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Understanding the Pathoecological Relationship between Ancient Diet and Modern Diabetes through Coprolite Analysis: A Case Example from Antelope Cave, Mojave County, Arizona

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| Karl J. Reinhard, Keith L. Johnson, Sara LeRoy-Toren, Kyle Wieseman, Isabel Teixeira-Santos, and Mónica Vieira Vieira | |
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Understanding the Pathoecological Relationship between Ancient Diet and Modern Diabetes through Coprolite Analysis

A Case Example from Antelope Cave, Mojave County, Arizona

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CA+ Online-Only Material: Supplements A-C

The elevated prevalence of non-insulin-dependent diabetes mellitus (NIDDM) in Native Americans of the Southwest has been explained by several authors in terms of a dietary change from preindustrial traditional foods to modern foods. Physiology adapted to traditional foods became deleterious during the process of modernization. Although several versions of this hypothesis exist, they all relate to the rise in modern NIDDM with change from prehistoric subsistence practices to modern dietary practices. This is especially true for the Southwestern desert tribes of Arizona and New Mexico. Coprolite analysts have been recovering the sort of data needed by diabetes researchers to explore the prehistoric dietary foundations for NIDDM. Diabetes researchers have missed these studies that are essential in understanding ancient diet. We are taking this opportunity to show how coprolite analysis of diet provides data relevant to understanding debates. Our case example comes from Antelope Cave, Mojave County, Arizona. There was a high reliance on fiber-rich plant foods with low glycemic indexes. However, these were not just famine foods as suggested by the original "thrifty gene" hypothesis. These were the foods eaten on a day-by-day basis during all seasons, in both feast and famine.

Southwestern Native American tribes, including the Akimel O'odham (Pima), Tohono O'odham (Papago), Quechan (Yuma), Ak-Chin (Maricopa), Haulapai, Mojave, and the Arizona and New Mexico Pueblo tribes, suffer high rates of diabetes. Specifically, the members of these tribes are susceptible to type 2, or non-insulin-dependent, diabetes mellitus

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(NIDDM). The elevated prevalence of NIDDM has been explained by several authors in terms of a "thrifty genotype" that was adaptive in ancient times but became deleterious during the process of modernization. Although several versions of the thrifty-gene hypothesis exist, they all relate to the rise in modern NIDDM with change from prehistoric subsistence practices to modern dietary practices. Neel (1962) first presented a thrifty-genotype hypothesis to explain why NIDDM occurs at such high rates in some modernizing tribal populations. It is hypothesized that this genotype evolved during thousands of years of feast or famine conditions. With food seasonally unavailable to hunter-gatherers, excessive caloric intake resulting in quickly elevated insulin secretion was adaptive. However, for populations that rapidly modernized, this genotype was maladaptive in the context of constant dependable sources of food.

The loss of high fiber in traditional diets has been linked to the emergence of NIDDM (Wolever et al. 1997). This has been demonstrated in experimental study. Williams et al. (2001) characterized "Indian," "Anglo," and "mixed" diets among the Akimel O'odahm and related these diets to diabetes prevalence. They found that complex carbohydrates, dietary fiber, insoluble fiber, vegetable proteins, and the proportion of total calories from complex carbohydrate and vegetable proteins were significantly higher in the Indian than in the Anglo diet. The risk of developing diabetes in the Anglo-diet group was 2.5 times higher than in the Indian-diet group.

Other researchers have sorted out the individual effects of the components of traditional diets. Hung et al. (2003) determined that the fiber content of carbohydrate food confers benefits in terms of diabetic control. Their work supports a dietary-fiber hypothesis proposed by Trowell (1975:764) that "dietary fiber depleted starchy foods are conducive to the development of diabetes mellitus in susceptible human genotypes." Also, these traditional diets were rich in food with low glycemic indexes (GIs). The GI is an indicator of carbohydrate's ability to raise blood glucose levels. Hung et al. (2003) found that for carbohydrates, the GI appears to be a better predictor of the metabolic effects of a diet than sugar content. Willett, Manson, and Liu (2002) summarized the value of the GI based on experimental studies of animals and humans. They stated that a high intake of carbohydrate foods with high GIs produces greater insulin resistance than intake of low-GI foods. Hu, van Dam, and Liu (2001) found that a low-GI diet with a high amount of fiber lowers the risk of NIDDM.

The traditional diets of the Tohono and Akimel O'odham have been studied with regard to NIDDM. However, the information concerning prehistoric traditional diet for Southwestern tribes in general is sketchy. This is similar to the knowledge of Aboriginal diet in Australia. As Gracey (2000: 1361) notes, "Most information about Aboriginal diets is anecdotal or semiquantitative. More effort needs to be invested in studies that more clearly and precisely define dietary pat-

terns in Aboriginal people, especially children, and how these patterns influence their growth, nutritional status, and health, prospectively." Gracey's comment is relevant to the study of Southwestern traditional diet.

The debates summarized above show that diabetes researchers have been actively searching for an effective method for reconstructing ancient diet. This is especially true for the Southwestern desert tribes of Arizona and New Mexico. Coprolite analysts have been recovering exactly the data desired by diabetes researchers for nearly 4 decades (Bryant 1974a, 1974b; Bryant and Dean 2006; Bryant and Williams-Dean 1975; Poinar et al. 2001; Rasmussen et al. 2009; Reinhard 1992; Reinhard and Bryant 1992, 2008). Diabetes researchers have missed these studies that are essential in understanding ancient diet. We are taking this opportunity to show how coprolite analysis of diet provides data relevant to the diabetes debates. Our case example comes from Antelope Cave, Mojave County, Arizona.

Martinson et al. (2003), followed by Reinhard (2007) and Reinhard and Bryant (2008), developed the concept of "pathoecology," which is the evaluation of environmental determinants of disease. The term was first used by Karl R. Reinhard (Karl J. Reinhard's father), an epidemiologist who worked on the history of disease development in the Arctic (Reinhard 1974a, 1974b). K. R. Reinhard applied this concept to emerging diabetes (Reinhard and Greenwalt 1975). K. J. Reinhard (1988) began pathoecological study of subsistence variation on parasitic disease. The connection between episodic malnutrition and parasitic disease with anemia as represented in skeletal remains was demonstrated later (Reinhard 1992, 2007). Also, coprolite data were linked to patterns of dental disease (Danielson and Reinhard 1998; Reinhard and Danielson 2005). Today, there is direct relevance of coprolite data to understanding the nutritional health, metabolic health, dental health, and level of parasite infection among prehistoric people. It was from this pathoecological perspective that Antelope Cave coprolites were analyzed.

Archaeological Background of Antelope Cave

Antelope Cave is located on the Arizona Strip in the northwest corner of Arizona, about 25 miles southeast of St. George, Utah. It is a large limestone grotto sunk into the semiarid rolling plains of the Uinkaret Plateau about 4,660 feet asl. The cave interior (fig. 1) measures 350 feet north-south by 150 feet east-west. Prehistoric Native Americans first lived in this subterranean cavern 4,000 years ago, and various Native groups occupied or visited the cave until AD 1150. The most abundant cultural materials recovered from the site were left behind by Ancestral Puebloan (Virgin Anasazi) peoples who lived there successfully, perhaps seasonally, for at least 450 years (AD 700–1150). It appears that the cave may have been a seasonal camp repeatedly visited to store artifacts such as sandals and nets, to gather local plants, and to hunt rabbits

to eat and then to prepare their hides for the manufacture of soft fur/skin robes.

Figure 1 is a plan drawing of Antelope Cave. The main entrance to the cave (now gated by the Bureau of Land Management) is on its southeast edge above large slabs of jumbled limestone. Once inside, a massive layer of broken limestone roof fall, beginning 10 feet below the entrance, extends over the eastern half of the cave. The dry and gray powdery midden deposit covers most of the western half. The surface of the midden slopes down rapidly from south to north, ending at a large secondary sinkhole, the bottom of which is 75 feet below the entrance to the cave.

Altschul and Fairley (1989) and Lyneis (1995) review the archaeology of the Arizona Strip, including the prehistoric Virgin Anasazi, some of whom were the primary occupants of Antelope Cave more than 1,000 years ago. We offer here a synopsis of their work and add information from Antelope Cave.

The Archaic Period (7000–300 BC) on the Arizona Strip is typified by distinctive styles of projectile points that represent early dispersed hunter-gatherer groups. Some Archaic-style points—Rocker side-notched, San Rafael side-notched, and Elko corner-notched—are reported from Antelope Cave. In support, charcoal samples from the cave yielded three calibrated ¹⁴C dates of 2028–1893 BC, 1891–1744 BC, and 1699–1444 BC (Janetski and Wilde 1989), further confirming the initial occupation of Antelope Cave by Archaic groups.

Basketmaker II (300 BC–AD 400) is the first Anasazi period to be recognized in the Southwest. Basketmaker II sites are typically pit structures or rock shelters and caves with slablined storage pits. Five are known on the Arizona Strip, including Antelope Cave, which has yielded a wooden atlatl (spear thrower) carbon dated at AD 100. Also, a small excavated area (fig. 1; University of California, Los Angeles [UCLA] excavation unit AC59-3, 4) near the west wall of the cave lacked ceramics but contained finely woven square-toed sandals and obsidian projectile points. This area is tentatively attributed to the Basketmaker II period (Johnson and Pendergast 1960:3).

Basketmaker III (AD 400–800) is represented by small pithouse villages, plain gray pottery, and cultivated beans to go with the growing of maize and squash. Probable Basketmaker III traits at Antelope Cave include cultivated beans, plain gray ceramics, and the wooden nock ends of arrows. In addition, three new radiocarbon dates from Antelope Cave indicate that the Ancestral Pueblo more than any other group made the greatest use of the site during late Basketmaker III through early Pueblo I times. The new dates are cal AD 680–890, cal AD 710–960, and cal AD 680–890. These new radiocarbon assays equate nicely with those from the Brigham Young University excavations in 1983 (Janetski and Hall 1983:40, 43).

The Pueblo I and Pueblo II periods (AD 800–1150) are characterized by the continuation and elaboration of Basket-maker III cultural traits. Pithouse villages are usually small and increase in number, and the houses may have benches

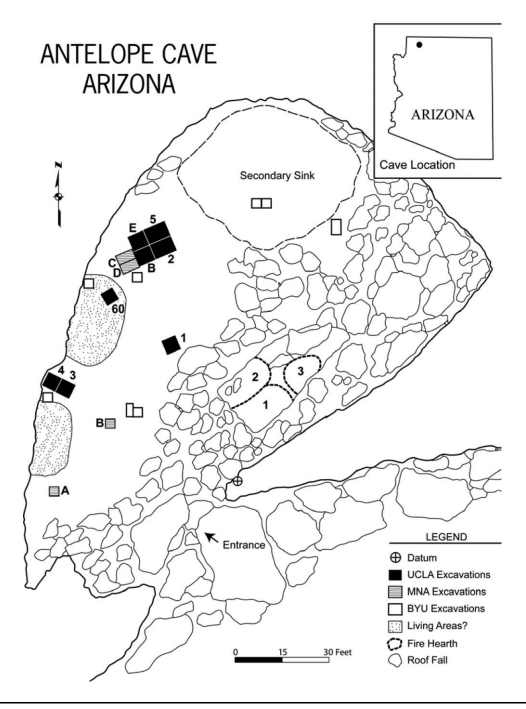


Figure 1. Antelope Cave, surface features and excavation units (University of California, Los Angeles [UCLA]; Museum of Northern Arizona [MNA]; Brigham Young University [BYU]).

and ventilators. Aboveground masonry structures first occur during this time period. Black-on-gray and black-on-white decorated pottery make their appearance early on. Corrugated pottery is a new style that debuts in the Pueblo II period (AD 1000–1150). Round- or pointed-toed sandals begin to replace square-toed sandals by Pueblo I times. At Antelope Cave, five Virgin Anasazi types of decorated ceramics as well as round-toed yucca sandals mark these two culture stages. However,

the seasonal use of Antelope Cave by Ancestral Puebloans was in decline by AD 1000. Of the 3,100 pottery sherds recovered during the UCLA excavations, only three were corrugated. This suggests little use of the cave during much of Pueblo II.

Several Anasazi sites have been dated to Early Pueblo III (AD 1150–1225) on the Arizona Strip. At this time, Antelope Cave had been pretty much abandoned by the Virgin Anasazi. Where they went or what happened to them is uncertain.

Prehistoric Southern Paiute groups began to move into Virgin Anasazi territory around AD 1000, and these newcomers eventually either absorbed or replaced the Ancestral Pueblo peoples on the strip. A large twined and pitch-covered Paiute water basket was recovered from the surface of Antelope Cave by Vilate Hardy of La Verkin, Utah, in the early 1950s. The location of this specimen in the cave provides a bit of evidence that either the Virgin Anasazi were in contact with the Southern Paiute or the Southern Paiute actually occupied the cave sometime after it was abandoned by the Pueblo people. More details of the archaeology of the region are presented in CA+ online supplement A.

Material and Methods

We analyzed 25 coprolites from Antelope Cave. Of these, 20 were human in origin and four were dog or some other canid in origin. One coprolite sample was actually consolidated cave sediment. Samples of 10 coprolites were submitted to two molecular biology labs, one at Fundação Oswaldo Cruz and one at the University of Oklahoma. The molecular results are not available at this time.

Initially, our goal was to select one coprolite from each Antelope Cave provenience. This involved identifying the most human-like coprolite from each provenience. However, four of the first 10 coprolites sampled in this way turned out to be of animal origin. Subsequently, the coprolite collection was examined with the goal of identifying human coprolites only. Therefore, those coprolites that had the most distinct human morphology and that did not contain obvious animal hair were included for study. The sampling followed the guidelines for distinguishing human from animal coprolites published by Chame (2003). The details of the analysis procedure are presented in CA+ online supplement B.

Results

The dietary results are presented in CA+ online supplement C. The tables in supplement C present definitions of terms, data for microresidues, and the macroresidue results. These results for the human coprolites are presented in table C1. Coprolites 6 and 22 are omitted from these results because analysis revealed that they were not human coprolites. Number 6 was an animal coprolite, and number 22 was consolidated cave sediment.

Most of the human coprolites analyzed for macroscopic and microscopic food remains appear to be from late summer and early fall depositions. At this time period, a diversity of wild-plant fruits and seeds were available for harvesting, and there would still be ambient pollen from sagebrush, grass, cheno-ams, and ragweed-type plants. Therefore, this coprolite series cannot be considered to represent a year-round diet.

There are four principal processed plant foods represented in the Antelope Cave coprolites. These are maize, wild-grass caryopses, sunflower seeds, and cheno-am seeds that have been ground to a fine flour. Much of the flour made from these plants passed through a 0.5-mm screen. Grinding was a significant part of preparing food at Antelope Cave. A caryopsis is a dry seed-like fruit produced by wild and cultivated cereal grasses. Maize and wild-grass caryopses consistent with dropseed were found in six coprolites each. They did not co-occur. Dropseed was very abundant in five coprolites, and maize was very abundant in six. Microscopically, maize starch occurred in seven coprolites. Thus, maize was slightly more important than wild-grass caryopses.

Sunflower achenes occurred in four coprolites and dominated three of these. An achene is a small, dry, indehiscent fruit with the seed distinct from the fruit wall. The composite family, among others, produces achenes. Microscopically, ground achene fragments were present in six coprolites. Therefore, it appears that sunflower achenes were significant to Antelope Cave users. Ground achenes, in our experience, are unique to Antelope Cave. Importantly, the entire achenes—shell and seed—were ground.

Flour was also made of cheno-am seeds and was found in three coprolites, but it was dominant in only one. In botanical archaeology terms, "cheno-am" is applied to fruits and pollen. These fruits look like small black seeds. They have a round starchy core surrounded by a thick seed coat. Cheno-am fruits come from plants in Chenopodium, Amaranthus, or related genera. Cheno-am pollen could be from either the family Chenopodiaceae or the genus Amaranthus. Cheno-am pollen is often ingested with cheno-am fruits. Cheno-am seeds are a common component of Southwestern diet in agricultural and preagricultural times. In our experience from coprolite analysis, the production of flour from cheno-am seeds is more an aspect of Archaic diet such as that represented at Dust Devil Cave, Utah (Reinhard, Ambler, and McGuffie 1985). The finely ground cheno-am flour at Antelope Cave is unusual for Ancestral Pueblo coprolites.

In addition to the four principal flour foods, a seed that could be four-winged saltbush was found in two coprolites. Unidentifiable seeds and wolfberry were observed in one coprolite each.

Following maize and wild grass, prickly pear pads were an important food source. Four coprolites included macroscopic remains of prickly pear, while 11 contain microscopic remains. Prickly pear was very abundant in two coprolites. However, in seven coprolites, prickly pear co-occurred with other foods. Therefore, prickly pear was an important standalone food, and it also supplemented other foods.

Discussion

There is no doubt that NIDDM has an evolutionary connection with ancient diet. Coprolite analysts provide the most insightful and empirical information about the nutritional content of traditional diets. The dietary data from the Antelope Cave coprolites can be directly applied to diabetes debates. We are using this site, combined with analysis with

other sites, to identify patterns of food use that formed an evolutionary pressure for the fixation of the thrifty gene.

Antelope Cave coprolite analysis highlights the nutritional nature of ancient foods. The analysis demonstrates the following points. First, prehistoric foods consistently had low GIs. Second, prehistoric diet was remarkably high in fiber. Third, there was a high reliance on hypoglycemic-effect foods. Fourth, inulin-rich foods were a consistent part of diet. This fourth aspect of prehistoric diet has already been established in the literature (Leach, Gibson, and Van Loo 2010; Leach and Sobilik 2010).

The GI measures the influence of carbohydrates on blood sugar levels. High-GI foods rapidly break down during digestion and rapidly release glucose into the bloodstream. Low-GI foods break down more slowly and release glucose gradually into the bloodstream. The GI is assigned to foods based on a standard of glucose that has a GI of 100. Low-GI foods have a value of less than 55. High-GI foods exceed a value of 70.

Some Antelope Cave food GIs can be approximated from modern counterparts. Prickly pear was a very important prehistoric food. It has a GI of 7, which is the lowest recorded for Southwestern plant foods and one of the lowest values for any recorded human food. Modern cultivated sunflower achenes have a GI of 10. Modern cultivated amaranth has a GI of 25. The prehistoric sunflowers and cheno-ams were uncultivated and probably had lower indexes than these. Thus, based on available GI values for modern versions of prickly pear, amaranth, and sunflowers, it appears that the wild-plant diet at Antelope Cave focused on very low-GI species. The total GI for stews made of wild-plant seeds and rabbit may have been close to the modern value of Traditional Akimel O'odham (Pima) acorn and venison stew, which is 23.

Looking for GI values of traditional Southwestern foods provides insight into the maize of Antelope Cave. Traditional Akimel O'odham maize hominy made from indigenous maize has a GI of 57. This was probably the highest GI for available foods at Antelope Cave.

As for the fiber content of the Antelope Cave diet, it was very high. Fiber appears in the form of xylem, phloem, epidermis, glumes, achene shells, seed testa, and fruit shells in Antelope Cave coprolites. In general, half of the weight of each coprolite was composed of fiber fragments larger than 0.25 mm. Microscopically, the concentrations of undigested plant fragments ranged from hundreds of thousands to millions of fragments per gram of coprolite. This fiber content is remarkable from the modern perspective but comparable to other Southwestern prehistoric coprolite series. By volume, about three-quarters of Antelope Cave coprolites are composed of insoluble fiber.

Prickly pear cactus pads were a common food in the prehistoric Southwest. Indeed, Reinhard (1992) identified prickly pear as one of the main dietary components in the region. Table 1 presents the frequency of coprolites positive for prickly

Table 1. Summary of phytolith finds from several Southwestern sites

| | | Орі | ıntia | Agavaceae | | |
|--------------------------|----|-----|-------|-----------|-----|--|
| Site | n | n | % | n | % | |
| Hunter-gatherer: | | | | | | |
| Hinds Cave, Texas | 14 | 13 | 93 | 14 | 100 | |
| Dust Devil Cave, Utah | 17 | 10 | 59 | 16 | 94 | |
| Bighorn Cave, Arizona | 17 | 14 | 82 | 16 | 94 | |
| Total | 48 | 37 | 77 | 46 | 96 | |
| Ancestral Pueblo: | | | | | | |
| Bighorn Sheep Ruin, Utah | 20 | 10 | 50 | 12 | 60 | |
| Antelope House, Arizona | 25 | 20 | 80 | 18 | 72 | |
| Salmon Ruin, New Mexico | 20 | 9 | 45 | 10 | 50 | |
| Antelope Cave, Arizona | 20 | 14 | 70 | 1 | 5 | |
| Total | 85 | 53 | 62 | 41 | 48 | |

Note. Hunter-gatherers consumed prickly pear (*Opuntia*), yucca, and/ or agave very commonly. Both yucca and agave are in the family Agavaceae. Consumption of Agavaceae food plants declined dramatically among the Ancestral Pueblo but was still common in some areas. However, prickly pear remained a common food in the Ancestral Pueblo diet.

pear as measured by the analysis of phytoliths from coprolites. Prickly pear pads contain distinctly shaped calcium oxalate crystals. By extracting phytoliths from coprolites, it is possible to assess the dietary reliance of ancient people on prickly pear pads. The data show that 77% of hunter-gatherer coprolites contain prickly pear phytoliths. For agricultural peoples, the frequency drops to 62%. This shows that prickly pear pads were a central part of the diet even after agriculture was established.

NIDDM researchers have long recognized the antidiabetic properties of prickly pear of the species Opuntia streptacantha, Opuntia ficus-indica, Opuntia engelmannii var. lindheimeri, and Opuntia fuliginosa (Frati-Munari et al. 1988, 1989a, 1989b, 1989c, 1989d; Meckes-Lozoya and Ibáñez-Camacho 1989; Meckes-Lozoya and Roman-Ramos 1986). Frati-Munari et al. (1988, 1989b, 1989c, 1991) demonstrated that broiled prickly pear pads had a hypoglycemic effect in diabetic humans. Various authors demonstrated the hypoglycemic effects of prickly pear on animals, including pancreatectomized rabbits (Ibáñez-Camacho and Roman-Ramos 1979), rats (Trejo-Gonzalez et al. 1996), pigs (Laurenz, Collier, and Kuti 2003), and other lab models (Ibáñez-Camacho, Meckes-Lozoya, and Mellado-Campos 1983). All of these lines of research demonstrated that prickly pear pads cause lower blood glucose levels in diabetic patients and animals. Meckes-Lozoya and Ibáñez-Camacho (1989) evaluated the hypoglycemic activity of prickly pear throughout the seasonal cycle of the plant and found no variation in the hypoglycemic activity of this plant.

Coprolite analysis focusing on consumption of desert succulent plants shows that prickly pear was a universal food source for prehistoric southwesterners (Reinhard and Danielson 2005). The discovery of prickly pear at Antelope Cave expands the known use of prickly pear to the Uinkaret Plateau. The consistent use of this plant through thousands of years of desert subsistence could have exerted part of an evolutionary pressure for the fixation of the thrifty gene.

Prickly pear pads were not the only desert succulents that were important in the Southwestern diet. Agave and vucca are desert succulents that were very commonly eaten by hunter-gatherer and agricultural peoples (table 1). They were exceptionally common in preagricultural diets. After the agricultural revolution in the Southwest, phytolith analysis indicates that about half of the human coprolites contain agave or yucca. Agave and yucca are fibrous desert succulents that provided several sources of food to prehistoric people (Leach and Sobolik 2010). Agave is a demonstrated dietary source of inulin. Yucca is not as well studied as agave, but it is also an inulin source. Inulin has minimal effect on blood sugar. For both agave and yucca, the buds and hearts of the plants were eaten. The hearts were available year-round. To collect the hearts, the plants had to be pried from the ground. The leaves were then cut from the base of the plant, leaving just the "heart" of leaf bases. The resulting structure looks like a very large artichoke heart. The hearts were then baked for up to 2 days in large underground rock-lined ovens. The cooked leaf bases were then pulled from the heart and chewed to extract carbohydrates. This resulted in a wad of fiber in the mouth called a "quid." The quids were swallowed or spit out. Fiber from quids is very commonly found in prehistoric coprolites and testifies to the high-fiber nature of yucca and agave. Both yucca and agave are fiber rich. Montonen et al. (2003), Salmerón et al. (1997a, 1997b), Brand-Miller et al. (2003), Pick et al. (1996), Guévin et al. (1996), Tabatabai and Li (2000), and Marlett, McBurney, and Slavin (2002) emphasized that fiber-rich foods were part of the traditional diets for tribal cultures that are experiencing rises in obesity and NIDDM.

Yucca in coprolites occurred only in traces at Antelope Cave. However, more than 300 yucca quids from the midden show that people did eat yucca there. Therefore, the Antelope Cave evidence is consistent with the general picture that desert succulents were important subsistence sources for Southwestern prehistoric people.

The growing dietary database from coprolite analysis supports the thrifty-gene hypothesis in general, but with a significant modification. There was a high reliance on highinulin fiber-rich plant foods with low GIs. However, these were not just famine foods, as suggested by the original hypothesis. These were the foods eaten on a day-by-day basis during all seasons, in both feast and famine. They continued to be eaten even after agriculture was developed. Antelope Cave coprolites show that this high-fiber diet was eaten during the warmer seasons of food abundance. Other sites, such as Antelope House in Canyon de Chelly, show that the reliance on high-fiber foods, especially yucca and prickly pear, was accentuated in winter and periods of ecological crisis (Sutton and Reinhard 1995). Therefore, we hypothesize that it was the consistent reliance on these foods that fixed the thrifty gene in Southwestern tribes. This hypothesis can be tested in future coprolite analyses and by review of existing data collected by coprolite analysts over the past decades.

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Supplement A from Reinhard et al., "Understanding the Pathoecological Relationship between Ancient Diet and Modern Diabetes through Coprolite Analysis"

(Current Anthropology, vol. 53, no. 4, p. 506)

Archaeological Background

Archaeological research on the Arizona Strip has been sporadic, and thus its prehistory is poorly known compared with other regions of the American Southwest. Altschul and Fairley (1989) offer a detailed overview of the environment and archaeology of the strip. Lyneis (1995) updates our knowledge of the prehistoric Virgin Anasazi, some of whom were the primary occupants of Antelope Cave over 1,000 years ago. The following synopsis is based on these two comprehensive publications plus additional information on Antelope Cave.

Evidence of Paleo-Indian presence on the Arizona Strip before 7000 BC is extremely minimal, being made up of one Clovis-like dart point and two Silver Lake point fragments.

The Archaic Period (7000–300 BC) also is typified by distinctive styles of projectile points that represent early dispersed hunter-gatherer groups. Some Archaic-style points—Rocker side-notched, San Rafael side-notched, and Elko corner-notched—are reported from Antelope Cave. In support, charcoal samples from the cave yielded three calibrated ¹⁴C dates of 2028–1893 BC, 1891–1744 BC, and 1699–1444 BC (Janetski and Wilde 1989), further confirming the initial occupation of Antelope Cave by Archaic groups. A few other sites (not lithic scatters) on the Arizona Strip have Archaic Period components, including Rock Canyon Shelter, approximately 7 miles north of Antelope Cave.

Basketmaker II (300 BC-AD 400) is the first Anasazi period to be recognized in the Southwest. In general, it is characterized by maize and squash cultivation, lack of ceramics, square-toed fiber sandals, human-hair cordage, slab-lined storage cists, pithouses, atlatl darts, coiled baskets, and rabbit-fur blankets. The question of whether the Basketmaker Puebloans evolved from Archaic hunter-gatherer groups is unresolved. Basketmaker II sites are typically pit structures or rock shelters and caves with slab-lined storage pits. Five are known on the Arizona Strip, including Antelope Cave, which has yielded an atlatl carbon dated at AD 100. Also, a small excavated area (fig. 1; UCLA excavation unit AC59-3, 4) near the west wall of the cave lacked ceramics but contained finely woven square-toed sandals and obsidian projectile points. This area is tentatively attributed to the Basketmaker II period (Johnson and Pendergast 1960:3).

Basketmaker III (AD 400–800) is represented by small pithouse villages, plain gray pottery, and cultivated beans to go with the growing of maize and squash. Bows and arrows begin to replace atlatls and darts during this period. Habitation sites are scattered over the Arizona Strip, and many are difficult to interpret because of the general lack of reliable dating. Probable Basketmaker III traits at Antelope Cave include cultivated beans, plain gray ceramics, and the wooden nock ends of arrows. In addition, three new radiocarbon dates from Antelope Cave indicate that the Anasazi more than any other group made the greatest use of the site during late Basketmaker III through early Pueblo I times. The ¹⁴C determinations are as follows.

| Beta 257788 | $1220 \pm 40 \text{ BP}$ | 680–890 cal AD (2σ) |
|-------------|---------------------------|---------------------|
| Beta 257787 | $1190~\pm~40~\mathrm{BP}$ | 710–960 cal AD (2σ) |
| Beta 257786 | $1230~\pm~40~\mathrm{BP}$ | 680–890 cal AD (2σ) |

Specimen Beta 257788 was run on a corncob (cat. 244–4835) from the 66–72-inch level of University of California, Los Angeles (UCLA) excavation unit AC59-5 (fig. 1). It represents the beginning of the Basketmaker III Anasazi occupation in this area of the cave. Beta 257787 dates a yucca quid (cat. 244-1490) from UCLA AC59-2, 24–30 inches below the surface. This sample was specifically selected to date the level containing a newly discovered tick that had been eaten by one of the cave's occupants (Johnson et al. 2008). The third date, Beta 257786, is from a yucca quid (cat. 244-658) in the top level (0–6 inches) of UCLA AC59-2. Unfortunately, the dated specimen appears to be from badly disturbed midden at the top of the excavation unit. These three new radiocarbon assays equate nicely with those from the Brigham Young University excavations in 1983 (Janetski and Hall 1983:40, 43).

The Pueblo I and Pueblo II periods (AD 800–1150) are characterized by the continuation and elaboration of Basketmaker III cultural traits. Pithouse villages are usually small and increase in number, and the houses may have

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benches and ventilators. Aboveground masonry structures first occur during this time period. Black-on-gray and black-on-white decorated pottery make their appearance early on. Corrugated pottery is a new style that debuts in the Pueblo II period (AD 1000–1150). Round- or pointed-toed sandals begin to replace square-toed sandals by Pueblo I times. At Antelope Cave, five Virgin Anasazi types of decorated ceramics as well as round-toed yucca sandals mark these two culture stages. However, the seasonal use of Antelope Cave by Ancestral Puebloans was in decline by AD 1000. Of the 3,100 pottery sherds recovered during the UCLA excavations, only three were corrugated. This suggests little use of the cave during much of Pueblo II.

Several Anasazi sites have been dated to Early Pueblo III (AD 1150–1225) on the Arizona Strip. The sites are still small, but some are larger than the settlements of previous periods. As usual, the family household remains the basic economic unit. At this time, Antelope Cave had been pretty much abandoned by the Virgin Anasazi. Where they went or what happened to them is uncertain.

Prehistoric Southern Paiute groups began to move into Virgin Anasazi territory around AD 1000, and these newcomers eventually either absorbed or replaced the Ancestral Pueblo peoples on the strip. A large twined and pitch-covered Paiute water basket was recovered from the surface of Antelope Cave by Vilate Hardy of La Verkin, Utah, in the early 1950s. The location of this specimen in the cave provides a bit of evidence that the Virgin Anasazi were either in contact with the Southern Paiute or the Southern Paiute actually occupied the cave sometime after it was abandoned by the Pueblo people.

Figure 1 is a plan drawing of Antelope Cave. The main entrance to the cave (now gated by the Bureau of Land Management) is on its southeast edge above large slabs of jumbled limestone. Once inside, a massive layer of broken limestone roof fall, beginning 10 feet below the entrance, extends over the eastern half of the cave. The dry and gray powdery midden deposit covers most of the western half. The surface of the midden slopes down rapidly from south to north, ending at a large secondary sinkhole, the bottom of which is 75 feet below the entrance to the cave.

Scientific investigation of the midden deposit began in 1954 when Robert Euler of the Museum of Northern Arizona (MNA) directed the excavation of four test pits (A, B, C, D). In 1983 and 1986, Brigham Young University (BYU) archaeologists under contract with the Bureau of land Management sampled six areas in the cave, including the secondary sink (Janetski and Hall 1983; Janetski and Wilde 1989). Their work provided excellent perishable and nonperishable cultural materials as well as important radiocarbon dates. Between the MNA and BYU excavations, archaeologists from UCLA carried out extensive investigations in the cave. In 1956 and 1957, under the direction of Robert Euler, UCLA crews from summer archaeology field schools in Utah dug units E and B adjacent to MNA pits C and D. Encouraged by Vilate Hardy and supported by a grant from the Department of Anthropology and Sociology at UCLA, three UCLA archaeologists spent 19 days in June of 1959 excavating units AC59-1 through 5 (Johnson and Pendergast 1960). AC59-1 is located close to the rock fall at the center of the cave. The midden became damp and rocky and was abandoned at 48 inches below the surface. AC59-3 and 4 were placed in a debris mound at the west edge of the cave. This area yielded early Basketmaker materials (Johnson and Pendergast 1960). AC59-2 and 5 were located in the culturally richest area of the midden near the north end of the cave, where the deposit is 72 inches deep. UCLA archaeologists returned to Antelope Cave for a few days in April 1960 to complete the excavation of AC59-5 and investigate two "living areas" abutting the west edge of the cavern. These possible living areas, the largest measuring 30 × 15 feet, were relatively flat and in 1959 appeared to be generally clear of surface cultural debris such as twigs, pottery fragments, corncobs, etc. Test unit AC60 was excavated to sample the subsurface of the larger "living area." The soil in the pit was brown rather than the typical gray midden of the site. A large limestone rock was exposed 2 inches below the surface, and two superimposed burned areas or fire hearths were encountered before rocks at 12 inches below the surface prevented further excavation. Among the rock fall on the eastern half of the site is a huge flat limestone slab. Three shallow fire hearths were noted on its surface by the UCLA crew in 1959, but they were not completely recorded until 2009. Fire hearth 3, the smallest of the three in diameter, is 2 inches thick and composed of very black charcoal fragments along with burned pine needles. Radiocarbon analysis of soil from hearth 3 (Beta 264019, 180 ± 40 BP, 1650–1950 cal AD [2σ]) indicates that visitors were still using the cave hundreds of years after it was abandoned by the Virgin Anasazi.

Prehistoric human coprolites from UCLA's work at Antelope Cave provide the basic data for this report. The excavations recovered 190 feces. They were found scattered in all excavation units with the exception of pits E and AC60, the latter of which may be in a family living area. The highest concentration of coprolites, 13, occurred in the 30–36-inch level of AC59-4, which yielded Basketmaker II materials. Twenty-five coprolites were selected for dietary analysis. Their provenience in Antelope Cave is listed in table A1 in the order of their assigned laboratory numbers.

Table A1. Major and minor food components listed for each coprolite based on the analyses summarized in supplementary results and tables

| Lab no. | Main association | Secondary association |
|---------|---|--|
| 1 | Roasted prickly pear pads | Dropseed seeds and pollen could be from a meal of caryopses without grinding or cooking |
| 2 | Not fully cooked stew of finely processed maize flour and rabbit | |
| 3 | Possibly a stew of highly processed sunflower and small-mammal meat and bone | High grass and cheno-am pollen concentrations result from earlier meals of these plants (probably seeds) |
| 4 | Fully cooked stew of maize and wolfberry fruit with small-animal meat and bone | Other foods are represented by prickly pear, sunflower, and other grass residues |
| 5 | Cooked stew of animal meat, bone, and ground dropseed flour | Prickly pear pads |
| 7 | Cooked stew of maize flour and small-mammal meat and bones | Prickly pear pads |
| 9 | Stew of small-mammal meat and bone and finely ground unidentifiable seed prepared with sagebrush | Prickly pear pads |
| 10 | Roasted prickly pear pads | |
| 11 | Crushed four-winged saltbush seeds, roasted prickly pear pads, stems or leaves of unidentifiable plant, and small-mammal meat and bone | |
| 12 | Stew of cooked maize and small-animal meat and bone | Prickly pear pads |
| 13 | Finely ground dropseed flour | Prickly pear pads, four-winged saltbush, sunflower, and other grass |
| 14 | Stew of sunflower and small-mammal meat and bone | Wild grass and cheno-am seeds |
| 15 | Parched and cooked plant food, source unidentifiable, with small-animal meat and bone | Prickly pear pads |
| 16 | Stew of sunflower and small-mammal meat and bone | |
| 17 | Stew or soup made from ground grass and small-mammal meat and bone | Prickly pear pads |
| 18 | Parched coarsely ground maize and roasted prickly pear pads | Termites probably from a previous meal |
| 19 | Roasted prickly pear pads and small-mammal meat and bone | Maize |
| 20 | Seed cake or stew composed of finely ground cheno-am fruits associ- ated with ground dropseed caryopses | Maize |
| 21 | Roasted prickly pear pads | Cooked maize and dropseed |
| 23 | A stew of roasted prickly pear pads, cheno-am seed, and small-mammal meat and bone | |

Note. The main association column lists the major foods found in each coprolite and an interpretation of the nature of the original foods. The secondary association is based on traces of the designated foods that may reflect vestiges of a separate meal or secondary foods from the same meal as the main association. The designation "cooked" or "completely cooked" is based on the presence of heat-altered starch or loss of starch completely. The designation "not fully cooked" is based on the presence of pristine, unaltered starch. The designation "processed" refers to ground foods. "Roasted" prickly pear is evidenced by the heat-altered white appearance of epidermal fragments. "Meat" is inferred by the find of bone and hair. Most identifiable bone and hair was either cottontail or jackrabbit. It is likely that all of the meat represented by bones is from rabbits with the exception of coprolite 4. Feather calami (quill bases) in 4 suggest that birds were eaten. "Parched" designation is based on association of carbon mixed with plant tissue. This could be from parching plant foods by swirling them with hot coals. "Stew" is a suggested method of preparing an association of foods applied to reoccurring associations.

Reference Cited Only in Supplement A

Johnson, Keith L., Karl J. Reinhard, Luciana Sianto, Adauto Araújo, Scott L. Gardner, and John Janovy Jr. 2008. A tick from a prehistoric Arizona coprolite. *Journal of Parasitology* 94:296–298.

Supplement B from Reinhard et al., "Understanding the Pathoecological Relationship between Ancient Diet and Modern Diabetes through Coprolite Analysis"

(Current Anthropology, vol. 53, no. 4, p. 506)

Material and Methods

Preliminary Steps

The specimens were assigned laboratory numbers and logged into a laboratory notebook by lab number and provenience (table B1). After photographing each specimen, we made specific observations about the shape, size, and content (as evident from the surface) of each coprolite. We also noted evidence of decomposition, such as the presence of larvae cases and beetle and arthropod holes. The coprolites were subsampled, and each subsample was weighed. Then the subsamples were rehydrated for 48 hours in 0.5% trisodium phosphate. *Lycopodium* tablets, each containing 12,500 spores, were added to the rehydrating samples. One tablet was added for each gram of coprolite with the exception of sample 15. One tablet was added to every 2 g of that sample. The rehydrated coprolites and dissolved *Lycopodium* tablets were disaggregated with a magnetic stirrer. The magnetic stirrer releases microfossils that might otherwise be trapped in the plant remains. The samples were then screened through a 150- μ m mesh. The fluid passing through the screen was collected in a large glass beaker and then centrifuged in 100-mL centrifugation tubes. The concentrated solid microscopic remains were then transferred to 50-mL tubes for microfossil extraction. The macroscopic remains on top of the screens were transferred to blotter paper and dried for analysis.

Macrofossil Dietary Analysis

The dried macrofossil remains were screened through 2.0-mm, 1.0-mm, and 0.5-mm geological sieves. The remains from each screening were examined with a binocular dissecting microscope. The remains were sorted by hand using forceps, tweezers, and wooden spatulettes. The macroscopic plant constituents were identified using the seed comparative collection and by published and online seed-image databases. In certain instances, wet mounts were made of plant tissue so that the cellular and phytolith arrangements could be examined with the compound microscope.

Microfossil Analysis

The key to microfossil quantification, analysis, and interpretation is the addition of Lycopodium spores. To calculate the concentrations of microfossils in samples of sediment, we added known number of Lycopodium spores into the samples (Reinhard et al. 2006). By adding a known number of exotic spores, we can estimate the amount or concentration of all types of microfossils. Concentrations of parasites, pollen grains, and starch grains are calculated with the following formula: microfossil concentration = $[(f/m) \times e]/v$, where f = microfossils counted, m = marker Lycopodium spores added, and m = marker Lycopodium spores counted.

For this analysis, *Lycopodium* spore batch 212761 was used. Previous analysis shows that approximately 12,500 spores are present in each tablet.

In order to retain Lycopodium spores for each type of analysis, a sequential microfossil analysis was done. First, parasitological scans were accomplished. Twelve microscope preparations were made and examined for parasite eggs and larvae at $250 \times$ with photographs taken at $400 \times$. The same slides were scanned for starch, calcium oxalate phytoliths, plant tissue, animal hair, and any other identifiable remains. This stage of analysis focuses on heavier mineral remains such as phytoliths.

The sediments were then treated with heavy-density zinc bromide solution. The solution is made with zinc bromide diluted to the desired specific gravity of 1.9 with 2% hydrochloric acid. The heavy-density solution was added to each centrifuge tube containing the microfossils. The tubes were centrifuged for 15 minutes to separate heavy from light remains. Three microscope preparations from the light fractions were analyzed for starch grains and parasite eggs. Then the analysis proceeded to palynology.

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The extraction of the pollen involved chemical destruction of silica, cellulose, and starch, leaving concentrated and stained pollen. The light fraction of microfossil remains were treated with hydrofluoric acid (40%). The acid was added to centrifuge tubes with the sediments. The tubes were placed in a hot water bath (95°–98°C) for an hour. The remains were then washed three times with distilled water and subsequently rinsed twice with glacial acetic acid preparatory to acetolysis. The acetolysis mixture of one part sulfuric acid and eight parts acetic anhydride was added to each tube. All tubes were then placed in a hot water bath for 15 minutes. Finally, the samples were washed with acetic acid and then several times with water until the supernatant was clear.

Following the chemical extraction, the residue was washed with 95% alcohol and transferred to small vials in alcohol. Microscope slides were prepared by pipetting a drop of residue onto a slide, allowing most of the alcohol to evaporate, and mixing in a drop of glycerol. A cover glass was placed on top and sealed with fingernail polish. A minimum count of 200 pollen grains was made for each sample at $400 \times$.

One important detail relates to identification of maize pollen. Many of the maize pollen grains were broken and torn by food processing. To prevent overcounting maize pollen, only fragments that exhibited an annulus were counted. Each maize pollen grain has only one annulus. Therefore, by counting only fragments with annuli, we prevented overcounting maize pollen.

Table B1. Coprolites analyzed from Antelope Cave by laboratory and field specimen (FS) numbers

| Lab no. | FS/excavation unit/depth in inch level | Weight (g) | Weight of macro residue (g) | No. <i>Lycopodium</i> tablets added |
|--------------------------------|--|------------|-----------------------------|-------------------------------------|
| Human coprolites: | | | | |
| 1 | 2487/AC59-4/30-36 | 3.66 | 1.65 | 4 |
| 2 | 1516/AC59-2/24-30 | 2.67 | 1.23 | 3 |
| 3 | 617/AC59-1/30-36 | 15.77 | 8.76 | 8 |
| 4 | 2302/AC59-4/6-12 | 6.39 | 3.55 | 6 |
| 5 | 2103/AC59-3/12-18 | 30.55 | 14.17 | 15 |
| 7 | 644/AC59-2/0-6 | 5.32 | 2.45 | 3 |
| 9 | 3172/AC59-5/12-18 | 7.0 | 3.42 | 4 |
| 10 | 3557/AC59-5/18-24 | 5.77 | 2.96 | 3 |
| 11 | 54/surface find | 2.56 | 1.59 | 3 |
| 12 | 153-294a/B/24-36 | 1.84 | .93 | 2 |
| 13 | 153-294b/B/24-36 | 2.51 | 1.55 | 3 |
| 14 | 617/AC59-1/30-36 | 8.02 | 4.58 | 4 |
| 15 | 1516/AC59-2/24-30 | 3.07 | 5.35 | 3 |
| 16 | 641/AC59-2/0-6 | 3.95 | 1.28 | 4 |
| 17 | 2103/AC59-3/12-18 | 29.27 | 16.32 | 15 |
| 18 | 2487/AC59-4/30-36 | 2.6 | 1.13 | 3 |
| 19 | 3557/AC59-5/18-24 | 2.97 | 1.53 | 3 |
| 20 | 3957/AC59-5/24-30 | 6.78 | 4.07 | 7 |
| 21 | 4874/AC59-5/36-42 | .2 | .06 | 1 |
| 23 | 244-000/surface find | 10.25 | 5.49 | 5 |
| Sediment sample, noncoprolite: | | | | |
| 22 | 244-4547/AC59-5/36-42 | 14.49 | ··· | 7 |
| Animal coprolites: | | | | |
| 6 | 153-294/B/24-36 | 7.54 | ••• | 4 |
| 8 | 2955/AC59-5/6-12 | 7.46 | | 4 |
| A1 | 153-262/B/12-24 | 8.18 | ••• | 4 |
| A2 | 4773/AC59-5/48-54 | 10.85 | | 5 |

Note. Weight and Lycopodium data are not available for Fundação Oswaldo Cruz 616 and 643 because dietary analysis of these samples is in progress.

Reference Cited Only in Supplement B

Reinhard, Karl J., Sherrian K. Edwards, Teyona R. Damon, and Debra K. Meier. 2006. Pollen concentration analysis of Ancestral Pueblo dietary variation. *Palaeogeography, Palaeoclimatology, and Palaeoecology* 237:92–109.

Supplement C from Reinhard et al., "Understanding the Pathoecological Relationship between Ancient Diet and Modern Diabetes through Coprolite Analysis"

(Current Anthropology, vol. 53, no. 4, p. 506)

Results

Preservation

The only decomposer insects discovered in the analysis were spider beetles, which were found in six coprolites. These never amounted to more than a trace of a gram, and never more than three were recovered from a coprolite sample. No invertebrate burrows were observed in the coprolites. No fly remains were found. Some mites were noted in microscopic analysis, but these were few and may have been ingested with plant food. No free-living nematodes were found. These observations attest to the excellent preservation of the Antelope Cave coprolites. Spider beetles prefer dry substrates. Flies and nematodes prefer moist substrates. The presence of a few beetles and the absence of flies and nematodes shows that the coprolites desiccated rapidly.

Diet

The dietary results are presented in the tables in this supplement. The definition of terms used in data tables is presented in tables C1 and C2. The data for microresidues is presented in tables C3–C8, and the macroresidue results are in tables C9–C13. The pollen counts and concentrations are presented in tables C14–C24. These results for the human coprolites are summarized below. Coprolites 6 and 22 are omitted from these results because analysis revealed that they were not human coprolites. Number 6 was an animal coprolite, and number 22 was consolidated cave sediment.

Coprolite laboratory number 1, FS 2487, is composed macroscopically of prickly pear pad fragments with traces of whole dropseed (*Sporobolus*) caryopses. The microfossils independent of pollen are exclusively from prickly pear. The pollen count is dominated by grass. This shows a meal of prickly pear pads, which were most likely roasted as evidenced by the heat-altered white appearance of epidermal fragments. The dropseed seeds and pollen could be from an earlier meal of caryopses eaten off of the plant without processing or cooking.

Coprolite laboratory number 2, FS 1516, is composed mostly of finely ground maize kernels, finely ground sunflower achenes, unknown plant epidermis and fiber, and bone. The bone is highly fragmented and eroded. The microfossils are dominated by maize starch with grass stem/epidermis fragments. There are traces of ring structures from prickly pear vascular bundles. The pollen has some cottonwood-type grains, but this type presents a problem because cottonwood can resemble many other spores and pollen from other taxa. This coprolite represents a meal of highly processed maize and probably rabbit apparently eaten together, perhaps in a stew, which would explain the erosion of the bone fragments. The maize was not extensively cooked because the maize starch is in a pristine form.

Coprolite laboratory number 3, FS 617, is composed macroscopically of a mix of finely ground sunflower achenes and bone. The bone is fragmented and eroded rabbit or rodent bone. Microscopically, sunflower fragments dominate the remains. The microfossil residue is rich in fibers, sunflower achene fragments, and seed coat fragments to the point that we could not estimate the actual numbers of them. Palynologically, the higher grass and cheno-am pollen concentrations could be influenced by earlier meals. This represents a meal of highly processed sunflower and fragmented small mammals, apparently eaten together, perhaps in a stew. The flour made of sunflowers would have been nearly inedible unless processed into a stew.

Coprolite laboratory number 4, FS 2302, is composed of feather calami (quill bases), whole wolfberry seeds from fruit, coarsely ground maize, and fragmented bone probably from rabbit. The nonpollen microremains are diverse. Round starch granules averaging 18 μ m in diameter with hila dominate the microscopic spectrum. These are probably from maize. Leaf epidermis fragment, grass epidermis, and xylem tracheids represent grasses and other vegetation. Prickly pear is represented by glochidia fragments and ring structures. Traces of sunflower achene fibers are also present. Palynologically, wild grass dominates the pollen spectrum. It may be that this coprolite represents a meal of bird and perhaps small-animal meat eaten with maize and wolfberry fruit. Wolfberry must be cooked to disperse poisonous compounds. This

1

may have been a stew, but it was not highly cooked because none of the 297 observed starch granules exhibit heat alteration. The prickly pear, sunflower, and other grass residues are from previous meals.

Coprolite laboratory number 5, FS 2103, is composed macroscopically of very finely ground plant residue, highly fragmented animal bone, finely ground dropseed, and an unidentified grass seed. The pollen is dominated by grass. The nonpollen microfossils are dominated by cactus calcium oxalate cactus druses and cactus glochidia. The druses and glochidia are from a type of cactus that is new to us. The absence of starch in this sample suggests extensive cooking in water, which would have destroyed the starch. Therefore, it may be that this was a stew of animal meat, bone, and ground dropseed.

Coprolite laboratory number 7, FS 644, macrofossils were dominated by fragmented bone and very finely ground maize. Microscopically, unidentifiable plant residue dominated the count. This sample appears to be derived from a combination of maize flour and small mammal, possibly eaten together in the form of a stew or soup. It appears that this was highly cooked in water because only one starch granule, a cooked maize grain, was observed. The pollen analysis revealed small amounts of wind-pollinated background types.

Coprolite laboratory number 9, FS 3172, is composed of fragmented bone and finely ground unidentifiable seed. The microscopic analysis shed no light on the origin of the seed. This is an enigmatic sample except that it is a repeat of the association of fragmented small-mammal bone with finely ground seed that is common in Antelope Cave coprolites. The pollen suggests the intentional use of sagebrush, *Artemisia*. There is a high concentration of sagebrush pollen and pollen aggregates of this taxon. Sagebrush is toxic to humans, but it is also medicinal. Treatments made of sagebrush taken internally kill intestinal worms and have an antibacterial effect. It was also used to treat internal bleeding (Tilford 1997).

Coprolite laboratory number 10, FS 3557, shows only prickly pear pad parts, both macroscopically and microscopically. The masses of fiber in this sample are probably from prickly pear. The prickly pear epidermis is whitened and made brittle by heat exposure and probably represents roasted prickly pear. The pollen analysis shows low concentrations of a diversity of pollen types but does not suggest economic use of these taxa.

Coprolite laboratory number 11, FS 54, is dominated by crushed seed, possibly four-winged saltbush with traces of bone fragments. Microscopically, there is an abundance of conductive vascular tissue from plants. Pollen analysis does not help identify the origin of the seed. Only small amounts of background types are present.

Coprolite laboratory number 12, FS 153-294a, is dominated macroscopically and microscopically by maize with traces of fragmented bone. The condition of maize starch shows that these foods were cooked. Of 220 observed maize starch granules, 219 show alteration due to cooking. Therefore, it appears that this is the result of eating a stew of maize and small-animal meat and bone. Interestingly, no maize pollen was recovered from this coprolite.

Coprolite laboratory number 13, 153-294b, is dominated macroscopically and microscopically by finely ground dropseed. There is also a lesser amount of crushed unknown seed similar to four-winged saltbush, probably from a previous meal. There is a high concentration of wild-grass pollen and aggregates of wild-grass pollen. This indicates that wild grass was consumed. There are traces of prickly pear in the form of microscopic glochidia, probably from a previous meal.

Coprolite laboratory number 14, FS 617, is dominated macroscopically and microscopically by ground sunflower achenes. There are also traces of bone and traces of cheno-am seeds. It is likely that the cheno-am seeds are from a previous meal, and a stew or soup of sunflower and small mammal was the meal most represented by this coprolite. Cheno-am pollen aggregates are present in this coprolite. Poaceae aggregates may be the residue of a previous meal of wild-grass seed.

Coprolite laboratory number 15, FS 1516, is a very difficult coprolite to interpret. There is fragmented small-mammal bone. However, the majority of the macroscopic and microscopic remains are of black granular material composed of carbon mixed with plant tissue. This could be from parching plant foods with hot coals. The pollen reveals one grass-pollen aggregate of two grains, but this is not significant.

Coprolite laboratory number 16, FS 641, is dominated macroscopically and microscopically by ground sunflower achenes. There is also fragmented small-mammal bone. This is a mixture of sunflower flour and crushed animal. Like coprolite 3, I believe these foods must have been a stew, because a flour made of sunflowers would have been nearly inedible. This coprolite is unique in that pollen was nearly absent. Extensive examination of several microscopic preparations reveal only one pollen grain. The absence of ambient pollen is very interesting. It might be that this coprolite was deposited in the cave at a time of low pollination, possibly winter.

Coprolite laboratory number 17, FS 2103, is dominated by fragmented small-mammal bone, jackrabbit claws, and extremely finely ground dropseed. Microscopically, there are hundreds of starch granules that are not birefringent. This appears to be a stew or soup made from ground grass and fragmented small mammal. There is a high concentration of wild-grass pollen with aggregates. This pollen was ingested with the seeds and inflorescences.

Coprolite laboratory number 18, FS 2487, is like coprolite 15. Fragmented bone appears with ash mixed with plant residue in a black granular substrate. The advantage with his coprolite is that there was some material liberated from the

aggregates. There was some coarsely ground maize. Microscopically, cactus glochidia and conductive plant tissue was clumped with ash with a few ring structures. This suggests that prickly pear pads were roasted, which resulted in the incorporation of ash or perhaps parched maize with prickly pear. Eleven thousand maize pollen grains per gram of coprolite were recovered. The majority of these are torn and fragmented from grinding. Three termites were found and may reflect dietary use of these insects.

Coprolite laboratory number 19, FS 3557, revealed macroscopic remains of fragmented small-mammal bone, tufts of jackrabbit hair, and aggregates of prickly pear epidermis with fiber and phytoliths. Microscopically, glochidia and prickly pear druses were the most common remains. This indicates that rabbit and prickly pear were eaten together. Wandsnider (1997) reviewed the method of cooking in Plains roasting pits and notes that plants and rabbits were roasted together. It appears that the composition of this coprolite, including hair, represents the preparation of rabbit and prickly pear together. Interestingly, 50,000 pollen grains of maize per gram of coprolite were evidenced by the pollen analysis. About half of these are torn. Thus, the pollen evidence shows that maize was eaten, probably independently and previously to the prickly pear and rabbit.

Coprolite laboratory number 20, FS 3957, is composed of finely ground cheno-am fruits associated with ground dropseed caryopses. Microscopically, remains of cheno-am and Poaceae dominate, although there are traces of sunflower. The pollen spectrum was dominated by wild grass, and many wild-grass pollen aggregates were noted. This appears to have been a seed cake or stew.

Coprolite laboratory number 21, FS 4874, contains a diversity of items and shows that analysis of even a small coprolite reveals a variety of information. Macroscopically, dropseed caryopses and prickly pear phytoliths dominate. Microscopically, there is a diversity of starch. Both cooked and uncooked maize starch is present. In addition there are two other starch forms from unknown plants and a variety of anatomical elements of prickly pear structures and grass. This represents as many as three dietary episodes of prickly pear, maize, and dropseed. The pollen spectrum was dominated by wild grass, and many wild-grass pollen aggregates were noted.

Coprolite laboratory number 23, FS 244-2256, is an association of fragmented small-animal bone and very finely ground cheno-am. Cheno-am seed coats and a variety of starch granules are evident microscopically. Nearly 70,000 cheno-am pollen grains were recovered per gram of coprolites, some of which were aggregates. Again, this appears to be an association of seed and meat in a stew or soup.

Table C1. Definition of terms applied to microscopic remains

| Exotic spores of arctic clubmoss added for quantification Animal hair Nonhuman hair Nonhuman hair Cac ₃ O ₄ druse Calcium oxalate phytolith probably from cactus Calcium oxalate phytolith probably from cactus Calcium oxalate phytolith probably from cactus Calcium oxalate phytolith from prickly pear Seeds that could be from plants in either the genera Chenopodium or Amaranthus or less likely Cycloloma Pollen grains that could be from either the family Chenopodiaceae or the genus Amaranthus Starch granules found within isolated plant cell Curious association of any type of plant remain, primarily from CaC ₂ O ₄ or silicified origins. They represent the most durable plant structures Stellate, microscopic, recurved spines associated with cactus are-oles and specific to cacti Glochidia are fragile and break into their component spines. These are individual spines Achenes are the simple fruits produced by the sunflower family and a few other families. These are consistent with very small sunflower fruits Fragments of exoskeleton of insects or other small arthropods Leaf epidermis fragment Segments of plant epidermis with stomata Isolated sieve-tube members from the phloem that conduct food materials in plants Segment of phloem with sieve-tube element, companion cells, and other phloem components Phytolith, unknown Phytolith that cannot be identified to plant or plant structure Plant epidermis, unknown Phytolith that cannot be identified to plant or plant structure Plant fiber Long tapered fibers from xylem or phloem Plant fiber Long tapered fibers from xylem or phloem Plant fiber Long tapered fibers arranged parallel in plant-tissue sections | Term | Definition |
|---|--|--|
| CaC ₂ O ₄ druse CaC ₂ O ₄ opuntioid druse Cheno-am macrofossils and nonpollen microfossils Cheno-am pollen Cheno-am pollen Cheno-am pollen Clump of mineralized plant structures in ash Clochidia Glochidia Glochidia Glochidia Felianthus achene fibers Helianthus achene fibers Arthropod fragment Leaf epidermis fragment Phloem sieve-tube element Phytolith, unknown Phytolith, unknown Plen Calcium oxalate phytolith from prickly pear Seeds that could be from plants in either the genera Chenopodium or Amaranthus or less likely Cycloloma Pollen grains that could be from either the family Chenopodiaceae or the genus Amaranthus Starch granules found within isolated plant cell Curious association of any type of plant remain, primarily from CaC ₂ O ₄ or silicified origins. They represent the most durable plant structures Stellate, microscopic, recurved spines associated with cactus areoles and specific to cacti Glochidia are fragile and break into their component spines. These are individual spines Achenes are the simple fruits produced by the sunflower family and a few other families. These are consistent with very small sunflower fruits Fragments of exoskeleton of insects or other small arthropods Segments of plant epidermis with stomata Isolated sieve-tube members from the phloem that conduct food materials in plants Phytolith, unknown Phytolith that cannot be identified to plant or plant structure Plant epidermis, unknown Phytolith that cannot be identified to plant or plant structure Plant epidermis, unknown morphology Plant fiber Long tapered fibers from xylem or phloem | Lycopodium | Exotic spores of arctic clubmoss added for quantification |
| CaC_O_4 opuntioid druse Cheno-am macrofossils and nonpollen microfossils Cheno-am macrofossils and nonpollen microfossils Cheno-am macrofossils and nonpollen microfossils Cheno-am pollen Cheno-am pollen Cheno-am pollen Cell containing starch Clump of mineralized plant structures in ash Clump of mineralized plant structures Clochidia Clump of mineralized plant structures Clochidia Clump of mineralized plant structures Clochidia Clump of mineralized plant structures in ash Curious association of any type of plant remain, primarily from CaC_O_O or silicified origins. They represent the most durable plant structures Clochidia are fragile and break into their component spines. These are individual spines Helianthus achene fibers Achenes are the simple fruits produced by the sunflower family and a few other families. These are consistent with very small sunflower fruits Segments of exoskeleton of insects or other small arthropods Segments of plant epidermis with stomata Phloem sieve-tube element Isolated sieve-tube members from the phloem that conduct food materials in plants Segment of phloem with sieve-tube elements, companion cells, and other phloem components Phytolith, unknown Phytolith that cannot be identified to plant or plant structure Epidermis, that shows unknown morphology Plant fiber Long tapered fibers from xylem or phloem | Animal hair | Nonhuman hair |
| Cheno-am macrofossils and nonpollen microfossils Cheno-am pollen Cheno-am pollen Cheno-am pollen Cheno-am pollen Cell containing starch Clump of mineralized plant structures in ash Cac ₂ O ₄ or silicified origins. They represent the most durable plant structures Glochidia Clochidia are fragile and break into their component spines. These are individual spines Achenes are the simple fruits produced by the sunflower family and a few other families. These are consistent with very small sunflower fruits Arthropod fragment Clochidia Clochidia Clochidia are fragile and break into their component spines. These are individual spines Clochidia are fragile and break into their component spines. These are individual spines Clochidia are fragile and break into their component spines. These are individual spines Clochidia are fragile and break into their component spines. These are individual spines Clochidia are fragile and break into their component spines. These are individual spines Clochidia are fragile and break into their component spines. These are individual spines Clochidia are fragile and break into their component spines. These are individual spines Clochidia are fragile and break into their component spines. These are individual spines Clochidia are fragile and break into their component spines. These are individual spines Clochidia are fragile and break into their component spines. These are individual spines Clochidia are fragile and break into their component spines. These are individual spines Clochidia are fragile and break into their component spines. These are individual spines Clochidia are fragile and break into their components Clochidia are fragile and break into their components Clochidia are fragile and break in | CaC ₂ O ₄ druse | Calcium oxalate phytolith probably from cactus |
| Cheno-am pollen Cheno-am pollen Pollen grains that could be from either the family Chenopodiaceae or the genus Amaranthus Cell containing starch Clump of mineralized plant structures in ash Cac ₂ O ₄ or silicified origins. They represent the most durable plant structures Glochidia Stellate, microscopic, recurved spines associated with cactus areoles and specific to cacti Glochidium fragment Glochidia are fragile and break into their component spines. These are individual spines Helianthus achene fibers Achenes are the simple fruits produced by the sunflower family and a few other families. These are consistent with very small sunflower fruits Arthropod fragment Fragments of exoskeleton of insects or other small arthropods Leaf epidermis fragment Phloem sieve-tube element Segments of plant epidermis with stomata Phloem with sieve-tube element Segment of phloem with sieve-tube elements, companion cells, and other phloem components Phytolith, unknown Phytolith that cannot be identified to plant or plant structure Plant epidermis, unknown Phytolith that cannot be identified to plant or plant structure Plant fiber Fragrent fibers from xylem or phloem | CaC ₂ O ₄ opuntioid druse | Calcium oxalate phytolith from prickly pear |
| aceae or the genus Amaranthus Cell containing starch Clump of mineralized plant structures in ash CaC ₂ O ₄ or silicified origins. They represent the most durable plant structures Glochidia Stellate, microscopic, recurved spines associated with cactus areoles and specific to cacti Glochidium fragment Glochidia are fragile and break into their component spines. These are individual spines Helianthus achene fibers Achenes are the simple fruits produced by the sunflower family and a few other families. These are consistent with very small sunflower fruits Arthropod fragment Arthropod fragment Fragments of exoskeleton of insects or other small arthropods Segments of plant epidermis with stomata Phloem sieve-tube element Segment of phloem with sieve-tube elements, companion cells, and other phloem components Phytolith, unknown Phytolith, unknown Phytolith that cannot be identified to plant or plant structure Plant epidermis, unknown Epidermis that shows unknown morphology Plant fiber Long tapered fibers from xylem or phloem | Cheno-am macrofossils and nonpollen microfossils | |
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| and a few other families. These are consistent with very small sunflower fruits Arthropod fragment Fragments of exoskeleton of insects or other small arthropods Leaf epidermis fragment Segments of plant epidermis with stomata Phloem sieve-tube element Isolated sieve-tube members from the phloem that conduct food materials in plants Phloem with sieve-tube element Segment of phloem with sieve-tube elements, companion cells, and other phloem components Phytolith, unknown Phytolith that cannot be identified to plant or plant structure Plant epidermis, unknown Epidermis that shows unknown morphology Plant fiber Long tapered fibers from xylem or phloem | Glochidium fragment | |
| Leaf epidermis fragmentSegments of plant epidermis with stomataPhloem sieve-tube elementIsolated sieve-tube members from the phloem that conduct food materials in plantsPhloem with sieve-tube elementSegment of phloem with sieve-tube elements, companion cells, and other phloem componentsPhytolith, unknownPhytolith that cannot be identified to plant or plant structurePlant epidermis, unknownEpidermis that shows unknown morphologyPlant fiberLong tapered fibers from xylem or phloem | Helianthus achene fibers | and a few other families. These are consistent with very small |
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| and other phloem components Phytolith, unknown Phytolith that cannot be identified to plant or plant structure Plant epidermis, unknown Plant fiber Epidermis that shows unknown morphology Long tapered fibers from xylem or phloem | Phloem sieve-tube element | |
| Plant epidermis, unknown Epidermis that shows unknown morphology Plant fiber Long tapered fibers from xylem or phloem | Phloem with sieve-tube element | |
| Plant fiber Long tapered fibers from xylem or phloem | Phytolith, unknown | Phytolith that cannot be identified to plant or plant structure |
| 8 mb | Plant epidermis, unknown | Epidermis that shows unknown morphology |
| Plant fiber bundle Fibers arranged parallel in plant-tissue sections | Plant fiber | Long tapered fibers from xylem or phloem |
| | Plant fiber bundle | Fibers arranged parallel in plant-tissue sections |

Supplement C from Reinhard et al., Ancient Diet and Modern Diabetes

Plant hair, mineralized

Plant hair

Small hair-like structures called trichomes derived from plant epi-

dermal cells

Poaceae epidermis Epidermis from grass
Poaceae leaf epidermis Grass epidermis with stomata

Poaceae long cell Long dendritic epidermal cell phytolith from grass

Raphide CaC₂O₄ needle-shaped phytolith found in several plant families

Raphide bundle Mass of raphides arranged in parallel

Ring structure with interference cross Birefringent doughnut-shaped structures arranged in columns

within tubes and here found only in cactus and specifically

prickly pear

Seed coat fragment Seed testa unidentifiable to plant taxon

Seed coat (cheno-am?) Seed coats from many species of the goosefoot family or some

species of the pigweed family

As above but mineralized into phytoliths

Seed testa, light color Seed testa perhaps from ground grass or maize Starch, indeterminate Very small starch grains with no distinct features

Starch, cheno-am Faceted granules 5 μ m in diameter with hila and found in

aggregates

Starch without interference cross Medium-sized spheroidal granules 10– $20~\mu m$ in diameter that are

not birefringent. All other starch are birefringent

Starch, 15 µm, round, no hilum Distinctive starch of an unknown source

Starch, 15 µm, round, monocolpate Distinctive starch of an unknown source that has a single groove

on the surface

Starch, 15 µm, faceted, with hilum

Distinctive starch of an unknown source
Starch, 18 µm, round, with hilum

Distinctive starch of an unknown source
Starch, 11 µm, round, with large hilum

Distinctive starch of an unknown source

Starch, Zea cooked Maize starch that is altered by heat. The stellate hila widen in

these examples, and some are partly destroyed

Starch, Zea uncooked Maize starch that retains pristine characteristics of irregular spher-

Mass of cooked maize starch

oid shape, stellate hilum, and birefringency

Starch, Zea cooked, large clump of 200+ granules

Starch, tuber aggregate

Aggregate of faceted starch granules of various sizes 5–15 μm

that are most consistent with starch from tubers

Xylem section Columns of conductive tissue with identifiable tracheids and ves-

sel elements. These are not identifiable to taxon. Their abundance reflects how much plant stem and leaves were eaten

Xylem tracheid Helical, often mineralized, structures. These conduct nutrients in

plants

Xylem tracheid, double helical As above except that the tracheids are paired in double helices

Opuntia cuticleFragments of the waxy coating covering prickly pear padsYucca phytolithWedge-shaped CaC_2O_4 phytoliths consistent with the Agavaceae

family of which *Yucca* is best represented in the Antelope

Cave area

Unidentified plant tissue Plant residue with no distinctive features

Table C2. Definitions of terms applied to macroscopic remains

| Component | Definition | | | | | |
|-----------------------------------|---|--|--|--|--|--|
| <.5-mm category | Very finely ground material | | | | | |
| >.5-<1.0-mm category | Finely ground material | | | | | |
| >1.0-mm category | Ground material | | | | | |
| Aggregates of macroscopic remains | Masses of consolidated seeds, bones, and fibers that did not disaggregate | | | | | |
| | during coprolite processing. The percent composition of the aggregates was estimated | | | | | |
| Ant | An ant head, probably a cave contaminant | | | | | |
| Ash mixed with plant residue | Residue of food preparation, possibly from the use of parching trays, which mixes food with ash | | | | | |
| Black granular material | Apparently charcoal | | | | | |
| Bone | Bone from human coprolites was highly fragmented and eroded from preparation and digestion | | | | | |
| Cactus glochidia | Stellate arrangements of recurved spines from cacti | | | | | |
| Cactus thorn | Modified leaf from a cactus | | | | | |
| Charcoal | Small charcoal fragment from tree or shrub | | | | | |
| Cheno-am | Seeds of the goosefoot family or pigweed genus. These seeds are most likely from | | | | | |

the goosefoot genus Chenopodium

Supplement C from Reinhard et al., Ancient Diet and Modern Diabetes

Claws, rabbit Claws from jackrabbit, not a contaminant

Cottonwood? Unusual white plant fiber attached to a woody matrix
Dropseed Seed morphology consistent with *Sporobolus* caryopses

Feather Down feather
Feather calami Bases of feather quills
Fiber Undistinct masses of fiber

Fiber, <.5 mm in smallest dimension Finely ground fiber, which indicates that plant stems were ground and eaten

Fur tuft, Lepus Microscopic examination shows that these tufts are from jackrabbit

Spider beetle From the Ptinidae family of beetles, which are general scavengers of dry substrates.

Found commonly in coprolites and mummies. They burrow into coprolites and leave long, narrow holes that allow for contaminants to enter the coprolite matrix

Grass stem A whole stem fragment

Hair Nonhuman hair Maize Corn kernel testa

Prickly pear Two or more prickly pear anatomical parts adhered together, including glochidia,

phytoliths, epidermis, and fiber

Prickly pear epidermis

Thick, brittle epidermis with classic pattern of CaC₂O₄ opuntioid druses in each cell.

Color ranges from light tan to white

Prickly pear phytoliths CaC₂O₄ opuntioid druses exceeding .25 mm in diameter

Sunflower seed Actually the ground achenes of sunflower or related genus including the outer fruit

wall and seed

Tick Dermacentor andersoni exoskeleton

Twig fragments Woody stem fragments

Unidentifiable seed Seed that is so finely ground that no identifiable morphology is visible to make an

identification

Unidentified arthropod Curved, spiny, exoskeleton similar to an isopod

Unknown seed Highly fragmented seeds similar to four-winged saltbush (Atriplex species, but this is

not a positive identification). May also have an arboreal origin

Whole dropseed Caryopses similar but smaller than sand dropseed Sporobolus cryptandrus

Whole wolfberry seeds Seeds from wolfberry fruits, probably *Lycium pallidum**Yucca fiber Fiber with distinct groove consistent with *Yucca*

Table C3. Microscopic counts from human coprolites

| | Lab no. | | | | | | | | |
|---|---------|---|----------|----|----|---|-----|----|----|
| Material | 1 | 2 | 3 | 4 | 5 | 7 | 9 | 10 | 11 |
| Lycopodium | 5 | 9 | 2 | 23 | 7 | 1 | 1 | 4 | 34 |
| Animal hair | | | | | | | | | 2 |
| CaC ₂ O ₄ druse | | | | | 5* | | | | |
| CaC ₂ O ₄ opuntioid druse | 93 | | | | | | 1 | | |
| Cell containing starch | | | | | | | | | |
| Clump of mineralized plant structures in ash | | | | | | | | | |
| Enterobius egg fragment? | | | | | | | | | 1 |
| Glochidia, mineralized | | | | | | | | 55 | |
| Glochid fragment, mineralized | | | | | 6 | 4 | | | |
| Helianthus achene fibers | | | ∞ | 5 | | | 54 | | |
| Insect fragment | | | | | | | | | |
| Leaf epidermis fragment | | | | 25 | | | | | 1 |
| Phloem sieve-tube element | | 1 | | | | | | 6 | 4 |
| Phloem with sieve-tube element | | | | | | 7 | | | 2 |
| Phytolith, unknown | | 3 | | | | | | | |
| Plant epidermis, unknown | | | | | | 1 | | | |
| Plant fiber | | | | | | | | | 78 |
| Plant fiber bundle | | | | | | | | | 12 |
| Plant hair | | | | 28 | | | | | 15 |
| Plant hair, mineralized | | | | | | | | | |
| Poaceae epidermis | | 3 | | 16 | 16 | | | | |
| Poaceae leaf epidermis | | | | | | | | | 3 |
| Poaceae long-cell phytolith | | 6 | | | | | | 2 | |
| Raphide | | | | | | | | | 4 |
| Raphide bundle | | | | | | | | | 1 |
| Ring structure with interference cross | 84 | 6 | | 2 | | | | | 1 |
| Seed coat fragment | | | ∞ | | | | | | |
| Seed testa, light color | | | | | | | 128 | | |
| Starch, 15 μ m, round, no hilum | | | | | | | 1 | | |

| Starch, 15 μ m, round, monocolpate | | | | | | | 1 | |
|---|----|----|-----|----|-----|---|----|----|
| Starch, 15–20 μ m, round, with hilum | | | 297 | | | | | 6 |
| Starch, 11 μ m, round, with large hilum | 1 | | | | | | | |
| Starch, Zea cooked | | | | | 1 | | | |
| Starch, Zea uncooked | | 22 | | | | | | |
| Xylem section | | 1 | | 28 | | 5 | 61 | 6 |
| Xylem tracheid | 56 | 10 | 16 | 2 | | 1 | 89 | 62 |
| Xylem tracheid, double helical | | | | | | | 2 | |
| Opuntia cuticle | | | | | | | | 6 |
| Yucca phytolith | | | | | | | | 1 |
| Unidentified plant tissue | | | | | 201 | | | |

Note. Samples 6 and 8 are excluded because they are canid in origin. $\infty =$ masses observed, too numerous to count; * = square plate projections on druses.

Table C4. Microscopic counts from human coprolites

| | Lab no. | | | | | | | | | | |
|---|---------|-----|-----|-----|-----|-----|----|-----|-----|-----|-----|
| Material | 12 | 13 | 14 | 15 | 16 | 17 | 18 | 19 | 20 | 21 | 23 |
| Lycopodium | 2 | 6 | 3 | 19 | 4 | 2 | 13 | 2 | 32 | 32 | 6 |
| Animal hair | 10 | | | 12 | | | | 2 | | | |
| CaC ₂ O ₄ opuntioid druse | | | | 2 | | 1 | | 14 | | | |
| Cell containing starch | 1 | | | | | | | | | | |
| Clump of mineralized plant structures in ash | | | | 137 | | | 92 | | | | |
| Glochid, mineralized | 1 | | | | | | 7 | 2 | | | |
| Glochid fragment, mineralized | | | | | | | 88 | 48 | | 2 | |
| Helianthus achene fibers | | | 180 | | 175 | | | | 7 | | |
| Insect fragment | | | | 1 | | | | | | | |
| Leaf epidermis fragment | | 1 | | | | | | | | | |
| Phloem sieve-tube element | | | | 2 | | | 5 | 6 | | | |
| Phloem with sieve-tube element | | | | 24 | | | 6 | | | 5 | |
| Plant epidermis, unknown | | | | 4 | | | 2 | | 4 | | 1 |
| Plant fiber | | | | | | 1 | | | | 10 | |
| Plant fiber bundle | | | | | | | | | | | 6 |
| Plant hair | | | | | | 3 | | | | | |
| Plant hair, mineralized | | 23 | | 13 | | | | | | | |
| Poaceae epidermis | | 177 | 5 | | | 5 | | | 14 | 57 | |
| Poaceae leaf epidermis | | 1 | | | | | | 10 | | | |
| Poaceae long-cell phytolith | | 2 | | | | | | 2 | 1 | 1 | |
| Raphide bundle | | | | 2 | | | 1 | | | 6 | |
| Ring structure with interference cross | | 3 | | 4 | | | 4 | | | 105 | |
| Seed coat fragment | | 5 | 29 | | | | | | | | |
| Seed coat, cheno-am? | | | 12 | | | | | | 10 | | 212 |
| Seed testa, light color | | | | | | | | | 173 | 1 | 1 |
| Starch, indeterminate | | | | 2 | | | | | | | 3 |
| Starch, cheno-am | | | | 1 | | | | | | | |
| Starch without interference cross | | | | | | 215 | | | | | |
| Starch, 10-15 μm, round, no hilum | | 1 | | | | | | | | | |
| Starch, $10-15 \mu m$, faceted, with hilum | | | | | | | 3 | | | 1 | |
| Starch, 18–22 μ m, round, with hilum | | | | | | | | | | 1 | |
| Starch, 10–15 μ m, round, with large hilum | | | | | 2 | | | | | | |
| Starch, Zea cooked | 219 | | | 1 | | | 1 | | | 1 | |
| Starch, Zea uncooked | 1 | | | | | | 1 | | 1 | 4 | |
| Starch, Zea cooked, large clump (200) | 1 | | | | | | | | | | |
| Starch, tuber aggregate | 2 | | | | | | | | | | |
| Xylem section | | | | 1 | | 7 | 5 | 10 | 6 | 8 | |
| Xylem tracheid | 1 | | 2 | 9 | | | 12 | | 2 | 18 | |
| Xylem tracheid, double helical | | | | | | | | 192 | | | |
| Opuntia cuticle | | | | 1 | | | | | | 1 | |

Note. The two tuber starch aggregates from sample 12 were composed of 11 and 13 individual starch grains.

Table C5. Microfossil concentration values from human coprolites

| | Lab no. | | | | | | | |
|---|---------|--------|----------|---------|--------|--|--|--|
| Material | 1 | 2 | 3 | 4 | 5 | | | |
| Lycopodium spores per gram of coprolite | 13,661 | 14,044 | 6,341 | 11,737 | 6,137 | | | |
| Lycopodium | 5 | 9 | 2 | 23 | 7 | | | |
| CaC ₂ O ₄ druse | | | | | 4,384 | | | |
| CaC ₂ O ₄ opuntioid druse | 253,164 | | | | | | | |
| Glochid fragment, mineralized | | | | | 5,260 | | | |
| Helianthus achene fibers | | | ∞ | 2,552 | | | | |
| Leaf epidermis fragment | | | | 12,758 | | | | |
| Phloem sieve-tube element | | 1,560 | | | | | | |
| Phytolith, unknown | | 4,681 | | | | | | |
| Plant hair | | | | 14,289 | | | | |
| Poaceae epidermis | | 4,681 | | 8,165 | 14,027 | | | |
| Poaceae long-cell phytolith | | 9,363 | | | | | | |
| Ring structure with interference cross | 229,504 | 9,363 | | 1,021 | | | | |
| Seed coat fragment | | | ∞ | | | | | |
| Starch, 15–20 μ m, round, with hilum | | | | 151,560 | | | | |
| Starch, 11 μ m, round, with large hilum | 2,732 | | | | | | | |
| Starch, Zea uncooked | | 34,330 | | | | | | |
| Xylem section | | 1,560 | | | 24,548 | | | |
| Xylem tracheid | 153,003 | 15,604 | | 8,165 | 1,753 | | | |

Note. ∞ = incalculably high.

Table C6. Microfossil concentration values from human coprolites

| | Lab no. | | | | | | | | |
|---|--------------------|---------|---------|--------|---------|--|--|--|--|
| Material | 7 | 9 | 10 | 11 | 12 | | | | |
| Lycopodium spores per gram of coprolite | 7,049 | 7,143 | 6,499 | 14,648 | 13,587 | | | | |
| Lycopodium | 1 | 1 | 4 | 34 | 2 | | | | |
| Animal hair | | | | 860 | 67,935 | | | | |
| CaC ₂ O ₄ opuntioid druse | | 7,164 | | | | | | | |
| Cell containing starch | | | | | 6,794 | | | | |
| Glochidia, mineralized | | | 88,841 | | | | | | |
| Glochid fragment, mineralized | 28,163 | | | | 6,794 | | | | |
| Helianthus achene fibers | | 386,845 | | | | | | | |
| Leaf epidermis fragment | | | | 430 | | | | | |
| Phloem sieve-tube element | | | 9,692 | 101 | | | | | |
| Phloem with sieve-tube element | 49,285 | | | 860 | | | | | |
| Plant epidermis, unknown | 7,041 | | | | | | | | |
| Plant fiber | | | | 33,546 | | | | | |
| Plant fiber bundle | | | | 5,161 | | | | | |
| Plant hair | | | | 6,451 | | | | | |
| Poaceae leaf epidermis | | | | 1,290 | | | | | |
| Poaceae long-cell phytolith | | | 3,231 | | | | | | |
| Raphide | | | | 101 | | | | | |
| Raphide bundle | | | | 430 | | | | | |
| Ring structure with interference cross | | | | 430 | | | | | |
| Seed testa, light color | | 914,304 | | | | | | | |
| Starch, 15 µm, round, no hilum | | 7,164 | | | | | | | |
| Starch, 15 μ m, round, monocolpate | | | 1,615 | | | | | | |
| Starch, 15–20 μ m, round, with hilum | | | | 2,580 | 143,555 | | | | |
| Starch, Zea cooked | 7,041 | | | | 6,794 | | | | |
| Starch, Zea uncooked | | | | | 6,794 | | | | |
| Starch, Zea cooked, large clump (200) | | | | | 13,587 | | | | |
| Xylem section | | 35,819 | 98,533 | 2,580 | | | | | |
| Xylem tracheid | | 7,164 | 143,761 | 26,665 | 6,794 | | | | |
| Xylem tracheid, double helical | | | 3,231 | | | | | | |
| Opuntia cuticle | | | | 2,580 | | | | | |
| Yucca phytolith | | | | 430 | | | | | |
| Unidentified plant tissue | 1.42×10^6 | | | | | | | | |

Table C7. Microfossil concentration values

| | | Lab no. | | | | |
|---|---------|---------|--------|---------|---------|--|
| Material | 13 | 14 | 15 | 16 | 17 | |
| Lycopodium spores per gram of coprolite | 14,940 | 6,234 | 12,214 | 12,658 | 6,406 | |
| Lycopodium | 6 | 3 | 19 | 4 | 2 | |
| Animal hair | | | 7,714 | | | |
| CaC ₂ O ₄ opuntioid druse | | | 3,214 | | 3,203 | |
| Clump of mineralized plant structures in ash | | | 12,214 | | | |
| Helianthus achene fibers | | 374,040 | | 553,788 | | |
| Insect fragment | | | 643 | | | |
| Leaf epidermis fragment | 2,490 | | | | | |
| Phloem sieve-tube element | | | 1,286 | | | |
| Phloem with sieve-tube element | | | 15,428 | | | |
| Plant epidermis, unknown | | | 2,571 | | | |
| Plant fiber | | | | | 3,203 | |
| Plant hair | | | | | 9,609 | |
| Plant hair, mineralized | 57,270 | | 8,357 | | | |
| Poaceae epidermis | 440,730 | 10,390 | | | 16,015 | |
| Poaceae leaf epidermis | 2,490 | | | | | |
| Poaceae long-cell phytolith | 4,980 | | | | | |
| Raphide bundle | | | 1,286 | | | |
| Ring structure with interference cross | 7,470 | | 2,571 | | | |
| Seed coat fragment | 12,450 | 60,262 | | | | |
| Seed coat, cheno-am? | | 24,936 | | | | |
| Starch, indeterminate | | | 1,286 | | | |
| Starch, cheno-am | | | 643 | | | |
| Starch without interference cross | | | | | 688,645 | |
| Starch, 10-15 μm, round, no hilum | 2,490 | | | | | |
| Starch, 10–15 μ m, round, with large hilum | | | | 6,329 | | |
| Starch, Zea cooked | | | 643 | | | |
| Xylem section | | | 643 | | 22,421 | |
| Xylem tracheid | | 4,156 | 5,786 | | | |
| Opuntia cuticle | | | 643 | | | |

Note. The two tuber starch aggregates from sample 12 were composed of 11 and 13 individual starch grains.

Table C8. Microfossil concentration values

| | Lab no. | | | | |
|---|---------|---------|--------|---------|---------|
| Material | 18 | 19 | 20 | 21 | 23 |
| Lycopodium spores per gram of coprolite | 14,423 | 12,626 | 12,906 | 62,500 | 6,098 |
| Lycopodium | 13 | 2 | 32 | 32 | 6 |
| Animal hair | | 12,626 | | | |
| CaC ₂ O ₄ opuntioid druse | | 88,382 | | | |
| Clump of mineralized plant structures in ash | 102,070 | | | | |
| Glochid, mineralized | 7,766 | 12,626 | | | |
| Glochid fragment, mineralized | 97,633 | 303,024 | | 3,906 | |
| Helianthus achene fibers | | | 2,823 | | |
| Phloem sieve-tube element | 5,547 | 151,512 | | | |
| Phloem with sieve-tube element | 4,555 | | | 9,766 | |
| Plant epidermis, unknown | 1,518 | | 1,613 | | 1,016 |
| Plant fiber | | | | 19,531 | |
| Plant fiber bundle | | | | | 6,098 |
| Poaceae epidermis | | | 14 | 111,328 | |
| Poaceae leaf epidermis | | 63,130 | | | |
| Poaceae long-cell phytolith | | 12,626 | 403 | 1,953 | |
| Raphide bundle | 759 | | | 11,719 | |
| Ring structure with interference cross | 3,036 | | | 205,078 | |
| Seed coat, cheno-am? | | | 4,033 | | 215,463 |
| Seed testa, light color | | | 69,773 | 1,953 | 1,016 |
| Starch, indeterminate | | | | | 3,049 |
| Starch, 10–15 μ m, faceted, with hilum | 2,277 | | | 1,953 | |
| Starch, 18–22 μ m, round, with hilum | | | | 1,953 | |
| Starch, Zea cooked | 759 | | | 1,953 | |

| Starch, Zea uncooked | 759 | | 403 | 7,813 |
|--------------------------------|-------|--------------------|-------|--------|
| Xylem section | 3,796 | 12,626 | 2,420 | 15,625 |
| Xylem tracheid | 9,109 | | 807 | 35,156 |
| Xylem tracheid, double helical | | 1.21×10^6 | | |
| Opuntia cuticle | | | | 1,953 |

Table C9. Macroscopic remains (weight in grams)

| Component | 1 | 2 | 3 | 4 |
|---|-------|-------|-------|------|
| Bone, <.5 mm | | | | |
| Bone, >.5-<1.0 mm | | | | |
| Bone, >1.0 mm | | .34 | Trace | .67 |
| Charcoal | | .02 | | |
| Cottonwood? | | Trace | | |
| Feather calami | | | | .01 |
| Ground dropseed, <.5 mm | | | | |
| Ground dropseed, >.5-<1.0 mm | | | | |
| Ground dropseed, >1.0 mm | .3 | | | |
| Ground maize, <.5 mm | | | | |
| Ground maize, >.5-<1.0 mm | | .77 | | 1.1 |
| Ground maize, >1.0 mm | | .07 | | 1.27 |
| Ground sunflower seed, <.5 mm | | | | |
| Ground sunflower seed, >.5-<1.0 mm | | | 7.51 | |
| Ground sunflower seed, >1.0 mm | | .01 | 1.24 | |
| Ground unidentifiable seed, <.5 mm | | | | |
| Ground unidentifiable seed, >.5-<1.0 mm | | | | |
| Ground unidentifiable seed, >1.0 mm | | | | |
| Ground unknown seed, <.5 mm | | | | |
| Ground unknown seed, >.5-<1.0 mm | | | | |
| Ground unknown seed, >1.0 mm | | | | |
| Prickly pear, >.5-<1.0 mm | | | | |
| Prickly pear, >1.0 mm | .94 | | | |
| Prickly pear phytoliths, <.5 mm | .4 | | | |
| Spider beetle | | Trace | | |
| Tick | | Trace | | |
| Twig fragments | | | Trace | |
| Whole dropseed | Trace | | | |
| Whole wolfberry seeds | | | | .5 |
| Yucca fiber | | Trace | | |

Table C10. Macroscopic remains (weight in grams)

| | Lab no. | | | | |
|-----------------------------------|---------|------------------|-----|-------|--|
| Component | 5 | 7 | 9 | 10 | |
| Aggregates of macroscopic remains | 4.06° | .81 ^b | | | |
| Bone, <.5 mm | | | | | |
| Bone, >.5-<1.0 mm | .19 | | .04 | | |
| Bone, >1.0 mm | 1.71 | .1 | .76 | | |
| Cactus glochidia | Trace | | | Trace | |
| Charcoal | Trace | | | | |
| Cottonwood? | | | | | |
| Fiber, >.5-<1.0 mm | | | | .85 | |
| Fiber, <.5 mm | | | | 1.82 | |
| Fiber, >1.0 mm | | | | .28 | |
| Fruit skin or corn testa | | | | Trace | |
| Ground dropseed, <.5 mm | 6.43 | | | | |
| Ground dropseed, >.5-<1.0 mm | 1.67 | | | | |
| Ground dropseed, >1.0 mm | | | | | |
| Ground maize, <.5 mm | | .56 | | | |
| Ground maize, >.5-<1.0 mm | | .42 | | | |
| Ground maize, >1.0 mm | | .28 | | | |

Supplement C from Reinhard et al., Ancient Diet and Modern Diabetes

Ground sunflower seed, <.5 mm Ground sunflower seed, >.5-<1.0 mm Ground sunflower seed, >1.0 mm .41 Ground unidentifiable seed, <.5 mm Ground unidentifiable seed, >.5-<1.0 mm .38 Ground unidentifiable seed, >1.0 mm 1.83° Ground unknown seed, <.5 mm Ground unknown seed, >.5-<1.0 mm Ground unknown seed, >1.0 mm Prickly pear, >.5-<1.0 mm Prickly pear, >1.0 mm Prickly pear epidermis .1 Prickly pear phytoliths, <.5 mm

Table C11. Macroscopic remains (weight in grams)

| | Lab no. | | | | |
|---|---------|-------|------------|-------|--|
| Component | 11 | 12 | 13 | 14 | |
| Aggregates of macroscopic remains | | | | | |
| Bone, <.5 mm | .05 | | | | |
| Bone, >.5-<1.0 mm | | | | Trace | |
| Bone, >1.0 mm | | .13 | | | |
| Cactus glochidia | | | | | |
| Crushed unknown seed, <.5 mm | .36 | | | | |
| Crushed unknown seed, >.5-<1.0 mm | .42 | | | | |
| Crushed unknown seed, >1.0 mm | .76 | | .12 | | |
| Fiber, >.5-<1.0 mm | | | | | |
| Fiber, <.5 mm | | | | | |
| Fiber, >1.0 mm | | | | | |
| Fruit skin or corn testa | | | | | |
| Grass stem | | | | Trace | |
| Ground cheno-am, <.5 mm | | | | | |
| Ground cheno-am, >.5-<1.0 mm | | | | Trace | |
| Ground cheno-am, >1.0 mm | | | | | |
| Ground dropseed, <.5 mm | | | .18a | | |
| Ground dropseed, >.5-<1.0 mm | | | 1.04^{a} | | |
| Ground dropseed, >1.0 mm | | | .21a,b | | |
| Ground maize, <.5 mm | | .06 | | | |
| Ground maize, >.5-<1.0 mm | | .07 | | | |
| Ground maize, >1.0 mm | | .67 | | | |
| Ground sunflower seed, <.5 mm | | | | .04 | |
| Ground sunflower seed, >.5-<1.0 mm | | | | .93 | |
| Ground sunflower seed, >1.0 mm | | | | 3.59 | |
| Ground unidentifiable seed, <.5 mm | | | | | |
| Ground unidentifiable seed, >.5-<1.0 mm | | | | | |
| Ground unidentifiable seed, >1.0 mm | | | | | |
| Prickly pear, >.5-<1.0 mm | | | | | |
| Prickly pear, >1.0 mm | | | | | |
| Prickly pear epidermis | | | | | |
| Prickly pear phytoliths, <.5 mm | | | | | |
| Spider beetle | | Trace | | Trace | |
| Tick | | | | | |
| Whole dropseed | | | | | |
| Wolfberry seeds | | | | | |
| Yucca fiber | | | | | |

^aComposed of ground seed, bracts, and stem.

^aMostly dropseed.

^bMostly maize.

^cUnidentifiable seed.

^bAggregates of ground seed, bracts, and stem.

Table C12. Macroscopic remains (weight in grams)

| | Lab no. | | | |
|---|---------|------------------|-------------------|-------|
| Component | 15 | 16 | 17 | 18 |
| Aggregates of macroscopic remains | | | | |
| Ash mixed with plant residue, <.5 mm | | | | .14 |
| Ash mixed with plant residue, >.5-<1.0 mm | | | | .17 |
| Ash mixed with plant residue, >1.0 mm | | | | .35 |
| Black granular material, <.5 mm | .14 | | | |
| Black granular material, >.5—<1.0 mm | .31 | | | |
| Black granular material, >1.0 mm | .9 | | | |
| Bone, <.5 mm | | | | |
| Bone, >.5-<1.0 mm | | .02 | 1.46 | |
| Bone, >1.0 mm | .24ª | | 2.73 | .17 |
| Cactus glochidia | | | | |
| Cactus thorn | | | Trace | |
| Charcoal | | | | |
| Claws, rabbit | | | .06 | |
| Cottonwood? | | | | |
| Feather | Trace | | Trace | |
| Fiber, <.5 mm in smallest dimension | .11 | | | .05 |
| Fur tuft, Lepus | Trace | | | |
| Grass stem | Trace | | | |
| Ground cheno-am, <.5 mm | | | | |
| Ground cheno-am, >.5-<1.0 mm | | | | |
| Ground cheno-am, >1.0 mm | | | | |
| Ground dropseed, <.5 mm | | | 5.9^{b} | |
| Ground dropseed, >.5-<1.0 mm | | | 1.24 ^b | |
| Ground dropseed, >1.0 mm | | | $3.7^{b,c}$ | |
| Ground maize, <.5 mm | .03 | | | .05 |
| Ground maize, >.5-<1.0 mm | .08 | | | .02 |
| Ground maize, >1.0 mm | 1.01 | | | .17 |
| Ground sunflower seed, <.5 mm | | .84 | | |
| Ground sunflower seed, >.5-<1.0 mm | | .39 | | |
| Ground sunflower seed, >1.0 mm | | .37 ^d | | |
| Ground unknown seed, <.5 mm | | | | |
| Ground unknown seed, >.5-<1.0 mm | | | | |
| Ground unknown seed, >1.0 mm | | | | |
| Hair | | | .05 | |
| Spider beetle | Trace | | Trace | Trace |
| Termites $(n = 3)$ | | | | Trace |
| Tick | Trace | | | |
| Unidentifiable | | | 1.16 | |

^aApparently juvenile or embryonic. ^bMixed seed, bracts, and stem. ^cWeight estimated from aggregates.

Table C13. Macroscopic remains (weight in grams)

| Component | Lab no. | | | | |
|-------------------------------------|---------|--------------------|----|------|--|
| | 19 | 20 | 21 | 23 | |
| Aggregates of macroscopic remains | | 1.12ª | | | |
| Ant | | | | | |
| Bone, <.5 mm | | | | | |
| Bone, >.5-<1.0 mm | | | | | |
| Bone, >1.0 mm | .11 | | | .13 | |
| Feather | | | | | |
| Fiber | .1 | | | | |
| Fiber, <.5 mm in smallest dimension | | .03 | | | |
| Fur tuft, Lepus | .19 | | | | |
| Grass stem | | Trace ^b | | | |
| Ground cheno-am, <.5 mm | | .65 | | 3.82 | |

dIn the form of aggregates.

| Ground cheno-am, >.5-<1.0 mm | | .15 | | .44 |
|--|------------------|-------|-----------|------|
| Ground cheno-am, >1.0 mm | | | | 1.03 |
| Ground dropseed, <.5 mm | | 1.43° | | |
| Ground dropseed, >.5-<1.0 mm | | .15 | $.01^{c}$ | |
| Ground dropseed, >1.0 mm | | | | |
| Ground maize, <.5 mm | | | | |
| Ground maize, >.5-<1.0 mm | | | | |
| Ground maize, >1.0 mm | | | | |
| Ground sunflower seed, <.5 mm | | | | |
| Ground sunflower seed, >.5-<1.0 mm | | | | |
| Ground sunflower seed, >1.0 mm | | | | |
| Ground unidentifiable seed, <.5 mm | | | | |
| Ground unidentifiable seed, >.5-<1.0 mm | | | | |
| Ground unidentifiable seed, >1.0 mm | | | | |
| Ground unknown seed, <.5 mm | | | | |
| Ground unknown seed, >.5-<1.0 mm | | | | |
| Ground unknown seed, >1.0 mm | | | | |
| Hair | | | | |
| Prickly pear, >.5-<1.0 mm | .27 ^d | | | |
| Prickly pear, >1.0 mm | .17 | | | |
| Prickly pear phytoliths with fiber, <.5 mm | .68 | | $.05^{e}$ | |
| Spider beetle | | | | |
| Tick | | | | |
| Unidentifiable | | .54 | | .07 |
| Whole dropseed | Trace | | | |

^aAggregates of crushed seed and fiber. ^bCut grass stem. ^cIncludes seed, bracts, and fibers. ^aPhytoliths and fiber.

Table C14. Pollen counts from coprolites

| | Lab no. | | | | | |
|--------------------------------|---------|-----|-----|-----|-----|--|
| Material | 1 | 2 | 3 | 4 | 5 | |
| Lycopodium spores | 134 | 876 | 123 | 84 | 35 | |
| Ambrosia type | 6 | 8 | 2 | 3 | | |
| Artemisia | 10 | 6 | 32 | 5 | 10 | |
| Celtis | | 5 | | | | |
| Cheno-am | 22 | 10 | 32 | 10 | 6 | |
| Ephedra sp. | | 1 | 1 | 2 | 1 | |
| Ephedra viridis | | | | | 1 | |
| Ephedra torreyana | 3 | | | | | |
| Eriogonum | | | | 1 | | |
| Fabaceae | 2 | | | | | |
| Helianthus type | | 4 | | | | |
| High-spine Asteraceae | | | 1 | 1 | | |
| Juniperus | 6 | 3 | 1 | | | |
| Low-spine Asteraceae | 7 | 7 | 18 | 5 | 1 | |
| Maize, torn | | 5 | | 1 | | |
| Maize, whole | 1 | 3 | | 1 | | |
| Onagraceae | | 1 | | | | |
| Pinus | 3 | 6 | 1 | 1 | 1 | |
| Poaceae | 140 | 10 | 123 | 167 | 173 | |
| Polygonum | | | | 1 | | |
| Populus type | 2 | 13 | | | | |
| Quercus | | 1 | | | | |
| Rhus | | 1 | | | 1 | |
| Salix | 1 | | 2 | | 1 | |
| Solanaceae type | 2 | 1 | | 1 | | |
| Unidentified | | 4 | 3 | | 3 | |
| Unknown | | 2 | | | 1 | |
| Unknown striate stephanoporate | 1 | | | | | |
| Yucca | | 1 | | | | |

eIncludes epidermis.

Table C15. Pollen counts from coprolites

| | Lab no. | | | | |
|-----------------------|---------|-----|-----|-----|-----|
| Material | 7 | 9 | 10 | 11 | 12 |
| Lycopodium spores | 159 | 27 | 106 | 100 | 100 |
| Ambrosia type | 1 | | 7 | 2 | |
| Artemisia | 2 | 182 | 23 | 1 | |
| Cheno-am | 4 | | 8 | | |
| Ephedra torreyana | | | | | 1 |
| Ephedra nevadensis | | | 1 | | |
| Fabaceae | | | 1 | | |
| Helianthus type | | | 16 | | |
| High-spine Asteraceae | | | 3 | | |
| Juniperus | | | 1 | | |
| Low-spine Asteraceae | | 1 | 2 | | |
| Maize, torn | | 1 | 1 | | |
| Maize, whole | 1 | | | | |
| Pinus | 5 | | 3 | | 1 |
| Poaceae | | 31 | 5 | 1 | 1 |
| Populus type | | | 1 | | |
| Rosaceae | 1 | | | | |
| Salix | | | 2 | | |
| Sarcobatus | | 1 | | | |
| Sphaeralcea | | | 1 | | |
| Unidentified | 1 | 9 | | | |

Table C16. Pollen counts from coprolites

| | | Lab no. | | | | |
|-----------------------|-----|---------|-----|-----|-----|--|
| Material | 13 | 14 | 15 | 16 | 17 | |
| Lycopodium spores | 17 | 102 | 150 | 100 | 35 | |
| Ambrosia type | 1 | 8 | | | | |
| Artemisia | 3 | 40 | 1 | | 7 | |
| Cheno-am | 1 | 57 | | | 2 | |
| Ephedra sp. | | | | 1 | | |
| Ephedra torreyana | | 1 | | | | |
| Ephedra nevadensis | | 1 | | | | |
| High-spine Asteraceae | | 3 | | | 1 | |
| Low-spine Asteraceae | | 10 | | | 1 | |
| Maize, whole | | | 1 | | | |
| Pinus | | 1 | 3 | | | |
| Poaceae | 306 | 80 | 1 | | 196 | |
| Quercus | | | 1 | | 1 | |
| Unidentified | | 3 | | | 2 | |
| Unknown | | | 1 | | | |

Table C17. Pollen counts from coprolites

| | Lab no. | | | | | |
|-----------------------|---------|----|-----|----|-----|--|
| Material | 18 | 19 | 20 | 21 | 23 | |
| Lycopodium spores | 259 | 25 | 140 | 93 | 18 | |
| Ambrosia type | | 2 | 1 | | 3 | |
| Artemisia | 1 | 6 | 1 | 2 | 1 | |
| Celtis | | | | | 1 | |
| Cheno-am | 3 | 1 | 3 | 2 | 206 | |
| Ephedra viridis | | | | 1 | | |
| Fabaceae | | 1 | | | | |
| Helianthus type | | 5 | | 1 | | |
| High-spine Asteraceae | | 2 | | | | |
| Juniperus | | | | | 1 | |
| Low-spine Asteraceae | 1 | | 1 | | | |

| Maize, torn | 172 | 40 | | | |
|--------------------------------|-----|-----|-----|-----|----|
| Maize, whole | 26 | 59 | | | |
| Pinus | | 2 | | | |
| Poaceae | 1 | 113 | 215 | 105 | 13 |
| Quercus | 1 | | 1 | | |
| Solanaceae type | 3 | | | | |
| Unknown striate stephanoporate | | 1 | | | |

Table C18. Pollen concentration values expressed in numbers of pollen grains per category per gram of coprolite

| | | | Lab no. | | |
|--------------------------------|--------|--------|---------|--------|--------|
| Material | 1 | 2 | 3 | 4 | 5 |
| Lycopodium spores per gram | 13,661 | 14,044 | 6,341 | 11,737 | 6,137 |
| Lycopodium counted | 134 | 876 | 123 | 84 | 35 |
| Ambrosia type | 612 | 128 | 362 | 419 | |
| Artemisia | 1,019 | 96 | 5,798 | 699 | 1,754 |
| Celtis | | 80 | | | |
| Cheno-am | 2,243 | 160 | 5,798 | 1,397 | 1,052 |
| Ephedra sp. | | 16 | 181 | 279 | 175 |
| Ephedra viridis | | | | | 175 |
| Ephedra torreyana | 306 | | | | |
| Eriogonum | | | | 140 | |
| Fabaceae | 204 | | | | |
| Helianthus type | | 64 | | | |
| High-spine Asteraceae | | | 181 | 140 | |
| Juniperus | 612 | 48 | 181 | | |
| Low-spine Asteraceae | 714 | 112 | 3,261 | 699 | 175 |
| Maize, torn | | 80 | | 140 | |
| Maize, whole | 102 | 48 | | 140 | |
| Onagraceae | | 16 | | | |
| Pinus | 306 | 96 | 181 | 140 | 175 |
| Poaceae | 14,273 | 160 | 22,284 | 23,334 | 30,337 |
| Polygonum | | | | 140 | |
| Populus type | 204 | 208 | | | |
| Quercus | | 16 | | | |
| Rhus | | 16 | | | 175 |
| Salix | 102 | | 362 | | 175 |
| Solanaceae type | 204 | 16 | | 140 | |
| Unidentifiable | | 654 | 544 | | 526 |
| Unknown | | 32 | | | 175 |
| Unknown striate stephanoporate | 102 | | | | |
| Yucca | | 16 | | | |

Table C19. Pollen concentration values expressed in numbers of pollen grains per category per gram of coprolite

| | Lab no. | | | | | |
|----------------------------|---------|--------|-------|--------|--------|--|
| Material | 7 | 9 | 10 | 11 | 12 | |
| Lycopodium spores per gram | 7,049 | 7,143 | 6,499 | 14,648 | 13,587 | |
| Lycopodium counted | 159 | 27 | 106 | 100 | 100 | |
| Ambrosia type | 44 | | 429 | 293 | | |
| Artemisia | 89 | 48,149 | 1,410 | 147 | | |
| Cheno-am | 177 | | 490 | | | |
| Ephedra torreyana | | | | | 136 | |
| Ephedra nevadensis | | | 61 | | | |
| Fabaceae | | | 61 | | | |
| Helianthus type | | | 981 | | | |
| High-spine Asteraceae | | | 182 | | | |
| Juniperus | | | 61 | | | |
| Low-spine Asteraceae | | 265 | 123 | | | |
| Maize, torn | | 265 | 61 | | | |

| Maize, whole | 177 | | | | |
|----------------|-----|-------|-----|-----|-----|
| Pinus | 222 | | 182 | | 136 |
| Poaceae | | 8,201 | 307 | 147 | 136 |
| Populus type | | | 61 | | |
| Rosaceae | 177 | | | | |
| Sarcobatus | | 265 | | | |
| Salix | | | 182 | | |
| Sphaeralcea | | | 61 | | |
| Unidentifiable | 177 | 2,381 | | | |

Table C20. Pollen concentration values expressed in numbers of pollen grains per category per gram of coprolite

| | | | Lab no. | | |
|----------------------------|---------|-------|---------|--------|--------|
| Material | 13 | 14 | 15 | 16 | 17 |
| Lycopodium spores per gram | 14,940 | 6,234 | 12,214 | 12,658 | 6,406 |
| Lycopodium counted | 17 | 102 | 150 | | 35 |
| Ambrosia type | 879 | 489 | | | |
| Artemisia | 2,636 | 2,445 | 81 | | 1,281 |
| Cheno-am | 879 | 3,484 | | | 366 |
| Ephedra viridis | | | | 127 | |
| Ephedra torreyana | | 61 | | | |
| Ephedra nevadensis | | 61 | | | |
| High-spine Asteraceae | | 183 | | | 183 |
| Low-spine Asteraceae | | 611 | | | 183 |
| Maize, torn | | | 81 | | |
| Pinus | | 61 | 244 | | |
| Poaceae | 268,920 | 4,889 | 81 | | 35,874 |
| Quercus | | | 81 | | 183 |
| Unidentified | | 183 | | | 366 |
| Unknown | | | 81 | | |

Table C21. Pollen concentration values expressed in numbers of pollen grains per category per gram of coprolite

| | Lab no. | | | | |
|--------------------------------|---------|--------|--------|--------|--------|
| Material | 18 | 19 | 20 | 21 | 23 |
| Lycopodium spores per gram | 14,423 | 12,626 | 12,906 | 62,500 | 6,098 |
| Lycopodium spores | 259 | 25 | 140 | 93 | 18 |
| Ambrosia type | | 1,010 | 91 | | 1,016 |
| Artemisia | 56 | 3,030 | 91 | 1,344 | 339 |
| Celtis | | | | | 339 |
| Cheno-am | 167 | 505 | 277 | 1,344 | 69,788 |
| Ephedra viridis | | | | 672 | |
| Fabaceae | | 505 | | | |
| Helianthus type | | 2,525 | | 672 | |
| High-spine Asteraceae | | 1,010 | | | |
| Juniperus | | | | | 339 |
| Low-spine Asteraceae | 56 | | 91 | | |
| Maize, torn | 9,578 | 20,202 | | | |
| Maize, whole | 1,448 | 29,797 | | | |
| Pinus | | 1,010 | | | |
| Poaceae | 56 | 57,070 | 19,820 | 70,566 | 4,404 |
| Quercus | 56 | | 91 | | |
| Solanaceae type | 167 | | | | |
| Unknown striate stephanoporate | | 505 | | | |

Table C22. Pollen aggregates counts

| | | | Lab no. | | |
|----------------------|--|---------------|-------------|-----------------------|----------------------|
| Material | 1 | 2 | 3 | 4 | 5 |
| Artemisia | | | | | |
| Cheno-am | | | | | |
| Low-spine Asteraceae | | | | (3) | |
| Maize | (2) | (2) | | | |
| Poaceae | 12(2), 6(3), (4), (5), (7), 2(8), (9), (12) | (14) | 7(2), 4(3), | 4(2), 2(3), (5), (10) | 7(2), 2(3), (6), (9) |
| Populus type | | (2), (3), (5) | | | |

Note. Each number in parentheses indicates one clump of the specified number of pollen grains; e.g., (6) = one aggregated clump of six pollen grains. A number in parentheses proceeded by a number without parentheses indicates several aggregated clumps of pollen of the specified number; e.g., 3(6) = three aggregates of six pollen grains each.

Table C23. Pollen aggregates counts

| | | Lab no. | | | | | | |
|--|----------------------|---------------------------------|------------------|-----|-----------------|--|--|--|
| Material | 9 | 13 | 14 | 15 | 17 | | | |
| Artemisia Cheno-am Low-spine Asteraceae Maize | 8(2), 2(3), (5), (7) | | (3) (4) | | (3) | | | |
| Poaceae | | 8(2), 5(3), (4), (5), (6), (12) | 17(2), 6(3), (4) | (2) | 11(2), (3), (5) | | | |

Note. Each number in parentheses indicates one clump of the specified number of pollen grains; e.g., (6) = one aggregated clump of six pollen grains. A number in parentheses proceeded by a number without parentheses indicates several aggregated clumps of pollen of the specified number; e.g., 3(6) = three aggregates of six pollen grains each. No aggregates were observed for coprolites 7, 10, 11, 12, and 16.

Table C24. Pollen aggregates counts

| | Lab no. | | | | | | |
|---|----------------|-----------------------------------|-----------------------------------|------------------------|----------------|--|--|
| Material | 18 | 19 | 20 | 21 | 23 | | |
| Artemisia Cheno-am Low-spine Asteraceae | | | | | 4(2), (3), (7) | | |
| Maize, torn Poaceae | (2), 2(3), (4) | 2(2), (6) 2(2), 2(3), (4), (6) | 7(2), 3(3), 2(4), (6), (10), (12) | 17(2), 3(3), 3(5), (6) | (3), (4), (9) | | |

Note. Each number in parentheses indicates one clump of the specified number of pollen grains; e.g., (6) = one aggregated clump of six pollen grains. A number in parentheses proceeded by a number without parentheses indicates several aggregated clumps of pollen of the specified number; e.g., 3(6) = three aggregates of six pollen grains each.

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