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Zoonotic and Human Parasites of Inhabitants of Cueva de los Muertos Chiquitos, Rio Zape Valley, Durango, Mexico

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ZOOONOTIC AND HUMAN PARASITES OF INHABITANTS OF CUEVA DE LOS MUERTOS CHIQUITOS, RIO ZAPE VALLEY, DURANGO, MEXICO

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ABSTRACT: We present the first reconstruction of the parasitoses among the people of the Loma San Gabriel culture, as represented by 36 coprolites excavated from the Cueva de los Muertos Chiquitos in Durango, Mexico. The coprolites date to approximately 1,400-yr-ago. Species identified based on eggs recovered include the trematode Echinostoma sp., the tapeworms Hymenolepis sp. and Dipylidium caninum, and the nematodes Ancylostoma duodenale, Enterobius vermicularis, and Trichuris trichiura. After rehydration and screening, 2 methods were used to recover eggs from these samples including spontaneous sedimentation and flotation. Samples were analyzed by 3 different laboratories for independent verification and comparison of methods. Spontaneous sedimentation resulted in the discovery of hymenolepidid eggs that were not found with flotation. Sedimentation was a more-sensitive indicator of prevalence as well. The modified method of flotation permitted estimation of egg concentration and resulted in the detection of a few specimens not found by sedimentation. The results of both methods showed that 19 (of 36) coprolites contained helminth eggs. Our results detected the presence of pathogenic helminths including hookworms and whipworms. The cestodes found do not cause severe pathology in humans. The early dates of hookworm and whipworm, relative to other findings in the southwest United States, indicate that these parasites arrived relatively late in prehistory in Arizona and New Mexico, probably moving into the area with travelers from Mesoamerica.

The first dietary reconstruction of Mesoamerican societies was based on the analysis of coprolites from the valley of Tehuacán (Callen, 1965, 1967; McNeish, 1967). This reconstruction addressed over 5,000 yr of Mexican indigenous cultural development, from egalitarian hunter-gatherers to complex civilizations. However, the study of parasites was not included in that research. After Callen’s death, the Tehuacán collection was moved to the Department of Anthropology at Texas A&M (Bryant, 1974, 1975). One of us (K. J. Reinhard, pers. obs.) examined the collection, but no unprocessed coprolites were preserved by Callen to be tested for parasites. Therefore, an opportunity to examine the effect of the rise of civilization on parasitism was lost. Such a study was completed on the Colorado Plateau on the margins of ancient Mesoamerican civilization (Reinhard, 1987). However, the geographic connection of parasitism on the Colorado Plateau with Mesoamerica could not be evaluated because coprolites were not studied from Mexico. The analysis of coprolites from the site of Cueva de los Muertos Chiquitos in central Durango, Mexico provided us with the first glimpse into the parasitological state of Mesoamerica. The site is also important because it is on the frontier of the American Southwest and Mesoamerica. Therefore, we can gain insight into the transfer of parasites between Mesoamerica and the Southwest in ancient times. The present study thus helps to fill the gap in our knowledge of Mesoamerican parasitology and parasite biogeography.

The archeological site of Cueva de los Muertos Chiquitos is located in the northern Durango region of el Zape in the transition zone between the greater Southwest and the northern-most edge of Mesoamerica (Kelley, 1956, 1971; Brooks and Brooks, 1980). Located about 18 km SE of Guanaceví, Durango, with a sub-humid climate, this rocky valley has a series of caves that have been shown to provide excellent preservation of coprolites. Aboriginal inhabitants of the region belonged to the Mesoamerican culture known as Loma San Gabriel and occupied the region of southern Chihuahua and northern Durango 1,200- to 1,400-yr-ago (Brooks et al., 1962; Foster, 1986). They have been characterized as sedentary villagers with the ability and knowledge to practice agriculture and also to pursue hunting and gathering (Kelley, 1956, 1971; Brooks et al., 1962; Foster, 1985).

Previous research has shown that, once human remains were deposited in the site, burials and middens were sealed with an adobe layer that prevented disturbance (Brooks et al., 1962; Brooks and Brooks, 1980). Skeletons discovered in this area include 7 children ranging in age from a few months to 5-yr-old. The burials included food offerings consisting of corn, beans, agave, piñón, and cucurbits as well as necklaces of marine shells, belts, and wooden plaques with turquoise inlays (Brooks et al., 1962). The objects from this part of the cave were dated to about 600 A.D. Additional burials of an adult and 1 infant were placed over the older tomb at some later date (Brooks et al., 1962; Brooks and Brooks, 1978, 1980). Evidence of skeletal lesions in the human remains is apparent (D. Martin, pers. comm.).

Several techniques were employed to recover the remains of parasites from a variety of archaeological materials, including mummy tissues (Allison et al., 1974; Guhl et al., 1997), latrine sediments (Warnock and Reinhard, 1992; Bouchet et al., 2003), and coprolites (Szidat, 1944; Pizzi and Schenone, 1954; Taylor, 1955; Samuels, 1965; Reinhard et al., 1988; Ferreira et al., 2000). The trisodium-phosphate rehydration technique has become the most widely applied method for the study of coprolites and mummy intestinal contents (Reinhard, 1992; Reinhard and Bryant, 1992, 2008). This method results in the recovery of dietary and parasitological material (Callen and Cameron, 1960) and has been widely applied in North America (Samuels, 1965; Reinhard et al., 1988; Reinhard, 1992; Ferreira et al., 2000; Reinhard and Bryant, 2008). Thus, there is a body of data for desert regions that allows for comparison of prehistoric infection across time, cultural adaption, and geography for the desert west
of North America (Reinhard 1992; Reinhard and Bryant, 2008). The analysis of the external appearance of stools also offers the testimony of some symptoms suffered by individuals under study, i.e., diarrhea. The analysis of their contents provides direct evidence of both feeding habits and parasitoses (Reinhard, 1992; Araújo and Ferreira, 2000; Araújo et al., 2003). Used in combination, the results of dietary and parasitological examinations may assist in the reconstruction of the interactions of ancient humans with their environment, settlement conditions, diet, hygiene, health (Araújo et al., 2003; Sianto et al., 2005), and trade routes (Ferreira et al., 1979; Araújo et al., 2003; Costa et al., 2009).

The present study offers insights on the parasites infecting people in El Zape valley in a site that predates other examined cultures in the Greater Southwest. Our analysis of coprolites followed 2 methods in 3 different laboratories. Therefore, our goals include (1) documenting the parasites present at the Cueva de los Muertos Chiquitos, and (2) providing definitive data on the relative efficacy of different methods of analysis of coprolites.

MATERIALS AND METHODS

Coprolites from different proveniences were collected in Cueva de los Muertos Chiquitos and placed into individual plastic and paper bags in the field by Sheilagh and Richard Brooks. The samples were stored in stable dry and temperate conditions; these were labeled by grid square and stratigraphic level of the excavation unit. We sampled one coprolite from each grid and excavation unit in order to maximize the chances of sampling a variety of the cave’s prehistoric inhabitants. In a positive-pressure, filtered-air laboratory (University of Nebraska, Lincoln, Nebraska), samples of 36 coprolites were placed in new plastic bags by gloved laboratory workers. We believe, based on diversification of samples, that we sampled 36 individuals. Each coprolite was described, photographed, and weighed. From each sample, 2 fragments of 2 cm each were removed with the aid of a clean spatula. One set of samples was processed at the University of Nebraska for parasite remains. A second set was submitted to the Molecular Anthropology Laboratories at the University of Oklahoma (Norman, Oklahoma) for DNA analysis (Tito et al., 2008). Molecular analysis of parasite DNA is on-going.

The samples were weighed and rehydrated in 0.5% trisodium phosphate for 48 hr. A tablet of Lycopodium (batch 212761, each containing 12,500 ± 400 Lycopodium spores, University of Lund, Sweden) was added to the rehydrated coprolites. This was done to aid in the quantification of parasite eggs from the coprolites (Warnock and Reinhard, 1992; Sianto et al., 2005). The rehydrated coprolites were then screened through a 250-μm mesh with distilled water. The water then was filtered and microscopic residues that passed through the screen were collected in a beaker and concentrated by centrifugation. The concentrated sediments were divided into 3 subsamples to be analyzed at 3 separate laboratories, i.e., the Archaeoeparasiology lab in the School of Natural Resources at the University of Nebraska–Lincoln, the Laboratório de Paleoparasitologia, Departamento de Biología, Universidad Nacional de Mar del Plata (Mar del Plata, Argentina), and the Laboratório de Paleoparasitologia da Escola Nacional de Saúde Pública (Rio de Janeiro, Brazil).

At the University of Nebraska, the sediments were centrifuged and the pellet (usually 1 ml) was resuspended in Sheather’s solution in 15-ml glass centrifuge tubes. The solution was added until it filled the tube. A coverslip was placed over each tube in direct contact with the solution. The tubes were spun at 1,000 g (Pritchard and Kruse, 1982; Gardner and Duszynski, 1990). The resulting coverslip was mounted on a slide and scanned with a compound microscope.

The concentration of eggs per g (epg) was estimated by dividing the number of eggs counted by the mass sample under the coverslip. Assuming uniform distribution of both spores of Lycopodium and parasite eggs, the mass sample under the coverslip was estimated by multiplying the number of spores of Lycopodium by the total weight of fecal sample used. The product was divided by the total number of Lycopodium spores (calculated by multiplying number of tablets used by the average number of spores).

Samples studied in the laboratories of paleoparasitology at Mar del Plata and FIOCRUZ were homogenized and allowed to spontaneously sediment (Lutz, 1919). Ten slides of each sample were prepared by mixing a drop (approximately 0.5 ml) collected from the bottom with a drop of glycerin. The slides prepared were scanned with a compound microscope. Eggs detected were measured and identified using specialized literature (Yamaguti, 1975; Lamotho-Argumedo and Garcia-Prieto, 1988; Anderson, 2000; Ash and Orihel, 2007).

RESULTS

From the 36 coprolites examined, 2 showed signs of diarrhea (coprolites 6 and 30), as indicated by a lamellar structure that resulted from the desiccation of watery stools. Five more displayed a flattened shape resulting from the desiccation of loose stools (coprolites 9, 12, 22, 26, 31).

The taxa found included digenetic trematodes and cestodes. Digenean eggs consistent in size and form with Echinostoma sp. were found (Fig. 1). Tapeworms were represented by Dipylidium caninum and a hymenolepidid species. The identification of D. caninum was based on the morphology of eggs found in packets. Only the embryophores were preserved from the hymenolepidid eggs, so diagnosis was possible only to Hymenolepididae. Digenean eggs were determined to be an unidentified species of the Echinostoma, based on the presence of an operculum and a thickening of the eggshell in the opposite pole to the operculum. We compared the archaeological eggs with eggs of Paragonimus and Fasciola species, but we eliminated the former taxon because the eggs we found lack the diagnostic operculum characteristic of species in this genus. Fasciola was also ruled out because of the larger size of eggs in fasciolid species. Therefore, we consider Echinostoma to be the best differential diagnosis.

The nematodes Ancylostoma duodenale (Dubini, 1843), Enterobius vermicularis (Linnaeus, 1758), and Trichuris trichiura (Linnaeus, 1771) were identified based on morphology and, in the case of T. trichiura, on the egg dimensions (Table I).
The use of the flotation technique in Sheather’s solution resulted in the detection of 10 positive coprolites. In these coprolites, 5 species of helminths were found (Table I). Eight of the coprolites (22%) were positive for the pinworm *E. vermicularis*. One coprolite was positive for *Echinostoma sp.*, *D. caninum*, *A. duodenale*, and *T. trichiura* (Table II).

The sedimentation technique revealed 14 positive coprolites and a total of 6 species of helminths (Table I). Individual eggs identified as human pinworm, *E. vermicularis*, were found in 12 coprolites (33%). *Echinostoma sp.* and *A. duodenale* were found in 3 coprolites (8%). One coprolite was positive for the hymenolepidid, *D. caninum*, and for *T. trichiura*.

The combined results of both techniques show that 19 coprolites were positive for 1, or more, of 6 helminth species. *Enterobius vermicularis* showed the highest prevalence, occurring in 16 (44%) of the samples. *Echinostoma sp.* and *A. duodenale* were found in 3 coprolites (8%). The hymenolepidid, *D. caninum*, and *T. trichiura* were found in 2 coprolites each (6%) (Table II).

The concentration of eggs per infected individual (in epg) was estimated as 2,702 ± 1,038 epg for hookworms, 1,127 epg for whipworms, and 8,848 epg for *Echinostoma sp.* Pinworms were detected in 8 fecal pellets and the concentration ranged from 158 to 2,254 epg, with an average of 823 ± 868 epg.

**DISCUSSION**

There are differences in results with variation in methods. For recovery and subsequent identification of helminth ova, the technique of spontaneous sedimentation appeared to be more sensitive, as it allowed the detection of 1 more species of parasite (*Hymenolepis*idae) and 5 more infected individuals (Table II). This is consistent with earlier assessment of methods (Reinhard et al., 1988). The simplicity of the technique of spontaneous sedimentation may have allowed a greater number of eggs present in the sample to concentrate toward the bottom of the test tube. This, in addition to the greater number of slides prepared per sample, may have increased the chances of detecting eggs, even if the egg output was low. The flotation technique may result in false negatives if the mass of sediment to be re-suspended exceeds a critical mass that may dilute the Sheather’s solution. False negatives may also result from the re-suspension of a small mass of sediment. The latter could be the case for coprolites 1, 2, 3, 5, 8, 11, and 12 (Table II), which were fiber rich. The presence of undigested mass in the coprolite reduces the proportional volume of digested contents and parasite ova.

Only 5 positive specimens were detected by both techniques (coprolites 2, 25, 27, 29, and 34; Table II) and resulted in the detection of the same species of parasite. The exception was coprolite 2, in which eggs of *Echinostoma sp.* were detected by means of the sedimentation technique. The method of flotation failed to detect parasites in 9 coprolites that were found to be positive by sedimentation (Table II). The latter method failed to detect eggs in 5 coprolites found infected by the former (Table II). Based on these observations, we recommend employing at least 2 different methods in subsequent studies.

The prevalence of parasites in coprolites from Cueva de los Muertos Chiquitos exceeds any published record from sites of the Greater Southwest in the United States (Reinhard et al., 1988; Reinhard, 1990, 1992) or from any site in the Americas in general (Gonzáles et al., 2003). The assemblage of parasites includes 6 species, 3 of which have a monoxenous and 3 a heteroxenous life cycle (Lamothe-Argumedo and García-Prieto, 1988; Anderson, 2000). The species showing direct patterns of transmission include *A. duodenale*, *E. vermicularis*, and *T. trichiura*. The rest of the species require at least 1 intermediate host to complete their life cycles. The assemblage of parasites suggest that the people from Cueva de los Muertos Chiquitos were exposed to a wide array of infections caused by parasites specific to humans and a set of zoonotic species. Inferences from the list may indicate that parasites could have been acquired via the ingestion of eggs, ingestion of food items carrying infective stages, and direct penetration.

Among the set of heteroxenous parasites detected in the present samples, all taxa may be associated with rodents and dogs, with humans serving occasionally as definitive hosts (Lamothe-Argumedo and García-Prieto, 1988). In archaeological sites, the
presence of parasites of rodents in human feces has been associated with the practice of storing agricultural goods in granaries (Reinhard, 1987; Reinhard et al., 2007; Reinhard and Bryant, 2008). Grain containers attract arthropods and granivorous rodents that may feed on the grains and the fauna associated with them. Coleopterans and rodents serve as intermediate and definitive hosts for certain species of hymenolepidid tapeworms (Gardner, 1985; Gardner and Schmidt, 1988) although 1 species of hymenolepidid, *Hymenolepis nana*, also has an alternate, direct life cycle. Uncertainty for the taxonomic identity of the hymenolepidid eggs prevents us from determining a pattern of transmission that may be compared against the agricultural practices of the people of Loma San Gabriel (Kelley, 1956, 1971; Brooks et al., 1962; Foster, 1985). *Dipylidium caninum* use fleas as intermediate hosts. It is most commonly a parasite of dogs, but humans can be infected by eating fleas or lice. The presence of *Echinostoma sp.* may be a result of the hunting habits of the people who inhabited this cave. *Echinostoma spp.* include several generalist species that may occur in birds and mammals. Among them, *Echinostoma revolutum* (Froelich, 1802) is a species that is found in cricetine rodents, carnivores, and birds (Yamaguti, 1975) and infects people in Mexico (Lamothe-Argumedo and García-Prieto, 1988). The infection is acquired by eating snails or frogs containing the metacercaria of this trematode (Yamaguti, 1975). The pathology resulting from the infection of these 3 species would depend on the number of worms occurring in the patient. Typically, they may include diarrhea, abdominal pain, and headaches (Lamothe-Argumedo and García-Prieto, 1988), although heavy infections may cause anemia and other complications.

Secondary contamination of the samples by rodents defecating on the coprolites can be ruled out because the sealing of the original midden prevented rodents from gaining access to the burials, evidenced by the fact that the burial food offerings remained intact until their excavation (Brooks et al., 1962; Brooks and Brooks, 1978, 1980).

The 3 nematode species infect their host via direct transmission. Among them, both the pinworm and whipworm are transmitted orally, and the hookworm is generally transmitted via direct penetration of the skin (Miller, 1979).

Until the present, Ancestral Puebloans have had the highest prevalence of pinworm in walled structures built in caves. They make a case that eggs become airborne in such environments and result in infection through the inhalation of eggs and egg-contamination of food and water. The high prevalence of pinworm at the Cueva de los Muertos Chiquitos is consistent with this finding.

Because of the characteristics of their life cycle (Anderson, 2000), eggs of pinworms are usually found in the perianal folds of people and are seldom found in feces. This characteristic has served as the foundation to infer that a prevalence of 5% may indicate the presence of this nematode in almost all the individuals of a population (Fry, 1977). Clearly, the people who lived at Cueva de los Muertos Chiquitos had a serious problem with pinworm infection. Heavy infections with pinworms may cause hyperemia in the blood vessels of the serosa surrounding the appendix, yet they do not induce a severe inflammatory response. The high prevalence of pinworms is consistent with the fact that people from Loma San Gabriel lived in crowded groups (Kelley, 1956, 1971; Foster, 1985, 1986), which can facilitate the transmission of the parasite.

Because of their direct pattern of transmission, both hookworm and whipworms are common parasites of people and occur in high prevalence worldwide (Brooker, 2010). Currently, both species are found in Mexico (Lamothe-Argumedo and Garcia-Prieto, 1988) and their remains had been documented in archaeological sites of the Greater Southwest and other parts of the continent (Allison et al., 1974; Reinhard, 1987; Gonçalves et al., 2003). In the present study, both hookworm and whipworm occurred with a prevalence of 8% and 6%, respectively. Although a precise estimation of the morbidity and mortality caused by these 2 parasites may not be possible, it has been estimated that severe trichuriasis affects 5% of infected children under the age of 15, and anemia-inducing infections by hookworm may affect 6% of infected individuals (Brooker, 2010).

Several species of whipworms are known to infect humans including the specific *T. trichiura* and the zoonotic *T. suis* (Schrank, 1788) (Lamothe-Argumedo and Garcia-Prieto, 1988; Anderson, 2000). These whipworm species may have evolved parallel to the evolution of humans (*T. trichiura*) and pigs (*T. suis*). The absence of pigs in the prehistoric New World indicates that *T. trichiura* infected people at Cueva de los Muertos Chiquitos. Severe trichuriasis affects children and may induce

**Table II. Prevalence of parasite remains of 36 individual coprolites associated with a population of the Loma San Gabriel Culture, located in Cueva de los Muertos Chiquitos, El Zape, Mexico. Average measurements of the ova are presented in μm. An estimation of the total prevalence, as well as the comparative results of both spontaneous sedimentation and flotation technique, is included.**

<table>
<thead>
<tr>
<th>Parasite</th>
<th>Average size ± SD</th>
<th>Prevalence flotation technique</th>
<th>Prevalence sedimentation</th>
<th>Total prevalence</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Echinostoma sp.</em></td>
<td>108 × 66</td>
<td>3%</td>
<td>8%</td>
<td>8%</td>
</tr>
<tr>
<td>Hymenolepididae</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Dipylidium sp.</em></td>
<td>22 × 25</td>
<td></td>
<td>3%</td>
<td>3%</td>
</tr>
<tr>
<td>Nematoda</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Ancylostoma duodenale</em></td>
<td>77 × 62 ± 4 × 6.2</td>
<td>3%</td>
<td>8%</td>
<td>8%</td>
</tr>
<tr>
<td><em>Enterobius vermicularis</em></td>
<td>58 × 30 ± 3.2 × 2</td>
<td>26%</td>
<td>35%</td>
<td>44%</td>
</tr>
<tr>
<td><em>Trichuris trichiura</em></td>
<td>62 × 35 ± 9.2 × 4.2</td>
<td>3%</td>
<td>3%</td>
<td>6%</td>
</tr>
</tbody>
</table>
dysentery, anemia, and rectal prolapse (Brooker, 2010) even before the worm reaches sexual maturity, which happens between 2 and 3 mo post-infection (Anderson, 2000).

The pattern of transmission of hookworms includes the hatching of the larvae in the soil and an active migration of the infective stage from the moist soil through the skin into the lungs and then the small intestine, with a prepatent period of about 43 days (Miller, 1979; Anderson, 2000). Infections by hookworms in children are known to be lethal in post-infection periods as short as 30 days (Miller, 1971) and are known to induce high mortality in children 1 to 5 yr of age in developing countries (Zimmerman, 1946; Bwimbo, 1970). We did not find any blood or a high concentration of watery mucus in the coprolites (Miller, 1971), which are indicators of acute infections. However, the concentration of 2,702 hookworm epg of feces found in 1 infected individual suggests that inhabitants of the cave would have been exposed to heavy infections. Heavy infections are known to cause severe pathology before worms reach sexual maturity (Miller, 1971, 1979). Mechanical damage of the lungs is caused when larvae transgress the pulmonary circulation to the alveoli, causing internal hemorrhaging, and induces death within 24–72 hr after experimental infection of puppies (Miller, 1971).

The presence of these parasites is an indicator of the poor sanitary conditions of the cave environs and the low hygiene levels of the cave inhabitants. The finding of hymenolepidid tapeworms, and D. caninum, suggests the presence of intermediate hosts in the settlements. The presence of whipworms and hookworms indicate defecation in, or in the vicinity of, the dwelling or work areas such as agricultural fields (Reinhard, 1992, 2008; Reinhard and Bryant, 2008; Walker et al., 2009). The latter observation could be associated with the high mortality observed in children 2 yr or younger. The infections caused by parasites may cause a lethal effect on children as a result of the combination of malnutrition and the recurrent uptake of nutrients (especially vitamin B12) by the parasites. The combined results could become exacerbated during periods of drought and famine (Walker et al., 2009), which could have resulted in the high and traumatic mortality levels of the children buried in Cueva de los Muertos Chiquitos (Brooks and Brooks, 1978).

In the archaeological record, increased pinworm prevalence is associated with sedentary, crowded villages (Reinhard, 1987, 1992; Santoro et al., 2003). Reinhard (1992) defined a positive, and significant, correlation between pinworm prevalence in coprolites and cranial lesions from anemia in skeletal populations at the same sites. Recognizing the fact that pinworms do not cause serious pathology such as anemia, Reinhard (1992) made a case that pinworms are a good proxy gauge of pathogens associated with poor sanitation. These include water- and food-borne pathogens as well as air-borne pathogens. The sites with the highest prevalence of lesions attributable to anemia also had a diversity of parasites. Applying this model to Cueva de los Muertos Chiquitos, it is likely that anemia was also a problem faced by the Loma San Gabriel people.

Our results are the first reconstruction of the parasitoses hosted by the people of the Loma San Gabriel culture. With the available results, we have detected the presence of disease-inducing species including hookworms, whipworms, and severalcestodes. It is not possible at this point to find a correlation between the death of the children from Cueva de los Muertos Chiquitos and the parasites discovered. However, future integration of our data with skeletal pathology may reveal connections between parasitoses and morbidity-mortality at the site. At our study area, skeletal remains and coprolites were found in the same cultural association; therefore, the parasite data are directly applicable to a skeletal series. This is due to the fact that a midden (trash deposit) containing coprolites was also used as an area for burial of the dead (Brooks and Brooks, 1978). The Cueva de los Muertos Chiquitos analysis suggests that some parasite species entered into the southwestern United States from people in Mesoamerica. Hookworm became established among agricultural peoples in the Colorado Plateau after 900-yr-ago. The finding of hookworm in Durango, Mexico, 500 yr earlier indicates that hookworm was well established in Mesoamerica at an earlier time. Araújo et al. (2008) presented the value of using parasites as markers or probes for ancient human migrations. The transfer of hookworm from Mesoamerica to the Southwestern United States indicates that interaction between prehistoric peoples in these areas occurred. Importantly, prehistoric hookworms in the southwest were found at sites within the sphere of influence of the Ancestral Pueblo culture of Chaco Canyon. The archaeology of Chaco Canyon is replete with Mesoamerican cultural characteristics, ranging from pottery styles to trade goods such as cacao. Archaeologists have increasingly recognized Mesoamerican influences in Chaco Ancestral Puebloans (Lekson, 2009). The discovery of a hookworm connection between the Mesoamerica and the Ancestral Pueblo culture is the first pathogen indicator of Mesoamerican presence in the southwest.

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LITERATURE CITED


