and physiological data*. *DeSIR: Discrimination Estimation in R*. PeerJ Preprints. <https://doi.org/10.7287/peerj.preprints.1950v3>
- Hermanson, J. W., & Woods, C. A. (2012). Anatomical specializations of capromyid rodents: functional and phylogenetic considerations. In R. Borotto-Páez, C. A. Woods, & F. E. Sergile (Eds.), *Terrestrial mammals of the West Indies: Contributions* (pp. 179–194). The Florida Museum of Natural History and Wacahoota Press.
- Hofman, C. L., & Hoogland, M. L. P. (2015). Investigaciones arqueológicas en los sitios El Flaco (Loma de Guayacanes) y La Luperona (UNIJICA). *Boletín del Museo del Hombre Dominicano*, 46, 61–74.
- Hooghiemstra, H., Olijhoek, T., Hoogland, M., Prins, M., van Geel, B., Donders, T., Gosling, W., & Hofman, C. (2018). Columbus' environmental impact in the New World: Land use change in the Yaque River valley, Dominican Republic. *Holocene*, 28, 1818–1835. <https://doi.org/10.1177/0959683618788732>
- Hopkins, J. B., & Ferguson, J. M. (2012). Estimating the diets of animals using stable isotopes and a comprehensive Bayesian mixing model. *PLoS ONE*, 7, e28478. <https://doi.org/10.1371/journal.pone.0028478>
- Howland, M. R., Corr, L. T., Young, S. M. M., Jones, V., Jim, S., Van Der Merwe, N. J., Mitchell, A. D., & Evershed, R. P. (2003). Expression of the dietary isotope signal in the compound-specific $\delta^{13}\text{C}$ values of pig bone lipids and amino acids. *International Journal of Osteoarchaeology*, 13, 54–65. <https://doi.org/10.1002/oa.658>
- Keegan, W. F., & DeNiro, M. J. (1988). Stable carbon- and nitrogen-isotope ratios of bone collagen used to study coral-reef and terrestrial components of prehistoric Bahamian diet. *American Antiquity*, 53, 320–336. <https://doi.org/10.2307/281022>
- Keeley, J. E., & Rundel, P. W. (2003). Evolution of CAM and C4 carbon-concentrating mechanisms. *International Journal of Plant Sciences*, 164, S55–S77. <https://doi.org/10.1086/374192>
- Kohn, M. J. (2010). Carbon isotope compositions of terrestrial C3 plants as indicators of (paleo)ecology and (paleo)climate. *Proceedings of the National Academy of Sciences of the United States of America*, 107, 19691–19695. <https://doi.org/10.1073/pnas.1004933107>
- Krigbaum, J., Fitzpatrick, S. M., & Bankaitis, J. (2013). Human paleodiet at Grand Bay, Carriacou, Lesser Antilles. *The Journal of Island and Coastal Archaeology*, 8, 210–227. <https://doi.org/10.1080/15564894.2012.756082>
- Kurle, C. M., Koch, P. L., Tershy, B. R., & Croll, D. A. (2014). The effects of sex, tissue type, and dietary components on stable isotope discrimination factors ($\Delta^{13}\text{C}$ and $\Delta^{15}\text{N}$) in mammalian omnivores. *Isotopes in Environmental and Health Studies*, 50, 307–321. <https://doi.org/10.1080/10256016.2014.908872>
- Laffoon, J., & Vos, B. (2011). Diverse origins, similar diets: An integrated isotopic perspective from Anse à la Gourde, Guadeloupe. In *Communities in Contact: Essays in archaeology, ethnohistory and ethnography of the Amerindian circum-Caribbean* (pp. 187–204). Sidestone Press.
- Laffoon, J. E., Hoogland, M. L. P., Davies, G. R., & Hofman, C. L. (2019). A multi-isotope investigation of human and dog mobility and diet in the pre-colonial Antilles. *Environmental Archaeology*, 24, 132–148. <https://doi.org/10.1080/14614103.2017.1322831>
- Laffoon, J. E., Plomp, E., Davies, G. R., Hoogland, M. L. P., & Hofman, C. L. (2015). The movement and exchange of dogs in the prehistoric Caribbean: An isotopic investigation. *International Journal of Osteoarchaeology*, 25, 454–465. <https://doi.org/10.1002/oa.2313>
- Laffoon, J. E., Rojas, R. V., & Hofman, C. L. (2013). Oxygen and carbon isotope analysis of human dental enamel from the Caribbean: Implications for investigating individual origins. *Archaeometry*, 55, 742–765. <https://doi.org/10.1111/j.1475-4754.2012.00698.x>
- Laffoon, J. E., Valcárcel Rojas, R., Weston, D. A., Hoogland, M. L. P., Davies, G. R., & Hofman, C. L. (2020). Diverse and dynamic dietary patterns in early colonial Cuba: New insights from multiple isotope

- analyses. *Latin American Antiquity*, 31, 103–121. <https://doi.org/10.1017/laq.2019.103>
- Lee-Thorp, J. A., Sealy, J. C., & van der Merwe, N. J. (1989). Stable carbon isotope ratio differences between bone collagen and bone apatite, and their relationship to diet. *Journal of Archaeological Science*, 16, 585–599. [https://doi.org/10.1016/0305-4403\(89\)90024-1](https://doi.org/10.1016/0305-4403(89)90024-1)
- LeFebvre, M. J., & deFrance, S. D. (2014). Guinea pigs in the pre-Columbian West Indies. *The Journal of Island and Coastal Archaeology*, 9, 16–44. <https://doi.org/10.1080/15564894.2013.861545>
- LeFebvre, M. J., deFrance, S. D., Kamenov, G. D., Keegan, W. F., & Krigbaum, J. (2019). The zooarchaeology and isotopic ecology of the Bahamian hutia (*Geocapromys ingrahami*): Evidence for pre-Columbian anthropogenic management. *PLoS ONE*, 14, e0220284. <https://doi.org/10.1371/journal.pone.0220284>
- LeFebvre, M. J., DuChemin, G., deFrance, S. D., Keegan, W. F., & Walczesky, K. (2019). Bahamian hutia (*Geocapromys ingrahami*) in the Lucayan Realm: Pre-Columbian exploitation and translocation. *Environmental Archaeology*, 24, 115–131. <https://doi.org/10.1080/14614103.2018.1503809>
- Linares, O. F. (1976). “Garden hunting” in the American tropics. *Human Ecology*, 4, 331–349. <https://doi.org/10.1007/BF01557917>
- Loiselle, B. A., & Blake, J. G. (1992). Population variation in a tropical bird community. *Bioscience*, 42, 838–845. <https://doi.org/10.2307/1312083>
- MacPhee, R. D. E., & Iturralde-Vinent, M. (1995). *Origin of the Greater Antillean land mammal fauna. 1, New Tertiary fossils from Cuba and Puerto Rico*. American Museum Novitates; no. 3141. American Museum of Natural History.
- Makarewicz, C., & Tuross, N. (2012). Finding fodder and tracking transhumance: Isotopic detection of goat domestication processes in the near east. *Current Anthropology*, 53, 495–505. <https://doi.org/10.1086/665829>
- Manójjina, N., & Abreu, R. (1990). Utilización de algunos mulcosos y reptiles en la dieta de la jutía conga. *Ciencias Biológicas*, 23, 127–129.
- Mickleburgh, H. L., & Pagan-Jimenez, J. R. (2012). New insights into the consumption of maize and other food plants in the pre-Columbian Caribbean from starch grains trapped in human dental calculus. *Journal of Archaeological Science*, 39, 2468–2478. <https://doi.org/10.1016/j.jas.2012.02.020>
- Minagawa, M., & Wada, E. (1984). Stepwise enrichment of ^{15}N along food chains: Further evidence and the relation between $\delta^{15}\text{N}$ and animal age. *Geochimica et Cosmochimica Acta*, 48, 1135–1140. [https://doi.org/10.1016/0016-7037\(84\)90204-7](https://doi.org/10.1016/0016-7037(84)90204-7)
- Monagle, V., Conrad, C., & Jones, E. L. (2018). What makes a dog? Stable isotope analysis and human-canid relationships at Arroyo Hondo Pueblo. *Open Quaternary*, 4, 6. <https://doi.org/10.5334/oq.43>
- Müller, J., Clauss, M., Codron, D., Schulz, E., Hummel, J., Kircher, P., & Hatt, J.-M. (2015). Tooth length and incisal wear and growth in guinea pigs (*Cavia porcellus*) fed diets of different abrasiveness. *Journal of Animal Physiology and Animal Nutrition*, 99, 591–604. <https://doi.org/10.1111/jpn.12226>
- Newsom, L. A., & Wing, E. S. (2004). *On land and sea: Native American uses of biological resources in the West Indies*. The University of Alabama Press.
- Odling-Smee, F. J., Laland, K. N., & Feldman, M. W. (1996). Niche construction. *The American Naturalist*, 147, 641–648. <https://doi.org/10.1086/285870>
- O’Leary, M. H. (1988). Carbon isotopes in photosynthesis: Fractionation techniques may reveal new aspects of carbon dynamics in plants. *Bioscience*, 38, 328–336. <https://doi.org/10.2307/1310735>
- Oliver, W. L. R., Herr, R. H., & Kelly, D. L. (1986). The Jamaican hutia *Geocapromys brownii* captive breeding and reintroduction programme. History and progress. *The Dodo, Journal of the Jersey Wildlife Preservation Trust*, 23, 32–58.
- Oswald, J. A., Allen, J. M., LeFebvre, M. J., Stucky, B. J., Folk, R. A., Albury, N. A., Morgan, G. S., Guralnick, R. P., & Steadman, D. W. (2020). Ancient DNA and high-resolution chronometry reveal a long-term human role in the historical diversity and biogeography of the Bahamian hutia. *Scientific Reports*, 10, 1373. <https://doi.org/10.1038/s41598-020-58224-y>
- Pagán-Jiménez, J. R., Ali, Z., Santiago-Marrero, C. G., & Hofman, C. L. (2020). Plantscapes of dwelling: Precolonial household mounds, phyto-cultural dynamics and the ensuing human ecosystems at El Flaco and El Carril (cal. AD 990–1450), northern Dominican Republic. *Review of Palaeobotany and Palynology*, 274, 104160. <https://doi.org/10.1016/j.revpalbo.2020.104160>
- Pagán-Jiménez, J. R., & Mickleburgh, H. L. (2022). Caribbean deep-time culinary worlds revealed by ancient food starches: Beyond the dominant narratives. *Journal of Archaeological Research*. <https://doi.org/10.1007/s10814-021-09171-3>
- Park, M. K., Min, S.-Y., Song, J. S., Lee, J.-H., Jung, H.-S., & Kim, S.-O. (2017). Estimated time of biomineralization in developing rat incisors. *Journal of Korean Academy of Pediatric Dentistry*, 44, 138–146. <https://doi.org/10.5933/JKAPD.2017.44.2.138>
- Pestle, W. J. (2010). *Diet and society in prehistoric Puerto Rico: An isotopic approach*. PhD dissertation. University of Illinois at Chicago.
- Pestle, W. J. (2013). Fishing down a prehistoric Caribbean marine food web: Isotopic evidence from Punta Candelero, Puerto Rico. *The Journal of Island and Coastal Archaeology*, 8, 228–254. <https://doi.org/10.1080/15564894.2013.797943>
- Pestle, W. J., Hubbe, M., Smith, E. K., & Stevenson, J. M. (2015). Technical note: A linear model for predicting $\delta^{13}\text{C}_{\text{protein}}$. *American Journal of Physical Anthropology*, 157, 694–703. <https://doi.org/10.1002/ajpa.22743>
- Pestle, W. J., & Laffoon, J. (2018). Quantitative paleodietary reconstruction with complex foodwebs: An isotopic case study from the Caribbean. *Journal of Archaeological Science: Reports*, 17, 393–403. <https://doi.org/10.1016/j.jasrep.2017.11.032>
- Phillips, D. L., Inger, R., Bearhop, S., Jackson, A. L., Moore, J. W., Parnell, A. C., Semmens, B. X., & Ward, E. J. (2014). Best practices for use of stable isotope mixing models in food-web studies. *Canadian Journal of Zoology*, 92, 823–835. <https://doi.org/10.1139/cjz-2014-0127>
- Quinn, R. L. (2019). Isotopic equifinality and rethinking the diet of *Australopithecus anamensis*. *American Journal of Physical Anthropology*, 169, 403–421. <https://doi.org/10.1002/ajpa.23846>
- Ramírez-Barajas, P. J., & Calmé, S. (2015). Subsistence hunting and conservation. In G. A. Islebe, S. Calmé, J. L. León-Cortés, & B. Schmook (Eds.), *Biodiversity and conservation of the Yucatán peninsula* (pp. 333–351). Springer International Publishing. https://doi.org/10.1007/978-3-319-06529-8_13
- Roberts, P., Blumenthal, S. A., Dittus, W., Wedage, O., & Lee-Thorp, J. A. (2017). Stable carbon, oxygen, and nitrogen, isotope analysis of plants from a South Asian tropical forest: Implications for primatology. *American Journal of Primatology*, 79, e22656. <https://doi.org/10.1002/ajp.22656>
- Schoeninger, M. J., & DeNiro, M. J. (1984). Nitrogen and carbon isotopic composition of bone collagen from marine and terrestrial animals. *Geochimica et Cosmochimica Acta*, 48, 625–639. [https://doi.org/10.1016/0016-7037\(84\)90091-7](https://doi.org/10.1016/0016-7037(84)90091-7)
- Schwarcz, H. P. (2002). Some biochemical aspects of carbon isotopic paleodiet studies. In S. H. Ambrose & M. A. Katzenberg (Eds.), *Biogeochemical approaches to paleodietary analysis* (pp. 189–209). Springer US. https://doi.org/10.1007/0-306-47194-9_10
- Schwarcz, H. P., Melbye, J., Anne Katzenberg, M., & Knyf, M. (1985). Stable isotopes in human skeletons of Southern Ontario: Reconstructing palaeodiet. *Journal of Archaeological Science*, 12, 187–206. [https://doi.org/10.1016/0305-4403\(85\)90020-2](https://doi.org/10.1016/0305-4403(85)90020-2)

- Sharpe, A. E., Emery, K. F., Inomata, T., Triadan, D., Kamenov, G. D., & Krigbaum, J. (2018). Earliest isotopic evidence in the Maya region for animal management and long-distance trade at the site of Ceibal, Guatemala. *Proceedings of the National Academy of Sciences*, *115*, 3605–3610. <https://doi.org/10.1073/pnas.1713880115>
- Shev, G. T., Ali, Z., Almonte Milan, J. N., Casale, S., Djakovic, I., & Hofman, C. L. (2021). Coastal-hinterland exchange and garden hunting practices prior to the European invasion of Hispaniola. *Environmental Archaeology*, *1*–22. <https://doi.org/10.1080/14614103.2021.1993693>
- Shev, G. T., Laffoon, J. E., Grouard, S., & Hofman, C. L. (2020). An isotopic and morphometric examination of island dogs (*Canis familiaris*): Comparing dietary and mobility patterns in the precolumbian Caribbean. *Latin American Antiquity*, *31*, 632–638. <https://doi.org/10.1017/laq.2020.58>
- Shev, G. T., Laffoon, J. E., & Hofman, C. L. (2021). Human and hutia (*Isolobodon portoricensis*) interactions in pre-Columbian Hispaniola: The isotopic and morphological evidence. *Journal of Archaeological Science: Reports*, *37*, 102913. <https://doi.org/10.1016/j.jasrep.2021.102913>
- Smith, B. D. (2001). Low-level food production. *Journal of Archaeological Research*, *9*, 1–43. <https://doi.org/10.1023/A:1009436110049>
- Smith, B. D. (2007). Niche construction and the behavioral context of plant and animal domestication. *Evolutionary Anthropology: Issues, News, and Reviews*, *16*, 188–199. <https://doi.org/10.1002/evan.20135>
- Smith, B. D. (2011). General patterns of niche construction and the management of 'wild' plant and animal resources by small-scale pre-industrial societies. *Philosophical Transactions of the Royal Society, B: Biological Sciences*, *366*, 836–848. <https://doi.org/10.1098/rstb.2010.0253>
- Smith, D. A. (2005). Garden game: Shifting cultivation, indigenous hunting and wildlife ecology in Western Panama. *Human Ecology*, *33*, 505–537. <https://doi.org/10.1007/s10745-005-5157-Y>
- Soto-Centeno, J., Simmons, N., & Steadman, D. (2017). The bat community of Haiti and evidence for its long-term persistence at high elevations. *PLoS ONE*, *12*, e0178066. <https://doi.org/10.1371/journal.pone.0178066>
- Sponheimer, M., Robinson, T. F., Ayliffe, L. K., Roeder, B. L., Hammer, J., Passey, B. H., West, A. G., Cerling, T. E., Dearing, D. M., & Ehleringer, J. R. (2003). Nitrogen isotopes in mammalian herbivores: Hair $\delta^{15}\text{N}$ values from a controlled feeding study. *International Journal of Osteoarchaeology*, *13*, 80–87. <https://doi.org/10.1002/oa.655>
- Stock, B. C., Jackson, A. L., Ward, E. J., Parnell, A. C., Phillips, D. L., & Semmens, B. X. (2018). Analyzing mixing systems using a new generation of Bayesian tracer mixing models. *PeerJ*, *6*, e5096. <https://doi.org/10.7717/peerj.5096>
- Stokes, A. V. (2008). *A biogeographic survey of prehistoric human diet in the West Indies using stable isotopes*. University of Florida.
- Sugiyama, N., Martínez-Polanco, M. F., France, C. A. M., & Cooke, R. G. (2020). Domesticated landscapes of the neotropics: Isotope signatures of human-animal relationships in pre-Columbian Panama. *Journal of Anthropological Archaeology*, *59*, 101195. <https://doi.org/10.1016/j.jaa.2020.101195>
- Sugiyama, N., Somerville, A. D., & Schoeninger, M. J. (2015). Stable isotopes and zooarchaeology at Teotihuacan, Mexico reveal earliest evidence of wild carnivore management in Mesoamerica. *PLoS ONE*, *10*, e0135635. <https://doi.org/10.1371/journal.pone.0135635>
- Turvey, S. T., Oliver, J. R., Storde, N., & Rye, P. (2007). Late Holocene extinction of Puerto Rican native land mammals. *Biology Letters*, *3*, 193–196. <https://doi.org/10.1098/rsbl.2006.0585>
- van der Merwe, N. J., & Medina, E. (1991). The canopy effect, carbon isotope ratios and foodwebs in Amazonia. *Journal of Archaeological Science*, *18*, 249–259. [https://doi.org/10.1016/0305-4403\(91\)90064-V](https://doi.org/10.1016/0305-4403(91)90064-V)
- von Fischer, J. C., & Tieszen, L. L. (1995). Carbon isotope characterization of vegetation and soil organic matter in subtropical forests in Luquillo, Puerto Rico. *Biotropica*, *27*, 138–148. <https://doi.org/10.2307/2388989>
- Wing, E. S. (2008). Pets and camp followers in the West Indies. In E. J. Reitz, S. J. Scudder, & C. M. Scarry (Eds.), *Case studies in environmental archaeology* (pp. 405–425). Springer New York. https://doi.org/10.1007/978-0-387-71303-8_21
- Wing, E. S. (2012). Zooarchaeology of West Indian land mammals. In R. Borotto-Páez, C. A. Woods, & F. E. Sergile (Eds.), *Terrestrial mammals of the West Indies: Contributions* (pp. 341–356). Florida Museum of Natural History and Wacahoota Press.
- Wiscovitch-Russo, R., Rivera-Perez, J., Narganes-Storde, Y. M., García-Roldán, E., Bunkley-Williams, L., Cano, R., & Toranzos, G. A. (2020). Pre-Columbian zoonotic enteric parasites: An insight into Puerto Rican indigenous culture diets and life styles. *PLoS ONE*, *15*, e0227810. <https://doi.org/10.1371/journal.pone.0227810>
- Woods, C. A., & Ottenwalder, J. A. (1992). *The natural history of southern Haiti*. Gainesville.
- Woods, R., Barnes, I., Brace, S., & Turvey, S. T. (2021). Ancient DNA suggests single colonization and within-archipelago diversification of Caribbean caviomorph rodents. *Molecular Biology and Evolution*, *38*, 84–95. <https://doi.org/10.1093/molbev/msaa189>
- Zavodny, E., McClure, S. B., Culleton, B. J., Podrug, E., & Kennett, D. J. (2015). Identifying Neolithic animal management practices in the Adriatic using stable isotopes. *Documenta Praehistorica*, *42*, 261–274. <https://doi.org/10.4312/dp.42.18>
- Zeder, M. A. (2006). Central questions in the domestication of plants and animals. *Evolutionary Anthropology: Issues, News, and Reviews*, *15*, 105–117. <https://doi.org/10.1002/evan.20101>
- Zeder, M. A. (2012). Pathways to animal domestication. In P. Gepts, T. R. Famula, R. L. Bettinger, S. B. Brush, A. B. Damania, P. E. McGuire, & C. O. Qualset (Eds.), *Biodiversity in agriculture* (pp. 227–259). Cambridge University Press. <https://doi.org/10.1017/CBO9781139019514.013>
- Zeder, M. A. (2015). Core questions in domestication research. *Proceedings of the National Academy of Sciences*, *112*, 3191–3198. <https://doi.org/10.1073/pnas.1501711112>

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