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Exploration of fine-scale recombination rate variation in the domestic horse

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Total genetic map length and local recombination landscapes typically vary within and across populations. As a first step to understanding the recombination landscape in the domestic horse, we calculated population recombination rates and identified likely recombination hotspots using approximately 1.8 million SNP genotypes for 485 horses from 32 distinct breeds. The resulting breed-averaged recombination map spans 2.36 Gb and accounts for 2939.07 cM. Recombination hotspots occur once per 23.8 Mb on average and account for ~9% of the physical map length. Regions with elevated recombination rates in the entire cohort were enriched for genes in pathways involving interaction with the environment: immune system processes (specifically, MHC class I and class II genes), responses to stimuli, and serotonin receptor pathways. We found significant correlations between differences in local recombination rates and population differentiation quantified by F_{ST} . Analysis of breed-specific maps revealed thousands of hotspot regions unique to particular breeds, as well as unique “coldspots,” regions where a particular breed showed below-average recombination, whereas all other breeds had evidence of a hotspot. Finally, we identified relative enrichment ($P = 5.88 \times 10^{-27}$) for the in silico-predicted recognition motif for equine PR/SET domain 9 (PRDM9) in recombination hotspots. These results indicate that selective pressures and PRDM9 function contribute to variation in recombination rates across the domestic horse genome.

[Supplemental material is available for this article.]

Meiotic recombination is a major driving force in shaping genetic variation and haplotype structure in sexually reproducing eukaryotes. In most species studied to date, recombination events are not uniformly distributed along chromosomes but tend to cluster in small regions termed “hotspots” (Petes 2001; Paigen and Petkov 2010). Fine-scale recombination rates and hotspots are poorly conserved across mammalian species and can differ significantly between even closely related species (Winckler et al. 2005; Smagulova et al. 2016; Stevison et al. 2016), a phenomenon primarily attributed to variations in the DNA-binding specificity of the zinc finger array present in the PR/SET domain 9 (PRDM9) protein (Baudat et al. 2010; Grey et al. 2011). PRDM9 binding results in histone H3 lysine 4 trimethylation, a modification associated with recombination initiation via formation of a double-strand break (DSB) (Buard et al. 2009). In species with functional copies of the PRDM9 gene, total genetic map length and local recombination rates vary between individuals, as well as across populations (Crawford et al. 2004; Coop et al. 2008; Keinan and Reich 2010; Ma et al. 2015; Petit et al. 2017), indicating that recombination landscapes are rapidly evolving in these species and may be subject to selective pressures.

Several factors are associated with a reduction in recombination. In humans, rates tend to be lower within transcribed regions of genes (Myers et al. 2005) and in regulatory domains (Liu et al. 2017), possibly because of increased mutation frequency associated with DSBs. It is also well known that chromosomal inversions result in suppressed regional recombination when heterozygous (Sturtevant 1921; Jaarola et al. 1998), a proposed mechanism by which speciation arises in the presence of gene flow (Noor et al. 2001; Rieseberg 2001). Regions showing little or no recombination

(“coldspots”) also occur in association with copy number variants (CNVs) in mice (Morgan et al. 2017). Finally, although population-based estimates of recombination rates are limited by the fact that genetic variation is necessary to detect historical recombination events, a downward bias in these rate estimates is often present in regions of the genome under selection (O’Reilly et al. 2008), an area of particular interest in the study of domestication and selective breeding of livestock.

The horse is a unique species for which domestication has resulted in numerous breeds specialized for the purpose of performing work or participating in athletic competition, and genomic resources have been under development to enable investigation of the genetic bases for such traits. Low-resolution linkage maps for the domestic horse were first created using microsatellite markers (Guérin et al. 1999; Swinburne et al. 2000, 2006; Penedo et al. 2005), paving the way for mapping coat color traits and several diseases of economic importance to the horse industry (Swinburne et al. 2002; Pielberg et al. 2005; Brunberg et al. 2006; Tryon et al. 2007; McCue et al. 2008). Since that time, linkage maps and microsatellite markers have given way to genome-wide SNP chip data and high-throughput sequencing, allowing an even closer look at genetic variation in the horse. Very little is known about fine-scale recombination and its determinants in the domestic horse, although several studies have reported relevant findings. Steiner and Ryder (2013) found evidence for positive selection in the zinc fingers of the PRDM9 gene across several species of equids. They also identified multiple PRDM9 alleles in the domestic horse and hybrids thereof, indicating that there may be

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population-specific differences in recombination rates and usage of recombination hotspots in this species. In addition, structural variation in the equine genome has long been a subject of interest to the equine genetics community, leading to the identification of thousands of CNVs (Doan et al. 2012; Metzger et al. 2013; Ghosh et al. 2014) with possible ties to phenotypes such as body size and sexual development, as well as a chromosomal inversion on equine Chromosome (ECA) 3 responsible for the Tobiano coat color phenotype (Brooks et al. 2007). Because the horse is a domestic species with several breeds possessing unique distinguishing phenotypes such as gaitedness and athletic performance, investigations of the genetic targets of natural and artificial selection in the horse have resulted in several discoveries providing new insights into selective breeding's functional consequences (Petersen et al. 2013b, 2014; McCoy et al. 2014; Avila et al. 2018). The impacts of the *PRDM9* gene, structural variants, and selective pressures on the local recombination landscape in the horse remain unknown.

The recent development of a 2-million-marker SNP array (Schaefer et al. 2017) allows for fine-scale investigation of recombination rate variation across the equine genome, as historical recombination rates can be estimated by coalescent analysis of linkage disequilibrium (LD) patterns in high-density SNP data (Auton and McVean 2007). These estimates can help identify putative recombination hotspots and compare local recombination rate differences between populations, as well as improve genotype imputation accuracy from lower- to higher-density SNP arrays. Here we report creation of fine-scale recombination maps and predicted hotspot locations for the domestic horse as well as 12 individual breeds. We assess relative enrichment for the *in silico* predicted DNA recognition motif for equine *PRDM9* in predicted recombination hotspots compared with genomic regions without evidence of hotspot activity. Finally, we investigate recombination rate distribution across the equine genome as it relates to genes and gene function and compare recombination rates across breeds.

Results

Recombination rate distribution across the equine genome

We estimated fine-scale population recombination rates ($4N_e r$) with the *LDhat* program *rhomap*, which uses a coalescent model-based approach to infer historical recombination rates between SNP markers from LD patterns (Auton and McVean 2007). Our data set consisted of 1,931,359 SNP genotypes for 485 horses from 32 distinct breeds (Schaefer et al. 2017). We constructed a breed- and sex-averaged recombination map, as well as 12 breed-specific maps for breeds in which we had genotype data for at least 18 individuals (i.e., Arabian, Belgian, Franches-Montagnes, French Trotter, Icelandic, Lusitano, Maremmano, Morgan, Quarter Horse, Standardbred, Thoroughbred, and Welsh Pony). To convert $4N_e r$ estimates to centimorgans (cM) for all maps, we estimated contemporary effective population sizes (N_e) for all equine chromosomes via gradient boosting regression using coalescent simulations modeling the demographic history of the horse. True and estimated N_e values had a Pearson's correlation coefficient of 0.85 (95% CI: 0.83–0.87; $P < 2.2 \times 10^{-16}$) (Supplemental Fig. S1) based on 1000-fold cross-validation. N_e estimates for the domestic horse data set ranged from 6843 for ECA11 to 9836 for ECA22, with a genome-wide average of 8460. Individual breed estimates varied from 1784 in the Thoroughbred to 6516 in the Quarter Horse (Supplemental Table S1).

Our fine-scale breed- and sex-averaged recombination map spans ~2.36 Gb and accounts for 2939.07 cM, making the average genome-wide recombination rate 1.24 cM/Mb (Table 1). Chromosomal recombination rates range from 1.06 cM/Mb on ECA15 to 2.13 cM/Mb on ECA12 (Table 1), with smaller autosomes generally showing higher rates. The Pearson's correlation coefficient for log of autosome size in megabases versus recombination rate (in cM/Mb) was -0.61 ($P = 0.0002972$). Approximately 75% of recombination occurs in 40% of the genome (Fig. 1A), illustrating the nonuniform nature of recombination distribution. Further, the probability of recombination events within transcribed regions of genes is relatively low, with a symmetrical increase in variability in recombination estimates as distance increases from the nearest start codon (Fig. 1B). We identified a significant weak correlation (linear regression $R^2 = 0.00314$; $P < 2.2 \times 10^{-16}$) between recombination rate and per-site nucleotide diversity. Supplemental Figure S2 shows that recombination rates along each equine chromosome vary considerably across chromosomes on a fine scale.

We used the *g:GOST* function of *g:Profiler* (Reimand et al. 2016) to assess functional enrichment of genes associated with Gene Ontology (GO) terms (The Gene Ontology Consortium 2000) in regions with both high and low recombination rates (Supplemental Table S2). The top 500 genes by recombination rate ranged from 4.77 to 15.70 cM/Mb with an average rate of 6.28 cM/Mb, whereas the bottom 500 genes ranged from 0.14 to 0.20 cM/Mb with an average rate of 0.19 cM/Mb. Genes with elevated recombination rates were enriched for immune system process terms, specifically major histocompatibility class I and class II

Table 1. Physical and genetic distance, mean recombination rate, and number of hotspots identified by chromosome

	Genetic length (cM)	Physical length (Mb)	Mean rate (cM/Mb)	Hotspots
Genome	2939.07	2363.42	1.24	56,299
ECA1	198.00	185.83	1.07	4080
ECA2	140.25	120.73	1.16	2662
ECA3	129.96	119.37	1.09	2731
ECA4	119.44	108.46	1.10	2511
ECA5	116.91	99.58	1.17	2375
ECA6	98.07	84.62	1.16	2112
ECA7	119.52	98.42	1.21	2320
ECA8	102.93	93.93	1.10	1973
ECA9	91.23	83.44	1.09	1974
ECA10	109.72	83.91	1.31	2042
ECA11	92.20	61.19	1.51	1361
ECA12	70.08	32.95	2.13	1054
ECA13	58.10	42.46	1.37	1055
ECA14	103.66	93.78	1.11	2132
ECA15	97.11	91.47	1.06	2146
ECA16	94.92	87.24	1.09	2033
ECA17	97.28	80.63	1.21	1967
ECA18	92.00	82.42	1.12	1927
ECA19	75.58	59.86	1.26	1359
ECA20	98.36	64.09	1.53	1690
ECA21	70.22	57.61	1.22	1410
ECA22	53.73	49.82	1.08	1219
ECA23	61.50	55.60	1.11	1337
ECA24	57.53	46.56	1.24	1060
ECA25	47.18	39.41	1.20	944
ECA26	65.00	41.76	1.56	1094
ECA27	50.44	39.85	1.27	1031
ECA28	53.55	46.06	1.16	1013
ECA29	50.87	33.56	1.52	888
ECA30	43.62	29.95	1.46	841
ECA31	34.48	24.87	1.39	678
ECA32	245.64	123.99	1.98	3280

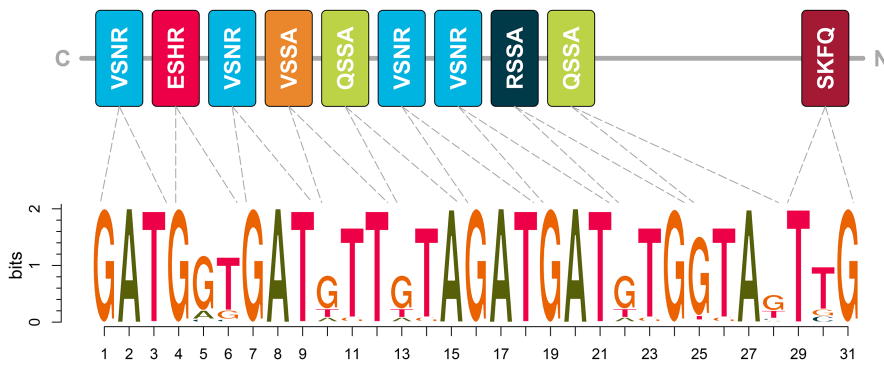


Figure 3. Zinc finger array for the equine reference PRDM9 protein (*top*) and corresponding in silico-predicted 31-mer DNA recognition sequence (*bottom*). Each colored box represents a single zinc finger, with distinct zinc fingers represented in different colors. Letters within the boxes signify DNA-contacting amino acids at the -1 , 2 , 3 , and 6 positions. Dotted lines lead to the portion of the DNA sequence motif recognized by each zinc finger.

Mantel test, establishing a significant positive correlation ($r=0.67$ – 0.84 ; $P=0.0001$ – 0.0016) between local recombination rates and population differentiation quantified by F_{ST} (Fig. 4B; Supplemental Fig. S3).

Analysis of breed-specific recombination hotspots revealed thousands of hotspot regions unique to a particular breed, as well as unique “coldspots,” or regions where one breed showed below-average recombination, whereas all other breeds showed evidence of hotspot activity (Supplemental Table S6; Supplemental Figs. S6–S17).

We calculated and compared breed-specific genic recombination rates (Supplemental Table S2) for functional enrichment using g:Profiler (Reimand et al. 2016). Most enriched GO terms are shared across all or many breeds for genes with both high and low recombination rates and reflect findings from the breed- and sex-averaged map. However, there were several GO enrichments that were unique to a particular breed (Supplemental Figs. S4, S5). Notable examples include enrichment of organic acid and anion transmembrane transporter activity in regions of elevated recombination rates in the Standardbred and negative regulation of metabolic processes in regions of low recombination rates in the Quarter Horse. We also found several breed-specific recombination hotspots and coldspots that overlap with regions previously reported as signatures of selection based on genetic divergence from other breeds in work by Petersen et al. (2013b) for nine breeds included in both studies (Supplemental Figs. S6–S17).

Discussion

Here we report the first look at fine-scale recombination rate variation in the domestic horse, a species for which genetic bases of health, disease, and performance traits are being increasingly discovered as new resources become available to study genome structure and function. We used SNP data from 1,931,359 loci in 485 horses to estimate effective population sizes, calculate population recombination rates, and predict recombination hotspots across the equine genome. To

investigate these results further, we analyzed known functions of the genes with the highest and lowest recombination rates across the genome. Finally, we identified differences in recombination rates and hotspot usage across 12 different breeds.

We estimate contemporary effective population size for the domestic horse at 8460. Our breed-specific N_e estimates are much higher than previously reported in many of the same breeds (Poncet et al. 2006; Corbin et al. 2010; Janssens et al. 2010; Campana et al. 2012; Vicente et al. 2012; Petersen et al. 2013a; Lee et al. 2014). This discrepancy is likely because of differences in methods used, which ranged from pedigree-based analyses to using genome-wide

SNP array data with varying sample sizes. N_e is a difficult parameter to estimate, sometimes leading to confidence intervals spanning multiple orders of magnitude, and estimates based on genetic markers are relative (Wang 2005; Waples 2006). Previous SNP arrays for the horse had much lower SNP density than the array used in this study and were developed with strong bias in favor of common SNPs with the overall goal of high informativeness (minor allele frequency [MAF] ≥ 0.05) in many breeds (McCue et al. 2012). Further, the SNPs included in these arrays have relatively poor informativeness in draft and pony breeds (McCue et al. 2012; Schaefer et al. 2017). These features are likely to lead to underestimation of genetic diversity across and within breeds and downward biases in SNP-based estimates of N_e . Similar discrepancies in N_e estimates between methods and marker sets for an animal breed can be seen in other agricultural species, in particular, Holstein cattle (Kim and Kirkpatrick 2009; Boitard et al. 2016) and Lacaune sheep (Kijas et al. 2012; Petit et al. 2017). Further, correction for N_e in this study results in reasonable genome-wide recombination rate figures both within and across breeds that are largely in agreement with existing equine linkage maps, as described below.

On the genome-wide scale, our recombination map length of 2939.07 cM falls between that of the two most recent existing

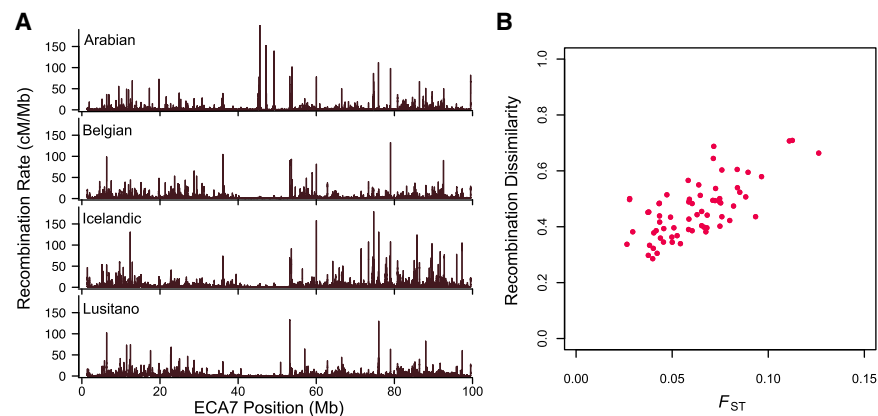


Figure 4. Fine-scale recombination rate variation on ECA7. (A) Recombination rate variation along ECA7 in five breeds. (B) Pairwise F_{ST} versus recombination dissimilarity ($1 - \text{Spearman's rank correlation coefficient}$) across ECA7.

linkage maps for the horse (2772 cM by Swinburne et al. [2006] and 3740 cM by Penedo et al. [2005]), an indication of relative agreement between these different methods despite the differences between studies in populations used and density of marker sets genotyped. The map created by Penedo et al. (2005) is a male-specific map, but it is longer than both the sex-averaged map created by Swinburne et al. (2006) and the map created as a part of this study. In mammals, genome-wide recombination rates are generally higher in females than in males (Kong et al. 2010; Stapley et al. 2017), with the notable exceptions of cattle (Ma et al. 2015) and sheep (Johnston et al. 2016). Although the biological significance of this discrepancy is unknown, one proposed mechanism is intense artificial selection on males of these species leading to indirect selection for higher recombination rates in males (Ma et al. 2015). Collectively, our results, in conjunction with the previous linkage maps, suggest that this explanation could also extend to the horse, a species subjected to strong sex bias during domestication and traditional focus on stallions in selective breeding programs (Lindgren et al. 2004; Wallner et al. 2013). Because our recombination rate estimates are based on LD patterns in SNP data from unrelated individuals, we are unable to estimate differences between male and female recombination rates in the horse using this data set.

Our average genome-wide recombination rate of 1.24 cM/Mb is comparable to findings in other mammals (Stapley et al. 2017). At the chromosome level, we found a significant correlation between autosome size and recombination rate, a result that has been reported in other species as well (Kaback et al. 1992; Jensen-Seaman et al. 2004; Johnston et al. 2016; Stapley et al. 2017). In the equine genome, ~75% of recombination occurs in 40% of sequence, resembling figures found in other mammals (Myers et al. 2005; Petit et al. 2017).

We identified 56,299 putative recombination hotspot regions in the equine genome. This is similar to the number of hotspots detected in other mammals with functional *PRDM9* genes (The International HapMap Consortium 2007; Brunshwig et al. 2012; Petit et al. 2017). We also found that hotspot regions account for 9% of the physical length of the equine genome. This proportion falls between that found in humans at 6% (The International HapMap Consortium 2007) and mice at ~13% (Brunshwig et al. 2012). Within these hotspot regions, we found significant enrichment for the *in silico* predicted DNA recognition motif for *PRDM9* compared with regions with below-average recombination rates, suggesting the possibility that *PRDM9* may play a role in hotspot determination in the horse.

The GO term analysis in regions with elevated recombination shows enrichment for genes associated with pathways involving interactions with the environment; in particular, immune system processes, responses to stimuli, and serotonin receptor signaling pathways. This is consistent with previous findings suggestive of historical diversifying selection in immune-related and olfactory receptor genes in the horse (Orlando et al. 2013). Genomic regions containing MHC complex genes have been shown in other species to be more genetically diverse and have relatively higher recombination rates and hotspot densities than the genome average (Jeffreys et al. 2001; Schaschl et al. 2006; The 1000 Genomes Project Consortium 2010; Fulton et al. 2016), reflecting the vital importance of maintaining within-species variability at these loci for antigen recognition and pathogen resistance. An animal's sense of smell serves a similar purpose to its immune system, helping it to recognize and distinguish thousands of different odorant molecules as a means of gathering information (Young and Trask

2002). Although analysis of our breed-averaged map did not show enrichment for olfactory receptors in highly recombining regions, the majority of individual breeds showed significant enrichment for genes responsible for sensory perception of smell. These results echo those found in a study of the genetic determinants of recombination in sheep (Petit et al. 2017). In contrast, genes with historically low recombination rates in the horse had fewer enriched GO terms but were primarily involved in intracellular processes and as parts of organelles. A similar pattern exists in humans (The International HapMap Consortium 2007). These findings may correspond to selection against recombination in regions of the genome containing more highly conserved genes. However, it is important to consider that our approach is limited by the fact that genetic variation and heterozygosity within a population are required in order to detect recombination events, so these regions of low recombination could potentially represent regions of low variation rather than regions with truly suppressed recombination.

We were able to identify trends in local recombination rate estimates across 12 different horse breeds. However, given the number of breeds and the complexity of the data presented here, a full detailed description of breed-specific recombination maps is beyond the scope of this study. Individual breeds showed positive correlations between recombination rate differences and population differentiation quantified by F_{ST} , similar to results found in humans (Keinan and Reich 2010; Laayouni et al. 2011). This variability may be in part a result of variation in demographic history between populations, as $4N_e r$ estimates are sensitive to these differences (Dapper and Payseur 2018). Icelandic horses had a much higher genome-wide recombination rate in this study than any of the other breeds analyzed. It is important to note that we generated both $4N_e r$ and N_e estimates in this breed based on SNP data from the smallest sample size ($n=18$) of all breeds for which a breed-specific map was made. As such, it is likely that the discrepancy is in part because of errors in the N_e estimate used to normalize recombination rates. We identified thousands of hotspots unique to a breed, illustrating significant within-species variation in hotspot usage in the horse. Significant GO terms associated with genes with low recombination rates unique to a breed as well as breed-specific coldspots may represent targets of selection resulting from downward bias in $4N_e r$ estimates because of local reductions in N_e .

Overall, our findings add to the growing body of knowledge surrounding fine-scale recombination rate variability within and across species. We have identified several areas warranting further investigation. Recombination rate estimates based on LD patterns represent historical recombination events and are influenced by demographic history, fluctuations in SNP density, and variability in N_e on a finer scale than just by chromosome. Large-scale pedigree-based recombination analyses using dense SNP data are necessary to resolve these possible confounding factors. This approach would also allow for the investigation of total map length and fine-scale recombination differences between male and female horses. Additionally, hotspots and coldspots presented here are computationally predicted and have not yet been experimentally validated to rule out false positives. We show that *PRDM9* is likely involved in recombination hotspot determination in the horse and that local recombination landscape varies considerably by breed, but it is not yet known whether population-specific *PRDM9* alleles exist that may influence hotspot usage at a breed level. Further characterization of this gene and its effects on recombination in a larger population would help resolve this question.

Methods

Data set

Genotype data from 485 horses (235 male; 281 female) representing 32 distinct breeds were previously generated as a part of the development of a high-density SNP array for the horse (Schaefer et al. 2017). Briefly, whole-genome sequence data compiled from 153 horses representing 24 separate breeds were used to discover approximately 23 million biallelic candidate SNPs. After quality control and filtration based on breed representation, even spacing across the genome, and probe design considerations, 2,001,826 SNPs were selected for the Affymetrix equine MNEc2M SNP array. A cohort of 332 horses from 20 breeds was genotyped using this array to serve as a reference population for genotype imputation and creation of a recombination map (Schaefer et al. 2017). Of these SNPs, 70,467 were located on unplaced contigs and therefore excluded from this study, leaving 1,931,359 SNPs for analysis. Whole-genome sequence SNPs from the variant discovery cohort (NCBI BioProjects PRJEB14779, PRJNA273402, and PRJEB10098) were masked to the MNEc2M SNP loci and combined with the array data (genotypes available at <https://www.animalgenome.org/cgi-bin/util/vcf>) to create a uniform set of SNPs on Chromosomes 1 through 31 and X from 485 total horses. Genotypes were phased with Beagle version 4.0 (Browning and Browning 2007) to fill in missing genotypes and predict haplotypes.

Recombination rate estimation and hotspot detection

One hundred random samples of 40 chromosomes from the data set described above were generated to estimate population recombination rates ($4N_e r$) using the *rhomap* program in the *LDhat* package (Auton and McVean 2007). Likelihood lookup tables for all analyses were generated from the table included with the software package, assuming a mutation rate (θ) of 0.001 per site per generation. Chromosomes were split into intervals of 2000 SNPs with 200-SNP overlaps between adjacent windows. Each analysis was run for 10,100,000 total iterations, sampling every 2000 iterations, with a 100,000-iteration burn-in period. To generate a breed- and sex-averaged recombination map, rates were averaged across the random samples. Breed-specific maps were generated following the above procedure in individual breeds with genotypes for 18 or more individuals (Arabian, Belgian, Franches-Montagnes, French Trotter, Icelandic, Lusitano, Maremmano, Morgan, Quarter Horse, Standardbred, Thoroughbred, and Welsh Pony). Rate estimates from the first 100 SNPs on the 5' end and last 100 SNPs on the 3' end of overlapping intervals were discarded to achieve continual progression between consecutive windows. *LDhot* (Auton et al. 2014) was used to perform likelihood ratio tests over 100-kb sliding windows, comparing a constant recombination rate model to a model with a 1-kb hotspot in the center of the window, with the null distribution for each window coming from up to 1000 coalescent simulations matched for sample size, SNP density, and background recombination rate. Hotspots were considered significant at $P < 0.001$, with a cutoff of 0.1 for joining adjacent windows.

Effective population size estimation

Effective population size (N_e) was calculated by gradient boosting regression for the domestic horse as well as for each of the 12 breeds for which breed-specific maps were made. Gradient boosting regression is an ensemble machine learning method that uses random sampling of classifiers with replacement over weighted data to train a predictive model. Coalescent simulations model-

ing the demographic history of the horse were performed using *cosi2* (Shlyakhter et al. 2014). Demographic parameters were randomly selected from uniform prior distributions ranging from a 100 to 20,000 contemporary effective population size (N_c), 10 to 1000 generations since domestication or breed formation bottleneck, and a 100 to 50,000 ancestral population size (N_a), including a rule that N_a must be greater than N_c . Recombination maps were simulated for chromosomes of random lengths matching those of equine chromosomes (i.e., 25–186 Mb), featuring varying recombination rates with gamma-distributed hotspots. SNPs generated from *cosi2* were used to calculate MAF proportions in bins (MAF ≤ 0.02 , 0.02–0.04, 0.04–0.06, 0.06–0.08, 0.08–0.10, 0.10–0.15, 0.15–0.20, 0.20–0.25, 0.25–0.30, 0.30–0.35, and 0.35–0.40) and pairwise LD measurements in the form of r^2 in 25-kb bins up to 200 kb for each simulation. Gradient boosting regression was performed using the Python *scikit-learn* machine learning module (Pedregosa et al. 2011). Hyperparameters were tuned via grid search, with `min_samples_leaf=100`, `max_features="sqrt"`, `sub-sample=0.8`, `learning_rate=0.1`, `n_estimators=575` identified as the best combination based on mean accuracy for this data set. The effectiveness of this method for estimating N_e from our simulated data was tested via 1000-fold cross-validation. Finally, N_e was estimated for the entire population and each breed 100 times per chromosome with different random seeds, and the mean of these estimates was used to convert $4N_e r$ estimates to centimorgans. An average genome-wide N_e estimate was determined for each breed and for the entire population by back-calculating N_e from the resulting total genetic map lengths in centimorgans and $4N_e r$.

Recombination rate distribution

A list of the 18,129 annotated equine genes with associated GO terms (The Gene Ontology Consortium 2000) was extracted using BioMart (Smedley et al. 2015). To prevent biased estimates of recombination rates because of differences in gene size, gene size was controlled for as in phase II of the human HapMap project (The International HapMap Consortium 2007). An average recombination rate was estimated for each gene over a region with 10 kb flanking either side of the midpoint of the transcribed region of the gene. Genes were subsequently ranked by recombination rate, and the top and bottom 500 genes were analyzed using the *g:GOST* function of *g:Profiler* (Reimand et al. 2016) as ordered queries.

PRDM9 motif analysis

Motif enrichment analysis was used to assess relative enrichment for the DNA recognition sequence for equine PRDM9. The amino acid sequence for horse PRDM9 (Steiner and Ryder 2013) was used for de novo prediction of the DNA-binding specificity position weight matrix (PWM) of its Cys₂His₂ zinc finger array. This was achieved with software developed by Persikov and Singh (2014), which predicts DNA-binding motifs by calculating contact energies between the amino acids of each Cys₂His₂ zinc finger and all 256 possible 4-bp sequence combinations using support vector machines trained with an experimental database derived from known examples of zinc finger protein-DNA interactions. Sequences from hotspot regions ($n = 56,299$) predicted from the recombination map were extracted from the EquCab2 equine reference genome (Wade et al. 2009). These sequences were matched with "coldspot" sequences of identical length from the same chromosome extracted from regions with recombination rates less than half the mean rate across the chromosome. AME (McLay and Bailey 2010) was used to determine whether the in silico predicted zinc finger DNA recognition motif for equine PRDM9 was

enriched in hotspot DNA sequences relative to regions with low recombination. Sequences were scored using AME's average odds score method, and a Fisher's exact test was used to determine motif enrichment.

Breed-specific comparisons

Recombination rates were compared between the 12 breed-specific maps to identify breed-level variation. Mean pairwise fixation index (F_{ST}) values (Weir and Cockerham 1984) were calculated with VCFtools (Danecek et al. 2011) for each breed pair on each chromosome. Recombination dissimilarity was calculated by subtracting the Spearman's rank correlation coefficient of recombination rates across each chromosome from one as described by Laayouni et al. (2011). For each chromosome, symmetric F_{ST} and recombination dissimilarity distance matrices containing comparisons of each breed to all other breeds were tested for correlation using a Mantel test (Mantel 1967) as implemented in the Python package scikit-bio (scikit-bio.org) using the Pearson method with P -values determined based on 10,000 matrix permutations. Hotspots were considered unique to a breed if there was no evidence of boundary overlap with a hotspot predicted in another breed. Unique coldspots were defined as regions of below-average recombination in one breed where hotspots occur in all other breeds tested. Recombination rates were averaged for genes using the method described above. The top and bottom 500 genes by recombination rate per breed were analyzed using the g:Cocoa function of g:Profiler (Reimand et al. 2016) as ordered queries for across-breed comparison of biological function of genes in high and low recombination regions.

Data access

Recombination maps and hotspots from this study are available in the Supplemental Material and have been submitted to <https://www.animalgenome.org/repository/pub/UMN2019.0219/>.

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