

## Supplementary Materials for

### The evolutionary history of dogs in the Americas

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**Other Supplementary Material for this manuscript includes the following:** (available at www.sciencemag.org/content/361/6397/81/suppl/DC1)

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#### **Materials and Methods**

#### Archaeological Site Descriptions

Here we describe all sites associated with pre-contact samples that were used in the analysis (see Table S1 and Fig 1a for more information). We include the sample number(s) assigned by the home depository along with our study sample number(s) (*e.g.*, depository; sample). Where only one set of numbers is given the depository number and sample number are the same.

#### Aachim Lighthouse (AM1, AM2; CGG10, CGG11)

The Aachim Lighthouse site is located on the Aachim Peninsula on the Eastern Siberian Sea coast. This site has the remains of sea mammals and stone tools, as well as the mandibles of two dogs that have been radiocarbon dated to 1,700 BP (27).

# Angel Mounds (AM310A, AM310B, AM310C, AM474; FS3305, FS3283.33, FS3283.22, FS2187)

Angel Mounds is located along the Ohio River just east of present day Evansville, Indiana. The site consists of eleven mounds, several plazas, and numerous residential neighborhoods surrounded by a large semi-circular palisade (28). More recent research suggests that site construction began just before AD 1050 and continued through AD 1450 with the primary occupation occurring during the Late Mississippian period between AD 1350-1450 (29). Extensive excavations were conducted at Angel Mounds, from the WPA era in the late 1930's through the 1960's, and several dog burials and dog skulls are recorded from excavations. Of the four dog burials listed in the Angel Mounds catalog, only one contains an articulated dog skeleton that can be considered an intentional and undisturbed dog burial. This dog, FS 3305 (AM310A), was photographed during WPA excavations and was recovered from the block known as X11B. There are also the remains of two dogs (which are mixed in the same context) recorded as "dog burial", FS 3283.33 (AM310B) and FS 3283.22 (AM310C), which comes from the same excavation block as the articulated dog; and FS 2187 (AM474), which comes from block W11A. These second and third "burials" consist of several disarticulated bone fragments representing incomplete remains, and do not have any hand-drawn or photographed in situ representations associated with them. FS 3283 includes a smaller dog's cranial and mandibular fragments (partially burned) as well as a large dog's limb bones and postcranial elements. Materials associated with FS 2187 include the cranium, without mandibles or postcranial elements, of what is an average-sized Mississippian dog along with the shaft portion of a single right humerus from a sandhill crane (Grus canadensis) from context W11A.

#### Anker (ISM21C; AL2705)

The Anker site (IAS CK 21) is a Mississippian village site on the north side of the Little Calumet River, just south of Chicago (30). Part of the site was destroyed from previous construction efforts, but a dome-shaped house, fire pits, storage pits, and more than 30 human burials were identified. The individuals interred at this site were likely wealthy, as they were buried with numerous grave goods, including animal skulls and copper pieces. The pottery that has been recovered is fairly homogenous, all jars in the

Oneonta style, which suggests that this site dates to AD 1400–1500. A small number of *Canis* bones were recovered from the site, including some dog teeth that were used as grave goods. The Anker sample included in this study was morphologically identified as *Canis lupus*.

#### Apple Creek (ISM070; AL2707)

The Apple Creek site is a Woodland village on the north bank of Apple Creek, four miles north of Eldred, IL (*30*). The site was excavated in 1962 and 1963, and has cultural components from the Hopewell and Woodland periods. The majority of faunal bones recovered from the site were white-tailed deer (*Odocoileus virginianus*), suggesting that they were an important food source for the area (*31*). Seven dog burials have been recovered from the site, as well as some additional isolated elements, which show evidence of consumption. The dog burials are likely from the Middle or Late Woodland periods (2,000-1,000 BP). The dogs are all terrier-sized, which is smaller on average than most other Woodland dogs. The dog used in this analysis, Burial 1, was a small mature male dog with more tooth wear on the right side of the maxilla and mandible than the left.

#### Baum, Ohio (OSU 13320; AL2748)

The Baum site is a prehistoric village settlement in Paint Creek River Valley, Ross County, Ohio. This is the type site for the Baum Phase of the Fort Ancient culture, dating between AD 950-1200 (32, 33). The village extends over 10 acres of ground, and consists of dozens of circular post structures 3-4 meters in diameter, surrounding a large central mound (34, 35). The mound has been known since the mid-19<sup>th</sup> century, and is interpreted as a large ceremonial pyramid structure with two levels, containing 17 human burials (32, 34, 36), and is perhaps the best-preserved mound in the region. In three seasons of excavations covering 2 acres, Mills uncovered 47 tepee structures, 127 burials, and 234 refuse pits, concluding that the site was occupied year-round by a minimum of several hundred occupants (34).

Abundant, well preserved faunal remains were found at the site within the refuse pits, and remains of domestic dogs were found ubiquitously throughout the village. These remains were described, in the early  $20^{th}$  century, by A.F. Lucas as belonging to Indian dogs of a size and proportion similar to the bull terrier (*34*). A total of 50 dog remains, including 7 crania, were collected. Some of the bones had evidence of cut marks, while others were broken consistent with other prey species, and some were made into ornaments. All crania collected were broken, which Mills suggested was to remove the brain (*34*). Comparisons with other prehistoric dogs suggest these were of the general Indian dog type found at contemporary sites in the region, as well as at prehistoric sites in Texas, and the old Pueblos (*34*, *37*).

While individual dog burials and associated ritual activity are known from contemporary sites at the time (e.g., (38, 39) the evidence from Baum suggests a more domestic role for these dogs. The breakage of dog bones similar to other food animals indicates that at least on some occasions dogs were used as food. Gnaw marks from dogs on other animal bones found throughout the middens suggests that dogs had access to food waste. Unlike earlier Archaic period sites where dogs were regularly afforded

special treatment indicated by their careful burial, these later prehistoric dogs from Baum appear to have been a utilitarian presence at the site, likely similar to modern feral or village dogs, with deposition predominantly occurring in refuse pits.

#### *Cox (1Jo176-66-6; Cox6)*

The Cox mound is located in Jackson County, Alabama, and had a number of flint tools and pot sherds associated with the mound (40). The site dates to the Middle Woodland period (1-500 AD). To the east of the mound, there were thirty human burials, many of which included grave goods such as shell beads and tempered bowls. Multiple dogs have also been recovered from the site, one of which is included in this analysis.

# *Channel Islands (CA-SNI-4, CA-SNI-21 133.21-A, CA-SNI-21 133.21-7, CA-SBA-27 F.492, CA-SRI-41, CAW2, CAO1; CISNI4, CINHA, CINH7, CIAS, CICVD, CAW2, CAO1)*

The Channel Islands are 20 to 98 km from the coast of Southern California, and have been occupied for 13,000 years (41). The peoples living on the island had a primarily marine diet and complex trade networks both between islands and with the mainland. Dogs have lived on the island for at least 6,000 years, and were almost certainly brought over by humans (42). They were likely not used as hunters, but may have been an occasional food source, and were found on all of the Channel Islands. Seven individuals from three islands, as well as one mainland archaeological site, have been used in this study. Two dogs were recovered from Santa Cruz Island, one from the Orizaba Cove area (SCRI-434) that dates to the Late Holocene, and the other from Willows Canyon. Three dogs are from San Nicolas Island, two of which come from the North Head site (SNI-21), which was occupied during the Terminal Early Period (5,000 BP) and the Middle Period (2,000 BP). The third dog is from site SNI-4, although its age is uncertain. One dog derives from Santa Rosa Island, at the Canada Verde site (CA-SRI-41), which dates to the terminal Early Period (4,000 BP). The dog from the mainland was recovered from the coastal Chumash town of Syuxtun, which is now a part of modernday Santa Barbara, and was important both for trade and for interactions between mainland populations and the people from Santa Cruz Island.

#### *Flint River (1Mo48 – 40-11; FR11)*

The Flint River site was a village with a large circular shell mound (2 m tall, 15 m in diameter) located on the Flint River near Courtland, Alabama (40). The mound is made of raw clay, with a large shell deposit. Over 200 human burials have been recovered from the site, as well as 19 dog burials (43). Dogs were interred throughout the period of occupation, and did not share graves with humans. Large numbers of artifacts were also recovered from the mound, including pottery and sandstone bowls, pipes, awls, and other tools. The majority of the components are from the Late Archaic period, however intrusive Mississippian human burials have also been recovered from the mound (44). The dog analyzed in this study may be Late Archaic or Mississippian.

#### *Grass Mesa (5MT23-16; 5MT316)*

Grass Mesa Village was a large village located east of the Dolores River in Colorado (45). It was occupied from AD 700 to the early 900s AD, and has one of the highest

artifact densities of any archaeological site in the Mesa Verde region, with a large ceramic assemblage spanning the whole length of human occupation. The earliest period of occupation consisted of dispersed homesteads clustered around pit structures, while later periods of occupation (starting around AD 760) saw the construction of a great kiva, as well as room blocks and more closely-spaced residential structures. Grass Mesa Village had its height of occupation (roughly 150 households) around AD 850, although areas of the village began to be abandoned not long after. The majority of faunal remains from the site were of multiple species of deer and rabbits, although there were also small numbers of dog and unidentifiable *Canis spp.* bones.

Janey B. Goode, Illinois (11S1232 98-1, 11S1232 34-2, 11S1232 1671-1, 11S1232 845-1, 11S1232 939-1, 11S1232 2601-1, 11S1232 1724-1, 11S1232 4109-2, 11S1232 2793-2, 11S1232 5267-1, 11S1232 5819-1, 11S1232 6287-1, 11S1232 7023-1, 11S1232 7892-1, 11S1232 F9 PP4407-1, 11S1232 I15 PP4747-1, 11S1232 L17 P 2-1, 11S1232 3993-2, 11S1232 Q25 PP4811-1; JBG1M, JBG5, JBG11, JBG12, JBG13, JBG17, JBG19, JBG21, JBG24, JBG26, JBG32, JBG35, JBG37, JBG41, JBG42, JBG43, JBG45, JBG48, JBG50)

The Janey B. Goode site (11S1232) is a settlement near Brooklyn, Illinois, that was occupied from AD 650 to 1400 and was excavated from 1998 to 2004 (46). This site includes a small Late Woodland component (AD 650-900), and is primarily composed of Terminal Late Woodland period features (AD 900-1050), which include storage pits and structures that were likely houses. A small Mississippian component (AD 1050-1400) has also been identified, and this site is only 8 km from Cahokia, the largest Mississippian mound site as well as the center of the Mississippian world. Interestingly, Janey B. Goode also has one of the largest numbers of dog burials of any archaeological site dogs have been recovered from 102 different features (47). The majority of these dogs have been recovered from the Terminal Late Woodland component. Many of these dogs were recovered as complete skeletons, and were found in storage pits or buried under the settlement structures. A small number of individuals may have been ritually sacrificed – six males were buried without skulls, suggesting they were beheaded prior to their burial. Cut marks on a Mississippian dog indicate that dogs may have been consumed during the Mississippian period, but dogs from the Late Woodland and Terminal Late Woodland periods show no evidence of consumption, but instead have vertebral fractures, which may be suggestive of their use as pack animals prior to the Mississippian period.

#### Koster, Illinois (ISM F2256A; AL2135)

The Koster site is a complex, highly-stratified site located in a tributary valley of the lower Illinois River in west-central Illinois, excavated between 1969 and 1978 by the Foundation for Illinois Archaeology, Northwestern University and the Center for American Archeology (48-51). Cultural deposits dated from the early Archaic through Mississippian periods are buried to depths of over 10m, providing a continuous record of Holocene human occupation in the region (52, 53). Given this record, the Koster site has become a valuable resource for investigating changes in the environment, settlement patterns, technology, and human adaptations to climatic and social variation (53-55); see (56) for overview).

The early Archaic layers at Koster (Horizons 11 and 12), dating from c. 9,900-9,000 calibrated BP (57), are some of the most well-studied, revealing dense layers of habitation, artifacts, middens, and hearths. Horizon 11 also included the burial of nine humans, four infants and five adults ((53), outside of the living area. Excavators also discovered the burial of four dogs, found as complete, articulated skeletons in shallow pits (58, 59). At the time of excavation the dogs were dated based on associated material to around 8,500 years ago, making them the oldest examples of intentional dog burials in the world and some of the oldest domesticated dogs ever identified; certainly the earliest in the New World. Our recent direct dating of two of the Koster dog burials (ISM\_F2256A and ISM\_F2357) dated them to between 10,130-9,680 calibrated BP respectively (2), making them among the earliest dated material from Koster and confirming their position as the earliest identified domesticated dogs in the Americas and the earliest dog burials in the world.

One of the specimen included in this study (ISM\_F2256A; AL2135) was found in a shallow, basin-shaped pit with a metate and mano placed near its cranium, though it is not clear whether this is associated with the burial. The skeleton is complete and the animal was buried lying on its side, with no evidence of intentional cut marks or other trauma. Given the presence of preserved bacula in the other dogs, the missing baculum here suggests the dog was a female, and it was an adult (58). Morey and Wiant (58):225-226) previously discussed the confirmed status of the Koster canids as domesticated dogs based on comparisons to modern and ancient wolves and coyotes, due to their small size. Particularly, they noted the wide palate and cranial vault of this specimen in comparison to coyotes, a finding that seem to be corroborated by our genetic analyses (see below).

#### *Little Bear (CT8 D-2 2-90; LB2)*

The Little Bear Creek site is an Archaic shell mound located at the mouth of the Little Bear Creek, in Colbert County, Alabama (60). The mound contained nearly 200 burials, some of which were cremated or interred with grave goods including shell beads. Other artifacts recovered from the mound include shell-tempered pottery and flint knives and projectile points. The dog remains from this site are all from the Late Archaic period, and were deliberately buried in the mound, similar to the human interments, but dogs and humans were not found buried together at this site (61).

#### Mayapan (M2: Q152 bag 1, M3: Q152 bag 1, M4: Q152 bag 1, M10: Q162 bag 4; May2, May3, May4, May10)

Mayapan was the largest Mayan city during the Postclassic period, and was occupied from its founding at 900 BP until its abandonment at 450 BP ((62). It is located in Yucatan, Mexico, and consisted of a monumental center containing temples and ritual buildings, surrounded by residential areas which contained both palaces for the elite and smaller houselots (63). Dog remains were highly concentrated in the Templo Redondo Group, located in the main plaza of the monumental center (Masson and Lope 2013). Over 2,000 dog bone fragments have been identified from the site (64). Young dogs were consumed regularly, and older dogs were often used in ritual contexts.

#### *McPhee Pueblo (5MT4475-20; 5MT520)*

McPhee Pueblo was part of McPhee Village, which is located 6 km northwest of Dolores, Colorado (65). It was occupied for 200 years during the Pueblo I and II periods (roughly AD 800-980). The pueblo consisted of housing clusters with both living and storage areas, clustered in groups with pit structures. McPhee Pueblo was the largest pueblo in the village at its height of occupation (AD 870-900), and was also the only part of the village to be occupied for the full range of occupation. Over 200 bones have been recovered from the site that are identified as *Canis spp.*, including coyotes, wolves, and dogs.

#### Modoc (ISML50, ISM090; AL2810, AL2706)

The Modoc Rock Shelter is at the edge of the Mississippi floodplain, located 2 miles SE of Prairie du Rocher, IL, and is one of the earliest sites of human occupation in Illinois (*66*). The lowest strata has been dated to 11,000-9,000 BP, and the site was occupied for 6,000 years. In the earliest strata, tools and projectile points are primarily all that has been recovered, but slightly later in time fire pits have been identified as well. Trash piles of vertebrate remains suggests that the individuals living in the rock shelter were hunters, and also ate large numbers of gastropods. Post molds have been identified starting at 8,000 BP, along with six human burials, but there is no trace of human occupation after 4,000 BP. Two complete dog burials were recovered from the same strata as the human burials (*67*).

#### Perry (1Lu25-Dog 35, 1Lu25-Dog 39, 1Lu25-Dog59; P35, P39, P59)

The Perry site is a large shell mound located in northern Alabama, on the Tennessee River (68). The shell mound contains hundreds of human burials that have been dated to the middle Archaic to Mississippian periods, based on the artifacts present with the burials, as well as a large assemblage of stone tools (69). Over 100 dogs have also been buried in the shell mound, some of them with humans. The age of these dog burials are largely unknown, as the majority of the dogs were buried without associated grave goods.

#### Port au Choix, Newfoundland (MU NP50A 1; AL3194)

The Port au Choix site is located on Newfoundland's northwest coast on the Port au Choix peninsula, projecting into the confluence of the Gulf of St. Lawrence and the Strait of Belle Isle. The area encompasses a number of well-preserved localities, including a Maritime Archaic burial ground (Port au Choix-3) with over 100 preserved burials (Port au Choix-3, Locus II), excavated from 1967-1969 by Memorial University of Newfoundland (70-72). The Maritime Archaic are defined as native groups in the Atlantic Provinces, dating from approximately 9,000-3,500 years ago, who were well-adapted to a coastal environment (70, 73). The burial ground at Port au Choix is thought to date to approximately 4,400-3,300 years ago (74).

The remains of four dogs were recovered from the Port au Choix-3 burial ground (for review see (75) – two complete and articulated associated burials (c.f. (76) accompanying human Burial 50 (Locus II), fragments of one dog from Locus I, and another from burial fill at Locus II (72). All the dogs appear to be of the Large or Common Indian dog size (c.f. (77):459), though there is some variation between them

which is within the range of sexual dimorphism (72). The dog analyzed here (MU\_MP50A\_1, AL3194) is an older male, likely weighing between 45-55 pounds, killed by a blow to the head, and included, along with another male dog, in a multi-human burial (72): 77-78). Direct radiocarbon dating of the dog resulted in a date of 4,402-3,912 calibrated BP (UCIAMS159456), indicating it comes from the earliest use of the burial ground within the Maritime Archaic period. Tuck ((72):78) proposed these dogs were likely used as companions, hunting aids, and occasionally as travois dogs due to their well-developed muscle attachments. Comparisons to Eskimo sled dogs were unfavorable, but the Port au Choix specimens were similar to other large breed native dogs seen at the prehistoric site of Frontenac Island, New York (78). Early descriptions of large breed native dogs described them as slender and wolf-like, with large erect ears and a long, pointed snout (77):461).

# Prince Rupert Harbour (D1-184, D9-985, D10-973, W5-74, W8-969; PRD1, PRD9, PRD10, PRW5, PRW89)

Prince Rupert Harbour is on the northern Northwest Coast of North America, just south of the Alaskan boundary. Two sites, GbTo-13 and GbTo-54, had large-scale excavations in 2012-2013 (79). The work used a novel method of electronic data collection where virtually all data was spatially linked to real-world coordinates (80). The larger excavation, at GbTo-54, had nearly 500 m<sup>3</sup> excavated. The oldest component here dates from 900 BC (calibrated radiocarbon) to about 250 BC; the main occupation was between 250 BC and AD 800; and there was a slightly less intensive occupation from about AD 900 to 1300. Some late pre-contact activity also occurred, but there were no European trade goods, suggesting no admixture with European dogs likely occurred in the population of dogs at this site (also suggested by our mtDNA analysis; see below).

GbTo-13 had smaller an occupation limited to AD 1000–1300.

Both sites were remarkable for the large numbers of exotic and high-prestige artifacts and animal bones found. These latter include an unprecedented amount of mountain goat (*81*) as well as grizzly and black bear, sea lion, northern fur seal, wolf, and other species otherwise rare in Northwest Coast assemblages. The mammal NISP was 5,810 for GbTo-54 and 499 for GbTo-13 (*81*). Rank-linked artifact and faunal distribution indicated that one house was a chief's residence, yet even the poorest houses appear to have had much more wealth than at almost all other sites in Prince Rupert Harbour, suggesting regional and intra-village ranking. One purposeful dog burial was found just outside the chief's house at GbTo-54. Some 300 dog bones and 21 wolf bones were recovered from GbTo-54, and 25 dog from GbTo-13. The dogs are smaller than the wolves in the area, and so the two species can be distinguished. Three dogs and two wolves were used for this analysis.

#### Reinhardt, Ohio (OSU F5607; AL2772)

The Reinhardt site (33PI880) is a middle period (AD 1200-1400) Fort Ancient village site in the Scioto Valley, Ohio. The site was first systematically excavated in 1988, and was identified as an approximately 91-meter diameter midden ring visible on the surface immediately adjacent to a terrace edge located 150 meters from the Scioto River (82, 83). More recent survey and excavations confirm that the site was a circular, or

arc shaped, village, with a small central plaza (82). Associated AMS dates from wood charcoal and a single human burial support the dating of the site to within the Middle Fort Ancient period (82), though bifaces from at least four earlier time periods were recovered during excavations.

Excavations in 2008 revealed a small number of canid remains, including two individual dog burials (39, 83). The specimen in this study is from a fully articulated, adult, male dog with heavily worn teeth, that was found buried under a layer of sand with a turkey bone awl (83). Significant pathologies to the dog were recorded, including vertebral pathologies representing either severe arthritis, or a healed infection, and an abscess in the right M2 (39). In their analysis of the two dog burials, Nolan and Sciulli (39) conclude that there was significant variation between dogs at the Reinhardt site, though their sample size was prohibitively small. Comparisons with other dogs from the region indicate little change in morphology between the Archaic and later prehistoric periods, despite this being a time of intense cultural transmission and changes in subsistence patterns and lifeways (39, 84).

The role of dogs in the prehistoric Eastern Woodlands is intimately connected with the prevalence of dog burials throughout the Archaic period, a tradition which continued to a lesser degree into the later prehistoric periods (38). Dogs in this region were known to serve a variety of purposes, such as for transportation, hunting, companionship and as a food source. Dogs also participated in ceremonial and ritual activity, and dog sacrifices were not uncommon (38, 85, 86). The individual burial of dogs at Reinhardt suggests a level of intimacy remained between the Fort Ancient people and at least some of their dogs, though isolated dog remains found throughout the site indicate that this intimacy was not afforded to all dogs equally.

#### Scioto Caverns (OSU\_4816\_1\_1, OSU\_4816\_2\_2, OSU\_4816\_2\_4, OSU\_4816\_2\_6, OSU\_4816\_2\_8, OSU\_4816\_3\_4, OSU\_4816\_3\_8; OSU611, OSU622, OSU624, OSU626, OSU628, OSU634, OSU638)

The Scioto Caverns site in Ohio are a series of three limestone caverns located near the Scioto River and Wright-Holder earthworks complex (87). The bones of 25 dogs have also been recovered from this deposit, including 5 nearly-complete skulls and 11 mandibles. The dogs are likely from the Hopewell period (200 BC - 500 AD), based on the nature of their burial, in which they were covered with limestone slabs. Based on their size, the dogs seem similar to Archaic dogs identified from Kentucky and Alabama, and are smaller than Woodland period dogs from the Midwest.

#### Simonsen Bison Kill (ISM 13CK61 7 2; AL2699/ISM172)

The Simonsen site is located in northwest Iowa, near Quimby, and its usage dates to the Paleoindian and Archaic periods (88). It has been suggested to be a bison kill site, given that the majority of remains at the site are of bison, and there are large numbers of points associated with the bison (89). Three zones at the site contained cultural material (88). The deepest has a small number of artifacts and the most bison bones, likely from *Bison occidentalis*. The middle horizon contains charcoal as well as a single artifact,

while the highest zone contains charred wood and ash suggestive of a firepit, as well as the ramus of a large dog, which was used in this analysis.

#### *Tizayuca (MT04\_1661, MT02\_707; AL2546, AL2552)*

Tizayuca (Hidalgo, Mexico) is located in the Basin of Mexico, at only 20 km of Teotihuacan, the largest urban center of Mesoamerica from 1,800-1,400 BP. Operations of rescue archaeology, between 2004 and 2009, allow the identification of at least three successive occupations during the Classic (c. 1,800-1,500 BP), Early Postclassic (c. 1,100-800 BP) and Middle/Late Postclassic (c. 800-500 BP) under the respective influence of Teotihuacan, Toltec and Aztec cultures (90). Over 800 dog bones have been identified, from neonatal to adults, representing at least 41 individuals. Cut marks and burning patterns suggest the consumption of dogs, but several individuals have also been buried in residential and ceremonial areas (91).

Sample AL2552 comes from a partial articulated skeleton buried in a Teotihuacan residential compound. The presence of a baculum indicates it was a male. Sample AL2546 comes from a disarticulated skeleton found in the basement of a domestic household related to Aztec occupation.

#### Uyak, Alaska (HMCZ 38342; AL3198)

The Uyak Site (KOD-145), also known as "Our Point", is a substantial prehistoric midden at least three meters deep, covering hundreds of acres and located on the western side of Kodiak Island, Alaska. The site was excavated by physical anthropologist Aleš Hrdlička from 1933-1936 for the United States National Museum (now the National Museum of Natural History), and consisted of house structures and hearths, stone and organic artefacts, human remains, and a large faunal assemblage (92). The stratigraphy of the midden deposits was divided into three layers by Hrdlička's team based on artifact type and deposit descriptions, and occupation dates from 2,000 BP to Russian contact in the mid-18th century were suggested (92-94). More recent AMS dating of a selection of dog and fox bones confirms that the site was occupied during this period (95). Due to the long occupation history of the site, and without direct dating, the cultural context of the study specimen can only be summarized to span both the earlier Kachemak phase (4,000-900 cal BP) and the later Koniag phase (900-200 cal BP), as it is clear dogs were present throughout the occupation (92, 96). Both cultures were maritime hunter/gatherers, though debate still surrounds whether the nature of the relationship between the two cultures was a direct ancestral relationship or a replacement event (92, 97).

Primarily concerned with the recovery of human remains, Hrdlička and Allen prioritized the collection of only well-preserved and intact canid remains (96), and noted that dogs were ubiquitous throughout the site. Hundreds of domestic dog remains were collected, and early metric analysis identified two types of dogs, classified as a "small" and a "large" type (96). While Hrdlička (92) originally suggested that the small type was restricted to the earliest deposits, later replaced by the large type, other studies have noted size variability among contemporary prehistoric canids throughout the Kodiak archipelago (98), and more recent work suggests the variation could be a result of sexual dimorphism (95). Excavation of the specimens was poorly recorded, so it is unknown what context the dog remains were found in, and whether they represent intentional burials, midden/refuse deposits or some combination of deposition types. This makes any interpretation of the role of dogs at the site difficult, though cut marks suggesting butchery were recorded on a portion of both mandibular and cranial elements (99). Dogs are a known food source, both preferentially or as a "fall-back food" during times of hardship, and additionally serve a variety of other functions in the Arctic, including assisting in prey acquisition, transportation, sanitation through food waste disposal, and even warmth (100-102). As the region's sole domesticate, it is likely the dogs at Uyak fulfilled several of these roles simultaneously.

#### Weyanoke Old Town, Virginia (DB49-LC1, DBU2-LM1; AL3226, AL3223)

The Weyanoke Old Town site, also known as the Hatch site, is located on a tributary of the James River on the coastal plain southeast of Richmond, Virginia. The site was excavated between 1975 and 1989 by the Virginia Foundation for Archaeological Research, Inc. (103, 104) and covers over 4,329 sq. meters (105). Based on associated artifacts, occupation at the site spans from at least the early Archaic through the early English Colonial period, when the Virginia Algonquians occupied the region ((103). The site was home to a Weyanoke (Weanoc tribe) village dating from the prehistoric through the early Colonial period (106, 107)).

Over 112 domestic dogs have been recovered from the site, one of the largest discoveries of domestic dogs in the Americas (85, 105, 108). The dog was the only domesticated animal of the Virginia Algonquians ((109)) and what little is known about the use of native dogs in the region is pieced together from early accounts in contact-period historic documents. There is no evidence that dogs at Weyanoke were used for food, as all dogs were recovered in isolated and associated burials (c.f. (76)), articulated with no evidence of cut marks or burning (105). Dogs of the Virginia Algonquians are documented partaking in small and large game hunting and as protection from predators like wolves and bears (110, 111). Like other regions of the Midcontinent and Eastern Woodlands (112–114), dogs, both puppies and adults, played an important role in burial with humans at Weyanoke. It is unclear whether dogs at Weyanoke were viewed as companions, ritual offerings, or both, but dog sacrifices were not uncommon among contact-period tribes (e.g., (115, 116).

Descriptions of the Virginian native dog often mention a wolf-like ((116–119) or fox-like ((120) appearance with an inability to bark and a propensity for howling. In an analysis of the Weyanoke Old Town skeletal material, Blick ((108)) noted the dogs there are clearly domesticated and not wolf-like in their cranial and postcranial skeletal morphology. He later proposed that admixture between local wolves and aboriginal dogs may explain the reported "wolf-like" behavior and appearance of Algonquian dogs ((105):6). They stood an average of 42 cm high and weighed approximately 10kg, similar to a medium-sized Algonquian native village dog depicted in a 1585 painting by John White (see (121):62, Plate 32). This painting, the earliest depiction of a Native American dog by a European, shows a knee-height, medium-sized yellow-brown dog with pricked ears and a long tail, similar to many modern village dogs or dingoes. In contrast to this, in 1602 John Brereton described dogs in coastal Massachusetts as "fox-like, black, and sharp-nosed" ((*120*):13). The Weyanoke canids are associated with Late Woodland period artifacts, confirming their native ancestry and lack of inbreeding with later introduced European breeds. This is supported by direct radiocarbon dating of one of the dog used for DNA analysis (AL3223) to 985-935 cal BP (OxA-35,516).

#### Yellow Jacket Pueblo (5MT501)

Yellow Jacket Pueblo was a large village in the Central Mesa Verde region that was occupied during the Late Pueblo II through the Pueblo III periods (roughly AD 1050-1260) (122). The village had a number of public and ceremonial structures, including room blocks, plazas, kivas, small towers, and what may be the a Chacoan Great House. A small number of canid remains have been identified (NISP = 27), including one bone that was specifically identified as belonging to a dog. Some of the canid bones show evidence of burning or cut marks, suggesting that these dogs were used as food.

#### Zhokhov (Zhokh2004-18, Zh-90-2, Zhokh2004-19, Zh-90-1, Zhokh2004-113, Zh-03-97, Zh-90-3, Zh-04-154, Zh-05-29; CGG1, CGG2, CGG3, CGG4, CGG5, CGG6, CGG7, CGG8, CGG9)

Zhokhov Island is located off of the northeastern coast of Siberia (123, 124). Human occupation of Zhokhov began in the Late Pleistocene period, with the earliest evidence for humans at the site dating to 9,000 years ago ((124) . House pits have been identified from this time period, as well as bone and antler fragments and abundant amounts of wood. Small blades and bone and ivory points have also been recovered from the site, as well as a fairly sophisticated sledge runner for the time period. Two fragmented dog mandibles, as well as a small number of postcranial bones, have been recovered from the site, and date to the earliest period of human occupation. They are smaller than wolf mandibles, and are similar in size to other ancient Arctic dog remains that have been recovered. Previous mitochondrial DNA sequencing has demonstrated that the dogs belong to Haplogroup A, and are genetically indistinguishable from modern domestic dogs in the hypervariable region of the mitochondrial genome (27).

#### Ancient DNA - University of Oxford

#### DNA extraction

DNA was extracted from teeth or bone samples (see Table S1) in a dedicated ancient DNA laboratory using the appropriate sterile techniques and equipment. Extraction was carried out following the Dabney extraction protocol (*125*) but with the addition of a 30 minutes pre-digest stage (*126*).

#### DNA Sequencing

Illumina libraries were built following (127), with the addition of a six base-pair barcode added to the IS1\_adapter.P5 adapter. The libraries were then amplified on an Applied Biosystems StepOnePlus Real-Time PCR system to check that library building was successful, and to determine the optimum number of cycles to use during the indexing amplification PCR reaction. A six base-pair barcode was used during the indexing amplification reaction resulting in each library being double-barcoded with an "internal adapter" directly adjacent to the ancient DNA strand and which would form the first bases sequenced, and an external barcode that would be sequenced during Illumina barcode sequencing. Libraries were sequenced on an Illumina HiSeq 2500 (Single End 80bp) sequencer at the Danish National High-Throughput Sequencing Centre and on a Illumina NexSeq 500 (Single End 80bp) at the Natural History Museum (London).

#### Ancient DNA - University of Illinois

Most of the DNA extractions were performed at the University of Illinois at Urbana Champaign, at the Carl R. Woese Institute for Genomic Biology, with methods described in Witt et al. (7), but a subset of extractions was performed at the Centre for GeoGenetics at the University of Copenhagen, with methods described in Allentoft et al. (128).

At the University of Illinois, genomic libraries were built for the extracts using the NEBNext DNA Library Prep Kit for Illumina (New England Biolabs). They were amplified twice, first using the NEBNext DNA polymerase and the associated index primers, to allow the samples to be pooled and sequenced together. The first amplification followed manufacturer instructions, and was repeated for twelve cycles. For the second amplification, Phusion High-Fidelity PCR Master Mix with HF buffer (New England Biolabs) was used, and four PCR reactions were made for each sample. Five uL of PCR product from the first amplification was added to each reaction, and the DNA was amplified according to manufacturer's instructions for 12 cycles. The four reactions for each sample were pooled and cleaned using Ampure XP (Beckman Coulter) and MagSi-DNA NGS<sup>PREP</sup> beads (MagnaMedics), using an 80% ethanol: sample ratio. The library was visually examined on an agarose gel, and quantitated using a Qubit 1.0 fluorometer (Thermo Fisher Scientific). Only libraries with concentrations of at least 20 ng/uL were used for capture.

To enrich for mitochondrial DNA, we developed a custom set of RNA baits as part of a myBaits kit (Arbor Biosciences) that covered the complete dog mitogenome with 4x tiling density (see Table S1 for details on the samples that were captured). Captures were performed using the myBaits manual version 3.01 at the University of Illinois, with a 60° C incubation for 28 hours. The heat elution step was skipped, and the capture was amplified using KAPA Hi-Fi polymerase, following manufacturer's instructions for 16 cycles. The PCR reaction was cleaned using MagSi-DNA NGS<sup>PREP</sup> beads (MagnaMedics), and the capture was visually examined on an agarose gel and quantitated using a Qubit 1.0 Fluorimeter, following the manufacturer's instructions. If the DNA concentration of the amplified capture was lower than 20 ng/uL, the capture was reamplified for 8 cycles using the KAPA polymerase prior to sequencing.

At the University of Copenhagen, genomic libraries were built from the extracts using the NEBNext DNA Library Master Mix Set 2 (New England Biolabs), with modifications. The End Repair mix was incubated for 20 minutes at 12° C and 15 minutes at 37° C. The Quick Ligation mix was incubated for 20 minutes at 20° C. The Fill-In mix was incubated for 20 minutes at 65° C and 20 minutes at 80° C. Each step was purified using a Qiagen MinElute PCR Purification Kit. The protocol was followed as directed except that a differing amount of EB Buffer was used for each mix (30 uL for End Repair, and 42 uL for Quick Ligation) and the column was incubated for 15 minutes at 37° C prior to elution. The finished libraries were amplified using Taq Gold in a mix that included 10 uL Taq Gold Buffer, 2.5 mM MgCl<sub>2</sub>, 0.8 mM uL BSA, 0.08 mM dNTPs, 0.2 µM of each of Illumina's Multiplexing PCR primer and a custom-designed index primer with a six-nucleotide index and 2 uL Tag Gold, in a total volume of 100 uL. gPCR was performed on the libraries to assess the quantity of DNA. The PCR conditions were followed according to manufacturer's directions, amplifying for 10-14 cycles, depending on the qPCR results. The PCR reaction was purified using the OIAOuick PCR Purification Kit, with elution in 30 uL EB Buffer and an incubation at 37 C for 10 minutes prior to the elution step. DNA concentration was assayed using a Oubit 2.0 Fluorimeter, following manufacturer's instructions. If the DNA concentration was less than 20 ng/uL, a second PCR amplification was performed using Phusion. The mix included 20 uL template DNA, 2 uL each of primers IS5 and IS6, 50 uL Phusion Master Mix, and 26 uL H2O. The PCR program followed manufacturer's instructions but for 6-10 cycles, and was purified with a QIAQuick PCR Purification Kit as described above. The capture procedure was performed following manual version 2.3.1 at the University of Copenhagen, with a 65 C incubation for 18 hours. The heat elution step was skipped, and the capture was amplified using KAPA Hi-Fi polymerase, following manufacturer's instructions for 16 cycles. The PCR reaction was cleaned using a QiaQuick PCR Purification Kit, eluting 30 uL of EB Buffer after a 15 minute incubation at 37 C. The capture was visually examined and quantitated using an Agilent 3300 Bioanalyzer. If the DNA concentration was lower than 20 ng/uL, the capture was re amplified using the KAPA polymerase.

Samples were pooled 8-10 individuals to a sequencing lane, and were sequenced on an Illumina HiSeq 2500 (80 or 100 bp). The samples captured at Copenhagen were sequenced at the Danish National DNA Sequencing Center, and the samples captured at the University of Illinois were sequenced at the Roy J. Carver Biotechnology Center at the University of Illinois.

#### Data processing - ancient DNA

Raw reads were filtered allowing one mismatch to the indices used in library preparation. Adapter sequences were removed using AdapterRemoval (129). Reads were aligned using Burrows-Wheeler Aligner (BWA) version 0.7.5ar405 (130) to canFam3.1, with default parameters apart from disabling the seed option ("1 1024") (131). FilterUniqueSAMCons (132) was then used to remove duplicates. BAM files from different sequencing lanes were merged using the MergeSamFiles tool from Picard v1.129 (http://broadinstitute.github.io/picard/). To accommodate the low coverage of the nuclear genome of our newly sequenced North American dogs, genotypes were called by randomly sampling a single read of 20 base pair minimum and with a mapping quality (MAQ) and base quality (BQ) of at least 30 at each covered position in the genome, excluding bases within 5bp of the start and end of a read (133–135). The Newgrange dog was genotyped similarly as modern data (except for 5bp at start and end of a read; see below)(16).

For the mtDNA we generated majority consensus (using reads with  $BQ \ge 20$  and  $MAPQ \ge 30$ ) sequence for all samples that had at least 3x average coverage (71 samples; Table S1) excluding bases within 5bp of the start and end of a read.

Molecular damage was assessed using MapDamage2.0 using default parameters (*136*) (Figure S1; Figure S2). Most samples display clear signs of deamination (Figure S1; Figure S2). Samples AL3231 and AL2696 display limited deamination patterns consistent with these being from a relatively recent time period (1000-1400 AD; Table S1).

#### Publically available data

Raw reads/bam files for 47 canid genomes (13–16) were downloaded from NCBI or DoGSD (137)(Table S2). Samples downloaded from NCBI were aligned to the CanFam3.1 reference genome using BWA mem (130). We computed depth of coverage (DoC) for each sample using bedtools (138). These genomes were chosen due to their high coverage and geographic spread covering North and South American, East Asian, and Western European (African, Indian and European) dogs, as well as Eurasian and American gray wolves and Coyotes and an outgroup (Lycalopex culpaeus; Andean fox). We also obtained data from two additional publically available CTVT genomes (20), genome data from an ancient wolf from the Taimyr peninsula (139) and from an ancient Irish dog (16).

Lastly we obtained data from 5,406 modern dogs that were genotyped on the semicustom CanineHD SNP array (~185K SNPs) developed by (9).

#### Genotyping

Dog samples

We used samtools 'mpileup' (0.1.19) (140) to call genotypes with default settings. Pileup files were further filtered, for each sample, using the following criteria:

- Minimum  $DoC \ge 6$
- Excluded all sites in region of high DoC (top 5%)
- Excluded all sites within 3bp of an indel
- Only bases with quality  $\geq 30$  within reads with mapping quality  $\geq 30$  were used.
- Minimum fraction of reads supporting heterozygous (variant allele frequency [VAF] >= 0.3) all sites that did not pass this criteria (0 < VAF < 0.3) were coded as missing (N).
- For high coverage ancient sample (Newgrange dog) we also discarded the first and last 5bp of each read for genotype calling, to avoid incorporating errors from deaminated sites (see above).

The Taimyr wolf was processed using the same random read approach used for the other ancient data (see above).

CTVT

Ancestry analyses were performed using data from two CTVT genomes that have been previously described, 24T and 79T. The goal was to determine the phylogenetic placement of CTVT within a cohort of modern and ancient dogs. CTVT genomes carry two types of genetic variation: germline variation inherited by the CTVT founder dog, and somatic variation acquired during the somatic evolution of the CTVT clone. The goal of this part of the analysis was to capture CTVT germline variation, and to use this to include CTVT in a phylogenetic analysis.

We generated a list of callable sites in CTVT using the criteria outlined in 'Genotyping - Dog samples'. Only regions that retained germline diploidy, as previously described (20), were considered. Sites were further filtered to retain only those sites in which the variant allele fraction (VAF) for a non-reference allele was  $\geq 0.1$  in at least one CTVT tumor, and for which no more than two nucleotides were detected at VAF  $\geq 0.1$ (i.e. multi-allelic sites were rejected). These sites were defined as single nucleotide variant (SNV) candidates

CTVT tumor biopsies contain both CTVT cells and host cells. The latter derive from stromal, immune and blood vessel components. Thus DNA derived from CTVT tumors is an amalgam of CTVT and matched host DNA. In order to identify and exclude SNVs derived exclusively from the matched host, as well as to correctly genotype alleles shared between CTVT and matched hosts, we took the following approach. Sites identified as SNV candidates (see above) were genotyped in genomes 24H and 79H, the matched hosts for tumors 24T and 79T(20). The genotypes of 24H and 79H were inferred using the following VAF thresholds:

homozygous reference: VAF < 0.2heterozygous: VAF = [0.2-0.8]homozygous alternative: VAF > 0.8.

Using the known host contamination fractions for 24T and 79T (20), we used the following VAF thresholds to genotype SNV candidates in CTVT cells. SNVs that were homozygous reference in the matched host were genotyped in CTVT using the following VAF thresholds:

Homozygous reference: VAF<0.2 Heterozygous: VAF=[0.2-0.6] Homozygous alternative:VAF>0.6

SNVs that were heterozygous in the matched host were genotyped in CTVT using the following VAF thresholds:

Homozygous reference: VAF<0.3 Heterozygous: VAF=[0.3-0.7] Homozygous alternative:VAF>0.7 SNVs that were homozygous alternative in the matched host were genotyped in CTVT using the following VAF thresholds:

Homozygous reference: VAF<0.4 Heterozygous: VAF=[0.4-0.8] Homozygous alternative:VAF>0.8

CTVT SNVs were further processed as described below (Ascertainment panel).

#### Ascertainment panel

All genotypes from all genome-wide samples were then merged using bedtools. Ascertainment was done without outgroups (but including Coyote). We selected all biallelic markers excluding sites that 1) were heterozygous in only one sample (Table S2) (required a minimum of two chromosomes in our set of samples to carry the derived allele; in the case of sites only variable in CTVT we required the two genomes (24T and 79T) to be homozygous to limit the inclusion of somatic mutations into the list of SNPs 2) sites that were not covered in our outgroup (Andean fox) 3) sites with more than 20% missing data across samples. All low coverage ancient samples were excluded from this step. As the two CTVT matched hosts, 24H and 79H, were not included in our ascertainment panel (Table S2), the genotypes of these two individuals were not taken into account when determining which CTVT SNPs were represented in other dog genomes. This resulted in ~6.21M high quality SNPs. We then excluded all sites that were outside of the germline diploid region in CTVT (*20*). This resulted in ~2.03M SNPs, including ~600K transversions.

#### mtDNA analysis

#### RAxML

We used all samples with at least 3x average coverage and consensus sequences with at least 80% coverage over the entire mtDNA genome were considered for further analysis. We further obtained ancient and modern mtDNA genomes from (*141*). This data set contains representative samples of all four major haplogroups (A, B, C, D) including 3 ancient American dogs. We aligned the data using mafft v7.2 (*142*, *143*). We built a maximum likelihood tree, with 100 bootstrap replicates using GTR+G model as implemented in RAxML (*144*).

All but one ancient American mtDNAs formed a monophyletic clade within haplogroup A, (bootstrap value=87; Figure S3). North American dogs further cluster with ancient sled dogs from the the island of Zhokhov in Eastern Siberia (*124*). Unsurprisingly, samples CGG10-11 (Aachim dogs) fall outside of the Zhokhov / precontact clade as these are recent sled dogs from Siberia (~1.5kya; Table S1). Lastly, one sample from British Columbia (Prince Rupert Harbour site; PRW89; ~1.5Kya) clusters with North American wolves. Wolves and dogs are poorly distinguished at this site (see above) so the sequence of this sample might be from a wolf - although interbreeding between wolves and dogs is also a possibility (see below). We then assessed whether previous studies that used control region of the mtDNA were able to identify the pre-contact monophyletic clade we have identified here (Figure S3). To do so, we extracted the control regions from all samples, that overlapped with the fragments analysed in (5) and (8) (605 bp in total), filtering out samples with more than 10% missing data and a ML tree with RAxML. The result of this analysis are presented in Figure S4. We found that while the control region has the power to distinguish between the major dog haplogroup (A, B, C, D) it did not possess the power to distinguish between pre-contact dog and other dogs within haplogroup A.

We expanded our mitogenome sample size to assess whether the mtDNA haplogroup that we had identified in pre-contact dogs exists in modern America dogs. To do so we used 942 additional mitogenomes from a worldwide sample of dogs, including CTVT and hosts genomes as well as 169 village and breed dogs that were sampled in North and South America (145-149). Description and accession number of all additional samples can be found in (145-149) (in Supplementary file 1 and 8 of . We combined this data with the mtDNA genome analysed above and built a maximum likelihood tree, with 100 bootstrap replicates using a GTR+G model as implemented in RAxML (144). Out of 667 modern domestic dog genomes analysed here we found only five modern samples with a pre-contact mtDNA haplogroup (Figure S5): 1) Terrier cross from San Juan del Sur, Nicaragua (Accession: KU291094), 2) Chihuahua (Accession: EU408262) 3) Japanese Spitz (Accession: EU789755) 4) non-breed dog from Shanxixian, China (Accession: EU789669) 5) non-breed dog from Laem Ngop, Thailand (Accession: EU789664).

Interestingly, two out of five of these sequences are from American dogs (Chihuahua and Nicaragua dog). Three, however, are from East Asia. This is surprising and suggests a very low frequency of the PCD haplogroup in East Asia (~2.5%). All five modern samples cluster together with ancient Mexican dogs from Mayapan (Figure S5). Multiple scenarios could explain the finding of East Asian dogs within the PCD clade: 1) the clade to which these East Asian samples belong diverged from PCD dogs prior to their introduction into the Americas 2) there was some back and forth migration of dogs between America and Asia after the flooding of the land bridge between Western and Eastern Beringia ~11,000 years ago (150) 3) these sequences were mislabelled.

#### BEAST

We used BEAST v1.8.4 (151) to calibrate the evolutionary rate of our canid data set. We restricted this analysis to sequences with at least 10x average coverage (Table S1). The mtDNA was partitioned into four categories (tRNA, rRNA, control region and coding sequence). We fitted a separate substitution model to each partition: tRNA (HKY+I), rRNA (TN93+G), control region (HKY+G+I) and coding sequence (SDR06) as selected by Akaike information criteria (AIC) using partitionfinder (152). The same tree was used for all four partitions. Age of archeological samples was used as prior (uniform distribution of tip age). We used a Bayesian Skyline prior (153)(group size parameter = 10) and a strict molecular clock as in (141) (uncorrelated clock was also tested and did not result in noticeable changes). We ran 50 million Markov chain Monte Carlo (MCMC) chains and sampled tree parameters every 5,000 iterations. Convergence was evaluated with Tracer v 1.6.0 (ESS for each parameter  $\geq 100$ ). Trees were summarized using Maximum Clade Credibility as implemented in TreeAnnotator v1.8.4 (10% burn-in).

BEAST retrieved the same topology as RAxML, with all pre-contact dogs forming a highly supported monophyletic clade (with a posterior probability of 0.99; Figure S6). We estimate that TMRCA of all pre-contact dogs is ~14666 years (95% HPD: 12965-16484) and that the time to most recent common ancestor (TMRCA) of all sampled ~15606 years ago (95% HPD:13739-17646).

We also inferred the age of the MRCA between the Mayapan dogs and the five modern dogs identified as monophyletic with PCD (Figure S5). We found that these five modern dogs diverged from the Mayapan dogs between 9,865 and 6,289 years ago (95% HPD) suggesting that their divergence postdates the flooding of the land bridge between Western and Eastern Beringia. Given this results, it is unlikely that the mtDNA haplotype of these dogs originated in Eurasia. These results instead suggest that dogs carrying PCD ancestry have been transported from Americas into East Asia. This most likely took place during recent times and could be linked to the creation of hairless dog breeds in Asia (154). However, more research is needed to further test these possibilities, especially given the possibility of mislabelling in databases such as GenBank.

#### Nuclear ancestry analyses

#### PCA

Using smartpca (155) we performed Principal Components Analysis (PCA) using various projections and data sets on our 2.03M SNPs:

- 1. All canids (including wolves and coyotes) PCD samples projected (Figure S7)
- 2. Only dogs (excluding wolves and coyotes) PCD samples projected (Figure S8)
- 3. Only dogs (excluding wolves and coyotes) PCD and CTVT samples projected (Figure S9)

For PCD we used all 7 samples for which we could call at least 10,000 sites (minimum number of sites suggested for ancient DNA analysis (128)). We used all available sites (sites covered in at least 1 ancient sample; ~1.5M SNPs) to compute the eigenvectors and then projected PCD onto that space. We also projected CTVT to ensure that their placement was not an artefact of somatic mutations.

Figure S7 shows that PCD are more closely related to dogs (except for one sample, AL2135 from Koster; see below) than wolves or coyotes. It also shows that dogs are less variable than wolves or coyotes. Figure S8 shows a distinction between Arctic, East Asian, and European dogs. Lastly, Figure S9 recapitulates the same results demonstrating that this result is not induced by somatic mutations in CTVT and also shows how PCD and CTVT are more closely related to each other, and fall in between Arctic dogs and all other dogs. A PCD sample (AL2135; Koster, Illinois) was projected in between dogs and wild canids (Figure S7). This suggests that this sample is admixed with wild canids (see

D-statistics analyses below). Its mtDNA haplotype, however, clusters with other PCD dogs (Figure S3).

#### *Neighbour joining tree*

We used plink v1.9 (156) to compute an Identity By State (IBS) matrix using all 2.03M SNPs. This matrix was used to build a neighbour joining tree (NJ) using the R package "ape" (157); (Figure S10). The tree recapitulates the deep split between East Asian and Western Eurasian dogs (16) and confirms that CTVT is more closely related to the PCD dogs than to any other dog population (bootstrap=100). It also shows that CTVT/PCD form a monophyletic group with Arctic breeds that fall outside of the rest of the dogs. The tree also confirms that PCD form a monophyletic clade with high support (bootstrap = 100).

Admixture analyses (see D-statistics below) show that all East Asian dogs, excluding Vietnamese Village dogs, are significantly admixed with European dog populations. Such disproportionate admixture could affect the topology of the tree. To test this we built a tree excluding all East Asian dogs except Vietnamese. This tree shows a different topology with Vietnamese still grouping with Dingoes, however, PCD, Arctic dogs and CTVT are now more closely related to Western dogs than to Asian Dogs (Figure S11).

#### **Bayesian** Tree

We built a phylogeny using nuclear genotypes with MrBayes 3.2 (158). To do so we used PGDSpider 2.0.9.2 (159) to build a Nexus file with discrete SNP format (0=reference, 1=heterozygous, 2=homozygous alternative). We used the Mkv model (160) implemented in MrBayes (Ordered character), which provides a likelihood framework for data sets that contain only variable characters. We also imposed a minimum distance of 10Kb between SNPs to limit the influence of linkage disequilibrium (LD) and lastly included only PCD samples with higher coverage (AL3194 and AL3223; Table S1; ~30K SNP total).

We ran two independent runs of four MCMC chains with two million samples. Trees were summarized discarding 25% as burnin. To limit biases from missing data we limited this analysis to transversions that were covered in 90% of our samples. Convergence was assessed by ensuring that average standard deviation of split frequencies was below 0.01 and that the potential scale reduction factor was close to 1 for all parameters. This analysis confirms that CTVT and PCD are monophyletic with high support (Posterior probability [PP] = 1; Figure S12) and the basal placement of the CTVT/PCD clade. However, this analysis suggests that modern Arctic dogs are more closely related to Eurasian dogs than to PCD. This is most likely due to the complex ancestry of Arctic dogs such as admixture from European dogs (see below; Table S4).

#### f3 statistics

We computed outgroup *f3*-statistics as f3(pre-contact dogs [PCD], X; outgroup) using ADMIXTOOLS (*161*) where X is any other dog population (see Table S2), to quantify the amount of genetic drift shared between pre-contact dogs and other dogs

using only transversions (Figure S13). For this analysis, we used only two PCD samples (AL3194 and AL3223; Table S1), with ~1.9x and ~0.5x coverage, respectively. Our results support our NJ tree (Figure S10) demonstrating that PCD is more closely related to CTVT and Arctic dogs than any other dog population. These results also support the observation that PCD/CTVT and Arctic breeds are equally related to all other dogs, except for Basenji and one Indian dog, which could be due to admixture from wolves into these two samples (e.g. Indian wolf or golden wolf).

#### D-statistics

We only used two PCD samples with ~1.9x and ~0.5x coverage (AL3194 and AL3223; Table S1) for these analyses, except when explicitly mentioned (e.g. Koster dog AL2135; see below). We used all 2.03M SNPs.

#### PCD is more closely related to CTVT and Arctic breeds

We computed D(Outgroup, PCD, Pop3, Pop4) using ADMIXTOOLS (*161*) where Pop3 was fixed as either European dogs, Asian dogs, Arctic dogs, or CTVT and Pop4 represented any possible other sample. We plotted, as box plots, the results of these combinations (Figure S14; Figure S15). Positive values imply that PCD shares more derived alleles with the population on the y axis, while negative values imply that precontact dogs are closer to the other dog populations. The results indicate that PCD do not share any more derived alleles with European dogs than they do with Asian dogs, suggesting that they are equally related to both. This result supports our f3-statistics and NJ tree finding that PCD and CTVT are equally related to European and Asian dogs (Figure S10; Figure S13). Arctic breeds also appear more closely related to PCD/CTVT than any other dog population (Figure S14; Figure S15; Figure S13).

#### Admixture between Coyote / North American wolves and PCD

We tested for admixture from wild North American canids into higher coverage PCD genomes (AL3194; AL3223). To do so we computed D(Outgroup, Coyote/North American Wolf, Pop3, Pop4) where Pop3/4 can be any dog genome. We found that in both cases Z values were mostly above 3 in most cases (Figure S16; Figure S17) for both AL3194 and AL3223 indicating admixture from Coyotes / North American Wolves in PCD samples. We also tested for extra admixture from wild canids into our higher coverage PCD genomes (AL3194; AL3223). To do so we computed D(Outgroup, Coyote, AL3194, AL3223) and D(Outgroup, American Wolf, AL3194, AL3223). We found no evidence of extra admixture from wild canids into these samples (Table S3).

We computed D(Outgroup, Coyote, CTVT, B) and D(Outgroup, American Wolf, CTVT, B) where B represented every possible pair of populations to determine whether there was any detectable admixture from Coyote and American wolf populations into the CTVT founder dog (Figure S16; Figure S17). We also found evidence that the CTVT founder dog shared more derived alleles with Coyotes than other non-PCD population suggestive of admixture. This is consistent with TreeMix and Qpgraph analyses (see below). We note, however, that this pattern could be consistent with admixture in both direction (dogs to wolves / wolves to dogs; e.g. see (23)).

#### Estimating Eurasian ancestry in Arctic dogs

Using D-statistics based on whole genome data we found evidence that all Arctic breeds are a mixture of the basal lineage (that leads to CTVT and PCD) and of the Eurasian dog lineage (Table S4).

#### Taimyr admixture into Arctic dogs, PCD and CTVT

We used whole genome data to assess Taimyr wolf admixture into PCD, Arctic dogs and CTVT (139). We found few values with |Z| > 3 (AL3194 and Alaskan malamute; Table S5). Lowering the threshold to |Z| > 2.5, we found admixture from the Taimyr wolf into PCD (both AL3194 and AL3223) as well as in all Arctic dogs (husky, Greenland sledge dog, and Alaskan malamute) and CTVT. We find no evidence for additional admixture from the Taimyr wolf into either CTVT, PCD or Arctic breed (Table S6). This suggests that Taimyr admixture into Arctic dogs suggested in (139) may have taken place after the PCD, Arctic dog and CTVT lineage diverged from Eurasian dogs but before the divergence of the Arctic dog and PCD/CTVT lineages.

#### Admixture from European dogs into East Asian dogs

We tested for admixture from European dogs into east Asian dogs. Following (9, 16) we used Vietnamese village dogs as the reference East Asian population to test for by computing D(Outgroup, Portugual, Vietnam, X). We found evidence of admixture in all East Asian populations test in this study (Table S7).

#### Potential evidence for Coyote admixture in Koster dog (AL2135)

Our PCA analysis suggests that AL2135 is admixed with wild canids. To test this hypothesis we computed D(Outgroup, North American canid / Taimyr wolf, AL2135, AL3194). We restricted this analysis to AL3194 as it is the highest coverage PCD dog available in this study. We found borderline significant results (|Z|>2; Table S8) suggestive of admixture from Coyote into AL2135. This sample, however, is very low coverage (only ~17K SNPs were called). Its placement on the PCA and this positive admixture signal might therefore be due to this low coverage.

#### Estimating pre-contact ancestry in modern North American and Arctic dogs

We used the SNP array data obtained from (9) to assess the degree to which modern dog populations found in North America retained ancestry from pre-contact dogs. This SNP panel contained 28 genotyped populations from North America, such as Peruvian village dogs, Alaskan village dogs or Carolina dogs (see Table S9 for the full list). We computed f4 ratios using ADMIXTOOLS (*161*, *162*) to estimate admixture proportion ( $\alpha$ ) from pre-contact dogs into these populations by computing:

 $\alpha = f4(A,O;X,C) \div f4(A,O;B,C)$ 

Where A is CTVT, O is the Andean fox (outgroup), B is PCD (AL3194 and AL3223), C is any European or East Asian population (see Table S9) and X is any American dog (see Table S9 and Figure S18). We computed  $\alpha$  for all combinations of European/Asian and modern North American populations (jackknifing was performed with a block sizes of 1 cM).

Besides the Alaskan Village dogs, we found no significant signal of pre-contact ancestry in modern North American populations ( $\alpha$  always <4% and Z always < 3; Table S10). Alaskan Village dogs, on the other hand, have ~17% (11-20% and Z always > 4.5; Table S10) ancestry derived from pre-contact dogs. We also used outgroup f3 statistics to assess the degree of shared drift between various populations available on the SNP array and PCD (Figure 2b).

To further assess this result we used ADMIXTURE (163) on a subset of the SNP array samples including all modern North American populations as well as Arctic dogs, "basal" breeds (164) and selected European and Asian populations (e.g. Boxer and Chow-Chow). K=4 was selected as the best K value based on 10 fold cross validation (Figure S19). This analysis support previous f4 ratio analysis showing that most modern North American dog populations have little pre-contact ancestry (<4%; Figure S20). ADMIXTURE, however, detects some evidence of limited PCD/Arctic ancestry in Carolina dogs ranging from 0-33% (Figure S20; population CD). Such signal might not have been detected by our F4 analysis as a result of the variable amount of ancestry in this population. This analysis also reveals an affinity between Chinook and PCD/Arctic breeds (12-15%; Figure S20). This is not surprising given that Chinook dogs are considered as Sledge dogs. With K=4, however, we cannot distinguish between PCD and Arctic dogs ancestry. This PCD/Arctic component in Carolina dogs and Chinook might therefore be the result of admixture with Arctic dogs rather than PCD. To test this we tried to separate PCD/Arctic ancestry with higher K values. Both K=10 and K=15, however, failed to differentiate PCD and Arctic dog ancestry (Figure S20) but instead differentiated New World Arctic dogs (Alaskan malamute and Greenland sledge dogs) from Old World Arctic dogs/PCD (Figure S20).

Alaskan Village dogs were the population of north American village dog with the most PCD admixture. This is not surprising as these are closely related to Arctic breeds (6, 9). The f4 ratio conducted above is thus not appropriate for this population (as it assumes close relatedness to Eurasian dogs; see Figure S18). Here we wanted to test whether these dogs have any pre-contact ancestry (interbred with pre-contact dogs). To do so we computed every possible combination of the same f4 ratio as above but using only Arctic breeds. We found that both Alaskan malamute and Greenland sledge dog have a significant amount of ancestry from PCD (~4-14%; Table S11). This however, might be the result of substructure among Arctic dogs (see below).

We assessed whether these results could be affected by the ascertainment of the SNP array by repeating the analysis above (PCD admixture fraction into Alaskan malamute and Greenland sledge dogs) using whole genome data. We used only transversions for this analysis (~600K SNPs). We found very little difference in admixture fraction (~7-14%; Table S11) indicating that the ascertainment of the array did not introduce much bias.

We also used D-statistics on genome-wide data to test for admixture from PCD into Arctic breeds since their MRCA. As for the f4 ratio we found that both Alaskan malamute and Greenland sledge dogs have a significant amount of ancestry from PCD (Table S12). We tested whether this signal could be due to admixture from Eurasian dogs into Siberian husky dogs (making derived alleles in Alaskan malamute and Greenland sledge dogs (GSD) match PCD more often; Table S13). We found evidence that the Siberian husky and Alaskan malamute genomes that we analysed here received gene-flow from European dogs (Table S13). However, we found no evidence that GSD received gene-flow since the MRCA of Arctic breeds. This suggests that Eurasian admixture did not affect our result. As stated above, this could also be the result of ancient substructure within Arctic dogs.

We found, however, no signal that either Alaskan malamute or Greenland sledge dogs shared an excess of derived alleles with PCD compared with each other (D(Outgroup, AL3194, Alaskan malamute, Greenland sledge dog)=-0.0001, sd=-0.010). This shows that these dog lineages did not receive additional gene-flow from PCD since their divergence from each other. This result suggests that the signal detected above (excess shared derived alleles between American Arctic dogs and PCD) is due to ancient substructure within Arctic dogs (*165*). More precisely, we hypothesise that the Eurasian Arctic dogs that were recently brought into the Americas, all the way to Greenland, originated from a population that was more closely related to PCD dogs than other Arctic dogs. The high degree of mtDNA divergence within ancient Eurasian Arctic dogs from Zhokhov (~9,000 BP; Figure 1b) suggests that ancient substructure with Arctic dogs is a plausible scenario.

#### TreeMix

In order to test the topology suggested by our phylogenetics and f3 statistics analyses we used Treemix (18) to build a tree with admixture edges. We only used 3 representatives from each major dog group:

- *Western Eurasian dogs* Portuguese village dogs (DEU), German Shepherd (DGS)
- *East Asian dogs* Vietnamese village dogs (DVN) because they lack admixture from European dogs (see above) and Tibetan village dogs (DTI)
- *Pre-contact dogs* (PCD), including both Port au Choix (AL3194) and Weyanoke Old Town (AL3223; Table S1)
- *Arctic dogs* Malamute (DMA) and Greenland dogs (DGL) because they seem least admixed with Western dogs (see above) and
- *CTVT* (79T and 24T)
- *Eurasian wolves* (WEU) from Spain and Portugal
- North American wolves (WAM) from Yellowstone
- *Coyotes* (COY) as an outgroup

We only used transversions in order to limit the effect of DNA damage on the analysis and only used sites that were covered in all samples (~60,000 SNPs).

The results of these analyses are presented in Figure S21, Figure S22, Figure S23 and Figure S24. The placement of the East Asian dog (DVN) population was affected by

adding admixture edges (Figure S22, Figure S23). With two admixture edges DVN outgroup PCD/CTVT and Arctic dogs but has strong admixture into DTI. This support results from our D-statistics analysis (Table S7) and NJ analysis (Figure S10; Figure S11) that suggests that DTI is mixed with European ancestry. We also found evidence for European ancestry in Arctic dogs, supporting our D-statistics analyses (Table S4). Lastly we also found admixture from COY into PCD/CTVT, consistent with D-statistics (Figure S16).

#### qpGraph

We used qpGraph (161) to fit admixture graphs to nine populations representing PCD, CTVT, and each of the three major dog groups, plus wolves and coyotes.

- *Western Eurasian dogs* Portuguese village dogs (DEU)
- *East Asian dogs* Vietnamese village dogs (DVN)
- *Pre-contact dogs* (PCD), including both Port au Choix (AL3194) and Weyanoke Old Town (AL3223; Table S1)
- Canine transmissible venereal tumor (CTVT), including C\_24T, C\_79T and C\_399T
- *Arctic dogs* Alaskan malamute (DMA)
- Eurasian wolves (WEU) from Spain and Portugal
- North American wolves (WAM) from Yellowstone
- Coyotes (COY) from California
- Andean Fox (OUT) as the outgroup

We only used transversions in order to limit the effect of DNA damage on the analysis. This resulted in 600,991 high quality SNPs.

To explore the space of all possible admixture graphs we implemented a heuristic search algorithm. Given an outgroup with which to root the graph, a stepwise addition order algorithm was used for adding leaf nodes to the graph. At each step, insertion of a new node was tested at all branches of the graph, except the outgroup branch. Where a node could not be inserted without producing f4 outliers (i.e.  $|Z| \ge 3$ ) then all possible admixture combinations were also attempted. If a node could not be inserted via either approach, that sub-graph was discarded. If the node was successfully inserted, the remaining nodes were recursively inserted into that graph. All possible starting node orders were attempted to ensure full coverage of the graph space.

As the number of possible graphs grows super-exponentially with each additional leaf node, we initially excluded CTVT from the search space and looked for models with fit the remaining eight populations. We fitted 480,166 unique admixture graphs for these 8 populations and recorded the 892 graphs that left no f4 outliers (i.e. |Z| < 3). We then fitted a further 309,525 unique models, testing all possible insertions of CTVT into the 892 eight-population graphs, and recorded the 1,655 graphs that left no f4 outliers.

Treemix analysis was also performed using the same nine populations, with six admixture edges (the maximum number seen in the qpGraph analyses). We chose the

most plausible qpGraph model (Figure S25) by comparing all fitted models to the Treemix tree with the same sampling (Figure S26), Neighbour joining tree (Figure S10), Bayesian tree (Figure S12) and D-statistics analyses (see above).

#### Phenotypic information

Considering the evidence of introgression between wild North American canids into the pre-contact domestic dog population (Table S3), we assessed the presence of a marker associated with melanism, which has introgressed from dogs into North American gray wolves and coyotes , in the higher coverage PCD genomes (Port du Choix sample: AL3194, ~2x; Weyanoke old town sample: AL3223, ~0.5x). We found no evidence for the CBD103<sup> $\Delta$ G23</sup> / K<sup>B</sup> mutation in either of these samples.

#### CTVT Mutation rate analysis

#### Overall rationale

Our goal was to estimate a lower bound for the CTVT somatic mutation rate and to use this to estimate an upper range for the time at which CTVT originated. To do this, we collected biopsies from a pair of CTVT tumors involved in a naturally occurring direct transmission event, and identified mutations that had arisen during the known transmission time interval to define a somatic mutation rate. We then estimated the number of somatic mutations in the entire CTVT lineage and applied our somatic mutation rate to estimate the time of CTVT origin.

Previous estimates of CTVT time of origin have relied on microsatellite mutation rates in mammalian germlines (*166*, *167*), or mutation rates in human cancer (*20*). These have led to estimates of 250 to 2,500 years since the most recent common ancestor of a group of globally dispersed tumors (*166*), or 6,500 to 65,000 years (*167*) and 10,179-12,873 years (*20*) since the origin of CTVT.

#### *Case histories*

Dog 609 was a mixed-breed free-ranging dog from the Gambia with an approximately 31cm<sup>3</sup> vaginal CTVT tumor. Her ten-month-old male puppy, Dog 608, had several CTVT tumors on the ventral skin. This unusual CTVT presentation in Dog 608 suggested that CTVT cells may have transmitted from mother to puppy during parturition.

#### Samples

This project was approved by the Department of Veterinary Medicine, University of Cambridge, Ethics and Welfare Committee (reference CR174). A 1-2 mm<sup>3</sup> biopsy was sampled from Dog 609's vaginal tumor (609T). A 1-2 mm<sup>3</sup> biopsy was sampled from one of Dog 608's skin tumors on the same day (608T). Biopsies were also collected from host tissues (ovary (609H) or testis (608H)). Genomic DNA was extracted using the Qiagen DNeasy Blood and Tissue extraction kit (Qiagen, Hilden, Germany). CTVT diagnosis was confirmed as previously described (*21*). Whole genome sequencing libraries were prepared with insert size 450 bp and sequenced with 150 bp paired end reads using the Illumina HiSeq X Ten platform (Illumina, San Diego, CA). Reads were aligned to CanFam3.1 using BWA-MEM (*130*). Average sequencing depth is reported in Table S2.

#### Variant Calling

#### Variant extraction and filtering

We used Somatypus (https://github.com/baezortega/somatypus), a Platypus (168) based variant calling and genotyping pipeline, to identify SNVs and small insertions and deletions (indels). In order to make an initial call, SNVs were required to have  $\geq$ 3 supporting reads in at least one of the four sequenced samples (608T, 608H, 609T, 609H). Indels were inputted to GATK Realigner Target Creator (169) for local realignment and SNVs were re-called from realigned genomes.

The following in-built Platypus flags were used to exclude SNVs at two stages, before and after genotyping: badReads, MQ, QD, strandBias, SC.

The following post-processing filters were also implemented:

- *Strand bias filter*. For each SNV, the total coverage, as well as forward and reverse strand read support were extracted. For low total coverage positions (≤10 reads across all four samples), we discarded calls with less than two supporting reads in either forward or reverse direction. For high total coverage positions (>10 reads across all four samples), we discarded calls with less than 20% support on either the forward or reverse sequencing strands.
- *Simple repeat filter*. SNVs within simple repeats, as defined by the UCSC table browser (CanFam3.1), were excluded.
- *Extreme depth filter*. SNVs within regions of high read depth were also excluded. To detect high read depth (HRD) regions we first generated BigWig coverage files from matched normal whole genome sequence data files (608H, 609H). We then identified areas with coverage 12 standard deviations higher than the mean read coverage, on a chromosome by chromosome basis. Common intervals between normal samples were identified using bedtools multiinter (*138*) and were merged using bedtools merge. The maximum allowed distance between regions to be merged was 250 bp. HRD regions spanning less than 500 bp were excluded. Any HRD region that overlapped with gene regions as defined by the UCSC table browser (CanFam3.1, Genes and Gene Predictions, Ensembl Genes) was excluded.
- *Low VAF filter*. SNVs with VAF>0 and VAF<0.2 in both 608H and 609H were discarded if (i) they were not detected in 608T or 609T or (ii) they were found with VAF>0 and VAF<0.1 in either or both of 608T and 609T.
- *Regions filter*. SNVs occurring in the mitochondrial genome or on unassigned scaffolds were excluded. In addition, to avoid problems caused by variable coverage in hosts, SNV analysis was restricted to autosomes.

#### Germline and consensus filtering

SNVs identified in 608T and 609T will belong to one or more of the following categories:

- (i) Contaminating germline SNVs from matched host
- (ii) Germline SNVs inherited by the CTVT founder dog
- (iii) Somatic mutation SNVs

In order to enrich for somatic mutations, we filtered our candidate SNVs against a panel of 28,812,954 canid germline SNVs. Specifically, we excluded any genomic site that was reported in any of the following variant catalogues:

• 608H and 609H (sites with  $\geq$ 5 reads coverage and  $\geq$ 2 reads supporting a non-reference allele were considered SNVs)

- The Variant and Systematic Error Catalogue (VSEC) (170)
- The CanineHD 170K SNP array (171)

• The ascertainment panel generated in this study prior to incorporating CTVT samples (*Genotyping - Ascertainment panel*)

• A complete genome from a Greenland sledge dog (14)(Table S2) included in the ascertainment panel was additionally genotyped. This provided additional SNVs beyond those in the ascertainment panel, as the ascertainment panel excluded SNVs that were found on only one chromosome; thus SNVs that were exclusively found in the Greenland sledge dog individual and CTVT would not have been included in the ascertainment panel that we filtered against (see previous bullet point), but were excluded in this step (*Genotyping - Dog samples*)(14)

Next, we further filtered the remaining SNVs using the following criteria:

• We retained only those SNVs that had that had  $\geq 2$  reads supporting the variant, all with minimum base quality of 20 and minimum mapping quality of 35, in at least one of the two tumors using the alleleCount package (http://cancerit.github.io/alleleCount/).

• We retained only those SNVs that were identified by GATK Haplotype Caller. The GATK engine was restricted processing candidate loci.

• We required that the matched host must have coverage of at least 20 reads total at the candidate SNV position. Candidate SNVs that did not reach this threshold in one or both matched hosts were discarded.

• We discarded SNVs where one or both matched hosts had  $\geq 10$  reads total (regardless of whether they supported the variant) with base quality <20 and mapping quality <35 in matched hosts

1,934,103 and 1,934,125 "tumor-only" SNVs remained in 608T and 609T respectively after these steps; 1,933,897 of these were shared by 608T and 609T. Of the SNVs in this set that mapped to genomic regions retaining both parental copies, almost all SNVs were heterozygous. Thus, the majority of these SNVs are likely to be somatic; however, some germline variation that was present in the CTVT founder dog, but that is not represented in the germline panel used here likely still remains. It is also likely that some somatic mutations, which occurred in the same sites as germline SNVs represented in our panel, have been removed.

#### Tumor-unique SNVs

We next filtered tumor-only SNVs, as defined above, for those unique to either 608T or 609T. In order to be considered unique to a single tumor, a variant was required to be present with  $\geq 2$  supporting reads in only one tumor, with minimum base quality of 20 and minimum mapping quality of 35 for those reads supporting the variant.

This method yielded 206 tumor-unique SNVs in 608T and 228 tumor-unique SNVs in 609T.

#### Mutational spectrum

Each tumor-only SNV was classified as one of six possible mutation types in the pyrimidine context (C>A, C>G, C>T, T>A, T>C, T>G). The immediate 5' and 3' sequence contexts for each mutation was extracted from the CanFam3.1 dog reference genome (*172*) yielding 96 mutation types. The mutational spectrum for the 1,933897, CTVT tumor-only SNVs (shared between 608T and 609T) is shown in Figure S27A.

#### Mutational signature fitting

We performed mutational signature fitting in order to estimate the number of mutations contributed by different exposures to the CTVT mutational spectrum.

Validated mutational signatures were obtained from the Catalogue of Somatic Mutations in Cancer (COSMIC; http://cancer.sanger.ac.uk/cosmic/signatures) database and renormalized to the CanFam3.1 dog reference genome (172). In addition, we generated a "Dog Germline" signature, from the germline mutational spectrum of a Greenland sledge dog (*Variant calling - Germline and consensus filtering*; Table S2).

Reference (20) previously showed that COSMIC mutational signature 1 (5-methylcytosine deamination), signature 5 (unknown etiology), and signature 7 (ultraviolet light exposure) are operative in CTVT, and that these three signatures are sufficient to describe the pattern of somatic substitutions observed in CTVT. We therefore fitted these three signatures, together with the Dog Germline signature (see above), to the CTVT tumoronly mutation spectrum (Table S14; Figure S27B and C). Results are similar to previous findings (20). Signatures were fitted using sigfit (https://github.com/kgori/sigfit). Simulations were run using 100 chains with 10,000 iterations each. Importantly, the Dog Germline signature accounted for only 5.5% of the tumor-only SNVs, suggesting that the majority of the SNVs in this set are indeed somatic.

#### N[C>T]G CTVT tumor-only SNVs

Mutational signature 1 is largely composed of 5'-N[C>T]G-3' mutations (where N is any base) (http://cancer.sanger.ac.uk/cosmic/signatures). Of the 1,933,897 tumor-only SNVs shared by 608T and 609T, 222,072 are N[C>T]G.

#### Copy-number analysis

Average mappability and GC-content were generated for the dog reference genome (172) with the generateMap, mapCounter, and gcCounter tools in the HMMcopy package (173). GC content and genomic mappability biases for read counts in non-overlapping 1kb windows were corrected using HMMcopy. Copy number estimation was then performed on GC- and mappability-corrected read counts using a bespoke copy number calling pipeline (https://github.com/ymk1/cnv\_pipeline.git)

Tumor purity in 608T and 609T was evaluated based on the VAF distribution of tumor-only SNVs. Tumor purity was estimated as follows:

#### Purity=2\*VAFmed

Where  $VAF_{med}$  is the median VAF value of tumor-only SNVs. Using this method, 608T was estimated to be 49.3% CTVT cells and 609T was estimated to be 65% CTVT cells.

#### *Identifying clonal mutations in tumor-unique variant sets*

We categorised tumor-unique SNVs in 608T and 609T as either clonal or subclonal, that is, present in all or a fraction of tumor cells within a sample, respectively. To do this, we first examined the VAF distributions of germline SNPs in 608T and 609T for each copy number (CN) state (CN1, CN2, CN3, CN4, CN6). We used a Gaussian mixture model (k = 2), implemented using the R package MCLUST (*174*), to define VAF clusters for heterozygous and homozygous SNPs. Next, we fitted this model to VAF distributions of tumor-unique SNVs. SNVs that fell below the 5% lower bound were defined as subclonal; all other SNVs were considered clonal. The results of this analysis are shown in Table S15.

#### *N*[*C*>*T*]*G CTVT tumor-unique clonal SNVs*

Of the 183 and 174 clonal mutations identified uniquely in 608T and 609T respectively, 27 and 23 were N[C>T]G in 608T and 609T respectively. 26/27 and 21/23 (609T) were validated using read alignment visualisation.

#### *CTVT* mutation rate

We have determined that 608T and 609T acquired 183 and 174 clonal mutations, and 27 and 23 clonal N[C>T]G mutations respectively since they diverged from their most recent common ancestor (MRCA).

In order to estimate the CTVT mutation rate, we need to know the time intervals during which the clonal tumor-unique mutations arose in 608T and 609T. These time intervals ( $i_{608T}$  and  $i_{609T}$ ) correspond to:

 $i_{608T} = t_{MRCA-608T} - t_{MRCA-608T/609T}$  and  $i_{609T} = t_{MRCA-609T} - t_{MRCA-608T/609T}$ 

where  $t_{MRCA-608T}$  and  $t_{MRCA-609}$  are time-points defining the MRCA cells of 608T and 609T respectively, and  $t_{MRCA-608T/609T}$  is the time-point defining the MRCA cell of both 608T and 609T.

We assumed that  $t_{MRCA-608T/609T}$  occurred during 609T tumor development; i.e. the clones that spawned the sampled 608T and 609T biopsies diverged in the period after infection of Dog 609 (the mother) but before transmission to Dog 608 (the son). Thus, the earliest time-point for  $t_{MRCA-608T/609T}$  would coincide with the time at which Dog 609 (the mother) was infected with CTVT, i.e. month 0.

We assumed that Dog 609 was infected during the heat cycle in which she conceived the puppy, Dog 608. Although we cannot be certain that this assumption is

valid, we observed that Dog 609's tumor appeared to be of a similar size to Dog 608's tumor. Unless CTVT tumors have large variation in growth rate, we believe that it is unlikely that Dog 609 was infected with CTVT in the heat cycle previous to that in which she conceived Dog 608. Given that the gestation period in domestic dogs can range from 57-72 days (175-177), we estimated that Dog 609 was infected with CTVT approximately 2 months prior to when her son, Dog 608, was born and infected. This implies that the latest time-point for  $t_{MRCA-608T/609T}$  is 2 months after Dog 609 was infected with CTVT. Thus  $t_{MRCA-608T/609T} = 0$  to 2 months, defining month 0 as the month at which Dog 609 (the mother) was infected with CTVT.

Assuming no polyclonal seeding,  $t_{MRCA-608T}$  either occurred within Dog 608, or was the cell that transmitted from Dog 609 to Dog 608, and thus  $t_{MRCA-608T} = 2$  to 12 months, where month 0 is the month at which Dog 609 (the mother) was infected with CTVT, and 12 months corresponds to the time of sampling.

 $t_{MRCA-609T}$  could have occurred at any time during 609T tumor development. Thus  $t_{MRCA-609T} = 0$  to 12 months months, where month 0 is the month at which Dog 609 (the mother) was infected with CTVT, and 12 months corresponds to the time of sampling.

Thus, we estimate both  $i_{608}$  and  $i_{609}$  to be 0 to 12 months.

We determined that 608T and 609T had acquired 183 and 174 mutations since their divergence from their MRCA (MRCA<sub>608T-609T</sub>) and before the MRCA of the clone biopsied in 608T (MRCA<sub>608T</sub>) and the MRCA and the clone biopsied in 609T (MRCA<sub>609T</sub>). These mutations will likely have arisen as part of clock-like ageing-associated mutational signatures 1 and 5 and possibly as part of mutational signature 7 (exposure to ultraviolet light). Signature 1 mutation rate is believed to be highly dependent on cell division (*178*). Due to the small number of tumor-unique mutations, signature fitting cannot give us an accurate estimate of the respective contributions of these signatures to tumor-unique SNV sets. As mutational signature 1 is largely composed of N[C>T]G mutations (where N is any base), we used N[C>T]G as a proxy for signature 1. Table S16 shows the number of N[C>T]G mutations unique to 608T and 609T, as well as the number found in the somatic lineage from the CTVT founder dog until MRCA<sub>608T-609T</sub>.

As 608T harbours more clonal N[C>T]G mutations than 609T (Table S16), we infer that MRCA<sub>608T</sub> existed more recently than MRCA<sub>609T</sub>. Assuming that  $i_{608T}$  is up to 12 months (see above), then the slowest rate at which N[C>T]G mutations accumulate is 27 N[C>T]G mutations / year or 12.56 N[C>T]G mutations / Gigabase (Gb) / year, using the callable dog genome size (excluding simple repeats and regions of high read depth) of 2.15 Gb (*172*). Applying this rate to the whole lineage (222,072 N[C>T]G mutations across the callable genome ~ 2.15 Gigabase pairs, see above), we obtain an upper bound of 8,225 years for the origin of CTVT. Our method cannot directly infer an upper bound for the CTVT mutation rate, and hence a lower bound for the age of CTVT. However, assuming that the disease described by Blaine in 1810 (*179*) was indeed CTVT, then CTVT must have arisen at least 200 years ago.

#### Comparison with mutation rates in human cancer

Signature 1 mutation rate varies between human cancer tissue types(*178*). The mutation rate lower bound that we have derived for CTVT N[C>T]G mutations (>12.56 mutations / Gb / year) is comparable to the N[C>T]G mutation rates found in human cancers(*178*). Cervical cancer was reported to have the highest estimated rate of accumulation of N[C>T]G somatic mutations of 36 human cancer types (16.61 N[C>T]G somatic mutations / Gb / year)(*178*). Applying the cervical cancer N[C>T]G mutation rate to the CTVT lineage would provide an estimate of 6,195 years since CTVT origin.

#### *Limitations of the approach*

The approach to deriving the CTVT mutation rate that we have presented here is based on a number of assumptions. These are outlined below.

• *No polyclonal seeding.* We have discounted the possibility of polyclonal seeding of Dog 609's tumor. If polyclonal seeding occurred, then  $t_{MRCA-608T/609T}$  may have existed in Dog 609 tumor's donor, rather than in Dog 609's tumor itself. If this is the case, then the mutation rate would be slower, leading to older estimates for CTVT time-of-origin.

• Age of Dog 608. Dog 608 was estimated to be ten months old at the time of sampling. Given that Dog 609 was approximately 14 days pregnant at the time of sampling, and given that dogs have heat cycles every six months, we believe that Dog 608's age estimate is likely accurate. However, an inaccurate age estimate could affect our mutation rate estimates.

• *Time of infection of Dog 609.* We assumed that Dog 609 (the mother) was infected with CTVT at the time of the heat cycle during which she conceived Dog 608. However, if Dog 609 was in fact infected with CTVT during a previous heat cycle, then  $t_{MRCA-608T/609T}$  may have existed at an earlier time point: this would lead to estimation of a slower mutation rate and older estimates for CTVT time-of-origin.

• *Estimation of total mutation burden.* We estimated the total mutation burden by filtering against a large panel of variation in normal dogs (see *Germline and consensus filtering*). If our set of tumor-only SNVs (see section *tumor-unique SNVs* above) is substantially over-filtered (i.e. somatic mutations were removed as they occur at the same site as germline SNVs), then CTVT could have arisen earlier than our estimates suggest. If, on the other hand, substantial numbers of germline SNVs remain in the set of total mutations, then we may have over-estimated the time of CTVT origin.

• *Back-mutation*. We discounted back-mutation as a significant factor in our estimates.

• *Mutation false discovery rate.* The expected proportion of false positive SNVs should be the same in the tumor-unique and tumor-only variant sets as we have used the same filters in both cases. Thus, this is unlikely to have substantially affected estimates.

• *Mutation opportunity and variable mutation rate*. We have discounted the effects of variable mutation opportunity from our estimates. Mutation opportunity may change over time due to (i) altered DNA methylation or chromatin states; (ii) decrease in number of available NCG sites over time; (iii) copy number alterations. Furthermore, although N[C>T]G is usually considered constant, a recent study has detected germline mutations in the *MBD4* gene which alter the rate of signature 1 mutation accumulation

(180). As we detected a number of mutations in MBD4, we cannot exclude the possibility that somatic alteration of this or other loci in CTVT may have caused variation in the rate of N[C>T]G mutation accumulation.

• *Sampling error. We* based our estimates of the CTVT mutation rate on one observation and did not account for sampling variation. Future studies can address this by measuring mutations in additional CTVT time intervals.

Despite these limitations, our analysis provides a plausible estimate of CTVT somatic mutation rate, and is comparable with clock-like mutation rates observed in some human cancers (*178*).



#### **Fig. S1.**

Per library C to T (red) and G to A (blue) frequency of mis-incorporation at 3' and 5' end of read for samples used in nuclear genome analyses.



## Fig. S2

Per library C to T (red) and G to A (blue) frequency of mis-incorporation at 3' and 5' end of read for samples used in mtDNA analyses. Lack of 5' damage in some libraries is due to library preparation protocol (see Ancient DNA - Illinois section).



Fig. S2 (continued)

Per library C to T (red) and G to A (blue) frequency of mis-incorporation at 3' and 5' end of read for samples used in mtDNA analyses. Lack of 5' damage in some libraries is due to library preparation protocol (see Ancient DNA - Illinois section).


# Fig. S2 (continued)

Per library C to T (red) and G to A (blue) frequency of mis-incorporation at 3' and 5' end of read for samples used in mtDNA analyses. Lack of 5' damage in some libraries is due to library preparation protocol (see Ancient DNA - Illinois section).



Maximum likelihood tree based on mtDNA data. The four major dog haplogroups are indicated: A (red; includes all but one pre-contact dogs), B (purple), C (yellow), D (green). Blue tip label represent newly sequenced samples (this study). Dark blue highlighted clade represents American dogs (monophyletic, bootstrap support value=87). Light blue highlighted clades (CGG1-10) represent Zhokhov Island samples (~9Kya sled dogs from Eastern Siberia; see Table S1). CGG10-11 (outside of the Zhokhov / pre-contact clade) are more recent sled dogs from Siberia (~1.5kya; Table S1). Node labels indicate bootstrap replicates.



Maximum likelihood tree based on 605bp of the control region. All samples starting with prefix HQ were obtained from (5) while all samples starting with prefix AY were obtained from (8). The four major dog haplogroups are indicated with different branch colours: A (red; includes all but one pre-contact dogs), B (purple), C (yellow), D (green). All pre-contact dogs from this study are highlighted in light blue. Node labels indicate bootstrap replicates.



Maximum likelihood tree based on mtDNA data including data from (150–155). Red branches represent ancient pre-contact dogs. Node labels indicate bootstrap replicates.



Bayesian tree (*BEAST*) of mtDNA data. Red, purple and green circle represent nodes with >0.9, >0.7 and >0.5 posterior probability respectively. Blue bar represent confidence interval of divergence time (scaled in year before present).



Principal Components Analysis (PC1 versus PC2) of 57 canid samples (including wolves and coyotes) based on 2,063,129 SNPs ascertained using the genome-wide data-set. All pre-contact dog samples were projected.



Principal Components Analysis (PC1 versus PC2) of 44 dog samples (excluding wolves and coyotes) based on 2,063,129 SNPs ascertained using the genome-wide data-set. All pre-contact dog samples were projected.



Principal Components Analysis (PC1 versus PC2) of 44 dog samples (excluding wolves and coyotes) based on 2,063,129 SNPs ascertained using the genome-wide data-set. All pre-contact dog and CTVT samples were projected.



Neighbour Joining (NJ) tree based on Identity By State (IBS). This figure is the same as in Figure 1c. Confirms that CTVT is more closely related to pre-contact dogs than any other dog population. Confirms that PCD form a monophyletic clade.



Neighbour Joining (NJ) tree based on Identity By State (IBS). Same as Figure S3 but without East Asian dogs that are admixed with European dogs i.e. excluding all East Asian dogs except Vietnamese. PCD, Arctic dogs and CTVT founder now appear more closely related to Western dogs.



Bayesian tree based on ~26K transversions. Confirms that CTVT and PCD are monophyletic with high support and supports the basal placement of the CTVT/PCD clade. Support values represent posterior probability.



Shared genetic drift measured by *f*<sub>2</sub>(Outgroup; Y, X) where Y is either Port au Choix dog (AL3194), Weyanoke Old town dog (AL3223), Alaskan Malamute (D\_Mal68), Greenland sledge dog (D\_Green) and X represents modern dog populations. Error bars represent 1 SE.



Box plot representing **a**. D-statistics and **b**. significance of D-statistics (Z) for every combination of D(Outgroup, AL3194[Port au Choix], Pop3, Pop4), where Pop3 is fixed and Pop4 represents any other genome. Positive values support a close relationship between Pop3 and PCD while negative values imply PCD are closer to other dog populations. If Pop3 is not admixed with PCD, we expect -4<Z<4 (x-axis).



Box plot representing **a.** D-statistics and **b.** significance of D-statistics (Z) for every combination of D(Outgroup, AL3223[Weyanoke Old town], Pop3, Pop4), where Pop3 is fixed and Pop4 represents any other genome. Positive values support a close relationship between Pop3 and PCD while negative values imply PCD are closer to other dog populations. If Pop3 is not admixed with PCD, we expect -4 < Z < 4 (x-axis).



Box plot representing **a**. D-statistics and **b**. significance of D-statistics (Z) for every combination of D(Outgroup, Coyote, Pop3, Pop4), where Pop3 is fixed (x-axis) and Pop4 represents any other genome. Positive values support a close relationship between Pop3 and coyotes while negative values imply coyotes are closer to other dog populations. If Pop3 is not admixed with coyotes, we expect -4 < Z < 4 (x-axis).



Box plot representing **a**. D-statistics **b**. significance of D-statistics (Z) for every combination of D(Outgroup, north American wolf, Pop3, Pop4), where Pop3 is fixed and Pop4 represents any other genome. Positive values support a close relationship between Pop3 and north American wolves while negative values imply NA wolves are closer to other dog populations. If Pop3 is not admixed with north American wolves, we expect -4 < Z < 4 (x-axis).



Schematic representation of the assumed phylogeny for the f4 ratio test used to estimate pre-contact ancestry into modern North American dogs. Alpha represents the degree of ancestry from pre-contact dogs.







ADMIXTURE results based on SNP array data for K=4, 10 and 15. Population code: AED=American Eskimo Dog, AM=Alaskan Malamute, APBT=American Pit Bull Terrier, AST=American Staffordshire Terrier, BEA=Beagle, BOX=Boxer, CBR=Chesapeake Bay Retriever, CC=Chow Chow, CD=Carolina Dog, CHI=Chihuahua, CLD=Catahoula Leopard Dog, COO=Chinook, CSP=Chinese Shar-pei, CTVT=CTVT, DAL=Alaskan Husky, DCH=Chinese Village Dog, DEU=European Village Dog, DGL=Greenland Sledge Dog, DHU=Husky, DLB=Lebanese Village Dog, DMA=Malamute, DME=Mexican Hairless Dog, DPU=Peruvian Hairless Dog, DSL=Siberian Laika, EUR=Eurasier, FS=Finnish Spitz, GSD=Greenland Sledge Dog, NEW=Newfoundland, NSDTR=Nova Scotia Duck Tolling Retriever, PCD=Pre-Colombian Dogs, PIO=Peruvian Inca Orchid, SAM=Samoyed, SH=Siberian Husky, VDB=Village Dog Belize, VDB2=Village Dog Brazil, VDC=Village Dog Colombia, VDCR=Village Dog Costa Rica, VDDR=Village Dog Dominican Republic, VDH=Village Dog Honduras, VDP=Village Dog Panama, VDPA=Village Dog Peru-Arequipa, VDPC=Village Dog Peru-Cusco, VDPI=Village Dog Peru-Ica, VDPL=Village Dog Peru-Loreto, VDPP=Village Dog Peru-Puno, VDPR=Village Dog Puerto Rico, VDUA=Village Dog US-Alaska, XOL=Xoloitzcuintli (see Table S9 for more information).



Admixture graph without migration edge and matrix of residuals, expressed as the number of standard errors, inferred using *TreeMix* (based on transversions). Western Eurasian dogs - Portuguese village dogs (DEU), German Shepherd (DGS), East Asian dogs - Vietnamese village dogs (DVN) and Tibetan village dogs (DTI), Pre-contact dogs (PCD), including both Port au Choix (AL3194) and Weyanoke Old Town (AL3223), Arctic dogs - Malamute (DMA) and Greenland dogs (DGL), CTVT - (79T and 24T), Eurasian wolves (WEU) from Spain and Portugal, North American wolves (WAM) from Yellowstone, Coyotes (COY) as an outgroup.



Admixture graph with a single migration edge and matrix of residuals, expressed as the number of standard errors, inferred using TreeMix (based on transversions). We see evidence for admixture from coyotes (COY) into the pre-contact dog lineage (PCD/CTVT), consistent with Figure S16. Western Eurasian dogs - Portuguese village dogs (DEU), German Shepherd (DGS), East Asian dogs - Vietnamese village dogs (DVN) and Tibetan village dogs (DTI), Pre-contact dogs (PCD), including both Port au Choix (AL3194) and Weyanoke Old Town (AL3223), Arctic dogs - Malamute (DMA) and Greenland dogs (DGL), CTVT - (79T and 24T), Eurasian wolves (WEU) from Spain and Portugal, North American wolves (WAM) from Yellowstone, Coyotes (COY) as an outgroup.



Admixture graph with two migration edges and matrix of residuals, expressed as the number of standard errors, inferred using TreeMix (based on transversions). Western Eurasian dogs - Portuguese village dogs (DEU), German Shepherd (DGS), East Asian dogs - Vietnamese village dogs (DVN) and Tibetan village dogs (DTI), Pre-contact dogs (PCD), including both Port au Choix (AL3194) and Weyanoke Old Town (AL3223), Arctic dogs - Malamute (DMA) and Greenland dogs (DGL), CTVT - (79T and 24T), Eurasian wolves (WEU) from Spain and Portugal, North American wolves (WAM) from Yellowstone, Coyotes (COY) as an outgroup.



Admixture graph with three migration edges and matrix of residuals, expressed as the number of standard errors, inferred using TreeMix (based on transversions). Western Eurasian dogs - Portuguese village dogs (DEU), German Shepherd (DGS), East Asian dogs - Vietnamese village dogs (DVN) and Tibetan village dogs (DTI), Pre-contact dogs (PCD), including both Port au Choix (AL3194) and Weyanoke Old Town (AL3223), Arctic dogs - Malamute (DMA) and Greenland dogs (DGL), CTVT - (79T and 24T), Eurasian wolves (WEU) from Spain and Portugal, North American wolves (WAM) from Yellowstone, Coyotes (COY) as an outgroup.



Qpgraph model with admixture fractions. *Western Eurasian dogs* - Portuguese village dogs (DEU), *East Asian dogs* - Vietnamese village dogs (DVN), *Pre-contact dogs* (PCD), including both Port au Choix (AL3194) and Weyanoke Old Town (AL3223), *Arctic dogs* - Malamute (DMA), *CTVT* - (79T and 24T), *Eurasian wolves* (WEU) from Spain and Portugal, *North American wolves* (WAM) from Yellowstone, *Coyotes* (COY) as an outgroup.



Admixture graph and matrix of residuals, expressed as the number of standard errors, inferred using TreeMix for the same population as in Figure S25. *Western Eurasian dogs* - Portuguese village dogs (DEU), *East Asian dogs* - Vietnamese village dogs (DVN), *Pre-contact dogs* (PCD), including both Port au Choix (AL3194) and Weyanoke Old Town (AL3223), *Arctic dogs* - Malamute (DMA), *CTVT* - (79T and 24T), *Eurasian wolves* (WEU) from Spain and Portugal, *North American wolves* (WAM) from Yellowstone, *Coyotes* (COY) as an outgroup.



**A**. CTVT mutation spectrum. 1,933,897 tumor-only mutations in CTVT are displayed by mutation type (in pyrimidine context) with immediate 5' and 3' context. Each of the 96 mutation classes is displayed on the horizontal axis. Mutation proportions are displayed relative to CanFam3.1 **B**. Fraction of CTVT tumor-only mutations attributable to COSMIC Signatures 1, 5, 7, and the dog germline signature, as estimated using sigfit. **C**.Reconstruction of CTVT tumor-only spectrum using COSMIC signatures 1,5 and 7 and the dog germline signature.

### Table S1.

Information about samples ancient samples sequenced in this study including provenance, age (radiocarbon or stratigraphic information), and sequencing statistics (endogeneous content etc.). N\_SNP\_call corresponds to the number of sites from our ~6M sites that were call in each sample. Analysed Nuclear and mtDNA Analysed columns corresponds to sample that were (1) and were not (0) analysed for nuclear and mtDNA analyses respectively. mtDNA capture column indicates whether mtDNA was enriched using target capture (0, no; 1, yes).

## Table S2.

Table containing information (coverage, accession etc.) of modern whole genomes used in this study.

# Table S3.

Table containing information (coverage, accession etc.) of modern whole genomes used in this study.

Pop 2	Pop 3	Pop 4	D	Z	BABA	ABBA	n
C_Cal	AL3194	AL3223	0.0152	1.460	5,681	5,511	285,638
C_Cal	AL3223	AL3194	-0.0152	-1.460	5,511	5,681	285,638
C_MidW	AL3194	AL3223	0.0035	0.277	3,443	3,419	181,671
C_MidW	AL3223	AL3194	-0.0035	-0.277	3,419	3,443	181,671
W_Mex1	AL3194	AL3223	-0.0130	-1.086	7,394	7,588	282,977
W_Mex1	AL3223	AL3194	0.0130	1.086	7,588	7,394	282,977
W_Yellow1	AL3194	AL3223	-0.0083	-0.785	7,501	7,626	281,684
W_Yellow1	AL3223	AL3194	0.0083	0.785	7,626	7,501	281,684
W_Yellow2	AL3194	AL3223	-0.0067	-0.622	7,513	7,615	282,189
W_Yellow2	AL3223	AL3194	0.0067	0.622	7,615	7,513	282,189
TAI	AL3194	AL3223	-0.0001	-0.005	3,026	3,026	116,212
TAI	AL3223	AL3194	0.0001	0.005	3,026	3,026	116,212

#### Table S4.

D-statistics for D(Outgroup, Asian or European dogs, PCD or CTVT, Arctic dogs). The Andean Fox was used as an outgroup for these analyses. *n* corresponds to the number of SNPs where all populations have data. Standard error for these statistics was obtained by performing a weighted block jackknife over 1Mb blocks. Statistics with Z>3 and Z<-3 are shown in bold.

Pop 2	Pop 3	Pop 4	D	Z	BABA	ABBA	n	
D_China8	AL3194	D_AHusky91	0.0752	7.65	47,384	40,759	879,166	
D_China8	AL3194	D_Green	0.0606	6.01	47,406	41,994	920,449	
D_China8	AL3194	D_Husky	0.0693	7.253	51,412	44,755	936,527	
D_China8	AL3194	D_Husky89	0.0825	8.407	41,900	35,513	768,218	
D_China8	AL3194	D_Mal68	0.0655	6.538	46,698	40,958	907,759	
D_China8	C_24T	D_AHusky91	0.0907	9.461	61,440	51,230	1,115,093	
D_China8	C_24T	D_Green	0.0753	7.523	61,012	52,472	1,167,919	
D_China8	C_24T	D_Husky	0.0819	8.781	66,255	56,230	1,189,650	
D_China8	C_24T	D_Husky89	0.0987	9.798	54,693	44,866	985,295	
D_China8	C_24T	D_Mal68	0.0796	8.128	60,682	51,739	1,151,059	
D_China9	AL3194	D_AHusky91	0.064	6.644	45,860	40,348	849971	
D_China9	AL3194	D_Green	0.0812	8.035	47,827	40,641	893422	
D_China9	AL3194	D_Husky	0.0684	7.323	50,481	44,022	905549	
D_China9	AL3194	D_Husky89	0.087	8.783	41,939	35,228	760651	
D_China9	AL3194	D_Mal68	0.065	6.504	45,784	40,195	877630	
D_China9	C_24T	D_AHusky91	0.0776	7.936	59,705	51,110	1,078553	
D_China9	C_24T	D_Green	0.0926	9.696	61,461	51,046	1,133912	

D_China9	C_24T	D_Husky	0.0799	8.709	65,471	55,786	1,150989
D_China9	C_24T	D_Husky89	0.1001	9.815	54,894	44,904	974829
D_China9	C_24T	D_Mal68	0.0743	7.814	59,571	51,330	1,113510
D_Port61	AL3194	D_AHusky91	0.0981	9.344	49,646	40,780	897394
D_Port61	AL3194	D_Green	0.0733	7.059	49,098	42,391	939,704
D_Port61	AL3194	D_Husky	0.0994	10.286	54,206	44,402	956,025
D_Port61	AL3194	D_Husky89	0.1006	9.791	43,556	35,593	784,550
D_Port61	AL3194	D_Mal68	0.1044	9.88	49,860	40,438	926,900
D_Port61	C_24T	D_AHusky91	0.1115	10.706	64,216	51,337	1,137,495
D_Port61	C_24T	D_Green	0.088	8.191	63,226	53,005	1,191,621
D_Port61	C_24T	D_Husky	0.1125	11.651	70,044	55,878	1,213,643
D_Port61	C_24T	D_Husky89	0.1181	11.113	57,125	45,064	1,005,524
D_Port61	C_24T	D_Mal68	0.1167	11.029	64,747	51,223	1,174,636
D_Portt71	AL3194	D_AHusky91	0.0949	9.146	48,273	39,904	877,094
D_Portt71	AL3194	D_Green	0.0814	8.348	48,157	40,909	918,351
D_Portt71	AL3194	D_Husky	0.0967	10.277	52,649	43,367	934,632
D_Portt71	AL3194	D_Husky89	0.1035	10.427	42,550	34,572	765,611
D_Portt71	AL3194	D_Mal68	0.1047	10.09	48,497	39,308	906,275
D_Portt71	C_24T	D_AHusky91	0.1122	10.895	62,687	50,038	1,112,724
D_Portt71	C_24T	D_Green	0.0994	9.888	62,205	50,964	1,165,580
D_Portt71	C_24T	D_Husky	0.1139	12.371	68,354	54,374	1,187,444
D_Portt71	C_24T	D_Husky89	0.124	11.829	55,911	43,573	982,083
D_Portt71	C_24T	D_Mal68	0.1214	11.729	63,196	49,518	1,149,527

#### Table S5.

D-statistics for D(Outgroup, Taimyr, PCD/Arctic dogs, European, Asian, or Arctic dogs). The Andean Fox was used as an outgroup for these analyses. *n* corresponds to the number of SNPs where all populations have data. Standard error for these statistics was obtained by performing a weighted block jackknife over 1Mb blocks. Statistics with Z>3 and Z<-3 are shown in bold.

Pop 2	Pop 3	Pop 4	D	Z	BABA	ABBA	n
TAI	AL3194	D_India168	-0.0274	-3.377	13,074	13,812	297,919
TAI	AL3194	D_Peru	-0.0267	-3.293	17,139	18,079	384,525
TAI	AL3194	D_India60	-0.0247	-3.254	17,383	18,264	372,784
TAI	AL3194	D_Na8	-0.0256	-3.199	16,614	17,489	366,726
TAI	D_Mal68	D_Na8	-0.0206	-3.127	19,530	20,350	459,311
TAI	AL3194	D_Port61	-0.0236	-3.006	17,398	18,239	387,777
TAI	D_Mal68	D_Peru	-0.0195	-2.856	20,182	20,983	482,639
TAI	D_Mal68	D_India60	-0.0189	-2.789	20,930	21,735	465,930
TAI	D_Mal68	D_Port61	-0.0204	-2.768	20,358	21,206	485,553
TAI	AL3194	D_Viet59	-0.0227	-2.753	13,212	13,824	308,811
TAI	D_Green	D_Na8	-0.0202	-2.674	20,383	21,224	465,676
TAI	D_Husky89	D_India168	-0.0205	-2.664	13,042	13,588	314,547
TAI	D_Green	D_India60	-0.0198	-2.607	21,505	22,372	472,462
TAI	D_Husky89	D_Port61	-0.0215	-2.571	17,284	18,043	413,104
TAI	D_Husky89	D_Peru	-0.0201	-2.525	17,191	17,896	410,031
TAI	D_Green	D_Port61	-0.0197	-2.506	21,204	22,058	492,659
TAI	AL3194	D_Husky	-0.0199	-2.498	16,444	17,111	394,720
TAI	AL3194	D_SLaika	-0.0194	-2.485	16,821	17,487	379,149

TAI	D_Green	D_India168	-0.0178	-2.481	15,728	16,296	377,110
TAI	D_Mal68	D_Na89	-0.0195	-2.481	12,195	12,680	289,545
TAI	D_Green	D_Peru	-0.0194	-2.455	20,782	21,605	488,730
TAI	D_Husky89	D_India60	-0.0186	-2.44	17,456	18,117	394,520
TAI	AL3194	D_Mex	-0.0193	-2.426	17,007	17,675	380,509
TAI	AL3194	D_Viet21	-0.0204	-2.419	13,088	13,634	303,723
TAI	AL3194	D_Tibet3	-0.0184	-2.41	16,776	17,403	375,015
TAI	AL3194	D_Portt71	-0.0185	-2.366	16,997	17,637	379,175

#### Table S6.

D-statistics for D(Outgroup, Taimyr, PCD or Arctic dogs, CTVT/Arctic dogs or PCD). The Andean Fox was used as an outgroup for these analyses. *n* corresponds to the number of SNPs where all populations have data. Standard error for these statistics was obtained by performing a weighted block jackknife over 1Mb blocks. Statistics indicate that there has been no extra admixture from the Taimyr wolf into other populations.

Pop 2	Pop 3	Pop 4	D	Z	BABA	ABBA	n
TAI	AL3194	C_24T	-0.0239	-2.258	10,176	10,674	380,938
TAI	AL3194	C_79T	-0.0267	-2.364	8,948	9,438	346,188
TAI	AL3194	D_AHusky91	-0.0126	-1.58	15,165	15,554	370,227
TAI	AL3194	D_Green	-0.0059	-0.681	15,468	15,650	387,439
TAI	AL3194	D_Husky	-0.0199	-2.498	16,444	17,111	394,720
TAI	AL3194	D_Husky89	-0.0076	-0.915	13,258	13,462	321,758
TAI	AL3194	D_Mal68	-0.0025	-0.312	15,225	15,301	382,193
TAI	C_24T	AL3194	0.0239	2.258	10,674	10,176	380,938
TAI	C_24T	C_79T	-0.0014	-0.654	5,721	5,737	441,374
TAI	C_24T	D_AHusky91	0.0045	0.587	19,419	19,247	461,923
TAI	C_24T	D_Green	0.009	1.179	19,495	19,147	483,404
TAI	C_24T	D_Husky	-0.0028	-0.381	20,900	21,018	493,025
TAI	C_24T	D_Husky89	0.0071	0.929	17,107	16,865	405,779
TAI	C_24T	D_Mal68	0.0102	1.321	19,467	19,075	476,690
TAI	С_79Т	AL3194	0.0267	2.364	9,438	8,948	346,188
TAI	С_79Т	C_24T	0.0014	0.654	5,737	5,721	441,374
TAI	C_79T	D_AHusky91	0.0041	0.539	17,112	16,972	416,326

TAI	C_79T	D_Green	0.0062	0.786	17,041	16,832	435,794
TAI	С_79Т	D_Husky	-0.0021	-0.279	18,489	18,566	446,651
TAI	С_79Т	D_Husky89	0.0081	1.039	14,690	14,454	358,654
TAI	С_79Т	D_Mal68	0.0113	1.419	17,079	16,698	429,798
TAI	D_AHusky91	AL3194	0.0126	1.58	15,554	15,165	370,227
TAI	D_AHusky91	C_24T	-0.0045	-0.587	19,247	19,419	461,923
TAI	D_AHusky91	C_79T	-0.0041	-0.539	16,972	17,112	416,326
TAI	D_AHusky91	D_Green	0.0083	1.034	17,900	17,607	470,340
TAI	D_AHusky91	D_Husky	-0.0081	-1.105	15,523	15,775	478,592
TAI	D_AHusky91	D_Husky89	0.0009	0.115	12,481	12,458	394,759
TAI	D_AHusky91	D_Mal68	0.0072	1.009	18,163	17,904	463,584
TAI	D_Green	AL3194	0.0059	0.681	15,650	15,468	387,439
TAI	D_Green	C_24T	-0.009	-1.179	19,147	19,495	483,404
TAI	D_Green	C_79T	-0.0062	-0.786	16,832	17,041	435,794
TAI	D_Green	D_AHusky91	-0.0083	-1.034	17,607	17,900	470,340
TAI	D_Green	D_Husky	-0.0137	-1.839	19,216	19,750	501,352
TAI	D_Green	D_Husky89	-0.0025	-0.296	15,501	15,578	416,390
TAI	D_Green	D_Mal68	-0.0002	-0.029	16,258	16,266	485,928
TAI	D_Husky	AL3194	0.0199	2.498	17,111	16,444	394,720
TAI	D_Husky	C_24T	0.0028	0.381	21,018	20,900	493,025
TAI	D_Husky	C_79T	0.0021	0.279	18,566	18,489	446,651
TAI	D_Husky	D_AHusky91	0.0081	1.105	15,775	15,523	478,592
TAI	D_Husky	D_Green	0.0137	1.839	19,750	19,216	501,352
TAI	D_Husky	D_Husky89	0.0123	1.655	14,207	13,860	419,790
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TAI	D_Husky	D_Mal68	0.013	1.913	19,767	19,258	494,030
TAI	D_Husky89	AL3194	0.0076	0.915	13,462	13,258	321,758
TAI	D_Husky89	C_24T	-0.0071	-0.929	16,865	17,107	405,779
TAI	D_Husky89	С_79Т	-0.0081	-1.039	14,454	14,690	358,654
TAI	D_Husky89	D_AHusky91	-0.0009	-0.115	12,458	12,481	394,759
TAI	D_Husky89	D_Green	0.0025	0.296	15,578	15,501	416,390
TAI	D_Husky89	D_Husky	-0.0123	-1.655	13,860	14,207	419,790
TAI	D_Husky89	D_Mal68	0.0043	0.587	15,803	15,667	408,271
TAI	D_Mal68	AL3194	0.0025	0.312	15,301	15,225	382,193
TAI	D_Mal68	C_24T	-0.0102	-1.321	19,075	19,467	476,690
TAI	D_Mal68	C_79T	-0.0113	-1.419	16,698	17,079	429,798
TAI	D_Mal68	D_AHusky91	-0.0072	-1.009	17,904	18,163	463,584
TAI	D_Mal68	D_Green	0.0002	0.029	16,266	16,258	485,928
TAI	D_Mal68	D_Husky	-0.013	-1.913	19,258	19,767	494,030
TAI	D_Mal68	D_Husky89	-0.0043	-0.587	15,667	15,803	408,271

### Table S7.

D-statistics for D(Outgroup, Portuguese Village Dogs, Vietnamese Village Dogs, Asian Dogs). The Andean Fox was used as an outgroup for these analyses. *n* corresponds to the number of SNPs where all populations have data. Standard error for these statistics was obtained by performing a weighted block jackknife over 1Mb blocks.

Pop 2	Pop 3	Pop 4	D	Ζ	BABA	ABBA	n
D_Port61	D_Viet21	D_China8	0.1029	11.106	52,952	43,074	934,713
D_Port61	D_Viet21	D_China9	0.0557	6.079	48,963	43,796	924,557
D_Port61	D_Viet59	D_China8	0.0965	10.469	52,395	43,172	941,437
D_Port61	D_Viet59	D_China9	0.0481	5.589	47,976	43,576	920,022
D_Port61	D_Viet21	D_Tibet3	0.0663	6.823	49,504	43,352	922,535
D_Port61	D_Viet21	D_Tibet4	0.0599	6.610	49,756	44,134	914,979
D_Port61	D_Viet59	D_Tibet3	0.0575	6.277	49,026	43,693	929,474
D_Port61	D_Viet59	D_Tibet4	0.0513	5.871	49,127	44,333	921,845
D_Port61	D_Viet21	D_Mastif4	0.0714	7.278	49,690	43,069	894,260
D_Port61	D_Viet21	D_Mastif5	0.0643	7.229	49,757	43,747	913,367
D_Port61	D_Viet59	D_Mastif4	0.0639	6.791	48,007	42,240	898,585
D_Port61	D_Viet59	D_Mastif5	0.0587	6.929	48,421	43,054	917,851

### Table S8.

D-statistics for D(Outgroup, North American canids (wolf or coyote) and Taimyr wolf, PCD (AL2135 or AL3194), PCD (AL2135 or AL3194)). The Andean Fox was used as an outgroup for these analyses. *n* corresponds to the number of SNPs where all populations have data. Standard error for these statistics was obtained by performing a weighted block jackknife over 1Mb blocks. Results suggest that the most likely source of admixture into the Koster dog (AL2135) is a Mid Western coyote.

Pop 2	Pop 3	Pop 4	D	Z	BABA	ABBA	n
W_Mex1	AL2135	AL3194	0.0690	1.024	112	98	2,762
W_Mex1	AL3194	AL2135	-0.0690	-1.024	98	112	2,762
W_Yellow1	AL2135	AL3194	0.0808	1.304	124	106	2,758
W_Yellow1	AL3194	AL2135	0.0808	-1.304	106	124	2,758
W_Yellow2	AL2135	AL3194	0.0792	1.248	120	102	2,764
W_Yellow2	AL3194	AL2135	-0.0792	-1.248	102	120	2,764
C_Cal	AL2135	AL3194	-0.0004	-0.006	87	87	2,794
C_Cal	AL3194	AL2135	0.0004	0.006	87	87	2,794
C_MidW	AL2135	AL3194	-0.1814	-2.034	43	62	1,863
C_MidW	AL3194	AL2135	0.1814	2.034	62	43	1,863
TAI	AL2135	AL3194	0.1284	1.217	53	41	1,098
TAI	AL3194	AL2135	-0.1284	-1.217	41	53	1,098

## Table S9.

Sample code used as population name for SNP array analysis.

# Table S10.

Results of f4 ratio analysis depicted in Figure S18 (see Table S9 for population code).

## Table S11.

Results of f4 ratio analysis used to estimate proportion of PCD ancestry in Arctic breeds. This table include results for both SNP array and Genome-wide analyses.

### Table S12.

D-statistics for D(Outgroup, PCD (AL3194), Arctic dogs, Arctic dogs). The Andean Fox was used as an outgroup for these analyses. *n* corresponds to the number of SNPs where all populations have data. Standard error for these statistics was obtained by performing a weighted block jackknife over 1Mb blocks. Statistics with Z>3 and Z<-3 are shown in bold.

Pop 3	Pop 4	D	Z	BABA	ABBA	n
D_Mal68	D_Green	0.0001	0.010	38,564	38,555	926,919
D_Husky89	D_AHusky91	0.0059	0.416	29,524	29,174	749,696
D_Husky	D_Husky89	0.0166	1.254	33,365	32,275	797,067
D_Husky	D_AHusky91	0.0176	1.324	37,461	36,164	913,221
D_Husky89	D_Green	0.0437	3.261	38,311	35,101	790,543
D_Husky89	D_Mal68	0.0455	3.384	39,192	35,781	775,433
D_AHusky91	D_Green	0.0470	3.555	44,116	40,158	897,276
D_AHusky91	D_Mal68	0.0450	3.596	44,737	40,883	884,940
D_Husky	D_Green	0.0619	5.021	48,948	43,246	955,940
D_Husky	D_Mal68	0.0609	5.050	49,177	43,528	942,665

### Table S13.

D-statistics for D(Outgroup, Asian or European Dogs, Arctic dogs, Arctic dogs). The Andean Fox was used as an outgroup for these analyses. *n* corresponds to the number of SNPs where all populations have data. Standard error for these statistics was obtained by performing a weighted block jackknife over 1Mb blocks. Statistics with Z>3 are shown in bold.

Pop 2	Pop 3	Pop 4	D	Z	BABA	ABBA	n
D_Portt71	D_AHusky91	D_Husky89	0.0000	0.001	37,394	37,394	954,618
D_Portt71	D_Mal68	D_Husky89	0.0005	0.040	47,184	47,141	988,232
D_Portt71	D_Husky	D_Mal68	0.0009	0.086	58,360	58,259	1,190,049
D_Port61	D_AHusky91	D_Mal68	0.0016	0.149	55,075	54,905	1,143,740
D_Port61	D_Mal68	D_Husky	0.0018	0.186	59,906	59,692	1,217,824
D_Portt71	D_Husky89	D_Husky	0.0026	0.250	42,139	41,923	1,014,889
D_Port61	D_Husky89	D_Mal68	0.0029	0.271	48,766	48,481	1,013,144
D_Portt71	D_AHusky91	D_Mal68	0.0046	0.412	53,778	53,282	1,117,362
D_Port61	D_AHusky91	D_Husky	0.0060	0.586	48,323	47,748	1,179,281
D_Portt71	D_AHusky91	D_Husky	0.0060	0.596	46,884	46,321	1,151,944
D_Port61	D_Husky89	D_AHusky91	0.0077	0.675	38,794	38,202	979,116
D_Port61	D_Husky89	D_Husky	0.0098	0.948	43,569	42,725	1,040,592
D_Portt71	D_Green	D_AHusky91	0.0197	1.780	53,880	51,794	1,133,450
D_Portt71	D_Green	D_Husky	0.0242	2.401	59,660	56,844	1,207,389

D_Portt71	D_Green	D_Mal68	0.0281	2.433	49,870	47,151	1,171,857
D_Portt71	D_Green	D_Husky89	0.0267	2.466	47,462	44,995	1,007,512
D_Port61	D_Green	D_Husky89	0.0283	2.625	48,938	46,241	1,033,241
D_Port61	D_Green	D_AHusky91	0.0301	2.628	55,922	52,657	1,160,778
D_Port61	D_Green	D_Husky	0.0341	3.348	61,488	57,440	1,236,161
D_Port61	D_Green	D_Mal68	0.0375	3.355	51,677	47,947	1,199,625
D_China8	D_Husky89	D_AHusky91	0.0017	0.158	37,346	37,218	959,372
D_China9	D_Mal68	D_AHusky91	0.0040	0.424	52,229	51,810	1,085,883
D_China9	D_Green	D_Husky89	0.0063	0.586	46,453	45,871	1,003,859
D_China8	D_Green	D_Mal68	0.0066	0.628	48,381	47,747	1,175,269
D_China8	D_Husky	D_AHusky91	0.0068	0.744	46,638	46,006	1,155,853
D_China8	D_Mal68	D_Husky	0.0071	0.780	58,261	57,441	1,193,177
D_China8	D_Husky	D_Husky89	0.0078	0.820	42,094	41,441	1,019,519
D_China9	D_Husky	D_Green	0.0081	0.846	57,408	56,484	1,178,134
D_China9	D_AHusky91	D_Husky	0.0085	0.957	45,878	45,108	1,120,029
D_China9	D_Mal68	D_Husky	0.0096	1.121	57,254	56,163	1,156,324
D_China8	D_Green	D_Husky	0.0124	1.296	58,444	57,014	1,211,379
D_China9	D_Husky	D_Husky89	0.0141	1.428	42,263	41,090	1,009,765

D_China9	D_AHusky91	D_Green	0.0152	1.493	52,310	50,746	1,106,207
D_China8	D_Mal68	D_AHusky91	0.0146	1.553	53,877	52,324	1,120,670
D_China8	D_Mal68	D_Husky89	0.0160	1.601	47,812	46,307	992,341
D_China8	D_Green	D_AHusky91	0.0193	1.872	53,594	51,565	1,137,595
D_China9	D_Husky89	D_AHusky91	0.0208	1.901	37,871	36,329	950,029
D_China9	D_Mal68	D_Green	0.0205	2.167	48,324	46,381	1,142,965
D_China9	D_Mal68	D_Husky89	0.0224	2.288	47,913	45,810	982,870
D_China8	D_Green	D_Husky89	0.0250	2.488	47,327	45,021	1,012,257

# Table S14

Fitting of COSMIC mutational signatures 1, 5, 7 and Dog Germline signature to 1,933,897 CTVT tumor-only SNVs shared between 608T and 609T.

Mutational signature	Fit	Number of SNVs ( n=1,933,897)
1	Mean (15.4%)	297,820.1
	Lower bound (15.2%)	293,952.3
	Upper bound (15.5%)	299,754
5	Mean (40.1%)	775,492.7
	Lower bound (40.0%)	773,558.8
	Upper bound (40.3%)	779,360.5
7	Mean (38.95%)	753,252.9
	Lower bound (38.9%)	752,285.9
	Upper bound (39%)	754,219.8
Dog Germline	Mean (5.5%)	106,364.3
	Lower bound (5.3%)	102,496.5
	Upper bound (5.7%)	110,232.1

## Table S15

Number of clonal tumor-unique SNVs in 608T and 609T. CN, copy number. "All" indicates the complete set of tumor-unique SNVs, "Clonal", only those that are clonal.

Sample	Copy number state	All	Clonal
608T	CN1	40	33
	CN2	140	125
	CN3	22	21
	CN4	3	3
	CN6	1	1
		206	183
609T	CN1	43	30
	CN2	159	125
	CN3	21	15
	CN4	5	4
		228	174

# Table S16

Number of clonal N[C>T]G mutations unique to 608T and 609T, as well as the number found in the somatic lineage from the CTVT founder dog until MRCA<sub>608T-609T</sub>.

Variant set	N[C>T]G mutations
Clonal 608T-unique	27
Clonal 609T-unique	23
CTVT (origin to MRCA <sub>608T-609T</sub> )	222,072

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