The role of canids in ritual and domestic contexts: new ancient DNA insights from complex hunter–gatherer sites in prehistoric Central California

Brian F. Byrd a,⁎, Anna Cornellas b, Jelmer W. Eerkens c, Jeffrey S. Rosenthal a, Tim R. Carpenter e, Alan Leventhall d, Jennifer A. Leonard b

a Far Western Anthropological Research Group, 2727 Del Rio Place, Suite A, Davis, CA 95618, USA
b Conservation and Evolutionary Genetics Group, Estación Biológica de Doñana (EBD-CSIC), Seville, Spain
c Department of Anthropology, University of California, Davis, USA
d College of Social Sciences, San Jose State University, San Jose, CA, USA
e ArchaeoMetrics, Woodland, CA, USA

A B S T R A C T

This study explores the relationship between the genus Canis and hunter–gatherers through a case study of prehistoric Native Americans in the San Francisco Bay–Sacramento Delta area. A distinctive aspect of the region’s prehistoric record is the interment of canids, variously classified as coyotes, dogs, and wolves. Since these species are difficult to distinguish based solely on morphology, ancient DNA analysis was employed to distinguish species. The DNA study results, the first on canids from archaeological sites in California, are entirely represented by domesticated dogs (including both interments and disarticulated samples from midden deposits). These results, buttressed by stable isotope analyses, provide new insight into the complex interrelationship between humans and canids in both ritual and prosaic contexts, and reveal a more prominent role for dogs than previously envisioned.

Keywords:
Ancient DNA
Stable isotopes
Canid interments
Dogs
Ritual
Hunter–gatherers

1. Introduction

The genus Canis occupies a unique and enduring niche within ancient human societies, often figuring prominently as totemic symbols and within ceremonial and ritual events and also, in the case of the dog, in an array of functional contexts (Cook, 2012; Russell, 2010; Serpell, 1995). Strong archaeological insight into the nature of these symbolic and prosaic relationships is often hampered by our inability to readily distinguish between canid species based on morphology alone. This study explores this topic by examining the interrelationship between canids and Native American hunter–gatherers in the San Francisco Bay/Sacramento Delta area of central California. This region has a long, well-dated sequence of prehistoric occupation and, during the Late Holocene, complex social organization and high population densities (comparable to the most sophisticated prehistoric hunter–gatherer societies documented worldwide; Lightfoot, 1997; Lightfoot and Luby, 2002; Milliken et al., 2007).

Animal interments, most notably of canids, are a noteworthy aspect of this region’s Late Holocene record and provide an important glimpse into prehistoric ceremonial activities. Based on skeletal morphology, canid interments in the region have been most often classified by faunal analysts as coyotes (Canis latrans), less frequently as dogs (Canis familiaris), and only occasionally as wolves (Canis lupus) (Cambra et al., 1996; Heizer and Hewes, 1940; Langenwalter, 2005; Simons, 2004). Disarticulated canid remains also occur in high frequencies within midden deposits at a number of sites (e.g., Simons, 1992), suggesting canids may have had a significant utilitarian role as well. However, these remains rarely can be classified to the species level.

The considerable morphological similarity between domestic dogs, wild coyotes, and wild wolves can make distinguishing even complete skeletons difficult (Crockford, 2000; Morey, 1992). As a result, differences of opinion have emerged between faunal analysts regarding species assignment (e.g. Heizer and Hewes, 1940; Langenwalter, 1996). This ambiguity hampers our insights

⁎ Corresponding author. Tel.: +1 530 756 3941.
E-mail addresses: brian@farwestern.com, bfbryd@gmail.com (B.F. Byrd).
into symbolic and economic trends associated with the rise and persistence of complex hunter–gatherers in this region. Canids clearly were an integral aspect of prehistoric ritual and symbolic activities, and the ethnographic record in central California suggests that these three species played very different roles within Native Californian cosmology.

In order to gain a fresh perspective into this long-standing problem, we have employed ancient DNA (aDNA) analysis, a technique recently used elsewhere to distinguish canid species in archaeological contexts (e.g., Brown et al., 2013; Horsburgh, 2008; Losey et al., 2011). It is also an approach advocated by Simons (2004:42) for addressing canid identification challenges in the San Francisco Bay area. Stable isotope analyses were also conducted on some canids to gain insight into their diet, and to facilitate comparison with contemporaneous humans in the region ( Bartelink, 2009; Rick et al., 2011). It should be noted that while morphologically wolves and dogs can be very difficult to distinguish directly from their skeletal remains, other lines of evidence (such as the nature of pit disturbances and faunal elements chewing and gastric weathering) may also be drawn upon to discern the presence of these species in archaeological contexts (e.g., Hudson, 1993; Kuznar and Jeske, 2006).

Prehistoric samples include interments previously classified as coyote, dog, and wolf based on morphology alone. Where positive results were obtained at the species level, the aDNA results are entirely represented by domesticated dogs (including interments and disarticulated samples from midden deposits). The results provide new insight into the complex interrelationships between humans and canids, and reveal a greater importance for the dog, the only prehistoric domesticated animal in California, than heretofore recognized.

In this paper we begin by providing a general context for the ritual and prosaic aspects of canids in the prehistory of western North America, focusing on central and southern California. Then we present the samples studied, analytical methods used, and results obtained. We conclude with a consideration of broader implications of these initial results, focusing on the implications for widespread presence of dog rather than wild canids for understanding complex hunter–gatherer practices.

2. Background

Some of the earliest evidence for the symbolic importance of canids is documented more than 12,000 years ago among the Natufian, complex hunter–gatherers of the Near East, when they were first buried with humans (Tchernov and Valla, 1997). Dogs were domesticated from wolves in Eurasia during the Late Pleistocene (Lindblad-Toh et al., 2005; Vila et al., 1997). These domesticated dogs are believed to have then traveled with some of the early immigrants into the New World (Leonard et al., 2002). Once there, they encountered wild New World wolves and coyotes, both of which were to play a prominent role in Native American cosmology.

Dogs, of course, fulfilled varied roles in Native American societies (Snyder and Leonard, 2006: 458). In some contexts they helped with hunting, functioned as early warning devices when strangers approached the village, were beasts of burden, and of course were companions. Along the northwest coast, they were even bred for their wool production. Dogs also sometimes served as a food source, a common occurrence worldwide in traditional societies (Simons, 1994:200–252). For example, among the Plains Indians ethnohistorically, dogs functioned as a famine food and were also sacrificed to honor visitors as a symbol of the importance of their friendship (Snyder, 1991).

Morphological analyses of canid remains from archaeological sites have been used to suggest that there may have been some inter-breeding between canid species (Lupo and Janetski, 1994; Snyder and Leonard, 2006). This possibility is supported by the observation of such crosses in extant populations (Adams et al., 2003; Godinho et al., 2011; Muñoz-Fuentes et al., 2010). This potential adds to the challenge of distinguishing between these three species in the archaeological record.

In general, prehistoric canid interments are most often dogs and less frequently wild coyotes or wolves (Morey, 2006; Snyder and Moore, 2006). Wolves are very rarely documented, with notable examples from two sites in Siberia including one with ochre and a human skull from the Early Holocene (Bazaliiskiy and Savelyev, 2003) and one with a human skull between its legs from the Middle Holocene (Losey et al., 2011). The latter wolf was confirmed using aDNA analysis.

The earliest canid interments (all classified anatomically as dogs) in North America appear to date to at least 8500 years ago at Koster in the Illinois River valley (Morey and Wiant, 1992). In a recent overview, Morey (2006) notes dog burials are emblematic of the Archaic period in North America, found typically as individual interments rather than buried together with humans. In western North America, animal interments of all types are frequent prior to the Late Holocene, and they are most common in the last 1000 years. Canids in the Great Basin are most often reported from caves or from wetland sites, primarily in the western portion where they are occasionally associated with human burials (Dansie, 1990; Janetski et al., 1992; Lupo and Janetski, 1994). Difficulties in distinguishing between canids are well-recognized, although most interments are classified as dogs. Other animal interments are rare or absent.

In the American Southwest, Hill’s (2000) synthesis of animal interments reveals a complex spatial—temporal record that includes significant representation of canids, birds (raptors, macaws/parrots, and turkeys), and occasionally bears. Canids comprise 55% of all animal interments (total sample size of 164 interments and 206 individuals), and have a much longer Late Holocene temporal span (starting around 2500 years ago) than bird interments (mainly between 1000 and 600 years ago). Although many are only classified as canids, when species identifications are provided they are entirely classified as dogs. Contextually, canid interments in the Southwest are highly varied (Hill, 2000: 386–387). They occur in clusters, as multiple interments, and individually; they also occur periodically with human burials. Most are young and many have either cranial traumas or were interred headless. Contextually, they were recovered from floors of ritual and domestic structures, in pits, and in ventilator shafts.

In California, invariably the most common animal interments are identified as Canis, although other animals such as foxes, birds, and bears also occur. Animal interments are best documented in Central California and coastal Southern California. Canid interments are also highly varied, with ritual dismemberment more common in central California and on San Clemente Island in southern California than elsewhere (Hale and Salls, 2000; Heizer and Hewes, 1940). In these two localities, canid interments commonly have associated esoteric and utilitarian artifacts, including shell ornaments.

On the mainland of southern California, canid interments are infrequently reported (e.g., Langenwalter, 2005; Winterbourne, 1967:43–53), with most (17 individuals from 11 interments) from a single site that post-dates 1000 years ago (Langenwalter, 1986). In contrast, canids are better documented on the Southern California Channel Islands; Vellanoweth et al. (2008:3119) state that 95 Canis
interments have been recovered from 41 sites. Although caution is often used (Hale and Salls, 2000), they are generally considered to be dogs, in large part because wolves and coyotes are not indigenous to the islands (Rick et al., 2008, 2011; Vellanoweth et al., 2008). Indigenous island fox interments, however, are well-represented on the Channel Islands.

The southern-most Channel Island, San Clemente, has yielded the most diverse and complex record of animal interments in southern California, and these are generally not associated with human burial grounds. Concentrations of ritual features dating to the last 1000 years were recovered from several sites (Hale and Salls, 2000; Hardy, 2000; Raab et al., 1994; Salls and Hale, 2000). These features included Canis, island fox, and bird interments. Some Canis interments were partially dismembered and then interred with a wide range of associated offerings (red ochre, quartz crystals, bifaces, pipes, abalone shells, and baskets with seeds).

For the San Francisco Bay-Delta area in central California, the only synthesis of animal interments was an early study by Heizer and Hewes (1940) that addressed the Delta area only. They documented 30 animal interments from 10 sites, all dated to the last 1000 years. Canis interments dominate (43%), along with bears, badgers, deer/jack, and antelope. Heizer and Hewes’s (1940:589-590) initial sample of 13 interments from five sites were not classified anatomically as coyotes. Many (42%) lacked hind quarters and the vast majority (77%) had associated cultural material, including clusters of carved abalone pendants, clam shell disk beads, stone rods, a spear point, a charnsest, and a bone awl.

Since then, many more animal interments, including occasional birds, have been reported in the San Francisco Bay-Delta area, as well as in the nearby Sierra Nevada of central California (e.g., Cambra et al., 1996; Haag and Heizer, 1953; Johnson, 1970; Jones, 2010; Langenwalter, 1996; Peak, 1976; Simons, 2004). These more recent studies have confirmed that canids are the most common animal interment, and that the vast majority date to the last 1000 years. Taxonomically, however, the canid record appears more complex. Notably, Langenwalter (1996:1) argues that the coyotes reported by Heizer and Hewes (1940) are actually dogs, and their work was “flawed by poor taxonomic identifications.” Langenwalter (1996) does document a coyote interment within a larger sample of dog interments at a single site in the Sierra Nevada foothills. Canid interments from sites in the Bay Area have been variously reported as dogs (e.g., Haag and Heizer, 1953; Hildebrandt and Mikkelsen, 1993; Leventhal, 1993:351), coyotes (e.g., Gerow and Roland, 1968:84; Wallace and Lathrap, 1975:51), wolves (Cambra et al., 1996), or simply canids (Simons, 2004).

A patchy ethnographic record (comprised of occasional first-hand accounts by early European explorers and colonizers and later anthropological writing based largely on memory culture) provides some insight into the role of canids in Native American populations residing in Central California around the time of Spanish contact (circa 1770 into the mid 1800s). Names for all three canids are present in every Native American language represented in Central California, consistent with mammalogists’ assessment that coyotes and wolves were locally indigenous.

Coyotes figure more prominently than other canids in ethnohistorical accounts of central California Native religious beliefs and practices. For some, such as the Miwok and the Yokuts, coyotes were totemic symbols for lineages or moieties (Driver, 1937; Gayton, 1948). They also appear in ceremonial events, with individuals dressing as coyote dancers or clowns. Gayton (1948:29), in discussing totem animal redemption (the rite of paying for a ritual animal that was killed) among the southern Yokuts, notes “a dead coyote was “redeemed” in the same way by people of the Nutchuw society,” but this was not, apparently, a public affair. Any Nutchuw person who wished to pay for the coyote, whose carcass would then be buried, its hide being kept as a talisman. Dogs, in contrast, are generally relatively minor figures in religious mythology, and wolves are infrequently mentioned.

Varied opinions have been offered regarding the symbolic meaning of canid interments in California. Langenwalter (2005) suggests they typically represent prosaic disposal of personal property and pets after the owner’s death. In contrast, Salls and Hale (2000) referred to the ritual dismemberment of canids on San Clemente Island as “the canid ceremony” and suggested it was part of annual mourning ceremonies that took place within ritual enclosures. A third perspective, first offered by Heizer and Hewes (1940), interpreted these interments as representing symbolic totem markings for moieties or lineages (Jones, 2010; Langenwalter, 1996). Most importantly, these views were invariably contingent upon the species assignment made by the researchers. Clearly, the challenge of distinguishing between species hinders our understanding of the nature of canid-related prehistoric activities.

3. Analysis

We analyzed mitochondrial DNA from a pilot sample of prehistoric canid remains recovered as interments and from generalized midden deposits in the San Francisco Bay-Delta region. This included 13 canid samples from seven prehistoric sites: one from ALA-329, one from MRN-5/1, one from SAC-21, three from SAC-99, three from SCL-732, one from SFR-4/H, and three from YOL-13 (Table 1; Fig. 1). These samples came from both old archaeological projects (housed at the Phoebe A. Hearst Museum of Anthropology, University of California Berkeley and San Jose State University) as well as from recent projects.

All samples are from contexts that date to the last 2000 years, and come from some of the most prominent sites in the region including several sites (SAC-21, SAC-99 and YOL-13) discussed initially by Heizer and Hewes (1940). Most importantly these samples include interments previously classified as coyote, dog, and wolf based on morphology alone. The samples selected for ancient DNA analysis were primarily individual teeth. Teeth are generally the most useful element to analyze because enough DNA can be obtained from the root so the crown can be returned intact for future morphological studies.

Stable isotope analysis was also conducted on a subset of the canid samples to gain insight into their diet, and to facilitate comparison with contemporaneous humans in the same region. The five samples were selected from sites with positive DNA results that had accessable skeletal elements appropriate for bone collagen extraction. They included one sample from ALA-329, three samples from SCL-732, and one sample from SFR-4/H.

3.1. Archaeological contexts of samples

Of the 13 samples, six from five separate sites (ALA-329, MRN-5/1, SCL-732, SFR-4/H, and YOL-13) yielded positive ancient DNA results. The archaeological context of the analyzed canid remains at each of these five sites is briefly summarized (Table 1). It should be noted that although the two sites with multiple samples yielded both positive and negative results (SCL-732 yielded one positive and two negative results, while YOL-13 yielded two positive results and one negative result), all samples from these sites are discussed below. The archaeological contexts of four samples taken from canids at two additional prehistoric mounds (SAC-21 and SAC-99) along the Sacramento River are not presented since ancient DNA was not successfully extracted. Table 2 presents the Late Holocene chronologial sequence for Central California used here, with periods and subdivisions within them based in large part on
changes in predominant shell bead types, referred to as shell bead-style horizons (Groza et al., 2011).

ALA-329, the Ryan Mound, lies near the Coyote Hills along the southeast margin of San Francisco Bay. The site is an anthropogenic mound, some 135 by 90 m in extent and almost 5 m high (Coberly, 1973; Leventhal, 1993:31). Extensive excavations anchored by 39 radiocarbon dates document that the majority of this multicomponent mound dates to the Late Period, 685—180 calendar years before present (cal BP), underlain by Middle/Late Transition and Middle Period occupation horizons extending back until at least 2050 cal BP (Groza et al., 2011; Leventhal, 1993:75—87; Wilson, 1993). Of the 283 burials recovered from the site more than 80% appear to date to the Late Period, based on associated artifacts and stratigraphic position. A loosely flexed canid interment was recovered near the base of the Late Period deposit, and direct dating of the left tibia revealed the interment dates to the Middle/Late Transition or possibly the Middle 4 Period (OS-96418, 963 to 800 cal BP two sigma age range). The skeleton was found lying on its left side and associated with a harbor seal baculum (Fig. 2). The canid interment was originally classified as a dog based on morphology (Leventhal, 1993:351). This canid interment was sampled for ancient DNA and stable isotopes.

MRN-5/H is situated along the northwest edge of San Francisco Bay. This site is a small shell mound (38 by 15 m) adjacent to the marsh of Richardson Bay, a small inlet immediately north of the Golden Gate. Recent excavations by Evans and Smith (2009) documented a rich, 1.5—m-thick midden that included a diverse range of artifacts and a faunal assemblage dominated by clams and fish, along with water fowl, terrestrial mammals and sea mammals. Excavations also documented eight human burials. A series of four radiocarbon dates places the site in the Late Period (685—180 cal BP) and the Middle/Late Transition (930—685 cal BP). A complete canid skull, classified morphologically as a dog (Carpenter, 2009), was recovered from the midden and subjected to DNA analysis.

SCLI-732, the Kaphan Umu'x (Three Wolves) site, is located along the west side of Coyote Creek in the Santa Clara Valley. Broad-scale excavations revealed an extensive prehistoric deposit, concentrated in a 230 by 90 m area, which included a structure, various hearth and platform-related features, 100 human burials, and several animal interments (Cambra et al., 1996). The human burial assemblage is well-dated to the Early/Middle Transition through the Middle 1 Period (2550—1530 cal BP). Many of the features, the structure, and at least one animal interment, however, are dated to the Late Period 2 (430—180 cal BP). The animal features, each separated by ~25 m, include a single canid interment, a double canid interment, and a deer/snake interment (consisting of a headless snake and only the hind legs of a deer).

The three canid interments were all classified as wolves based on their morphology (Cambra et al., 1996) and each was sampled for DNA and stable isotopes as part of this study. The single canid interment (110—1) was loosely flexed, lying on its right side (Fig. 3). This interment dates to the Late Period 2 based on two radiocarbon dates on canid bones (WSU-4604, 311 to ~6 cal BP; OS-96402, 293 to ~2 cal BP; two sigma age ranges) and one date on charcoal from the pit (WSU-4605, 503 to 253 cal BP two sigma age range). In contrast, the double canid interment included a larger (111—1) and a smaller individual (111—2) buried on top of each other in a single pit, each positioned in a loosely flexed position. The larger, lower canid had a single charred three-strand braided rope, or possibly a net, "wrapped around the neck region and the hind quarters" (Cambra et al., 1996:71). Dating results for these two canids are unsatisfactory: modern dates have been obtained on bone samples from each canid (OS-96379 and OS-96386), while a 5301 to 4729 cal BP date (two sigma range, WSU-4539) was obtained on the

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Summary of site contexts with positive results from the ancient DNA analysis of canids.</th>
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<tbody>
<tr>
<td>Site</td>
<td>Setting</td>
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<tr>
<td>-------</td>
<td>--------</td>
</tr>
<tr>
<td>ALA-329</td>
<td>SouthEast San Francisco Bay</td>
</tr>
<tr>
<td>MRN-5/H</td>
<td>Northwest San Francisco Bay</td>
</tr>
<tr>
<td>SCL-732</td>
<td>Santa Clara Valley</td>
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<tr>
<td>SRK-A/H</td>
<td>Yelapa Bomen Island</td>
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<tr>
<td>YCI-13</td>
<td>Sacramento River/Feather River junction</td>
</tr>
<tr>
<td>SAC-21</td>
<td>SAC-99 not included as DNA was preserved.</td>
</tr>
</tbody>
</table>

This table provides a summary of the contexts with positive results from the ancient DNA analysis of canids.
In addition, a sequin bead (M1 type), with an age range of 1400 to 465 cal BP (Groza et al., 2011: Table 5) was recovered from the pit fill. Based on context, isotopic results, and ancient DNA results, these canids are most certainly indigenous and not modern, and most likely date to the latter portion of the prehistoric sequence, consistent with the Late Period dates on other features at the site.

SFR-4/H is situated on the east side of Yerba Buena Island within San Francisco Bay (Morgan and Dexter, 2008). The site includes a 1.3-m-thick shell midden well-dated to the Middle and Late periods, underlain by Early Period human burials within dune deposits (based on 27 radiocarbon dates; Rosenthal, 2008). For much of the sequence the faunal assemblage is dominated by sea mammals, water fowl, and fish. In the Late 1 period, however, terrestrial mammals greatly increase and represent 20% of the large mammal assemblage (Byrd, 2008). Most of these Late 1 Period (685–430 cal BP) terrestrial mammal remains were classified morphologically as canids, including the DNA sample submitted for this study (Simons, 2008). This sample was also analyzed for stable isotopes. The Late 1 Period is also noteworthy as it includes the only animal interment from the site: a golden eagle (Aquila chrysaetos) situated adjacent to a human burial (radiocarbon dated to 500–450 cal BP, two sigma age range, Beta-172582).

Yol-13, the Mustang Mound, is situated southwest of the confluent between the Feather and Sacramento rivers (Taggart and
Jackson, 2005). This Sacramento Valley mound was 2.4 m high and measured 70 by 45 m, with radiocarbon dates (n = 9) from the site revealing occupation concentrated in the Middle/Late Transition, with some earlier (Middle 4 Period) and later use (Late 2 Period) (Breschini et al., 1996; Groza et al., 2011). Initial excavations took place in the 1930s, and Heizer and Hewes (1940:590) report that one canid interment associated with a large, diagonally notched spear point and a charnstone was recovered in Middle/Late Transition contexts (930–685 cal BP). This canid was classified morphologically as a coyote. More extensive excavations took place from 1958 to 1961, yielding 112 human burials and 12 features, including additional canid interments, dating primarily to the Middle/Late Transition Period (900–700 cal BP) (Olsen, 1995). Three samples from canid interments (here designated Canids 1–3) were subject to DNA analyses.

### Table 2

<table>
<thead>
<tr>
<th>Period</th>
<th>Bead-style horizon</th>
<th>Time range (cal BP)</th>
<th>Duration (years)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Historic/Mission</td>
<td>Historic/Mission</td>
<td>180–115</td>
<td>65</td>
</tr>
<tr>
<td>Late</td>
<td>Late 2</td>
<td>430–180</td>
<td>250</td>
</tr>
<tr>
<td>Middle/Late transition</td>
<td>Late</td>
<td>685–430</td>
<td>255</td>
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<tr>
<td>Middle</td>
<td>Middle 4</td>
<td>1200–930</td>
<td>270</td>
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<tr>
<td></td>
<td>Middle 3</td>
<td>1355–1200</td>
<td>155</td>
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<td></td>
<td>Middle 2</td>
<td>1530–1355</td>
<td>175</td>
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<td>Middle 1</td>
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<td>620</td>
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<td>Early–Middle transition</td>
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<td>2550–2150</td>
<td>400</td>
</tr>
<tr>
<td>Early</td>
<td>Early</td>
<td>4000–2550</td>
<td>1450</td>
</tr>
</tbody>
</table>

*After Groza et al. (2011), using Dating Scheme D.*

### 3.2. Analysis methods

#### 3.2.1. Ancient DNA

All pre-amplification steps were performed in a dedicated ancient DNA laboratory at Estación Biológica de Doñana under the direction of co-author Leonard. To prevent possible contamination, this laboratory is physically isolated, has a private air-handling system, including positive pressure and HEPA (High-Efficiency Particulate Air) filters on all vents. Strict rules control access to the lab, and require proper clean attire under full overalls with head cover, dedicated shoes, face masks, and gloves. No materials from the main lab or storage areas can enter the ancient DNA lab; all materials are bought separately and go directly into the ancient DNA lab. All surfaces and materials are cleaned regularly with bleach and ultra violet lights. The lab is divided into two rooms, one for handling teeth and bones (which in some cases may generate dust) and a separate higher pressure room for handling chemicals, extracting DNA, and setting up other reactions.

The correct extraction method depends on the particularities of the site where the sample originated, so some tests of the different methods on samples from each site were performed. A silica-based column extraction method and phenol–chloroform extraction method, followed by column purification and concentration were tested in two individuals. All further extractions were with the phenol–chloroform method as in Losey et al. (2011). Each batch of extractions included a small number of samples and multiple negative controls were carried through the amplification step.

Canid-specific primers which target about 425 of the 5’ end of the mitochondrial control region and that have been developed for ancient DNA and extensively tested were used, in the pairs ThrL/ddi5; ddi1s/ddi3; ddi4/DLHc; ThrL/dog1R; dog2F/ddi5 (Leonard et al., 2002, 2005, 2007; Losey et al., 2011; Muñoz-Fuentes et al.,

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*Fig. 2. Canid interment from ALA-329.*
These primer sets amplify short fragments, which are much more likely to be preserved in ancient material than longer fragments. These short fragments amplify a region that overlaps with adjacent fragment(s) as an additional control against constructing chimeric sequences. Negative controls were included in every polymerase chain reaction (PCR).

Ancient DNA can yield apparent changes in sequences due to degradation of the DNA since the animal died, and not to the presence of a mutation in the animal. The type of degradation that yields these apparent mutations should be random, so to identify these changes and remove them from the data set each fragment of each individual was replicated through multiple PCRs. The various sequences obtained from a single individual were aligned in the program Sequencher (GeneCodes). If the sequence matched, it was considered confirmed. If a mismatch was identified, additional amplifications and sequencing were performed. The base pair at such sites was determined based on the most frequent base pair sequenced from three or more independently amplified products. Amplification was attempted a minimum of six times for each sample. Samples which yielded no positive reaction were discarded after 6–15 negative reactions, and further effort was put into the remaining samples which yielded at least one positive reaction.

A large number of coyote, wolf and dog sequences are published and publically available (i.e. Hailer and Leonard, 2008; Koblmüller et al., 2012; Leonard et al., 2002, 2005, 2007; Vilà et al., 1997, 1999). Sequences identified in the ancient material from this study were compared to all available sequences by BLAST search (http://blast.ncbi.nlm.nih.gov/Blast.cgi) to determine if the exact sequence has been previously identified in other animals. The sequences were then included in a data set of representative dog, wolf, and coyote sequences which was aligned with Clustal W v 1.83 (Larkin et al., 2007). This was used to construct a phylogeny using maximum likelihood with the web-based program RAXML (Stamatakis et al., 2008) and then visualized in Dendroscope (Hudson et al., 2007). Coyotes were assigned outgroup status to add directionality to the tree. If any of the unknown canids were coyote, this would not affect their placement with the coyotes in the phylogeny. This phylogeny enables individual canids to be assigned to species, and, for dogs, to clade.

3.2.2. Stable isotopes

Five bone samples were also analyzed for carbon and nitrogen stable isotopes from bone collagen (C and N) and carbon and oxygen from bone apatite (C and O) in the Stable Isotope Facility and Stable Isotope Lab at UC Davis, respectively. The surface of each bone was cleaned by abrasion with a drill bit and washed and sonicated in deionized water. For collagen extraction, samples were demineralized by immersing them in 0.5 M HCl at 1 °C for five to 15 days. The HCl solution was replaced every one–two days until the sample no longer visibly reacted and was spongy in texture. Samples were then treated with 0.125 M NaOH for 24 h to remove humic contaminants, rinsed with dH2O, immersed in pH = 3 water, and placed in an oven at 80 °C for 24 h to solubilize the collagen. Solubilized samples were freeze-dried. Collagen δ13C and δ15N was measured by continuous-flow mass spectrometry (PDZ Europa ANCA-GSL elemental analyzer interfaced to a PDZ Europa 20–20 isotope ratio mass spectrometer) at the Stable Isotope Facility at UC Davis. All samples yielded ample collagen (greater than 7% volume by weight) indicating excellent bone preservation. The atomic C/N ratio is also reported because it is a useful indicator of sample quality (DeNiro, 1985; van Klinken, 1999; Weber et al., 2005). Collagen from four samples (one from ALA-329 and three from SCL-732) was also submitted for AMS radiocarbon dating.

Apatite from sections of samples was also prepared for carbon and oxygen isotope analyses. Cleaned bone fragments were treated with 1.5% sodium hypochlorite to remove organic components (especially collagen), rinsed thoroughly with deionized water, and immersed in an acetic acid solution to remove soluble contaminants. δ13C and δ18O were measured using a GVI Optima Stable Isotope Ratio Mass Spectrometer at UC Davis.

4. Results

4.1. Ancient DNA

Of the 13 canid samples, six from five separate sites yielded positive ancient DNA results, the first on canids from archaeological sites in California (Table 3). These include four interments (ALA-329, SCL-732, YOL-13 Canid 2 and YOL-13 Canid 3) and two disarticulated samples from generalized midden deposits (MRN-5/H.
and SFR-4/H). Overall, the ancient DNA analysis documented the presence of only dogs in the Central California archaeological record, despite some of these canid interments being previously classified as coyotes and wolves (Fig. 4).

Three of the six samples yielded complete or near complete replicated sequences for the 425 base pair targeted fragment. All three of these samples (MRN-5/H, SFR-4/H, and YOL-13 Canid 2) have been genetically confirmed to be dogs. Previously these samples had been anatomically classified as unspecified canid (SFR-4/H), coyote (YOL-13 Canid 2), and dog (MRN-5/H). A fourth sample yielded about 70% coverage for the 425 base pair targeted fragment. This sample (SCL-732 Canid 111-1) is genetically assigned a dog with high confidence. Based on morphology, this skeleton (as well as the smaller skeleton, 111-2, in the same pit) had been previously classified as a wolf.

The remaining two positive DNA results yielded short fragments of 106 and 107 base pairs, and were not successfully replicated. The fifth positive DNA sample (YOL-13 Canid 3) yielded a short fragment of 106 base pairs of sufficient resolution to be classified as dog. Previously, this YOL-13 interment had been morphologically classified as a coyote. Finally, a sixth positive DNA sample (ALA-329) yielded a 107 base pair fragment that is less variable and only can be determined with high confidence to be a dog or wolf but not a coyote. Previously, the interment from ALA-329 had been morphologically classified as dog.

One of the dogs (SFR-4/H) was assigned to clade IV and all the others to clade I (following Vila et al., 1997). This pattern, with most individuals belonging to clade I and a small number to clade IV, was also observed in pre-contact American dogs, primarily from Latin America (Leonard et al., 2002). In that study a high percentage of the dogs formed a group within clade I, referred to as clade A, which likely evolved in the Americas and apparently went extinct after contact with Europeans (Castroviejo-Fisher et al., 2011). None of the dogs sequenced here fell into that ancient America-specific clade.

4.2. Stable isotopes

Collagen stable isotope analysis was conducted on five samples: two samples confirmed by the aDNA study to be dogs (SFR-4/H and SCL-732 Interment 111-1) and one sample (ALA-329) where the ancient DNA results revealed the interment was either a dog or a wolf (but not a coyote), and two interments from SCL-732 (111-2 and 110-1) that lacked ancient DNA. Collagen C and N isotopes are controlled mainly by protein sources (Ambrose and Norr, 1993; Froehle et al., 2009; Kellner and Schoeninger, 2007; Tiezen and Fagre, 1993). The results reveal a fairly tight distribution with respect to $\delta^{13}$C, ranging from $-16.7$ to $-19.8\%$ and $\delta^{15}$N ranging from $8.6$ to $9.3\%$, indicating a very similar diet (Table 4). Moreover, the aDNA-confirmed dog interment from SCL-732 (111-1) has an almost identical isotopic signature to the second interment in this feature (111-2, which lacked aDNA preservation), and the canine interment from ALA-329.

Our desire was to compare these values to those from humans from the same sites, and to prehistoric wild canids from the region. Unfortunately, isotopic data from humans are available for only some of the same sites as the canids from this study. Bartelink (2006) published isotope results on humans from ALA-329. No data are available from humans from SCL-732, however, data are available for humans from two nearby sites, SCL-134 (unpublished data from one of the authors, JWE) and SCL-287 (Bartelink et al., 2010) (see Fig. 1). Further, published isotopic data on prehistoric wild canids from the San Francisco Bay region are, to our knowledge, lacking. To place our results in context, Fig. 5 plots collagen isotope data from modern South American caves and dogs (Rick et al., 2011), and Pleistocene dire wolves and coyotes from the Los Angeles Basin (Coltrain et al., 2004).

The domesticated dogs from the California Channel Islands had a marine diet and so are completely unlike the canids in this study, and so are not illustrated in Fig. 5 with the other stable isotope values. Humans from the Bay area, plotted as smaller symbols, tend to fall along a diagonal, while wild canids tend to fall to the left and above the humans. Food webs available to the wild canids and humans plotted in Fig. 5 are not identical, making direct comparison difficult. However, overall this result suggests that wild canids in California are consuming protein more from terrestrial environments versus marine environments (less enriched $\delta^{13}$C) and from higher trophic levels (more enriched $\delta^{15}$N) than Bay area humans. The five canids sampled for stable isotope studies in this study are nearly identical in $\delta^{15}$N, ranging between 8.6 and 9.3%. This range is significantly less than that observed among humans (F-test, two tailed, $p = 0.01$) and the prehistoric (F-test $p = 0.007$) and modern (F-test $p = 0.01$) wild canids. This suggests the five canids from archaeological contexts were gaining their dietary protein from a very consistent source (in terms of trophic level). Similarly, $\delta^{13}$C in the five archaeological canids is more varied than $\delta^{15}$N, but still less variable than humans (F-test $p = 0.29$) and wild canids (F-test $p = 0.23$), again suggesting a consistent source for dietary protein. Absolute values for $\delta^{15}$N and $\delta^{13}$C for the dog from SFR-4/H and

<table>
<thead>
<tr>
<th>Site (CA)</th>
<th>Sample context &amp; catalog # (accession #)</th>
<th>Element*</th>
<th>Lab # (JAL)</th>
<th>Species (based on morphology)</th>
<th>DNA species assignment</th>
</tr>
</thead>
<tbody>
<tr>
<td>ALA-329</td>
<td>Canid interment, pit 60N, 60W (54° west, 21° south, 62° deep)</td>
<td>Upper left canine</td>
<td>5358</td>
<td>Dog</td>
<td>Dog or wolfb</td>
</tr>
<tr>
<td>MRN-5</td>
<td>Feature 1</td>
<td>Upper left M1</td>
<td>5362</td>
<td>Dog</td>
<td>Dog</td>
</tr>
<tr>
<td>SAC-21</td>
<td>Canid interment, 1-84570 (Ac# 759)</td>
<td>Lower right premolar (P3)</td>
<td>5353</td>
<td>Coyote</td>
<td>No result</td>
</tr>
<tr>
<td>SAC-99</td>
<td>Canid interment, L-17789</td>
<td>Lower left canine</td>
<td>5354</td>
<td>Coyote</td>
<td>No result</td>
</tr>
<tr>
<td>SAC-99</td>
<td>Canid interment, L-17790</td>
<td>Lower right canine</td>
<td>5355</td>
<td>Coyote</td>
<td>No result</td>
</tr>
<tr>
<td>SCL-732</td>
<td>Canid interment, L-17791</td>
<td>Upper (7), canine #2</td>
<td>5356</td>
<td>Coyote</td>
<td>No result</td>
</tr>
<tr>
<td>SCL-732</td>
<td>Canid interment, feature 110-1</td>
<td>Lower left canine</td>
<td>5357</td>
<td>Wolf</td>
<td>No result</td>
</tr>
<tr>
<td>SCL-732</td>
<td>Canid interment, feature 111-2 (larger canid)</td>
<td>Lower left canine</td>
<td>5360</td>
<td>Wolf</td>
<td>Dog</td>
</tr>
<tr>
<td>SCL-732</td>
<td>Canid interment, feature 111-2 (smaller canid)</td>
<td>Lower right canine</td>
<td>5361</td>
<td>Wolf</td>
<td>No result</td>
</tr>
<tr>
<td>SFR-04/H</td>
<td>Unr 24N, 3E, 30–40 cm (cat #1660.06)</td>
<td>Metatarsal #3, left</td>
<td>5359</td>
<td>Undeterminable</td>
<td>Dog</td>
</tr>
<tr>
<td>YOL-13</td>
<td>Canid interment 1, 1-213685 (Ac# 1295)</td>
<td>Lower, right premolar, #3</td>
<td>5363</td>
<td>Coyote</td>
<td>No result</td>
</tr>
<tr>
<td>YOL-13</td>
<td>Canid interment 2, 1-213685 (Ac# 1295)</td>
<td>Lower, left incisor, #3</td>
<td>5364</td>
<td>Coyote</td>
<td>Dog</td>
</tr>
<tr>
<td>YOL-13</td>
<td>Canid interment 3, 1-213687 (Ac# 1295)</td>
<td>Lower, right premolar #2</td>
<td>5365</td>
<td>Coyote</td>
<td>Dogb</td>
</tr>
</tbody>
</table>

* Identified by Tim Carpenter.
b Result not replicated.
## Table 4
Stable isotope results from canid bone collagen.

<table>
<thead>
<tr>
<th>Sample ID</th>
<th>Element</th>
<th>Collagen yield (%)</th>
<th>C:N ratio</th>
<th>$\delta^{13}$C</th>
<th>$\delta^{15}$N</th>
<th>$\delta^{13}$C apatite</th>
<th>$\delta^{18}$O apatite</th>
<th>Collagen–apatite spacing</th>
</tr>
</thead>
<tbody>
<tr>
<td>SFR-4/H, unit 24N 3E, 30–40 cm</td>
<td>Metatarsal 3</td>
<td>19.9</td>
<td>3.2</td>
<td>$-18.9$</td>
<td>9.3</td>
<td>$-14.9$</td>
<td>$-3.3$</td>
<td>$-4.0$</td>
</tr>
<tr>
<td>ALA 329 interment, pit 60N, 60W</td>
<td>Left tibia</td>
<td>21.5</td>
<td>3.2</td>
<td>$-16.7$</td>
<td>9.1</td>
<td>$-12.1$</td>
<td>$-4.4$</td>
<td>$-4.5$</td>
</tr>
<tr>
<td>SCL-732 interment, feature 110-1</td>
<td>Left radius midsections</td>
<td>7.4</td>
<td>3.3</td>
<td>$-19.8$</td>
<td>9.0</td>
<td>$-15.1$</td>
<td>$-5.2$</td>
<td>$-4.7$</td>
</tr>
<tr>
<td>SCL-732 interment, feature 111-1</td>
<td>Left radius</td>
<td>14.2</td>
<td>3.2</td>
<td>$-16.8$</td>
<td>8.6</td>
<td>$-10.4$</td>
<td>$-5.5$</td>
<td>$-6.4$</td>
</tr>
<tr>
<td>SCL-732 interment, feature 111-2</td>
<td>Left tibia</td>
<td>7.0</td>
<td>3.2</td>
<td>$-16.9$</td>
<td>8.8</td>
<td>$-11.5$</td>
<td>$-6.3$</td>
<td>$-5.4$</td>
</tr>
</tbody>
</table>

C: carbon, N: nitrogen, $\delta$: delta.

### Fig. 4
Maximum likelihood phylogeny of canids from California archaeological sites in this study (highlighted in bold) in relation to previously published domestic dogs (prefix D) from the four main clades, gray wolves (prefix Lu) and coyotes. Partial sequences are marked with an asterisk.
one of the canids lacking aDNA from SCL-732 overlap the bay area humans, suggesting a similar diet for these dogs and people. The remaining three samples comprise a tight cluster somewhat removed in isotopic space from the other samples, indicating diets close to the humans, but quite different than wild canids. The enriched carbon in these three samples suggests elevated input of marine carbon, as C4 and CAM plants are rare in this part of California. As well, depleted δ15N indicates relatively low-trophic-level sources of protein. Together, this suggests these three canids were consuming significant levels of low-trophic level marine foods, perhaps in the form of small fish or shellfish remains.

**Fig. 6** plots apatite δ13C (−10.4 to −15.1‰) and δ18O (−3.3 to −6.3‰) for the five samples in this study relative to the same set of Bay area humans (isotopic values for apatite were not available for the wild canids plotted in **Fig. 5**). Again, isotopic values for the samples in this study are narrow in their spread, especially δ18O. Significantly, the canids overlap with the values from the humans. Because δ18O is mostly controlled by water sources and δ13Capa by complete diet (Froehle et al., 2009; Kellner and Schoeninger, 2007), these data suggest the canids in this study were drinking water and consuming carbon from a similar source as humans.

In sum, collagen and apatite stable isotope values from the dogs are similar to recent stable isotopic studies of Late Holocene human remains from sites in the San Francisco Bay area (Bartelink, 2006; Beasley, 2008). For example, Late Period human burials (n = 8) at ALA-329 yielded collagen δ13C = −16.4 to −18.6‰ and δ15N = 8.8–11.8‰ (Bartelink, 2009: Table 1). Similarly, Middle 1 Period occupation (circa 2150–1530 cal BP) at SCL-287 yielded δ13C = −17.1 to −19.3‰ and δ15N = 6.5–14.4‰ (Bartelink n.d.:5–5 to 5–6). This indicates that the diet of these prehistoric Native Californians was largely terrestrial with only modest contributions of marine foods. As well, the protein component of diets was dominated by lower trophic-level foods, such as plants and herbivores.

The isotope results suggest that the canids in this study had a diet similar to humans. An analogous trend was documented by Rick et al. (2011) for dogs and humans on Santa Rosa Island in southern California, revealing a commensal relationship between humans and domestic dogs. Indeed, this pattern has been found in archaeological studies globally where collagen has been analyzed from dogs and humans from the same site (e.g., Cannon et al., 1999; Germonpre et al., 2009; Guiry, 2012). Such findings are consistent with the notion that domesticated dogs were fed by humans and, as a result, have a similar dietary composition. Overall, the isotope results from this study are consistent with the DNA results; namely that these canids are domesticated dogs and not wild wolves or coyotes.

### 5. Discussion

Our results are consistent with broader prehistoric trends that suggest that canid interments are most often dogs. The results also correspond well with the prevalence of dogs in interments on the Channel Islands, a setting where native wolves and coyotes were absent. The dogs identified in this ancient DNA study include three from prehistoric habitation sites along the edge of San Francisco Bay, one from the Santa Clara Valley, and two from the junction of the Sacramento and Feather rivers. All are from contexts dating to the last 1000 years associated mainly with the Middle/Late Transition and Late Period.

It is interesting to note that these sites with ancient DNA evidence of domestic dogs fall within the ethnographic territories of the Ohlone, Coast Miwok, and Patwin (see **Fig. 1**). Notably, this is the first definitive evidence that late prehistoric populations in the Coast Miwok and Patwin ethnographic territories had dogs. Kroober (1941:6–7) had previously noted that the ethnographic record indicated that dogs were rare or absent north and east of San Francisco Bay. This is in contrast to neighboring groups, such as the Yokuts in the Central Valley and Sierra Nevada, that were reported to have bred dogs and traded puppies to nearby groups including the Miwok and possibly the Ohlone (Barrett and Gifford, 1933:70; Driver, 1937; Gayton, 1936, 1948; Gifford, 1926, 1955; Kroober, 1941).

While being cautious not to over-generalize from these initial results, it is worthwhile to discuss potential implications if further analyses reveal that most of the canid interments and
disarticulated midden remains in the San Francisco Bay-Delta region are also dogs. Hill (2000: 363–364), in the context of the American Southwest, provides a useful conceptual orientation and pragmatic approach to unraveling the role of animal interments in ancient societies, focusing on ritual behaviors expressed in the archaeological record rather than beliefs about animals. She distinguishes three types of animal interments: animal sacrifice and disposal as “ceremonial trash” often after portions of the body (such as wing feathers or hides) were retained for use in subsequent ritual activities (see also Walker, 1995); dedicatory interment as an offering during a commemorative function (such as the founding or abandonment of a ceremonial structure); and simple interments/expedient disposal lacking perimortem trauma or contextual association. This interpretive approach stresses the importance of distinguishing context, the cause of death, and what portions of the body were interred to understand the roles that animals played in ritual practices.

If dogs were mainly interred in the San Francisco Bay-Delta area, how does this impact insight into the underlying reasons? First, the consistent presence of esoteric artifacts and the occasional postmortem removal of the hind quarter suggests that these interments may not just represent the disposal of the deceased personal property as often suggested (Langenwalter, 2005: 25). Second, the earlier interpretation that canine interments represent lineage or moiety totems appears only applicable if the interments were wolves or coyotes (e.g. Heizer and Hewes, 1940: 602; Langenwalter, 1996). If one can apply direct historical analogy to the problem, then the dearth of dog totems ethnographically in the region reduces the likelihood that this interpretation readily explains most dog interments.

Instead, it appears likely that many of these dogs may have been ritually killed as part of specific prehistoric ceremonial activities. These activities would then have been followed by ceremonial disposal as an interment. Dog interments may also represent offerings of food to the dead as part of ceremonial events such as annual mourning rituals. The removal of the hind quarters from some interments may suggest that ritual consumption also occurred. There is often a complex interplay between ceremonial events and subsistence activities, and in a number of other contexts ethnographically in North America dogs figured prominently in ceremonial and feasting events.

Central California ethnographies provide anecdotal support for the role of dogs in various ceremonies. For example, Kroeber (1932: 328) states that as part of the Kukus initiation ceremony among the Patwin, a bandage was used to cover the initiation wound, and “This bandage has been dipped in the blood of a dog previously killed.” The Pota ceremony among the Central Miwok conducted to appease family members whose relative was killed by violence or where witchcraft was suspected, on occasion included the ceremonial killing of a dog. Dancers with bows and arrows ran through the village and shot dogs who entered a clearing within the village where the ceremony was taking place (Gifford, 1926: 397, 1955: 195–196).

Gayton (1948: 154, 290) notes that the northern Yokuts may have eaten dogs at ceremonial events such as the ghost dance as well as at festivals. Similarly, in the fall of 1819 the Spanish soldier Estudillo witnessed and reported in his diary an annual mourning ceremony among the southern Yokuts when upland groups congregated at the foothill village of Chishcas and conducted a ceremonial mock attack where dogs were killed, the dogs’ owners compensated, and the dogs fed to the visitors: “The men arrived making skirmishes with their bows and arrows, killing dogs and chickens [gallinas] with permission from the Chishcas, and afterward paid the latter with beads. Then they commenced to eat them with great pleasure” (translation in Gayton, 1936: 75). Similarly, speaking of the Yokuts, Powers (1877: 379) states: “Dogs are reared (or were) largely for the flesh which they supply, which is accounted by them a special dainty, and which comes well in play, like the farmer’s yellow-legged chicken, when other meat is scarce.”

These central California ethnographic examples demonstrate that dogs were periodically used to feed guests/visitors, and in some contexts dogs may have been eaten on a more regular basis as well. In contrast, there are no indications in central California ethnographies that coyotes and wolves were eaten, and among some groups they were taboo foods (Gayton, 1948; Kroeber, 1932).

Ancient DNA results from our prehistoric interment and generalized midden samples also raise the possibility that disarticulated canid remains within generalized midden deposits in the region are mainly dogs. In many sites along the San Francisco Bay margin, canids represent an unusually high frequency of medium-large mammals, that at times rival the frequency of ungulates or sea mammals. For example, in an overview of vertebrate faunal assemblages, Simons (1992: Table 4.5) documents that five of the 11 assemblages had Canis remains representing 20–32% of identifiable large- and medium-sized mammals. If these high frequencies of canid remains are primarily dogs, then they may represent a periodic food source that served either as a famine food or were consumed at ceremonial events.

The suggestion that dogs were a food source in Central California is consistent with broader patterns in prehistory. As Snyder and Leonard (2006: 458) note: “In some areas of the Americas, dogs served as a more regular meat source. The use of dogs as food is or was a nearly worldwide phenomenon (Simoons, 1994: 200–252), and in the Americas this practice was widespread (Driver and Massey, 1957: 182, map 8).” The importance of dogs as a food resource in North America appears to have included contexts where they were a staple, where they were eaten in times of scarcity, and where they were only eaten in the context of ritual meals. A number of scholars have commented that the complex roles that dogs played in traditional societies often resulted in an ambivalent attitude (Russell, 2010). Serpell (1995: 248, 254) has noted: “...dog-eating is often associated with more complex psychological contortions” and “In symbolic terms, the domestic dog exists precariously in the no-man’s-land between the human and non-human worlds.”

As discussed by Snyder (1991: 370–374) and Cail (2011) in studies of Plains and Canadian Plateau Indians, dogs have a high fat content compared to other food sources, and unlike wild animals, their fat content varies little between seasons since they scavenge human food remains. As such, dogs would have been an attractive food source during the winter and early spring when wild food resources were lean. Hayden (1995) has argued that feasts and ceremonial events invariably involved foods that were rare, tasty, or had high fat content. Dogs fit his prediction of likely foods to be consumed at feasts such as those documented ethnohistorically in Central California. Whether such activities took place only at the very end of the prehistoric record (between 1000 and 170 cal BP) — the time frame of the dogs samples documented in this study — requires more in-depth research into the nature of the canid faunal record. A profitable focus of future investigations would be to explore variation in canid-related activities between site types (such as at major mounds — potential ceremonial centers — versus smaller seasonal settlements) and within sites (i.e., domestic midden areas versus specialized contexts such as features and public areas).

6. Conclusion

Our results provide further indication of several trends with respect to ritual and symbolic activities associated with prehistoric
animal interments in a slightly broader regional context in western North America. First, the record is primarily from the Late Holocene, especially the last 1000 years. Though other animals are well-represented in Central California, the Channel Islands, and the Southwest, Canis interments are invariably the most ubiquitous. Interments of Canis are also highly varied, with ritual dismemberment most common in central California and San Clemente Island in southern California. In these two localities, Canis interments commonly have associated artifacts. These trends suggest that most were sacrificed and then ceremonially interred, although some appear to represent dedicatory offerings. For San Clemente Island, Salls and Hale (2000) suggest that ritual dismemberment was an aspect of mourning ceremonies that took place in ritual enclosures.

In summary, the ancient DNA and isotopic results provide new insights into the relative importance of dogs, coyotes, and wolves in the archaeological record of the San Francisco Bay-Delta area. The complete dominance of dogs in this pilot study points toward the importance of this domestic species and provides the opportunity to construct a fresh framework for understanding their importance in ceremonial and prosaic contexts. These results also highlight the difficulty of morphologically distinguishing species of Canis in the archaeological record, and so in cases where specific identification is important to the hypothesis, identification by ancient DNA analysis should be employed, if possible.

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