Molecular and Evolutionary History of Melanism in North American Gray Wolves

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Morphologic diversity within closely related species is an essential aspect of evolution and adaptation. Mutations in the Melanocortin 1 receptor (Mc1r) gene contribute to pigmentation diversity in natural populations of fish, birds, and many mammals. However, melanism in the gray wolf, Canis lupus, is caused by a different melanocortin pathway component, the K locus, that encodes a beta-defensin protein which acts as an alternative ligand for the Mc1r. We show that the melanistic K locus mutation in North American wolves derives from past hybridization with domestic dogs, has risen to high frequency in forested habitats, and exhibits a molecular signature of positive selection. The same mutation also causes melanism in the coyote, Canis latrans, and Italian gray wolves, and hence our results demonstrate how traits selected in domesticated species can influence the morphologic diversity of their wild relatives.

The correspondence between coat color and habitat is often attributed to natural selection but rarely is supporting evidence provided at the molecular level. In North American gray wolves, coat color frequencies differ between wolves of forested and open habitats throughout western North America (1), including Denali National Park (2) and the Kenai Peninsula in Alaska (3), and much of the Canadian Arctic (4, 5). These differences are especially dramatic between wolves of the high tundra that are migratory and follow barren ground caribou to their breeding areas, and wolves that are year-round residents in the neighboring boreal forest and hunt non-migratory prey. Dark-colored wolves are extremely rare in the tundra, but increase in frequency along a Southwest cline towards forested areas (Fig. 1A). The potential selective value of dark vs. light coat color has been suggested to include concealment during predation, and/or indirect effects due to pleiotropy, but remains unresolved because the underlying gene(s) have not been identified (5–7).

In many vertebrates, natural pigmented variation is controlled by the Agouti—Melanocortin 1 receptor (Mc1r) pathway, a ligand receptor pair that modulates the amount and type of pigment, red/yellow phaeomelanin or brown/black eumelanin, produced by melanocytes in skin, hair, or feathers. Gain-of-function Mc1r mutations are well-recognized causes of melanism in many domestic and laboratory animal species (8, 9), as well as in several natural populations of birds (10), rodents (11, 12), and canids (13). Recently, we found that pigment-type switching in domestic dogs involves an additional component of the melanocortin pathway, the K locus, which encodes a beta-defensin protein, CBD103 (14, 15).

Coat color in Canadian wolves is genetically complex, with phenotypes ranging from white to gray to black, and is also confounded by an independent effect of graying with age (Fig. 1B). However, in Yellowstone National Park, where a small number of founder animals from Canada were recently reintroduced (16, 17), gray and black coat colors segregate as a Mendelian trait. We surveyed molecular variation in Agouti, Mc1r, and CBD103 in wolves from North America and identified several Mc1r and Agouti polymorphisms. However, none of these were predicted to affect gene function, and did not associate with black coat color (table S1). By contrast, in a 14 member, 3 generation kindred from Yellowstone, we observed complete co-segregation between black coat color and markers at the K locus (LOD = 4.21 at θ = 0, Fig. 1C), which is unlinked and lies on a different chromosome from Agouti and Mc1r.

In dogs, the ancestral CBD103 allele (k) confers normal Agouti and Mc1r gene action, whereas a 3 bp deletion...
(CBD103\textsuperscript{4G23} or \(K^b\)) suppresses Agouti gene action, leading to dominant inheritance of a black coat (14, 15). We observed the same 3 bp deletion in 102/104 black-colored wolves from Yellowstone, and 9/9 from the Canadian Arctic. Conversely, CBD103\textsuperscript{4G23} was absent from 120/120 gray-colored wolves from Yellowstone, and 22/22 white-colored wolves from the Canadian Arctic (Table 1). We also found CBD103\textsuperscript{4G23} in 6/10 gray-colored wolves from the Canadian Arctic, suggesting that gray coat color can result either from the absence of CBD103\textsuperscript{4G23} and a modified Agouti phenotype (in which individual hairs contain both cream-colored phemeomelanin and dark eumelanin), or from secondary factors such as age that dilute pigmentation of hairs that contain only eumelanin. [Additional genealogy studies of the Yellowstone population (17) together with the paucity of McIr variation in wolves (table S1) suggests that black coat color reported for the 2 \(k^b/k^b\) Yellowstone wolves is likely to reflect phenotypic ambiguity or misclassification at the time of sampling.] Allele frequencies for CBD103\textsuperscript{4G23} in tundra and forest wolves overall were estimated at 0.02 and 0.19, corresponding to phenotype frequencies of 2% - 33% and 33% - 64% for dark wolves in tundra and forest populations, respectively (Fig. 1A) (4).

To investigate the evolutionary history of the melanistic K allele, we sequenced 8 single-copy non-coding segments distributed across a ~150 kb region centered on CBD103 in 32 Arctic and 15 unrelated Yellowstone wolves, as well as in 12 domestic dogs, 6 \(k^b/k^b\) (akita, basenji, boxer, bulldog, Doberman pinscher, great dane) and 6 \(K^b/K^b\) (curly-coated retriever, Dalmatian, great dane, Labrador retriever, poodle, Portuguese water dog). We identified 52 biallelic polymorphisms across all canids (36 in wolves), and estimated haplotype structure (tables S3 and S4, Fig. 2B, and fig. S2). The rate of polymorphism among all wolf amplicons was 1 SNP per 510 bp (Watterson's estimator, \(\theta_w = 1.96 \times 10^{-3}\)), similar to genome-wide measurements of polymorphism between the Boxer and the gray wolf (1/580 bp) and the coyote (1/420 bp) (18). However, partitioning our data according to \(K\) locus genotype and proximity to CBD103 revealed little or no polymorphism among \(K^b\)-bearing chromosomes close to CBD103, rising to levels at or above those observed in \(k^b\)-bearing chromosomes in the 75 kb spanning either side of the locus (Fig. 2A). This pattern, and the analogous one for nucleotide diversity (\(\pi\), fig. S1), is also reflected in a significant difference in haplotype diversity between \(K^b\) (8 unique of 22 total) and \(k^b\) (59 unique of 72 total) chromosomes (\(\chi^2 = 14.2, p < 0.001\)). Together with the correlations between coat color and habitat (2–5), the combination of low diversity and high frequency suggest that \(K^b\) has been under positive selection in North American forest wolves.

Overall, the patterns of linkage disequilibrium (LD) across 150 kb surrounding the \(K\) locus were similar to comparisons between different breeds of domestic dogs (18), with relatively small haplotype blocks, including a ~4 kb CBD103 core region within which there is no evidence for historical recombination (Fig. 2C). However, different evolutionary histories for the Arctic wolf \(K^b\) and \(k^b\) alleles were apparent when the SNP patterns (Fig. 2B) were depicted as haplotype bifurcation diagrams (Fig. 2D), which highlight a central region of ~60 kb devoid of polymorphism among wolf \(K^b\) haplotypes. This characteristic, and the corresponding difference between \(K^b\) and \(k^b\) chromosomes, were represented quantitatively by the EHH (extended haplotype homozygosity) statistic (19), which is the empirical probability that two chromosomes chosen at random remain identical at progressively increasing distances from CBD103. As depicted in Fig. 2, E and F, the distribution of EHH was considerably broader for \(K^b\) compared to \(k^b\) chromosomes in wolves, whereas the distributions were nearly identical for \(K^b\) compared to \(k^b\) chromosomes in dogs. Together with additional analyses of genome-wide SNP data (SOM text, fig. S3), these observations suggest that \(K^b\) has risen to high frequency by a selective sweep.

As with black dogs and melanistic wolves, CBD103\textsuperscript{4G23} was associated with coat color in 67 coyotes (6 black and 61 gray, Table 1 and table S2). These findings suggest three possible evolutionary histories. First, the 3 bp deletion may be relatively old, having occurred in a canid ancestor more than 1 million years ago prior to the divergence of coyotes from wolves. Second, the 3 bp deletion may have occurred more recently in one of the species, followed by introgression into the others. Finally, the 3 bp deletion may represent a mutational hotspot, having recurred independently in coyotes, wolves, and dogs. To distinguish among these possibilities, we ascertained and compared coyote haplotypes (6 \(K^b\) and 18 \(k^b\)) with those from the North American wolf and dog.

The pattern of haplotype diversity for all 3 canids was similar to that observed in wolves alone, and showed significantly less diversity among \(K^b\) (15 unique of 40 total) relative to \(k^b\) (66 unique of 102 total) chromosomes (\(\chi^2 = 9.7, p = 0.003\)). Of the 15 unique \(K^b\) haplotypes, one haplotype was observed in 3 coyotes and 6 dogs, and a second haplotype was observed in 2 coyotes and 13 wolves (Fig. 3A). However, none of the 66 unique \(k^b\) haplotypes were observed in more than one species (fig. S2).

Reconstruction of a phylogenetic network for the entire 150 kb region is complicated by historical recombination between extant \(K^b\) and \(k^b\) chromosomes (e.g. arrows in Fig. 2B), and lack of a suitable approach for inferring accurate gene genealogies in the presence of recombination (20). However, by focusing on the 4 kb CBD103 core region (Fig. 2C) a simple neighbor joining tree was constructed for 18
core region haplotypes representing 142 (94 wolf, 24 dog, and 24 coyote) chromosomes (Fig. 3B). In this tree, all the \(K^B\) chromosomes define a 2 haplotype cluster, whereas the remaining 16 haplotypes (which represent all the \(k^B\) chromosomes) are more dispersed. Furthermore, many of the \(k^B\) chromosomes cluster by species (9/12 of the dog, and 44/72 of the wolf), unlike the \(K^B\) chromosomes. This contrasting phylogenetic pattern suggests that the \(k^B\) mutation occurred in a single species, and was later distributed among dogs, wolves, and coyotes by interspecific hybridization. [The 24 \(k^B\) haplotypes from coyotes are no closer to each other than to \(k^B\) haplotypes from wolves or dogs (Fig. 3B), which is consistent with their history of hybridization with other canids (21)].

To gain additional insight into how \(K\) locus variation in dogs and wolves arose, we estimated coalescent time to the most recent common ancestor (TMRCA) as a function of cumulative distance from \(CBD103\) for \(k^B\) and \(K^B\) chromosomes from wolves, dogs, and both groups together. We applied a molecular clock approach to sequencing data from individual amplicons across the entire 150 kb region (Fig. 2), which assumes that mutations occur at the same constant rate at all sites in wolves and dogs, and integrates the effects of both recombination and demography (22). Close to \(CBD103\), TMRCA estimates were near zero for all \(K^B\) subsets (Fig. 3C) because there is little or no polymorphism in this region (Fig. 3A). However, at greater distances from \(CBD103\) (10 – 50 kb), estimates for dog chromosomes are similar to those of dog and wolf chromosomes considered together, regardless of genotype. This suggests that \(K^B\) in dogs is sufficiently old to have undergone extensive recombination with \(k^B\) chromosomes, and that the recombination history includes hybridization between dogs and wolves. However, in the same 10 – 50 kb range, TMRCA estimates for wolf \(K^B\) chromosomes were considerably less than those from dog \(k^B\) chromosomes (or from dog and wolf \(K^B\) chromosomes considered together), suggesting that \(K^B\) was introduced into North American wolves from dogs, not vice versa.

Introggression of \(K^B\) from dogs into North American wolves is also supported by geographical and ecological considerations. \(K^B\) is widely distributed among domestic dogs, including ancient breeds originating in Asia and Africa. In wolves, however, melanism has only been reported outside North America in Italy, where it is associated with molecular and/or morphologic evidence of recent hybridization with free-ranging dogs (23). Indeed, we also examined 22 samples from the Italian Apennines, and observed \(K^B\) in 6 of 7 black “wolves” (including 1 previously classified to be a dog-wolf hybrid) but 0 of 15 gray wolves. By contrast, genome-wide SNP analysis of 10 \(K^B/k^B\) and 10 \(k^B/k^B\) North American wolves showed no evidence for recent dog-wolf hybridization (SOM text, fig. S3B).

The dog was domesticated between 15,000 – 40,000 years ago in East Asia from gray wolves (24, 25), and we estimate that \(K^B\) is at least 46,886 years old (95% confidence limit: 12,779 - 121,182), therefore we cannot distinguish whether \(K^B\) arose before or after domestication. However, if \(K^B\) arose in Old World wolves prior to domestication, our data indicate that it must have been lost from the gene pool and reacquired in North America, perhaps from Native American dogs that accompanied humans across the Bering Strait 12,000 – 14,000 years ago (26) (Fig. 3D).

The wolf in the United States faces grave threats, in some cases by eradication, and in others, by hybridization, such as in the Great Lakes region (27). However, apparent selection for the \(K^B\) locus in North American gray wolves shows how genetic diversity —preserved by humans in domestic dogs— may flourish in wild wolf populations. As the available tundra habitat declines due to development and/or global warming, the frequency of the \(K^B\) mutation may increase further in northern latitudes. Thus, introduction of genetic diversity into a natural population from a mutation originally selected in domesticated animals may, ironically, provide a mechanism to adapt to a changing environment. Interspecific hybridization has been widely observed between other domesticated species of animals and plants (28–30). Our results imply that variants which appear under domestication can be viable in the wild and enrich the genetic legacy of natural populations.

References and Notes
31. Supported by grants from the National Institutes of Health (G.S.B.), the National Science Foundation (R.K.W.), and the Swedish Research Council (J.A.L.).
32. H. Chen and S. Schmutz for advice, to H. Manuel for technical assistance, and to members of the USDA Wildlife Services and private citizens for assistance with sample collection. Sequences generated in this study are deposited in GenBank accessions FJ609634 - FJ609641.

Supporting Online Material
www.sciencemag.org/cgi/content/full/1165448/DC1
Materials and Methods
Figs. S1 to S4
Tables S1 to S5
References

3 September 2008; accepted 15 January 2009
Published online 5 February 2009; 10.1126/science.1165448
Include this information when citing this paper.

Fig. 1. Distribution of melanism and K locus genotypes in North American gray wolves. (A) Location and coat color phenotype of Canadian samples used here and as described (4). (B) Age-related graying and the associated difficulty of inferring genotype from phenotype in gray animals. Each pair of photos shows the same individual at different ages (10 months and 10 years) and documents an increasingly gray appearance at 10 years reflecting dilution of eumelanin and pheomelanin in the k'/k individual. Images courtesy of Monty Sloan, Wolf Park, Battle Ground, Indiana. (C) Co-segregation of Kβ and black coat color in a three generation pedigree from the Leopold pack in Yellowstone National Park (17). ΔG indicates the dominant Kβ allele, while + indicates the wild type allele, k'.

Fig. 2. Polymorphism and haplotype structure of the K locus in North American gray wolves [(A) to (E), 1 Kβ/kβ, 20 Kβ/k'/k' and 26 k'/k'] and domestic dogs [(F), 6 Kβ/kβ, and 6 k'/k']. (A) Polymorphism (Watterson’s theta, θw, +/- sd) as a function of distance from CBD103. (B) Wolf haplotype structure was inferred on the basis of 36 SNPs; each row represents a Kβ- or k'-bearing chromosome, blue and yellow squares represent the major and minor alleles, respectively, gray squares represent missing data. Red and black arrows indicate examples of haplotypes likely to represent historical recombination and 5' and 3' ends of the locus, respectively. (C) Pairwise linkage disequilibrium (LD) values (expressed as D') for all wolf chromosomes; red outline indicates a core region (as in Fig. 3) unlikely to have undergone historical recombination. (D) Haplotype bifurcation diagrams for Kβ- or k'-bearing chromosomes, in which the central blue dot represents CBD103, branches represent haplotype divergence, and the thick of the line is proportional to the number of chromosomes. Extended haplotype homozygosity (EHH) for Kβ- or k'-bearing chromosomes in wolves (E) and dogs (F) as a function of distance from CBD103.

Fig. 3. Evolutionary relationships and history of the K locus in canids. (A) Kβ haplotype structure in wolf-like canids based on genotypes defined by 52 SNPs. Each row represents a Kβ- or k'-bearing haplotype found in coyotes (C), dogs (D), or wolves (W) listed with their respective frequencies on the right, and colored as Fig. 2B. (B) Inferred genealogic relationships of the core region (Fig. 2C) haplotypes (with bootstrap values from 500 replicates shown next to branches). Each branch represents 1 of 18 different haplotypes with the number of chromosomes for each haplotype indicated underneath according to species. (C) Time to most recent common ancestor (TMRCA) estimates for indicated chromosome subsets calculated according to a molecular clock (22), and expressed as a fraction of the divergence time for all wolf-like canids. Individual points represent sets of chromosome segments whose relative TMRCA increases as a function of distance from CBD103, presumably due to ancient hybridization and recombination. (D) Timeline scenario for K locus evolution in dogs and wolves, in which ancestral k' chromosomes are indicated in orange, derivative Kβ chromosomes in gray, and recombinant chromosomes as an orange-gray checked pattern. The k' to kβ mutation may have overlapped or even predated domestication, but
introgression of $K^B$ into North American gray wolves is more recent.
Table 1. Distribution of CBD103 alleles in wolves and coyotes.

<table>
<thead>
<tr>
<th>Animal and location</th>
<th>Total no.</th>
<th>No. carrying $K^B$</th>
<th>White</th>
<th>Gray</th>
<th>Black</th>
</tr>
</thead>
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<tr>
<td>Forest wolves*</td>
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<td></td>
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<td>2</td>
<td>7</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>0</td>
<td>1</td>
<td>7</td>
</tr>
<tr>
<td>Tundra/taiga wolves*</td>
<td></td>
<td></td>
<td>10</td>
<td>8</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>0</td>
<td>5</td>
<td>2</td>
</tr>
<tr>
<td>Yellowstone wolves</td>
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<td>0</td>
<td>120</td>
<td>104</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>N/A</td>
<td>0</td>
<td>102</td>
</tr>
<tr>
<td>Coyotes‡</td>
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<td></td>
<td>0</td>
<td>61</td>
<td>6</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>N/A</td>
<td>0</td>
<td>6</td>
</tr>
</tbody>
</table>

*Forest and tundra/taiga wolves are from the Canadian Arctic (Fig. 1A). The overall frequency of dark (gray or black) wolves is 62% and 7% in the forest and tundra/taiga, respectively (4), and the genotype distributions shown do not represent population-based frequencies. All forest and tundra/taiga wolves carrying $K^B$ were $K^B/k^y$; in the Yellowstone population, 10 were $K^B/K^B$ and 92 were $K^B/k^y$. †This categorical designation of phenotypes, as defined at sample collection, does not fully capture the spectrum of normal coat color variation as indicated in Fig. 1B. ‡Gray coyotes surveyed were from Nebraska (30) or West Virginia (30); black coyotes were from Minnesota (2) or West Virginia (4).
A map showing the distribution of phenotypes in different regions of Canada. The map indicates the presence of dark and light phenotypes in areas such as Tundra/Taiga and Boreal Coniferous Forest.

B. Images of two wolf phenotypes:
- **CBD103 ΔG/+ (K^B/k^Y)**: 10 mo (dark) and 10 yrs (light).
- **CBD103 +/+ (k^Y/k^Y)**: 10 mo (light) and 10 yrs (light).

C. LOD score calculation for Yellowstone National Park:
- LOD score = 4.21 at θ = 0.