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Roan, ticked and clear coat patterns in the canine are associated with three haplotypes near usherin on CFA38

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Summary

White coat patterning is a feature of many dog breeds and is known to be coded primarily by the gene microphalmia-associated transcription factor (MITF). This patterning in the coat can be modified by other factors to produce the attractive phenotypes termed 'ticked' and 'roan' that describe the presence of flecks of color that vary in distribution and intensity within otherwise 'clear' white markings. The appearance of the pigment in the white patterning caused by ticking and roaning intensifies in the weeks after birth. We applied genome-wide association to compare English Cocker Spaniels of roan phenotype (N = 34)with *parti-color* (non-roan) English Cocker Spaniels (N = 9) and identified an associated locus on CFA 38, CFA38:11 057 040 ($P_{\text{raw}} = 8.9 \times 10^{-10}$, $P_{\text{genome}} = 2.7 \times 10^{-5}$). A local case– control association in English Springer Spaniels comparing 11 ticked and six clear dogs identified indicative association with a different haplotype, CFA38:11 122 467G>T $(P_{raw} = 1.7 \times 10^{-5})$ and CFA38:11 124 294A>C $(P_{raw} = 1.7 \times 10^{-5})$. We characterize three haplotypes in Spaniels according to their putative functional variant profiles at CFA38:11 111 286C>T (missense), CFA38:11 131 841-11 143 239DUP.insTTAA (using strongly linked marker CFA38:11 143 243C>T) and CFA38:11 156 425T>C (splice site). In Spaniels, the haplotypes work as an allelic series including alleles (t, recessive clear; T, dominant ticked/parti-color; and T_R, incomplete dominant roan) to control the appearance of pigmented spots or flecks in otherwise white areas of the canine coat. In Spaniels the associated haplotypes are t (CCT), T (TCC) and $T_{\rm R}$ (TTT) for SNP markers on CFA38 at 11 111 286C>T, 11 143 243C>T and 11 156 425T>C respectively. It is likely that other alleles exist in this series and together the haplotypes result in a complex range of patterning that is only visible when dogs have white patterning resulting from the epistatic gene Micropthalmia-associated transcription factor (the S-locus).

Keywords coat-color, pigmentation, USH2A, usherin

Introduction

Domestic dogs exhibit a wide range of coat colors and coat patterns, including white patterning that may occur with or without pigmented spots or interspersed hairs. The broader absence of pigmentation in defined regions of the coat (white patterning) is primarily affected by the activity of the

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micropthalmia-associated transcription factor (MITF) gene (Karlsson et al. 2007). White patterning on dogs may range from as little as a white chest spot on an otherwise selfcolored (or solid-colored) dog to an entirely white dog (known as extreme white). The extent of the white patterning is increased by the presence of a small interspersed nuclear element and affected by a length polymorphism, both in the vicinity of the promoter region of the MITF gene. The longer the length polymorphism, the more extensive is the white patterning on the dog (Körberg et al. 2014).

Two well-described variations of white patterning are roan (Fig. 1a) and ticked (Fig. 1b), which are both

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characterized by the presence of pigmented hairs in otherwise clear white patterned areas of the coat. Pups that will exhibit the pattern variants as adults are born with clear white patterning and pigmentation develops in the first weeks of life (Fig. 1c). Roan and ticked are described by various names across dog breeds, including roan in the English Cocker Spaniel (ECS) and German Short-haired Pointer, blue in the Australian Cattle Dog and Belton in the English Setter. Colloquially, ticked is regarded as the appearance of pigmented spots in white areas of the body. Pigmented spots from *ticking* range in size from only a few hairs to coin-sized, and are most dense on the muzzle and legs (Fig. 1b). In contrast, pigmentation in the white patterning associated with roan is a mixture of pigmented and white hairs throughout the areas of the coat that would otherwise be white (Little 1953), with the phenotype dispersed relatively evenly across the body. ECSs that have visible white markings lacking roan are known as parti-color. The white-patterned areas in parti-color ECSs frequently contain pigmented spots (are *ticked*), but may be *clear*.

Interestingly, the spots that are characteristic of the Dalmatian dog breed share two features with roan: the appearance of pigmented spots in white-patterned areas in the first weeks after birth and the dispersal of the pigmentation evenly across the body. Unlike roan, however, Dalmatians have well-defined pigmented spots in the white-patterned areas of the coat rather than interspersed

Figure 1 Coat patterning in Spaniel breeds: (a) *roan*; (b) *ticked*; and (c) appearance of pigmentation in white-patterned area of coat is not present in pups before loss of birth coat. The image demonstrates a disparity in ticked phenotype between pups and their dam (images under license from iStockPhoto.com).

pigmented and white hairs. The size and definition of the spots in the Dalmatian breed are predicted to be controlled by a locus that is in LD with the hyperuricosurea locus on CFA3 (Bannasch *et al.* 2008).

In classic literature, the two phenotypes of *ticked* and *roan* have been presumed to be controlled by two separate loci (Little 1953); however, even in the early characterization of the trait it was accepted that the phenotypic observations might equally have been explained by a multiallelic single locus. The accepted phenotypic dominance hierarchy places *roan* as dominant over *ticked* and both dominant over 'clear' (no pigmented spots or few spots). However, the range of the phenotypes displayed by individuals suggests a complex inheritance.

The hue of the pigmented hair associated with *ticked* or *roan* pigmentation patterns is controlled by many separate coat color loci (Little 1953). When individuals have red/ yellow pigmented base color, it is sometimes difficult to discern light *roan* from lightly *ticked* individuals, particularly when the dogs are young. Interestingly, pigmented skin spots in the absence of corresponding pigmented fur appear in *non-ticked* dog breeds, although the inheritance of this characteristic is as yet unexplored.

In the current study, we apply within-breed genome-wide association mapping to identify a locus that is strongly associated with the appearance of pigmentation in white patches of the coat in the ECS and English Springer Spaniel



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(ESS). Spaniels are a group of dogs in the sporting group that includes dogs bred for indicating, flushing and retrieving small game (American Kennel Club 2020). As the name implies, Spaniel breeds are derived from dogs that were taken from Spain to Britain in the Middle Ages, where the modern breeds were developed (Australian National Kennel Council 2006). Alongside other Spaniel breeds such as the American Cocker Spaniel and the Cavalier King Charles Spaniel, the ECS and ESS are broadly described as small to medium-sized short-legged dogs with large, drooping ears. Although they are closely related, the ECS can be distinguished from the ESS by its smaller stature.

Materials and methods

Samples

Information relating to dogs used in the various analyses follows.

Samples used in genome-wide association

Forty-three dogs of the ECS breed – 34 roan (19 male and 15 female) and nine *parti-color* individuals (seven female, two male) – were analyzed by genome-wide association. All were privately owned dogs that were sampled on presentation at Norwegian veterinary clinics for health checks, behavioral consultation or medical procedures unrelated to their coat color. All samples were collected as part of diagnostic procedures and remains were stored with the owner's consent. Coat colors were visually phenotyped at the time of sample collection. *Self-colored* ECSs were excluded from the association analysis because the segregation of *roan* and *parti-color* cannot be observed on this genetic background – although the patterns are expected to segregate freely in HWE in the absence of white overlay.

Samples used for haplotype refinement

Limited genotyping with a denser group of markers was used to identify haplotypes segregating throughout the region of association with the *roan* coat pattern. The analysis used nine of the *roan* and nine *parti-color* ECSs from the same group as used in the genome-wide association. To test whether regional association separated *ticked* and *clear* individuals, a cohort of privately owned coat-phenotyped ESS dogs (six *clear* and 11 *ticked* with ticking observed by photograph ranging from light to heavy) was assayed using the same markers.

Samples used for representative haplotype variant calling

Electronic data comprising a variant call file were available from a 590 dog resource published by the Dog Biomedical Variant Consortium (DBVC) (Jagannathan *et al.* 2019). The DBVC data were used for identifying putative functional variants on haplotypes observed in the homozygous state among the phenotyped ECSs and ESSs with the denser marker set.

Samples used to validate roan-associated variants

Blood samples for all dogs were collected in ethylenediaminetetraacetic acid-coated tubes and genomic DNA was isolated using DNeasy Tissue kit (Qiagen GmbH). All nucleic acid measurements and quality assessments were performed with a NanoDrop 1000 spectrophotometer (Thermo Fisher Scientific Inc.). Samples were collected under institutional ethics guidance (approval numbers N00/9-2009/3/5109, 2015/902 and 2018-1449).

Samples for the determination of ancestral alleles

Electronic data from wolf samples (N = 3) from the DBVC data (Jagannathan *et al.* 2019) and from Plassais *et al.* (2019) (N = 48) were assessed at loci represented in Table S2 to determine allelic history.

Genotyping and analysis

Genome-wide association analysis

DNA samples for within-breed genome-wide association analysis were purified according to Affymetrix (Affymetrix) sample preparation guidelines and genotyped using the Affymetrix Version 2 (49 663 markers) Custom Canine SNP array (Affymetrix). They were called using Affymetrix's snp5-geno-qc software.

The within-breed (ECS) genome-wide association analysis was conducted using standard case–control association analysis in PLINK (Purcell *et al.* 2007). SNPs with a genotyping rate of <10%, with strong deviation from Hardy–Weinberg expectation (P < 0.0005) and MAF of lower than 0.05 were removed. Genome-wide significance was ascertained by Bonferroni correction using the adjusted option in PLINK (Purcell *et al.* 2007).

Haplotype refinement analysis

To define haplotypes, the primary regions of the association were assayed by 53 SNPs, including nine markers from the array and 44 new variants in the region. The samples were genotyped using the iPLEX Gold Mass ARRAY (Sequenom; Gabriel *et al.* 2009). The genotype data were analyzed using the TYPER 4.0 ANALYZER software (Sequenom) for cluster analysis. SNPs with a call rate of greater than 75% (--geno 0.25) and a MAF of at least 5% were included in each analysis. Samples with a call rate of less than or equal to 75% (--mind 0.25) were excluded from further analysis.

Genotypes from two dog breeds (ECS and ESS) were visually phased to identify haplotypes in the region of the most associated markers. To prevent ambiguity of haplotype calling, phasing was restricted to animals that were homozygous throughout the region of association. Tagging variants were identified to enable clear definition of observed homozygous haplotypes (Table S2).

Representative haplotype variant calling analysis

Coat patterns related to the mapped locus were not expected to affect animal genetic fitness (reproduction and longevity), thus the causal mutations for observed major phenotypes were expected to segregate in many breeds. This assumption enabled us to use haplotype representatives from general canine whole-genome sequencing panels as proxies for homozygous ECSs and ESSs to identify potential functional variant sites in strong LD with the ECS/ESS observed haplotypes (Table S3).

The TABIX tool (Li 2011) was used to extract called variants from the relevant region of the DBVC variant call file (vcf) (Jagannathan *et al.* 2019). From the ECS/ESS haplotype tag genotypes, individual DBVC dogs representing the relevant observed homozygous haplotypes throughout the region of association were identified. PLINK (Purcell *et al.* 2007) was used to convert the DBVC region to a standard tped file and to reduce the data to include only the homozygous haplotype representative dogs. Data were reconverted to vcf format to enable phasing of functional variants to individual haplotypes.

Variant call files from the representative individuals from across the region of association were uploaded as custom tracks onto the UCSC genome browser (genome.ucsc.edu). The VARIANT ANNOTATION INTEGRATOR (VAI) function in the UCSC genome browser (Hinrichs *et al.* 2016) was used to annotate potential functional SNP and small insertiondeletion variants within the associated region for each individual.

Structural variants were visualized by plotting nucleotide mean variant cover from the DBVC data. Dogs were encoded and grouped according to their homozygosity for representative haplotypes shown in Table S3. Other dogs and canids were coded as unknown.

Validation analysis

Seven putative functional variants in the critical region were genotyped in 10 ESSs (various ticking density) and six ECSs [*parti-color* (4), *roan* (2)] by Sanger sequencing (Australian Genome Research Facility, Sydney, Australia) employing capillary separation on an AB 3730xl machine. Primer sequences are provided in Table S4.

The presence of an observed structural variant was further investigated by PCR in 72 individuals [ECS *roan* (23), ECS *parti-color* (5), ECS *self-color* (black) (7), ESS *ticked* (6), ESS *clear* (4), Dalmatian (23), Australian Cattle Dog (2) and German Short-haired Pointer (2)]. Primers were designed using Primer3 (Untergasser *et al.* 2012) to amplify the intervening product between the tandem duplicates. The PCR reaction was performed in a total volume of 20 μ l using AmpliTaq Gold 360 Master Mix (Applied Biosystems). Each reaction contained 25 ng of template DNA and 1 μ M of each primer (forward TCAACATCCATACTTCTCATGCTT; reverse TGCTCAACCTAAATGTGCTAA) at an annealing temperature of 55°C. PCR conditions were heat activation for 15 min at 95°C followed by 35 cycles of 45 s at 95°C, 45 s at 55°C and 1 min 30 s at 72°C and terminated with a final elongation step at 72 °C for 10 min. PCR products were visualized on a 1% agarose gel.

Results

Genome-wide association analysis

In the ECS genome-wide association analysis 26 014 SNPs were included after filtering. A total of 10 695 SNPs failed missingness, 16 067 failed the frequency test and 132 were excluded owing to deviation from HWE.

The whole-genome association, Q–Q distribution of probabilities and multidimensional scaling plots are shown in Fig. 2. An interval of genome-wide significant association with roan coat color spans CFA38:10 268 280–11 341,635 (CANFAM3.1). The 10 most strongly associated variants from genome-wide association are shown in Table S1.

Haplotype refinement

Association over the denser marker set revealed the strongest association at CFA38:11 069 051G>A ($P_{\text{raw}} = 8.0 \times 10^{-8}$) for *roan* vs. *parti-color* in ECSs. Among ESSs, two markers demonstrated indicative association with *ticked* vs. *clear*: CFA38:11 122 467G>T ($P_{\text{raw}} = 1.7 \times 10^{-5}$) and CFA38 11 124 294A>C ($P_{\text{raw}} = 1.7 \times 10^{-5}$).

Haplotype refinement analysis of both ECS and ESS data revealed the presence of three major haplotypes (*t, clear*; *T, ticked*; and $T_{\rm R}$, *roan*). Other minor haplotypes exist in heterozygous animals (Table S2). ECSs genotyped as homozygous *ticked* but phenotyped as *parti-color* exhibited the same haplotype at associated markers as some ESSs determined as *ticked*. Animals homozygous for the *parti-color* haplotype did exhibit pigmented spotting in their whitemarked coat areas. Typically, these spots were more common at the muzzle and paws.

The defined area of investigation was restricted to a region where markers for ESSs with the *clear* haplotype agreed with the reference genome (CANFAM3.1 – Boxer breed) markers (known to be *clear* phenotype). This encompassed markers between and exclusive of CFA38:11 089 995–11 176 811 (Table S2). Two markers in this region (CFA38:11



Figure 2 (a) Negative log of probability of wholegenome association of *roan* vs. *non-roan* in the English Cocker Spaniel (nine *non-roan* and 34 *roan*) (b) Q–Q plot of association. (c) Multidimensional scaling plot of first two components of variation in samples. (d) Negative log of probability of local association of *roan* vs. *non-roan* in English Cocker Spaniels and *ticked* vs. *clear* in English Springer Spaniels.

163 111 and CFA38:11 163 250) did not differentiate the observed haplotypes and were unrepresented in the public data, suggesting that they were monomorphic.

ESSs that had been determined as *clear* had few (if any) spots. Homozygosity for the reference genome haplotype was not observed in the ECS samples analyzed. *Roan* ECSs were homozygous for a haplotype that diverged from the

reference. One heavily marked ESS was homozygous for a fourth haplotype that was unobserved in the DBVC data but that was most similar to the *roan* haplotype (marked as TR* in Table S2). Genotyped markers did not clearly differentiate heavy and light ticking in the ESS haplotype refinement analysis, but did clearly differentiate the *ticked* from *clear* phenotypes in homozygous individuals.

Representative haplotype functional variant calling

Based on the variant calls available for DBVC haplotyperepresentative animals, six putative functional variants were identified using the VAI (Hinrichs *et al.* 2016) that segregated among the three representative haplotypes in the DBVC data. DBVC homozygous representatives of haplotypes observed in the ECSs and ESSs are identified by their allelic haplotype in Table S3. Across all dogs, many additional haplotypes and putative functional variants exist that were not observed to segregate in either the ECS or ESS. Heterozygous dogs or dogs with haplotypes unobserved in the ECS or ESS are shown as *unknown* in Table S3 (N = 558).

Below we describe the putative functional variant profiles for three alleles within an allelic series based on proxy haplotype. The characterized alleles are described in phenotype order from least to most pigmented within the whitepatterned areas of the coat (Fig. 3). Information relating to the functions of putative functional variants in the region as determined by ENSEMBL (Yates *et al.* 2020) is shown in Table S5. Other neutral variants segregate with the haplotypes and these are shown in Table S3.

In Spaniels, the reference genome haplotype *clear* (*t*) is differs from both the *ticked* and *roan* haplotypes by one missense SNP in *usherin* (*USH2A*; CFA38:g.11 111 286C>T) where the *clear* allele is C (Table 1). All observed *ticked* and *roan* haplotype representative animals have at least one copy of the T allele. Four of nine *parti-color* ECSs used in haplotype characterization were heterozygous for the *t* haplotype; the remainder had no copies of the *t* haplotype. Two of six *clear* ESSs were homozygous for the *t* haplotype and four were heterozygous.

Spaniels with the *ticked* (*T*) haplotype exhibit the alleles A and C at missense variant CFA38:g.11 156 276T>A and splice-site variant CFA38:g.11 156 425T>C respectively, whereas both *roan* and *clear* Spaniels have the T allele at both sites (Table 1). In addition, *ticked* Spaniels differ from the *clear* haplotype at CFA38:g.11 111 286C>T. One *particolor* ECS was homozygous for haplotype *T*.

The roan ($T_{\rm R}$) haplotype is diverged from the reference/ clear haplotype at four putative functional sites. Two are missense variants CFA38:g.11 085 443G>A and CFA38: g.11 111 286C>T (Table S3). The variant at CFA38:g.11 085 443G>A lies proximal to the genomic region that clearly distinguishes the three haplotypes based on concordance of the *clear* and reference haplotypes. The $T_{\rm R}$ haplotype has two additional variants that affect the secondary structure of the DNA in the region (see the next section).

Structural variants

By analysis of regional coverage in representative individuals, we identified a duplicated DNA segment (11 398 bp) lying within the 67th intron of USH2A and spanning CFA38:11 131 841–11 143 239 (CANFAM3.1), which is unique to the $T_{\rm R}$ haplotype (Fig. 2; Table S6). Subsequent testing for the presence of the PCR product capturing the junction between the duplicated segments identified the product in *roan* ECSs, all Dalmatians (typically described as evenly spotted), German Short-haired Pointer (typically *roan*) and Australian Cattle Dogs (typically *roan*). The PCR product was not captured in *parti-color* ECSs (*ticked*) nor

Figure 3 Illustration of allelic series hypothesis in the English Cocker Spaniel (ECS) and English Springer Spaniel (ESS). The *roan* allele is presumed to be codominant, whereas the *ticked* allele is codominant with *roan* but *dominant* in relation to clear. The following genotypes were observed in our haplotype refinement analysis: t/t (ESS = 2), T/t (ECS = 3, ESS = 1), T/T (ECS = 2, ESS = 1), T/T_R (ECS = 1, ESS = 1), T_R/T_R (ECS = 8). No dogs were identified with the t/T_R genotype.



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Table 1	Description of variants on canine CFA38 segregating with haplotypes that represent observed coat phenotypes with pu	tative functional
variants	s within the region segregating among three homozygous haplotypes in bold type.	

Position	Gene	Variant source	Putative function	Coding change	SIFT	Reference	Alternate
11 057 040		hr ¹ /wga ² /DBVC ³	Neutral	None		С	т
11 106 067		hr/wga/DBVC	Neutral	None		А	G
11 085 443	USH2A	DBVC	Missense	P/S	0.00	G	А
11 111 286	USH2A	hr/wga/DBVC	Missense	E/K	0.95	с	т
11 122 467		hr/DBVC	Neutral	None		G	Т
11 123 271		hr/DBVC	Neutral	None		Т	G
11 123 896		hr/DBVC	Neutral	None		А	G
11 124 294		hr	Neutral	None		С	А
11 143 243	USH2A	DBVC	Duplication edge	None		с	т
11 149 787		hr/DBVC	Neutral	None		G	А
11 154 628		hr/DBVC	Neutral	None		G	А
11 154 832		hr/wga/DBVC	Neutral	None		А	С
11 156 276	USH2A	DBVC	Missense	Q/L	0.33	Α	т
11 156 425	USH2A	DBVC	Splice-region	Unknown		т	с
11 167 847	USH2A	DBVC	Splice-region	Unknown		GA	GAA/DEL
11 169 445	USH2A	DBVC	Missense	P/S	0.45	G	А
11 176 811		hr	Neutral	None		А	С

A further putative functional variant in strong linkage with the *roan* haplotype but outside of the main analysis region (at 11 085 443) is shown also. ¹Haplotype refinement.

²Whole-genome association.

³Dog Biomedical Variant Consortium.

Australian ESS (*clear* or *ticked*). All animals identified with the PCR product had pigmented spots or interspersed pigmented hairs in their white markings. One of 23 coatphenotyped *roan* ECSs tested for the product reported a negative result based on owner-reported color. All other of 72 results were concordant with expectation based on coat phenotype (Table S6). Roan is a common phenotype in ECSs and is not observable when dogs do not have white patterning from the activity of *MITF*. In such dogs, the duplication is expected to freely segregate in accordance with Hardy–Weinberg expectation. In accordance with this expectation, black ECSs (where *ticked* and *roan* are unobserved) segregated the product.

The duplication allele is in strong LD with a variant at CFA38:11 143 243C>T (reference allele is C) that lies within 5 bp of the distal duplication edge but outside of the duplication. The variant acts as a proxy for the presence of the segmental duplication on the roan haplotype and enables the assessment of heterozygosity for the duplication. The dosage of the T-allele at the marker SNP is completely concordant with the expected copy number as ascertained by sequence coverage of the region of the duplication in the $T_{\rm R}$ dogs with the WGS (DBVC animals showing the $T_{\rm R}$ haplotype in Table S3). For DBVC breeds with observable ticking (i.e. breeds that commonly display white markings), the variant co-occurs with roan patterning in the affected breeds (i.e. Australian Cattle Dog, Dalmatian, German Wire-haired Pointer). Animals assessed as homozygous for the duplication according to this proxy have even dispersion of pigment throughout the white markings, i.e. the pigmentation favors neither the feet nor the muzzle.

DBVC coverage data (Fig. S1) show the presence of a VNTR mutation at CFA38:11 112 907–11 113 501. This variant lies close to an intron–exon boundary (ENSCAFG00000010731.4 exon 62/71) and it may affect the phenotypic expression of the *USH2A* gene. Ambiguity in the reference assembly at this site precludes effective assay of the variant at this time.

Validation analysis

Two variants of seven sequenced putatively functional variants (Table S8) segregated perfectly with roan phenotype (duplication bridge product and missense CFA38:g.11 169 445G>A) in the Australian genotyped Spaniels. Missense variant CFA38:g.11 169 445G>A segregates more broadly in the wider population and the roan-associated A allele is present as a homozygote in observable non-roan breed representatives, e.g. SID00101, Cavalier King Charles Spaniel, and SRR2094385, Beagle (Table S3). Sequencing shows that the Spaniel roan differs from the roan proxy haplotype at variants CFA38:11 085 443 (deleterious missense) and CFA38:11 156 276 (tolerated missense), where Spaniels have the reference and derived alleles respectively. Further, deleterious missense variant CFA38:11 085 443G>A that seemed strongly associated with roan in homozygous dogs from Table S3 did not segregate in Spaniels, showing that these variants are not causative for the roan phenotype. Together the results suggest that the segmental duplication variant (CFA38:11 131 841-11 143 239DUP.insTTAA) is the most likely causative variant for roan. The duplication associated marker CFA38:11 143 243C>T is homozygous in one ESS without the duplication, showing that linkage with the duplication is imperfect in Spaniels. Ticking density among dogs without the *roan* haplotype is not clearly associated with any of the genotyped variants. However, no animals homozygous for the *clear* haplotype were sampled in this analysis.

Determination of ancestral alleles

Wider data for the Plassais cohort (Plassais *et al.* 2019) are represented in Table S9. All loci shown in Table S3 were polymorphic in the wolf samples other than CFA38:g.11 085 443G>A and CFA38:g.11 143 243C>T, which describe the wider *roan* haplotype (T_R , wolf monomorphic for reference alleles G and C respectively; Table S10; Jagannathan *et al.* 2019; Plassais *et al.* 2019).

Discussion

The purpose of the initial study was to identify the genetic basis of the coat-patterning phenotype *roan*. Mapping of this phenotype without regard to the presence or absence of white patterning in ECSs misidentifies the causative locus as the white-patterning locus (*MITF*; data not shown) because frequently ECSs with white patterning are also *roan*. Restricting the analysis to only animals that have white patterning from the activity of *MITF* correctly identifies the causative locus as being in LD with *USH2A* on CFA38. *Parti-color* in the ECS is relatively rare and only five animals of 35 sampled in Australia had this phenotype (Table S7).

In the course of exploring the genomic association with *roan* phenotype, we asked whether *ticked* might be controlled by a related mechanism using ESSs which segregate both *ticked* and *non-ticked* phenotypes. The analysis including ESSs revealed that the most common *non-roan* haplotype in the ECSs was shared with the *ticked* ESSs and that homozygous *non-ticked* ESSs had a third haplotype in the same region that was concordant with the reference genome sequence (from a *clear* Boxer).

We have identified a region on canine CFA38 that segregates the *roan* and *parti-color* phenotypes in ECSs with genome-wide significance. Hair length, which is controlled by the gene *fibroblast growth-factor* (*FGF5*; Cadieu *et al.* 2009), can also affect our perception of color dispersion in the coat with shorter hair resulting in spots with precise edges whereas spots in long hair appear as diffuse. This may explain the discrepancy of one ECS in validation that is probably *ticked* based on inspection of a photograph.

On CFA38, the Affymetrix array marker most associated with *roan* in the ECS was CFA38:11 057 040 ($P_{\rm raw} = 8.9 \times 10^{-10}$). Further examination of genetic markers in the same region distinguished *clear* from *ticked* ESSs with the most associated markers at CFA38:11 122 467 and CFA38:11 124 294 (both $P_{\rm raw} = 1.7 \times 10^{-5}$) based on a denser

marker set with a limited number of phenotyped dogs. Observation of variant call files from dogs of various breeds with representative haplotypes in public-domain WGS data across the region of association identified further putative functional variants that segregate in LD with the three common haplotypes identified in coat-phenotyped ECSs and ESSs. We have labelled these haplotypes as *t* (recessive *clear* and reference genome haplotype), *T* (*ticked*) and $T_{\rm R}$ (*roan*). The phene has been allocated in OMIA as: coat color, ticked in *Canis lupus familiaris* (002264-9615).

Although our evidence strongly implicates the involvement of one gene (*USH2A*), the authors do not suggest that there is classic monogenic inheritance of these phenotypes. Rather, the three haplotypes of this allelic series together with others yet to be characterized result in a complex range of *clear*, *ticked* and *roan* patterns in the two phenotyped breeds.

The haplotypes that were characterized in our phenotyped Spaniels ignore other haplotypes present in domestic dogs, including rarer haplotypes in ECSs and ESSs. An illustration representing our current understanding of the most simplistic interaction of genotype and phenotype in ECSs and ESSs at this locus is shown in Fig. 3. Results from validation (Tables S2 and S8) suggest that in Spaniels the *ticked* phenotype can result from a large number of haplotypes in the region, whereas *roan* and *clear* phenotypes each seem to be clearly concordant with single haplotypes.

Sampled wolves did not segregate the $T_{\rm R}$ (*roan*) haplotype, suggesting that this haplotype evolved in domestic dogs. Other variants associated with the *clear* and *ticked* haplo-types did segregate in wolves.

A number of putative functional variants occur throughout the region of association in proxy animals; however, those with the strongest LD on the characterized haplotypes consist of four missense variants in the gene *USH2A* (CFA38:g.11 085 443G>A, CFA38:g.11 111 286C>T, CFA38:g.11 156 276A>T and CFA38:g.11 169 445G>A), two splicing variants in *USH2A* (CFA38:g11 156 425T>C and CFA38:g.11 167 847GA>GAA/DEL) and two structural variants (CFA38:11 112 907–11 113 501VNTR and CFA38:g11 131 841–11 143 239DUP.insTTAA).

The roan haplotype ($T_{\rm R}$) segregates strongly with the duplication of 11 kb located within the 67th intron of *USH2A* (CFA38:g11 131 841–11 143 239DUP.insTTAA). Evidence of the duplication is reported elsewhere (Ensembl), although not in relation to any particular phenotype (Decker *et al.* 2015). Puzzlingly, homozygous dogs from two breeds with disparate phenotypes (Dalmatian and Australian Cattle Dog) share a common haplotype with other *roan* breeds throughout the wider region of association. Their phenotypic divergence may be affected by the VNTR structural variant (CFA38:11 112 907–11 113 501VNTR) or variants elsewhere in the genome. Coverage data (Table S6) suggest polymorphism for copy number between breeds at the VNTR locus. Australian Cattle Dogs

in both public data resources (which are most always *roan*) demonstrated either homozygosity or heterozygosity for the $T_{\rm R}$ haplotype, supporting a proposed (incomplete) dominant mode of inheritance of the *roan* phenotype. One identified variant (CFA38:g.11 156 425T>C) differentiating the *T* (*ticked*) allele from *roan* and *clear* alleles in the cohort used for haplotype characterization affects gene splicing and this may alter somatic expression of the *USH2A* transcript.

The USH2A gene encodes the Usherin-2A protein and is found in the basement membrane of the extracellular matrix (ECM). Usherin-2A comprises a pentaxin domain with multiple laminin epidermal growth factor and many fibronectin type III motifs (Stelzer et al. 2016). In turn, laminin and fibronectin are proteins of the ECM and interact with other ECM proteins such as collagen. All variants that are associated with the coat patterning phenotypes examined in this research occur in the fibronectin region of the gene. Public data from the canine (Hoeppner et al. 2014) suggest that there are two alternatively spliced isoforms of USH2A that differ according to the presence or absence of 69 an exon between exons and 70 (ENSCAFG00000010731.4). The missing exon is represented in canine EST DN750099 (Genbank).

In humans, variants within USH2A are associated with Usher Syndrome type IIA (OMIM entry no. 276901), an autosomal recessive disorder characterized by varying severities of aural, visual and neurological impairment (Drever et al. 2008). The skin, eye and inner ear originate from the migration of neural crest cells. USH2A-associated hearing impairment is believed to occur as a result of disruptions to hair bundle formation and stereocilia differentiation within cochlear hair cells of the inner ear, whereas retinitis pigmentosa, also commonly associated with Usher Syndrome type IIA, is caused by defects in the retinal pigment epithelium (made from melanin and lipofuscin). To the authors' knowledge, the prevalence of ocular and/or aural disorders in roan or ticked Spaniel dogs has not yet been investigated but may be of interest given the role of USH2A in aural and visual phenotypes. Further, the mechanisms by which the USH2A variants functionally influence ticked or roan canine coat are at this time unclear.

Two breeds with the $T_{\rm R}$ haplotype (Dalmatian and Australian Cattle Dog) have some individuals that exhibit congenital sensorineural deafness (Stritzel *et al.* 2009; Sommerlad *et al.* 2014). The two breeds additionally have high frequency of the extreme white coat color at the white markings locus (*MITF*). The interaction of variants between these two loci in deafness-affected breeds remains under investigation.

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Conflict of interest

The authors have no conflicts to declare.

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Supporting information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

Figure S1. Regional coverage by representative haplotype clearly showing the presence of the segmental duplication and VNTR expansion on the $T_{\rm R}$ haplotype: (a) haplotype t (no structural variant, gray) vs. haplotype $T_{\rm R}$ (dark blue); (b) haplotype T (no structural variant, light blue) vs. haplotype $T_{\rm R}$ (dark blue).

 Table S1. Segregation of the 10 most associated variants

 from within-breed whole-genome-wide association analysis

for *parti-color (non-roan)* and *roan* coat patterns in the English Cocker Spaniel.

Table S2. Haplotype defining genotypes for English Cocker Spaniels (ECSs) and English Springer Spaniels (ESSs) with coat phenotypes and showing homozygous haplotypes.

Table S3. Dog Biomedical Variant Consortium Dogs showing breed and haplotype representation (homozygous individuals only).

 Table S4. Primers for capture of putative functional variants segregating among identified haplotypes.

Table S5. Putative functional variants in *usherin* (*USH2A*) as predicted by the variant table of ENSEMBL (Yates *et al.* 2020) for CANFAM3.1.

Table S6. Summary of variant calls and cover by representative haplotype (DBVC data: *t*, *clear*; *T*, *ticked*; T_R , *roan*, unknown?).

Table S7. Positive or negative result for structural variant bridge segment in coat phenotyped Australian dogs defined by positive capture of fragment in PCR using primers (forward, TCAACATCCATACTTCTCATGCTT; reverse, TGCTCAACCTAAATGTGCTAA) and an annealing temperature of 55°C.

 Table S8. Genotyping of identified putative functional variants in five ECSs and 11 ESSs.

Table S9. Genotypes for 722 dogs and canids derived from Plassais *et al.* (2019) for variants segregating with three coat phenotype haplotypes: *t* (*clear*), *T* (*ticked*) and $T_{\rm R}$ (*roan*) in ESSs and ECSs.

Table S10. Genotypes for wolves derived from Plassais *et al.* (2019) for variants segregating with three coat phenotype alleles *t* (*clear*), *T* (*ticked*) and $T_{\rm R}$ (*roan*) in ESSs and ECSs.