Natural Selection and Origin of a Melanistic Allele in North American Gray Wolves

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Abstract

Pigmentation is often used to understand how natural selection affects genetic variation in wild populations since it can have a simple genetic basis, and can affect a variety of fitness-related traits (e.g., camouflage, thermoregulation, and sexual display). In gray wolves, the K locus, a b-defensin gene, causes black coat color via a dominantly inherited K allele. The allele is derived from dog-wolf hybridization and is at high frequency in North American wolf populations. We designed a DNA capture array to probe the geographic origin, age, and number of introgression events of the K allele in a panel of 331 wolves and 20 dogs. We found low diversity in K, but not ancestral ky, wolf haplotypes consistent with a selective sweep of the black haplotype across North America. Further, North American wolf K haplotypes are monophyletic, suggesting that a single adaptive introgression from dogs to wolves most likely occurred in the Northwest Territories or Yukon. We use a new analytical approach to date the origin of the K allele in Yukon wolves to between 1,598 and 7,248 years ago, suggesting that introgression with early Native American dogs was the source. Using population genetic simulations, we show that the K locus is undergoing natural selection in four wolf populations. We find evidence for balancing selection, specifically in Yellowstone wolves, which could be a result of selection for enhanced immunity in response to distemper. With these data, we demonstrate how the spread of an adaptive variant may have occurred across a species' geographic range.

Key words: CBD103, gray wolf, allele age, adaptive introgression, melanism, sequence capture.

Introduction

Understanding how polymorphisms are generated and maintained is a key goal of evolutionary biology. One of the most distinct polymorphic traits in nature is coloration (Wright 1917), which is generally determined by the spatial distribution of pigmented hairs, feathers, or scales across the body and by pigment type (Wright 1917; Protas and Patel 2008; Manceau et al. 2010). In mammals, two main types of pigments control coloration, eumelanin (black/brown) and pheomelanin (red/yellow) (Protas and Patel 2008). Switching between these two pigment types is often controlled by the Agouti (ligand)–Mc1r (receptor) pathway (Protas and Patel 2008). Coloration can serve several functions, including camouflage, intraspecific and interspecific signaling, and thermoregulation (Wright 1917; Little 1958; Protas and Patel 2008), and can be subject to positive selection (e.g., Hoekstra and Nachman 2003). Additionally, melanin-based coloration genes are associated with physiological and behavioral traits through pleiotropic effects (Ducrest et al. 2008). Although the evolution of color polymorphism is inherently interesting for the varied functions mentioned above, it is also key to understanding evolutionary processes in general (Svensson 2017). If the causative alleles for phenotypic variation are known, the adaptive function of such alleles could...
potentially be determined in a natural evolutionary context (e.g., Barrett and Hoekstra 2011).

Melanism in gray wolves (Canis lupus) represents a novel system for understanding how demography and selection can change the patterns of adaptive variation across the entire geographic range of a species. Although many melanistic phenotypes are caused by mutations in Agouti or Mc1r, in the gray wolf melanism is manifested by a novel mutation in the β-defensin CBD103 gene (also referred to as the K locus). This gene encodes an alternative ligand for Mc1r that outcompetes a functioning Agouti ligand if present (Candille et al. 2007; Kerns et al. 2007; Anderson et al. 2009). The K<sup>d</sup> allele contains a 3 bp deletion that confers a dominantly inherited black (melanistic) coat color phenotype, whereas the wild-type K<sup>a</sup> allele confers a gray (agouti) coat color in homozygotes (Anderson et al. 2009). A previous study of 47 Arctic and Yellowstone wolves from forest and tundra/taiga habitats by Anderson et al. (2009) used a data set of 52 SNPs to show that the K locus experienced a strong selective sweep in black wolves after an initial introgression event from dogs >500 years ago (Leonard et al. 2002; Anderson et al. 2009). The evidence for a selective sweep, detailed in Anderson et al. (2009), consisted of a near absence of polymorphism within 50 kb of the K<sup>d</sup> allele in wolves, yet the wild-type K<sup>a</sup> allele was highly polymorphic. Additionally, the authors compared coalescent times, using a molecular clock approach, as a function of increasing distance from the K locus in dogs and wolves, and showed that the K<sup>d</sup> allele was introduced into North American wolves from dogs (Anderson et al. 2009). Research over the last several years has clarified the genetic basis and potential fitness advantage of melanism in wolves (e.g., Anderson et al. 2009; Coulson et al. 2011; Stahler et al. 2013; Cassidy et al. 2017), yet details of the evolutionary history, such as the number, timing, and geographic origin of introgression events, remain unknown.

In wolves, the target of selection for black coat color seems unrelated to camouflage, intraspecific signaling, or thermoregulation (e.g., Anderson et al. 2009; Coulson et al. 2011). Specifically, wolf coat color varies from black to white, with tawny variations, and with higher frequencies of lighter phenotypes in the high Arctic where the background vegetation is paler (Musiani et al. 2007). However, gray and black individuals often exist within the same populations and at similar frequencies over time, suggesting that mechanisms such as fitness tradeoffs maintain the polymorphism (Anderson et al. 2009; Coulson et al. 2011; Hedrick et al. 2016). In Yellowstone National Park, where wolf coat color has been intensively studied, the estimated frequency of black wolves in the population remained constant over a 19-year period (1996–2014) at 0.45 (Hedrick et al. 2016). The maintenance of these frequencies has been explained by a strong heterozygous advantage for the K<sup>d</sup> allele (Coulson et al. 2011; Hedrick et al. 2014), and recent evidence suggests strong disassortative mating for coat color (Hedrick et al. 2016). Demonstrated fitness differences among color genotypes in the Yellowstone population includes highest annual survival, lifespan, and lifetime reproductive success for black heterozygotes and lowest survival, reproductive success, and frequency for black homozygotes (Coulson et al. 2011). In contrast, gray homozygous females show higher annual reproductive success (Stahler et al. 2013). Such fitness variation among all three genotypes suggests pleiotropic effects of variants at the coat color locus (e.g., Ducrest et al. 2008) and that coat color may not be the proximate object of selection given the difference in fitness between the heterozygous and homozygous black coat color genotypes. In this regard, the K locus is a member of the β-defensin family of antimicrobial peptides (Pazgier et al. 2006) and may be involved in adaptive immune response (Yang et al. 1999). K locus mRNA is expressed in canine skin and respiratory tract (Candille et al. 2007; Erles and Brownlie 2010), and mRNA levels are correlated with increased antimicrobial activity against Bordetella bronchiseptica, a respiratory pathogen (Erles and Brownlie 2010). Thus, fitness tradeoffs involving the costs and benefits of immunity under varying environmental conditions (Coulson et al. 2011; Stahler et al. 2013) and the fitness of specific mating pairs (Hedrick et al. 2016) may determine the frequency of each genotype. Additionally, the frequency of a SNP in linkage disequilibrium (LD) with the 3 bp deletion is significantly and positively associated with maximum temperature of warmest month across North American wolf ecotypes (Schweizer, vonHoldt et al. 2016), and temperature and pathogen prevalence are positively related in a multitude of species (Allen et al. 2002; Guernier et al. 2004; Dionne et al. 2007), further supporting the notion that selection within the K locus likely involves immunity.

Initial studies of the K locus have raised further questions concerning the geographic origin, number, and age of introgression events that could not be addressed previously due to limited genomic and geographic sampling. Consequently, we designed a custom hybridization capture array to perform extensive resequencing of a five megabase (Mb) region surrounding the K locus, as well as neutral and control genomic regions to assess genomic changes due to demography alone. We sequenced these regions in a large geographic sample of North American wolves and 20 domestic dog breeds. We assessed patterns of nucleotide and haplotype diversity, population-specific decay in linkage disequilibrium (LD), and hierarchical patterns of genetic divergence among populations, and used a novel method to determine when the K<sup>d</sup> allele likely originated in each of our sampled populations. With these data, we reconstructed the evolutionary history of the black genotype and find a single origin, followed by a rapid spread across North America. We hypothesize that this spread may have been accelerated by the presence of dogs and mesocarnivores as reservoirs for pathogens to which the black locus provided enhanced immunity. Lastly, our population genetic simulations demonstrate that coat color is currently undergoing natural selection in four different wolf populations.

**Results**

**Targeted Capture Sequencing of Dogs and Wolves**

We designed a custom hybridization capture array to assay sequence variation surrounding the K locus, and in neutral, genic, telomeric, and nontelomeric regions (fig. 1, see
Methods and Supplementary Material online). These presumed neutral, telomeric and nontelomeric regions provide an empirical background for summary statistics, since the K locus is located close to the telomere on chromosome 16 and may show a positional bias in genetic diversity. After target enrichment, sequencing, and alignment to the CanFam3.1 dog reference, all individuals sequenced (n = 382) retained approximately 60% of the target regions with at least 25 coverage (mean ± 1SD: 150 ± 61×) (supplementary table S1 and fig. S1, Supplementary Material online). After genotype filtering and quality control, 10,922,248 positions with 162,815 variants and 12,774 insertions/deletions were retained. Based on data from 109 individuals previously genotyped on the Affymetrix Dog SNP array, we calculated 99.7% genotype concordance for heterozygotes (supplementary table S2, Supplementary Material online). After additional quality control filters (see Supplementary Material online), 353 samples were used for further analysis (table 1 and supplementary fig. S2, Supplementary Material online). Pedigree data from 203 Yellowstone wolves was used to aid phasing of haplotypes among all 353 samples using SHAPEIT software (supplementary fig. S3 and table S3, Supplementary Material online).

To further assess sequencing quality and to characterize genetic patterns, we partitioned our sequence data into six genomic categories: 1) neutral; 2) the K locus core (200 kb region); 3) genic (all exons and introns); 4) exonic; 5) telomeric; and 6) nontelomeric regions (supplementary table S3, Supplementary Material online). Heterozygosity values measured across all six partitions (fig. 1B and supplementary table S4, Supplementary Material online) were similar to those based on SNP array data and expectations based on population histories (Gray et al. 2009; vonHoldt, Pollinger, et al. 2010; vonHoldt et al. 2011). Dog heterozygosity values were lower than those in many wolf populations (fig. 1B) which is consistent with a bottleneck in the history of dog domestication (e.g., vonHoldt, Pollinger, et al. 2010; Freedman et al. 2014). Moreover, among wolves and dogs, heterozygosity was higher in telomeric regions than in nontelomeric regions (Ho, telomeric: 0.176% ± 0.031%; Ho, nontelomeric: 0.121% ± 0.03%; P value < 1e-15), which is consistent with telomeric regions in dogs and humans having higher recombination rates and higher diversity (Myers et al. 2005; Auton et al. 2013). Compared with other genic regions, the 200 kb K locus in wolves and in dogs showed high heterozygosity (Ho: 0.229% ± 0.103%; P value < 1e-15), which is unexpected for a selective sweep scenario at the K locus. However, the 200 kb region includes several genic and nongenic regions, which may elevate heterozygosity relative to purely genic regions.

Haplotype Diversity of K^b versus K^v in Dogs and Wolves

K^b haplotypes appeared less variable than K^v in all wolf populations, and variability was diminished in K^b wolves relative to K^v dogs (supplementary fig. S5, Supplementary Material online). Yellowstone wolf haplotypes appeared to be largely composed of a single K^b haplotype, with very little haplotype

![Fig. 1. Resequencing strategy and heterozygosity values within sequenced regions.](https://example.com/fig1.png)
Table 1. Sampling Information for Wolves and Dogs Sequenced on the Capture Array.

<table>
<thead>
<tr>
<th>Origin</th>
<th># Sequenced</th>
<th># Used for Analyses</th>
<th>(k^B) Haplotypes</th>
<th>(k^R) Haplotypes</th>
<th>Black</th>
<th>Gray</th>
<th>White</th>
<th>Other</th>
<th>Unknown</th>
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<td>Gray wolf</td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
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<td>117</td>
<td>544</td>
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<td>234</td>
<td>103</td>
<td>365</td>
<td>94</td>
<td>129</td>
<td>0</td>
<td>0</td>
<td>11</td>
</tr>
<tr>
<td>Yellowstone</td>
<td>34</td>
<td>33</td>
<td>8</td>
<td>58</td>
<td>6</td>
<td>16</td>
<td>0</td>
<td>0</td>
<td>11</td>
</tr>
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<td>212</td>
<td>201</td>
<td>95</td>
<td>307</td>
<td>88</td>
<td>113</td>
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<td>0</td>
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<td>7</td>
<td>25</td>
<td>0</td>
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<td>4</td>
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<td>0</td>
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<td></td>
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<td></td>
<td></td>
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<tr>
<td>Newfoundland</td>
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<td>7</td>
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<td>14</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>2</td>
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<td>Northwest Territories</td>
<td>32</td>
<td>32</td>
<td>9</td>
<td>55</td>
<td>4</td>
<td>4</td>
<td>13</td>
<td>0</td>
<td>11</td>
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<tr>
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<td>0</td>
<td>56</td>
<td>0</td>
<td>0</td>
<td>9</td>
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<td>0</td>
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<td></td>
<td></td>
<td></td>
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<td>0</td>
<td>14</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>7</td>
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<tr>
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<td>4</td>
<td>0</td>
<td>8</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>4</td>
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<tr>
<td>Yukon</td>
<td>7</td>
<td>6</td>
<td>2</td>
<td>9</td>
<td>0</td>
<td>0</td>
<td>0</td>
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<tr>
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<td>8</td>
<td>2</td>
<td>14</td>
<td>2</td>
<td>2</td>
<td>0</td>
<td>0</td>
<td>4</td>
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<tr>
<td>Domestic dog</td>
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<td></td>
<td></td>
<td></td>
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<tr>
<td>Breeds fixed for (k^B)</td>
<td>21</td>
<td>20</td>
<td>16</td>
<td>24</td>
<td>5</td>
<td>0</td>
<td>1</td>
<td>14</td>
<td>0</td>
</tr>
<tr>
<td>Breeds fixed for (k^R)</td>
<td>8</td>
<td>8</td>
<td>16</td>
<td>0</td>
<td>5</td>
<td>0</td>
<td>0</td>
<td>3</td>
<td>0</td>
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<tr>
<td>Breeds variable for (k^B) and (k^R)</td>
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<td>12</td>
<td>0</td>
<td>0</td>
<td>6</td>
<td>0</td>
<td>0</td>
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<tr>
<td>Breeds with previously unknown (K) alleles</td>
<td>4</td>
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<td>0</td>
<td>3</td>
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<tr>
<td>Total</td>
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<td></td>
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<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Admixed Italy</td>
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<td>4</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>2</td>
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</tbody>
</table>


Consistent with a domestication bottleneck in the latter. Telomeric regions had higher values of \(\pi\) and \(\theta_w\), perhaps because recombination rate increases towards telomeres and as a result diversity in those regions is higher (Auton et al. 2013; table 2). In dog and wolf \(k^R\) haplotypes, diversity was significantly higher than that of the other telomeric regions (e.g., \(\pi_{dog, k^R}: 0.002 \pm 0.0013; \pi_{dog, Tel}: 0.00139 \pm 0.0009; P\) value: 0.00059; table 2 and supplementary table S5, Supplementary Material online), whereas diversity in the \(K^B\) haplotypes was consistently lower than the other telomeric regions, although this pattern was more pronounced in wolves than in dogs (table 2 and supplementary table S5, Supplementary Material online). These results suggest that the patterns of diversity measured at the \(K\) locus, especially with regards to the \(K^B\) haplotype, are unusual, even in comparison to similarly positioned and sized regions in other chromosomes. In aggregate, these results suggest that the low levels of diversity found within \(K^B\) haplotypes, especially in wolves, are not characteristic of telomeric regions in general, and could not have been predicted solely from the genomic location of the \(K\) locus.

Extended Haplotype Homozygosity of the \(K\) Locus

Using the extended haplotype homozygosity (EHH) score, which ranges from 0 (no homozygosity) to 1 (complete homozygosity) (Sabeti et al. 2002), we found that the haplotype containing the derived \(K^B\) allele had more extensive homozygosity in wolves than in dogs (blue lines in fig. 3A and B;
supplementary fig. S7A, Supplementary Material online). These data suggest a recent and dramatic selective sweep for the $K^B$ allele in wolves. These patterns are not driven by uneven sampling of the $K^B$ haplotype among wolf populations or by relatedness among individuals, as the patterns remain when the Yellowstone wolf population (comprising over half our samples) is subsampled to only 35 unrelated individuals (supplementary fig. S8, Supplementary Material online), and when all wolves are subsampled to be unrelated (supplementary fig. S8, Supplementary Material online).

Comparison of Genetic Variation among Wolf Populations Containing the $K^B$ Allele

Using population-wide summary statistics, we inferred the geographic origin of the $K^B$ allele. The region of origin should have the highest nucleotide and haplotype diversity, and the highest proportion of ancestral haplotypes (e.g., Tishkoff et al. 2001; Gray et al. 2010) because more time has elapsed for new mutations and recombination events to occur (e.g., Shannon et al. 2015). Additionally, populations moving outside of the origin may be expected to carry a subset of the diversity of the ancestral population (Ramachandran et al. 2005). Given that Alberta and Newfoundland each only had a single $K^B$ haplotype, we focused on relative diversity in Alaska, the Northwest Territories, Yellowstone, and Yukon 200 kb $K$ locus regions. For each of the four populations, the diversity of the $K^B$ haplotype was significantly lower than the diversity of the telomeric regions (fig. 4 and table 2). We found that $K^B$ haplotypes from Yukon had the greatest diversity, followed by Northwest Territories (fig. 4). Yellowstone wolves showed...
the lowest $K^B$ haplotype diversity, despite including the largest
e number of sampled $K^B$ haplotypes (table 1). Importantly,
the Yellowstone population is otherwise genetically diverse
(fig. 1B), having been derived from three distinct populations
from Montana, Alberta, and British Columbia. Values of
Tajima’s $D$ for $K^B$ haplotypes in Yellowstone wolves were
negative ($-1.13885 \pm 0.86024$; table 2), and Alaska $K^B$
haplotypes also had a slightly negative Tajima’s $D$ ($-0.50833 \pm 1.22007$; table 2). Although the sample size of
Yukon wolves was too low to calculate $D$, the values of $\pi$ and
$\theta_w$ indicate the highest $K^B$ diversity was found in Northwest
Territories or Yukon suggesting that region as an origin, with
lower diversity in Alaska and Yellowstone.
Levels of LD were much higher for $K^B$ haplotypes than $K^C$
haplotypes for the $K$ locus region (fig. 5 and supplementary
fig. S9, Supplementary Material online), which had overall
higher levels of LD than those of the 10 parallel telomeric
regions (supplementary fig. S10, Supplementary Material
online). For the $K^D$ haplotypes (fig. 5A), most populations had LD
below $r^2 = 0.2$ for the 5 Mb region at a distance extending
from 20 kb to 1 Mb from the 3 bp deletion (supplementary
table S6, Supplementary Material online). Patterns of LD in
the Northwest Territories suggest that it was one of the
most diverse populations that also contained $K^B$ haplo-
types. For $K^B$ haplotypes (fig. 5B), only dogs, and wolves
from the Northwest Territories and Alaska, ever reached
$r^2 = 0.2$ within the sampled interval (supplementary table
S6, Supplementary Material online). Similar patterns are
observed in heatmap triangle plots of pairwise LD (sup-
plementary fig. S9, Supplementary Material online). In
contrast, LD decayed rapidly over the $\sim 200$ kb reference
telomeric regions, with LD being only slightly higher in
dogs than in wolves (supplementary fig. S10, Supplementary Material online).

### Extended Haplotype Homozygosity of the K Locus among Wolf Populations

For each wolf population that contained individuals with $K^B$
haplotypes, we calculated the EHH statistic to assess the
decay of homozygosity (fig. 3C–F and supplementary table S7,
Supplementary Material online), and found that Alaska and
Yellowstone wolves had the most extensive EHH in $K^B$
haplotypes (blue lines in fig. 3C and D). In contrast, Northwest
Territories (fig. 3E) and Yukon (fig. 3F) populations had EHH
scores that declined more rapidly (fig. 3 and supplementary
table S7, Supplementary Material online). The nonsymmet-
rical decay of EHH on either side of the deletion, especially in
the Northwest Territories population, may reflect fewer re-
combination events downstream of the $K$ locus. These uneven
patterns upstream and downstream of the $K$ locus may
be a result of selection at neighboring loci, although sequence
data did not show any fixed, functional differences between
$K^B$ or $K^D$ haplotypes in dogs or wolves other than the 3 bp
deletion (Supplementary Material online).

To explore whether patterns of EHH within the reintro-
duced Yellowstone population were a result of limited diver-
sity within the founders or a subsequent sweep, we measured
EHH solely within 12 founders, of which five were $K^D/k^D$ and
seven were $k^D/k^D$. Decay of EHH surrounding the core
mutation occurred more rapidly in Yellowstone founders (sup-
plementary fig. S11, Supplementary Material online) than in the
Yellowstone population (fig. 3D and supplementary table S7,
Supplementary Material online). Over the 5 Mb region, EHH
decayed to zero in $K^B$ haplotypes $\sim 1$ Mb upstream

### Table 2. Haplotype Diversity within Dogs and North American Wolves, for Each of Four Types of Region within the Genome.

<table>
<thead>
<tr>
<th>Region Type</th>
<th>Group</th>
<th>$\pi$</th>
<th>$\theta_w$</th>
<th>Tajima’s $D$</th>
</tr>
</thead>
<tbody>
<tr>
<td>K locus Core</td>
<td>Dog $K^A$</td>
<td>0.00117 (0.000105)</td>
<td>0.00115 (0.00096)</td>
<td>0.02901 (0.91231)</td>
</tr>
<tr>
<td></td>
<td>Dog $K^C$</td>
<td>0.0002 (0.0013)</td>
<td>0.00174 (0.00095)</td>
<td>0.51471 (1.32856)</td>
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<tr>
<td></td>
<td>Wolf $K^B$</td>
<td>0.00035 (0.00045)</td>
<td>0.00093 (0.00068)</td>
<td>-1.30547 (0.59812)</td>
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<tr>
<td></td>
<td>Alaska</td>
<td>0.00035 (0.00085)</td>
<td>0.00039 (0.0009)</td>
<td>-0.50833 (1.22007)</td>
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<td></td>
<td>NWT</td>
<td>0.00094 (0.00142)</td>
<td>0.00066 (0.00097)</td>
<td>1.66771 (0.82527)</td>
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<tr>
<td></td>
<td>YNP</td>
<td>0.00011 (0.0003)</td>
<td>0.00033 (0.00047)</td>
<td>-1.13885 (0.86024)</td>
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<tr>
<td></td>
<td>Yukon</td>
<td>0.00119 (0.0015)</td>
<td>0.00119 (0.0015)</td>
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<td></td>
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<tr>
<td></td>
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<td>0.00197 (0.00098)</td>
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**Note:** Italics text indicates values for the $K$ locus region that are significantly different from Telomeric regions ($P < 0.05$ with Wilcoxon rank sum test). See supplementary table S5, Supplementary Material online for $P$ values. $\theta_w$, Watterson’s theta; $\pi$, nucleotide diversity.

*Summary statistics were calculated from 1 kb nonoverlapping windows (K locus Core: 103, Nontelomeric: 96, Telomeric: 75) or 10 kb nonoverlapping windows (Neutral: 491), and the mean and standard deviation are provided.
Supplementary Fig. S11 and Supplementary Table S7, Supplementary Material online), whereas in the current Yellowstone population the EHH score remained above zero even C24 upstream (supplementary fig. S7 and supplementary table S7, Supplementary Material online). Given that the Yellowstone population is not inbred (vonHoldt et al. 2008; vonHoldt, Stahler, et al. 2010), these results suggest that an advantageous haplotype rose in frequency subsequent to the foundation of the population. This pattern, in association with results from neutral regions, suggests that selection may be especially strong at the K locus in Yellowstone wolves relative to other wolf populations.

Simulations of Allele Frequency Change in Yellowstone Wolves
We sought to determine whether the present-day allele frequency of the K\textsuperscript{B} allele (22%) in Yellowstone wolves could be explained simply by genetic drift and negative selection or whether other nonneutral processes would have to be considered. We chose to focus our simulations

on the Yellowstone population, since its demographic history is known and can be incorporated into models to test for selection. We performed forward simulations based on the initial allele frequency (20%) and demography of the wolves, adjusting \( N_e \) for nonrandom mating (see Materials and Methods). Our null model included a lethal recessive condition such that \( K^B \) homozygotes would not survive. We found that the final allele frequency of 22% was unlikely to be seen under this null model (\( P = 0.018; \) fig. 6A). This result suggests that a nonneutral process (e.g., positive selection, balancing selection) has altered the frequency of the \( K^B \) allele since the founding of the Yellowstone population in 1995.

To better understand the magnitude and nature of selection, we used approximate Bayesian computation (ABC) to infer the dominance coefficient (\( h \)) of the \( K^B \) allele. We used a uniform prior over \((-2, 2)\) and rejection sampling, retaining simulations where the final allele frequency in each simulation replicate was within 1e-3 of the final observed allele frequency. We found that the mean of the posterior distribution was \(-0.179\) (95% HPD: \(-0.483, -0.0040; \) fig. 6B), suggesting that balancing selection maintains variation at this locus. Importantly, the posterior distribution excludes 0, further arguing against the \( K^B \) allele being a simple recessive lethal.

**Hierarchical Patterns of Divergence among Populations**

Neighbor joining (NJ) trees generated using pairwise nucleotide divergence values from \( K \) locus haplotypes were compared with those based on the neutral data, since the former should provide insight into the specific history at the selected \( K \) locus, and the latter should reflect population history (vonHoldt, Pollinger, et al. 2010; vonHoldt et al. 2011; Pilot et al. 2014; Schweizer, vonHoldt, et al. 2016). NJ trees of the neutral regions showed groupings that were concordant with major geographic or taxonomic differences (supplementary fig. S12, Supplementary Material online). For instance, dogs formed a single grouping near Italian wolves, which is concordant with likely European or Asian origins of domestication (Savolainen et al. 2002; Thalmann et al. 2013; Freedman et al. 2014; Fan et al. 2016). Next, we generated NJ trees using data from a 26 kb region surrounding the \( K \) locus where there was no evidence of recombination (fig. 7A and supplementary fig. S13, Supplementary Material online; see Methods). North American \( K^B \) haplotypes form a single cluster, suggesting a single introgression event (fig. 7A). This cluster also contains a North American dog breed, the Newfoundland, as well as Labrador Retrievers, German Shorthaired Pointer and Portuguese Water Dog, and a Siberian Husky \( k^p \) haplotype was sister to this group (fig. 7A). A separate cluster of \( K^B \) haplotypes from Italian wolves are related to European dog breeds, suggesting a separate introgression event. NJ trees generated using pairwise nucleotide divergence between larger regions were not informative due to recombination events further upstream and downstream of the \( K \) locus, most notably observed in Northwest Territory wolves (results not shown).

We built haplotype networks using the same nonrecombining 26 kb region surrounding the \( K \) locus (see Methods; fig. 7B and supplementary fig. S14, Supplementary Material online). Among \( k^p \) haplotypes, most sampled dog breeds lie on a single branch of the network whereas North American and Italian wolf haplotypes are represented on other branches of the network (supplementary fig. S14, Supplementary Material online), which is consistent with the evolutionary history of these populations. Among the \( K^B \) haplotype network, North American wolves mostly form a single cluster, with a few dog haplotypes nested within the cluster (supplementary fig. S14, Supplementary Material online). Italian wolf \( K^B \) haplotypes group most closely to dog haplotypes rather than wolf haplotypes, which again suggests a different introgression event than that which occurred in North American wolves (supplementary fig. S14,
A closer examination of the wolf populations and dog breeds represented by the $k^b$ haplotypes (fig. 7B) shows that the Newfoundland dog haplotype (haplotype VI) is close to the most common haplotype (haplotype I) in North American wolf populations.

**Age of Introgression in North American Wolves**

To understand the timing and spread of the dog $k^b$ allele into wolf populations in North America, we estimated time to the common ancestor (TMRCA) for the deletion in each of the four populations containing more than one $k^b$ haplotype.
We used a new method (Smith et al. 2018) which leverages both the decay in LD between the selected allele and nearby sites, as well as the accumulation of new mutations on the selected allele’s ancestral haplotype. We find that the oldest TMRCA estimates are found in the Yukon with posterior means ranging from 1598 to 7248 ya, depending on the mutation rate used (fig. 8 and table 3). The youngest TMRCA values are consistently found in Yellowstone, ranging from 202 to 1942 ya. Estimates for Alaska, Northwest Territories, and two Yellowstone posteriors have significant overlap and all fall roughly between 694 and 2135 ya.

In some cases, the choice of different mutation rates did not affect the variability of time estimates (table 3). Among the Alaska and Northwest Territories samples, mutation rates from 0.4 \times 10^{-8} to 0.45 \times 10^{-8} impose similar time estimates. The same is true for the Yellowstone samples at mutation rates of 0.3 \times 10^{-8} and 0.4 \times 10^{-8}. However, all estimates using a mutation rate of 0.3 \times 10^{-8} did result in younger TMRCA values.

Demographic Inference and Simulations of Selection in Alaska, Northwest Territories, and Yukon Wolves

To gain insight into the processes that maintain variation at the K locus, we performed demographic inferences from the neutral genetic variation data in the Alaska, Northwest Territories, and Yukon wolf populations. We used a diffusion approximation based method (Gutenkunst et al. 2009) and inferred a single population size change demographic model for each population. After incorporating ancestral sequence misidentification into the model, we fit the site frequency spectra across all frequency bins (supplementary fig. S15, Supplementary Material online). We infer that all three populations underwent a contraction at some point in the past, with the Yukon population undergoing the most severe contraction (supplementary table S8, Supplementary Material online).

Armed with this information, we sought to understand the patterns of selection at the K locus more closely. Specifically, we tested whether a null model with complex demography and a recessive lethal mutation introduced through recent admixture could be compatible with the data. In contrast to the Yellowstone population, we do not know the timing of introgression nor the frequency of introgression. Therefore, we used ABC to infer the unknown parameters (time of introgression and introgression frequency). We then asked whether these inferred values are sensible given the other known information. We found that in order to match the final allele frequency data, introgression must have occurred very recently and at nearly 100% frequency in every population (fig. 9). This timing is not supported by our TMRCA estimates (fig. 8), which indicates the allele is much older than a few generations ago. Further, the unreasonably large introgression frequency suggests that the null scenario of complex demography combined with introgression and purifying selection on a recessive lethal allele is unlikely. Instead, our results strongly favor some type of balancing selection maintaining the frequency of the K^B allele in these populations.

Discussion

Melanism in gray wolves is caused by a dominantly inherited 3 bp deletion at the K locus, and the region surrounding the mutation shows signatures of strong selection within K^B haplotypes (Anderson et al. 2009). Using a resequencing approach (fig. 1A), we demonstrate that haplotypes containing the K^B allele showed signals of a selective sweep in North American wolves more so than dogs.

![Fig. 6](https://example.com/fig6.png)

**Fig. 6.** (A) Null distribution of allele frequencies based on the Yellowstone National Park wolf reintroduction demographic history. The dashed red line denotes the observed value, which corresponds to a P value of 0.018. (B) Prior and posterior distributions of the dominance (h) coefficient for the K^B allele. The mean posterior value was −0.179 (95% HPD: −0.483, −0.0040).
We also find support for a single introgression event, likely in Yukon or Northwest Territories. These results are corroborated by the finding that the $K^b$ allele age is oldest in the Yukon wolf population (fig. 8).

Italian wolves that contain the $K^b$ mutation represent a separate introgression with European dog breeds that we speculate may be more recent based on patterns of genetic diversity and linkage disequilibrium.
Selection on the K\textsuperscript{B} Allele in Wolves

Using multiple diversity statistics, LD decay, and EHH, we find compelling evidence of a partial selective sweep of the K\textsuperscript{B} haplotypes and diversity differences between wolves and dogs point to a recent selective sweep within wolves (figs. 2 and 3; supplementary table S5, Supplementary Material online). Given slightly lower diversity in the dog K\textsuperscript{B} haplotype than the dog K\textsuperscript{H} haplotype, it is possible that the dog K\textsuperscript{B} allele underwent an older selective event, perhaps due to selective breeding by humans or natural selection. The target of natural selection on the K\textsuperscript{B} allele may not be melanistic coat color, but rather immunological differences between alleles, due to the previously identified role of \textbeta-defensin genes in innate and adaptive immunity (Yang et al. 1999). In dogs, the K locus demonstrates antimicrobial activity against respiratory pathogens (Erles and Brownlie 2010), and there is ample evidence for differential selection in Yellowstone wolves (Coulson et al. 2011; Hedrick 2015). Black heterozygote wolves have a significantly higher overall survival rate than black homozygote wolves (which are rare in the population), implying that the selective effects of a K\textsuperscript{B} allele may have to do with disease rather than pigmentation, since both genotypes have the same coat color (Anderson et al. 2009; Coulson et al. 2011). In fact, black heterozygote wolves of Yellowstone National Park showed higher annual survivorship following the three documented distemper outbreaks than gray homozygous wolves (Yellowstone National Park Wolf Project, unpublished data). Given that multispecies carnivore populations are a reservoir for canine distemper (Almberg et al. 2010), there is a possibility that dogs provide the resistance allele to distemper common in the dog population and in other carnivore species.

With the exception of the Yellowstone wolves, the demographic history of the populations we sampled is uncertain, but population specific demographic events may have reduced haplotype variation that could be confused with selective sweeps. Consequently, we sampled \(~3.74\text{ Mb}\) genic sequence, \(~86.9\text{ kb}\) sequence in similar telomeric positions as the K locus, and \(~5.03\text{ Mb}\) neutral sequence (supplementary table S3, Supplementary Material online), to determine baseline genome wide and genic patterns of variation. Our results show that the K locus core has dramatically reduced variation relative to other genomic regions suggesting this pattern is due to selection, as selection is expected to be locus specific whereas demographic effects should be manifest genome wide. In fact, our specific demographic modeling of the Yellowstone population indicates that selection can explain even changes in haplotype frequency of \(<10%\) from an initial founding value (fig. 6). Additionally, population genetic modeling of the three populations with complex demography shows that patterns of diversity cannot be solely explained by inferred demography and purifying selection alone, suggesting balancing selection is required to explain observed diversity (fig. 9).

A Single Origin of Black Coat Color in North American Wolves

Wolves in North America are unique for having a high frequency of black coat color, approaching 50% of individuals in some populations except those in the high Arctic (Anderson et al. 2009). In the Old World, black wolves are rare and generally restricted to regions where dogs and wolves frequently hybridize (but see Galaverni et al. 2017). The high abundance of black wolves throughout North America suggests that a selective sweep has occurred. Our extensive geographic sampling of wolves from 10 states and provinces within North America implies that the admixture between dogs and wolves first occurred in Northern Canada, and most likely in the Yukon Territory. The ancestral population with regards to K\textsuperscript{B} introgression is predicted to have had the most time for recombination to shorten haplotype blocks and therefore generate greater sequence and haplotype diversity. Wolves within the Northwest Territories demonstrated some of the highest levels of K\textsuperscript{B} haplotype diversity (fig. 4), the lowest LD (fig. 5), and extent of haplotype homozygosity (fig. 3). Wolves from Yukon demonstrate similar patterns of diversity, and so are also candidates for K\textsuperscript{B} origin in wolves, especially given estimated K\textsuperscript{B} allele ages (see below). Additional sampling of wolf populations and dog breeds, especially of Arctic breeds and village dogs, may further elucidate the geographic origin and timing of introgression. Samples from Alaska and Yellowstone lack genetic diversity at the K\textsuperscript{B} haplotype, have high levels of LD within K\textsuperscript{B} haplotypes, and have long tracts of extended homozygosity, which suggest the presence of K\textsuperscript{B} allele is more recent in these populations. The diversity of K\textsuperscript{B} haplotypes and their grouping in phylogenetic trees and network is consistent with a single, ancient origin from Native (pre-Columbian) dogs.

Higher levels of haplotype variation and lower levels of LD suggest that the K\textsuperscript{B} allele was first transferred to wolves in Yukon or Northwest Territories. However, this result is not confirmed by phylogenetic analysis which may be underpowered given the very recent transfer of the K\textsuperscript{B} allele to wolves which we estimate occurred between 1,598 and 7,248 years ago (fig. 8 and table 3). Essentially, few if any phylogenetically informative mutations have appeared subsequent to this event that might provide a well-resolved phylogeny. Therefore, we follow previous analyses which have used patterns of LD and haplotype variation to infer origin of introgression events (e.g. Tishkoff et al. 2001; Gray et al. 2010). Conceivably, demographic events might have occurred in some wolf populations which lowered K\textsuperscript{B} diversity and obscured an origin of introgression there. However, genic and genome-wide patterns of variation outside the K locus do not show lower variation in other populations outside Yukon and Northwest Territories. For example, Yellowstone wolves have high genome-wide diversity but the lowest K\textsuperscript{B} diversity (fig. 18), suggesting the patterns of variation at the K locus are not due to demography.

Several studies exploring geographic origins of Native American dog breeds have suggested that Arctic dog breeds, such as the Inuit sled dog, the Canadian Eskimo dog and the Greenland dog, have archaic mitochondrial DNA haplotypes and show evidence of ancient admixture with wolves (Brown et al. 2013; van Asch et al. 2013). Furthermore, dogs within these populations represent the only living dog breeds with mitochondrial haplotypes that are unique to New World
wolves, and black coat color in these breeds are determined by mutations at the \textit{K} locus. Based on these findings we hypothesize that the original introgression event likely occurred in the High Arctic regions of Northern America, where dogs and native people first coexisted with large wolf populations that shared similar prey (e.g., caribou). Future efforts focused on increasing the sampling of dog breeds may yield insight into the specific dog breed or group of breeds involved in the introgression event. Within our data set, it is notable that the dog breed with the closest \textit{KB} haplotype, the Labrador Retriever, was established in the high latitudes of North America, and may share some ancestry with native North American dogs that were related to the first black dogs which successfully crossed with gray wolves.

The most ancient \textit{K}^{B} allele occurs in Yukon wolves, regardless of the mutation rate used (fig. 8). Using the most current estimates of dog mutation rates (0.3–0.4 \times 10^{-8}; Skoglund et al. 2015; Frantz et al. 2016) would place the \textit{K}^{B} origin in Yukon population at around 1,598–3,500 years ago. This date is much more recent than the arrival of the first dogs 8,500–12,000 years ago (Leonard et al. 2002), and on the more recent end of previously estimated range of 500 – 14,000 years ago (Anderson et al. 2009). Contact from Europeans was limited before the early 19th century, so it is likely that Yukon wolves may have interacted exclusively with dogs of First Nations people. Humans, and likely their dogs, were at relatively low population density, so that encounter rates between dogs and wolves may have been infrequent such that only a single introgression event was successful.

**Limited Genetic Diversity of the K Locus in Yellowstone Wolves**

We found genetic evidence for balancing selection in Yellowstone wolves based on multiple summary statistics and simulations. The Yellowstone population was originally founded in 1995 with 31 wolves from two discrete populations in Alberta and British Columbia, and 10 additional wolves from Montana added the following year (Bangs and Fritts 1996). Given this history, we would expect an increase in heterozygosity and diversity of \textit{KB} haplotypes relative to a single wild population from southwestern Canada. However, the Yellowstone wolf \textit{KB} haplotypes demonstrate very low levels of diversity (supplementary fig. S5, Supplementary Material online and fig. 4), high LD (fig. 5B), and an almost completely monomorphic core region (fig. 3D). This pattern might reflect lower diversity in the founding stock, but we found that other genomic regions of the Yellowstone population had average levels of heterozygosity (fig. 1B). We also show that the few Yellowstone wolves
founders that we sequenced had substantially higher diversity than the current population (supplementary fig. S7, Supplementary Material online), and consequently, diversity within the K locus region has been lost relative to other genomic regions. Results from our simulations show that the K locus is under balancing selection in the Yellowstone wolf population. Yellowstone may be unique among North American wolf populations in that it surrounded by a high density of carnivore groups, such as mustelids, canids, Ursids, felids, and raccoons, that are hosts for disease (Almberg et al. 2010). These other carnivore species are potentially a reservoir for canine disease such as distemper which causes substantial mortality in the Yellowstone wolf population (Stahler et al. 2013) and could be the cause underlying current balancing selection in Yellowstone wolves. Elsewhere, as represented by our wolf populations, the density of host species is lower and the encounter probabilities less.

Conclusions

Using population genetic simulations, we demonstrate evidence of selection at the K locus in North American wolf populations, and confidently rule out genomic location or sampling effects as explanation of previous results. We find that there was likely a single successful introgression event in the Canadian Arctic, that thereafter spread rapidly across the entire continent leaving other wolf populations with lower diversity at the K^6 haplotype. We apply a new approach using both recombination and mutation rate to estimate the time of introgression, and find that the event is very recent, suggesting one of the most rapid spread of an adaptive variant known in vertebrates. This spread was likely facilitated by the great dispersal abilities of gray wolves, which commonly disperse over 50 km before establishing new territories (Mech and Boitani 2003). Further, although the specific focus of selection is uncertain, the canonical function of the K locus is in immunity, and hence the presence of dogs and other carnivores as reservoirs of disease may have augmented the selective coefficient at this locus. Conceivably, the K^6 allele may have first evolved in dogs and persisted because of enhanced immunity in dense dog populations where disease thrives, and then was preadapted for selection in adjoining wolf populations that may have also been infected by diseases from a carnivore reservoir. This hypothesis requires confirmation with functional tests, such as with a CRISPR modified cell line (Shalem et al. 2014). Finally, consistent with this hypothesis, we detect a signature of balancing selection since the founding of the Yellowstone National Park population, which has been known to suffer from recurrent zoonotic disease. These zoonotics include introduced disease such as mange, and also distemper, a disease common in sympatric dog and wild carnivore populations with which wolves may interact in open range country surrounding the park.

Materials and Methods

Sample Selection and Array Design

We selected wolf samples to maximize the following parameters: 1) the geographic distribution of samples across regions and ecotypes (Schweizer, Robinson, et al. 2016; Schweizer, vonHoltz, et al. 2016); 2) the number of samples with known coat color phenotype and life history data; and 3) the quality and quantity of DNA. A total of 382 samples were sequenced, and, after genotype and sample quality filtering, 353 samples were retained for analysis (table 1).

The capture array was designed to reconstruct the origin and spread of the K^6 allele. We extracted putatively “neutral” regions from the dog reference genome (CanFam3.1; fig. 1A) for background estimates of individual relatedness and population demography. Details of the design of these regions have been described elsewhere (Freedman et al. 2014; Schweizer, Robinson, et al. 2016) and follow guidelines set by previous studies in humans (Wall et al. 2008). To characterize genetic variation around the K locus, we designed an extensive resequencing approach for almost 5 Mb surrounding the 3 bp causative deletion on chromosome 16. This design included the 200 kb “core” region partially sequenced previously (Anderson et al. 2009), plus 1 kb segments spaced every 10 kb, extending to the end of the chromosome ~560 kb downstream of the mutation (the K locus is near the telomeres) and 4.2 Mb upstream of the mutation. To test how the proximity of the K locus to the end of the

<table>
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<th>Population</th>
<th>Mutation Rate (/bp/gen.)</th>
<th>TMRCA (years)</th>
<th>95% Credible Interval</th>
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chromosome impacts decay of LD and diversity statistics, we sequenced ten 1 kb fragments spaced on a 200 kb segment in nontelomeric regions for five larger and five smaller chromosomes, plus chromosome 16. Given that dog and wolf chromosomes are acrocentric, we chose the midpoint for each nontelomeric region. Finally, as an empirical background for selection at the \( K \) locus, we assessed decay of LD in other telomeric regions by similarly designing ten 1 kb fragments spaced on a 200 kb segment in telomeric regions for five larger and five smaller chromosomes. Each telomeric region began at the same relative distance from the end of the chromosome as the \( K \) locus mutation on chromosome 16. In total, we captured approximately 10 Mb of sequence from each individual. To do this, 105,000 120 bp RNA baits were developed by MYcroarray (Ann Arbor, Michigan) so as to maximize specificity of bait hybridization and unique mapping within the genome. These baits covered approximately 91% of the regions we aimed to capture.

Library Prep, Target Enrichment, and Sequencing

Samples were prepared as previously described (Schweizer, Robinson, et al. 2016). Briefly, we extracted genomic DNA from blood or tissue, and sheared DNA of high quality and quantity using a Biorupter NGS Sonication System (Diagenode). DNA samples were sheared to approximately 300–450 bp, and before sequencing were randomized with respect to extraction, library prep and enrichment dates to prevent batch effects. Preparation of sequencing libraries followed the with-bead library preparation protocol of Faircloth (2015), and each sample was barcoded with a unique 6 bp index sequence during adapter ligation so as to enable pooling of 24–25 individuals per lane (Faircloth and Glenn 2012). Subsequent to library preparation, samples were target enriched following the manufacturer’s protocol. To check that libraries were enriched for target regions, we performed enrichment qPCR on all samples prior to sequencing (see Supplementary Material online). Enriched libraries were quantified, pooled in equimolar amounts, and sequenced on a HiSeq 2000 with 100 bp paired-end reads by the QB3 Vincent J. Coates Genomics Sequencing Laboratory (Berkeley, California, USA).

Sequence Alignment, Processing, and Genotype Filtration

Sequence alignment and processing followed the Broad Genome Analysis ToolKit “Best Practices” pipeline (https://www.broadinstitute.org/gatk/guide/best-practices), with the specifics of our processing protocol published elsewhere (Schweizer, Robinson, et al. 2016). In short, demultiplexed fastq reads passing the Illumina filter were trimmed for remaining adapter sequences, then forward and reverse reads were aligned and mapped to the reference boxer genome (CanFam3.1) using bwa aln and bwa sampe (Li 2014). Alignment of wolf sequences to CanFam3.1 produces high quality genotype calls and minimal reference bias due to the very short sequence divergence of wolves and dogs (~0.1%; Freedman et al. 2014). A recently published de novo wolf genome might provide a unique future reference for our analysis (Gopalakrishnan et al. 2017). However, the development of this genome is at early stage relative to the dog genome that we used, which was based on Sanger sequencing and minimale level information and annotation. After duplicate removal with samtools rmdup (Li et al. 2009), local realignment with GATK, and fixing mate information with picard (http://broadinstitute.github.io/picard/), we ran GATK Base Quality Score Recalibration using a previously generated set of “known” variant sites (Schweizer, Robinson, et al. 2016).

FIG. 9. Posterior densities for introgression time (in generations) and introgression frequency in null simulations of (A) Alaska, (B) Northwest Territories, and (C) Yukon wolf populations. Red indicates higher probability, whereas blue indicates lower probability (see key for each plot). Our null model includes negative selection and a demography estimated from the data. Under this null model, there must be very recent introgression at a very high frequency, providing support for an alternate model with positive selection.
The GATK UnifiedGenotyper algorithm was used to call SNPs and indels (insertions and deletions) over the capture array intervals with a padding of 1,000 bp. Variant positions identified by the Unified Genotyper were filtered with GATK VariantFilter using ten annotation values, as recommended by the Broad Best Practices pipeline. We also removed variant positions with genotype quality (GQ) < 30 and all positions with depth of coverage (DP) < 10.

Data Quality Control
Genotype concordance was assessed for 109 individuals and 204 sites that overlapped between the Affymetrix dog SNP array v2 and the capture array target intervals (Schweizer, vonHoldt, et al. 2016). Using the vcftools package (Daneczek et al. 2011), we calculated the sequence-wide heterozygosity, transition/transversion ratio, site missingness and individual missingness for each set of regions. Significant differences in heterozygosity among region types were tested in R using a Mann–Whitney test. The K locus indel genotype had previously been determined for 235 individuals using either Sanger sequence (Anderson et al. 2009) or high resolution melt curve analysis (Coulson et al. 2011; Schweizer & B.M. vonHoldt, unpublished data). We used these data for quality control in our sequence samples.

We performed principal components analysis (PCA) within the smartpca package of eigenstrat (Price et al. 2006). First, using neutral sequence data, we verified that samples grouped according to that expected based on previous studies (vonHoldt et al. 2011; Pilot et al. 2014; Schweizer, vonHoldt, et al. 2016). Samples that did not group according to their expected population or species were dropped from further analysis, as they likely represent bad sequencing runs or poor quality samples. For this analysis, we generated a set of LD-pruned SNPs using the “--indep-pairwise 50 5 0.5” option in PLINK (Purcell et al. 2007). We also generated a subset of unrelated individuals using PRIMUS (Staples et al. 2013) and a maximum identity-by-descent of 0.1, as in (Schweizer, Robinson, et al. 2016).

Summary Statistics on Phased Haplotypes
To phase haplotypes, we used SHAPEITv2 since it can handle varying levels of relatedness among individuals (O’Connell et al. 2014). We filtered sites for a 100% call rate in all individuals, a DP ≥10 and variants with GQ ≥30. The remaining 7,761,114 sites were phased by individual chromosome with the following parameters: --burn 10 --prune 10 --main 20 --states 200 --window 0.1 --rho 0.001 --effective-size 20000 --duohmm. Pedigree information from Yellowstone wolves was used by SHAPEIT to improve the accuracy of phasing (O’Connell et al. 2014). We also phased genome data from a Kenyan golden jackal (Canis aureus) for use as the ancestral state (Koepfl et al. 2015).

Using the phased haplotype data, we calculated summary statistics to investigate patterns of polymorphism and divergence within the K locus core and surrounding regions, neutral regions, genic regions, parallel telomeric, and nontelomeric regions. These statistics included the following: π (the average pairwise differences), Watterson’s theta (θw, the average number of segregating sites [Watterson 1975]), and Tajima’s D (a measure of the skew in the site frequency spectrum, which is an indicator of both selection and population history [Tajima 1989]). These statistics were calculated in nonoverlapping windows of 1 kb (for telomeric, nontelomeric, and K locus) or 10 kb (for neutral), in combinations of each sequence type, population, and K^d versus K^y haplotype. All calculations were performed within the Python EggLib package (De Mitia and Siol 2012), and significance of differences in means was tested using a Wilcoxon signed rank test or a Wilcoxon rank sum test, with continuity correction, in R.

Comparison of Selection Signals in K^B versus K^Y
We used two methods to test for signals of selection in dogs or wolves, and whether the partial sweep occurred on the K^d or K^y containing haplotypes. First, we visualized large-scale patterns of variation in haplotypes by generating haplotype structure plots in R 3.1.3 (http://www.R-project.org) with the rehh package (Gautier and Vitalis 2012) that showed the ancestral and derived state of each polymorphic position. These plots were also useful for visualizing regions where recombination events may have occurred. Next, we implemented the extended haplotype homozygosity (EHH) test, which measures the relationship between the frequency of an allele of interest and the amount of LD surrounding it (Sabeti et al. 2002). Once a core mutation is identified (here, the K^d mutation), increasingly distant SNPs are used to measure the decay of LD from the core haplotype. This measure of decay is called the EHH score and provides the probability that two randomly chosen chromosomes out of a population are identical between the core haplotype and the increasingly distant SNP. The phased Kenyan golden jackal sample was used to infer the ancestral state for each variant, and EHH tests were implemented with the rehh package (Gautier and Vitalis 2012).

Comparison of Diversity among Populations Containing the K^B Allele
Using EggLib, we calculated LD decay by measuring the square Pearson’s correlation coefficient (r^2) between each polymorphic SNP, after filtering for a minimum allele frequency of 0.05 in each population. Genomic distances were binned by 30 kb for telomeric regions, and by exponentially increasing distances for the surrounding 5 Mb region, and the decay of r^2 was plotted in R. Given that sample size can affect LD decay, we performed calculations using a random subset of at most eight haplotypes for a few populations with small sample sizes, there were fewer than eight haplotypes (see Results). LD in the K locus core and parallel telomeric regions was calculated separately.

Forward Single-Locus Simulations for Yellowstone Wolves
We used historical census data to obtain a demographic model from which to simulate. We modeled alpha-mating as a reduction in N_e using values suggested in (vonHoldt et al. 2008). We assumed that K^d homozygous individuals did not
survive and used a custom Wright–Fisher forward simulator, sampling the allele frequency from a binomial distribution every generation. Specifically, we simulated eight generations from the founding till the present day. We matched the initial allele frequency and sampled the frequency each generation, retaining the allele frequency from the final generation. We ran 10,000 simulations and used a one-sided test to determine the probability of observing an allele frequency as high or higher than the observation from the data. We used approximate Bayesian computation to infer the dominance coefficient at this locus. We set $s = -1$ and drew $h$ from a uniform distribution over $(-2, 2)$ to include over and under-dominance. We used a rejection sampling approach and kept 500 simulations where the final allele frequency was within 1e-3 of the observed value.

Patterns of Genetic Divergence among Populations
To identify a region showing no signal of past recombination events, we calculated the minimum number of recombinations using the four gamete test of Hudson and Kaplan (1985), implemented in EggLib, in sliding windows of 1,000 bp with 10 bp step size. We visually confirmed that there were no recombinant haplotypes within this region. We used neighbor-joining trees to infer the hierarchy between haplotypes from different geographic localities and the subsequent implications for introgression history. We used EggLib to calculate the pairwise number of differences between each haplotype for the nonrecombining 26 kb $K$ locus haplotype and neutral data sets. We constructed neighbor-joining trees with 5,000 bootstraps using the package ape 3.1-2 in R (Paradis et al. 2004). A haplotype network of the nonrecombining region was constructed within the pegas package of R (Paradis 2010) separately for $K^A$ and $K^B$ containing haplotypes.

Estimating Age of the $K^A$ Allele among North American Wolves
To describe the relative timing of the $K^A$ allele’s spatial spread and/or timing of selection among wolf populations in North America, we estimated time to the common ancestor (TMRCA) for the deletion in each of the four samples. We used a recent method (Smith et al. 2018) which leverages both the decay in LD between the selected allele and nearby sites, as well as the accumulation of new mutations on the selected allele’s ancestral haplotype. If assumptions of the model are reasonably approximated by the data, this approach leads to dramatic gains in accuracy relative to more commonly used Approximate Bayesian Computation methods. Specifically, the method assumes a “star-shaped” genealogy among sites linked to the selected allele’s ancestral haplotype. This is a reasonable assumption in cases where the focal allele is subject to strong positive selection; which for our case is appropriate given the results of reduced polymorphism and Tajima’s D, and elevated EHH scores observed at the selected locus.

A second assumption of this method involves the specification of an appropriate reference panel of haplotypes that do not have the selected allele. This reference panel should approximate the background haplotypes with which the selected haplotype recombined during its increase in frequency. Simulation results show that a reference panel which is too similar to the selected haplotype can lead to underestimates of the true TMRCA while a misspecified reference panel that is too diverged from the true reference panel will overestimate the TMRCA. For this reason, we excluded the sample of dogs for which a suitable reference panel was not available. We used the local reference panel of haplotypes without the selected allele in each of the four samples from natural populations of North American wolves (see Results).

TMRCA estimates can vary depending on the mutation and recombination rates used. To account for locus specific variation in recombination rates across the $K$ locus region, we used a recombination map inferred for dogs based on patterns of LD (Auton et al. 2013). This recombination map, however, does not include the entire sequenced region downstream of the selected site. We predicted the unobserved recombination rates at these sites using the adjacent 4 Mb of observed recombination rates using the smooth spline function in R with the smoothing parameter set to 0.95.

Several estimates of the per basepair per generation mutation rate have been inferred using different approaches. Skoglund et al. (2015) use ancient DNA from a 35,000 year old wolf and to infer a rate of $0.4 \times 10^{-8}$; Frantz et al. (2016) calibrate a molecular clock using radiocarbon dating on an ancient dog to infer a similar mutation rate between $0.3 \times 10^{-8}$and $0.45 \times 10^{-8}$. We report TMRCA estimates using these values in addition to one higher rate of $1 \times 10^{-8}$ (as used in Freedman et al. 2014).

Demographic Inference and Simulations of Selection in Alaska, Northwest Territories, and Yukon Wolves
To further test whether patterns of genetic diversity at the $K$ locus are unusual, we inferred the demographic history of the three remaining wolf populations where black wolves are found in our sampling (i.e., Alaska, Northwest Territories, and Yukon). We used ANGSD v 0.920 (Korneliussen et al. 2014) to generate site frequency spectra for each population from 5,055,985 neutral sites, using the Kenyan golden jackal as the ancestral reference sequence (Koepfl et al. 2015). We used $\delta a i o$ (Gutenkunst et al. 2009) to set up a model where a population undergoes a size change $T$ generations ago to size $N$. We included a parameter for ancestral state misidentification and ran the inference seven different times to ensure we are not optimizing to local minima. We converted time to generations using a mutation rate $0.4 \times 10^{-8}$ (Skoglund et al. 2015) and used 5,055,985 sites as the sequence length.

Next, we simulated data under these demographies using forward simulations as described above for the Yellowstone population. Since we do not know the initial introgression frequency (i.e., the starting frequency of the $K^A$ allele) nor the timing of the introgression, we estimated these parameters using an ABC rejection sampling approach. We drew these parameters from uniform prior distributions and kept simulations where the error was under 0.1, defined as
error = \frac{|F_o - F_s|}{F_o}.

where \(F_o\) is the observed present day allele frequency and \(F_s\) is the simulated final allele frequency. We then simulated data using a forward simulation algorithm that samples the allele frequency for each generation using binomial sampling. We apply selection every generation, assuming a recessive lethal condition for the derived allele. We retained 100 pairs of posterior parameter estimates for each population and plotted their joint distribution using a two-dimensional kernel density estimate (Venables and Ripley 2002).

Supplementary Material
Supplementary data are available at Molecular Biology and Evolution online.

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Author Contributions

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