A Comprehensive Linkage Map of the Dog Genome

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ABSTRACT

We have leveraged the reference sequence of a Boxer to construct the first complete linkage map for the domestic dog. The new map improves access to the dog's unique biology, from human disease counterparts to fascinating evolutionary adaptations. The map was constructed with ~3000 microsatellite markers developed from the reference sequence. Familial resources afforded 450 mostly phase-known meioses for map assembly. The genotype data supported a framework map with ~1500 loci. An additional ~1500 markers served as map validators, contributing modestly to estimates of recombination rate but supporting the framework content. Data from ~22k SNPs informing on a subset of meioses supported map integrity. The sexaveraged map extended 21 Morgans and revealed marked region- and sex-specific differences in recombination rate. The map will enable empiric coverage estimates and multipoint linkage analysis. Knowledge of the variation in recombination rate will also inform on genome-wide patterns of linkage disequilibrium (LD), and thus benefit association, selective sweep, and phylogenetic mapping approaches. The computational and wet-bench strategies can be applied to the reference genome of any non-model organism to assemble a de novo linkage map.

INTRODUCTION

Genomics has broadened the exploration of natural variation, helping to bring the naturalist perspective back into the fold of modern biology. The dog is an example of a non-model organism with much to offer in the way of natural phenotypic diversity. Since Darwin (1883), the domesticated dog has been recognized as a model of mammalian evolution, with striking variation in form and function. Much of the phenotypic diversity in the dog appears to have evolved rapidly, accelerated by the intense pressure of artificial selection and the force of genetic drift through population bottlenecks. Selective breeding over centuries has served as the preliminary experiment in canine genetics: the mutants have been screened and the strains have been established. The architectures of domestic traits are likely enriched for genes of large effect, given that artificial selection can act only on discernible phenotypic differences (Andersson, 2001; Neff and Rine, 2006). It follows that the diversity in size, shape, and behavior in the dog is genetically tractable. Moreover, this 'adaptive' variation can be couched in the developmental context of an extant progenitor, the wolf (Vilà et al., 1997), to illuminate the morphological and behavioral antecedents from which breed-defining traits have been derived. In addition to propagating purposefully bred traits, managed breeding has had unintended consequences. The dog suffers many of the same diseases as man, from Mendelian defects (e.g., deafness) to complex diseases (e.g., cancer susceptibilities) (Ostrander et al., 1993). These health issues follow breed predilection, implying that ancestral mutations have been trapped within the now-closed gene pools, presumably from founder effects at the inception of breed registries.

Variation in the dog, both adaptive and maladaptive, can be understood genetically through several phenotype-driven approaches (reviewed in Karlsson and Lindblad-Toh, 2008).

Linkage analysis of Mendelian traits and monogenic diseases is particularly powerful owing to large sib-ships and pedigree ascertainment through breeders. Genome-wide association mapping leverages the strengths of population structure, and is driven by a breed-based LD that is 20- to 100-fold more extensive than that observed in human populations (Sutter *et al.*, 2004; Lindblad-Toh *et al.*, 2005). Less conventional methods for detecting genotype-phenotype correlations, such as *in silico* association (Grupe *et al.*, 2001; Jones *et al.*, 2008) and selective sweep (Pollinger *et al.*, 2005) mapping, also hold promise in the dog, especially for understanding hallmark traits that have been bred to fixation, thereby presenting a 'segregation problem' to geneticists.

Each of these approaches to genetic mapping is more powerfully applied with a thorough understanding of recombination rate. This includes a relevance to LD mapping that may be particularly important in canine genetics. In man, LD tends to extend over a kb-scale, for which genetic and physical distances are reasonably correlated. In contrast, LD in dog breeds extends to a Mb-scale (Karlsson *et al.*, 2007); understanding local variation in recombination rate (cM/Mb) could facilitate interpretations of LD.

Given the strengths of canine genetics, a principal unmet need is a comprehensive description of genome-wide variation in recombination rate. The current linkage map is built upon the success of previous efforts (Neff *et al.*, 1999; Langston *et al.*, 1999; Dolf, 1999; Mellersh *et al.*, 1997; Lingaas *et al.*, 1997), however those maps were constructed with less than 600 markers and lack the coverage, precision, and accuracy needed to optimize linkage analysis and subserve LD mapping (Dukes-McEwan and Jackson, 2002). A comprehensive map would enhance genetic analysis in several ways. It would allow assessment of empiric coverage during linkage scans, provide essential information for multipoint and fine-resolution mapping, and

serve as a foundation for interpreting LD across the genome. The map can also serve as a comparative resource for continuing to improve our understanding of the basic processes governing meiotic recombination and faithful chromosome segregation.

Toward this end, we have leveraged the high quality draft sequence (7.6x) of the dog genome (Lindblad-Toh *et al.*, 2005) to assemble a *de novo* linkage map. The reference sequence afforded an abundance of molecular polymorphisms, which were computationally 'cloned' for even spacing. The physical scaffold aided map assembly by providing a first approximation of marker order that was ultimately tested with genetic data. Several measures of map integrity indicated that our estimates of recombination rate were both accurate and reasonably precise.

Thus a comprehensive map will join a list of important canine resources, from high resolution radiation hybrid maps (Guyon *et al.*, 2003; Breen *et al.*, 2001; Vignaux *et al.*, 1999) to the high quality draft sequence, annotated with 2.4 million SNPs. We discuss the biological implications of this new canine map, and provide online resources to facilitate its application.

MATERIALS AND METHODS

Samples and markers: The familial resources used in this study are summarized in Table 1. The Cornell Families (CF) were a genetically admixed research colony inclusive of six breed backgrounds (Neff *et al.*, 1999). The F₂ family was developed previously as an experimental intercross of a Border Collie (sire) and a Newfoundland (dam) (Neff *et al.*, 1999). The extended Silken Windhound family was a privately managed pedigree provided for this study. A panel of 36 purebred dogs was assembled and surveyed to test marker performance across breed populations (Supplemental Table 1). DNA was prepared from blood, tissue, or buccal swab samples using previously described methods (Bell *et al.*, 1981; Oberbauer *et al.*, 2003).

Microsatellite loci were computationally mined from canFam1 and canFam2 (Lindblad-Toh et~al., 2005) by exploiting the RepeatMasker track of the UCSC Genome Browser (www.genome.ucsc.edu). Perfect (CA)_n microsatellites (n >12) were chosen for even spacing across the genome (~750 kb steps). PCR primers were designed with Primer3 using standardized parameters (Supplemental Table 2). Candidate oligo sequences were screened against the genome sequence using BLAST to ensure unique primer binding sites. Markers with a primer sequence having $\geq 95\%$ identity to a secondary annealing site were replaced by a suitable adjacent locus. Supplemental Table 3 lists the autosomal molecular markers developed in this study. An electronic file with the full marker set is also available (Supplemental File 2).

Microsatellite genotyping: Microsatellite markers were multiplexed in sets of 3-9 loci based on differential dye labels and expected differences in PCR product size. Marker loci were amplified according to a published method that uses dye-specific M13 oligonucleotides to incorporate fluorescent label during the PCR (Oetting *et al.*, 1995). In addition, a leader sequence (GTTTCTT) was appended to the 5' end of a subset of reverse primers to catalyze the non-templated enzymatic addition of a nucleotide to the 3' end of the labeled product strand (Brownstein *et al.*, 1996). Final PCR concentrations for forward (40-mers) and reverse primers (27-mers) were 0.18 and 1.8 μ M, respectively. M13 primers for labeling product were included in the PCR at a final concentration of 0.36 μ M.

PCRs were performed in 17 μ l reaction volumes with 50 ng genomic DNA template or 2 μ ls buccal swab extract, and final reaction conditions of 2.5 mM MgCl₂, 1x buffer, 200 μ M each dNTP (GenScript), and 0.15 units of Taq DNA Polymerase (ABGene). Reactions were covered with inert Chill-Out® (BioRad) to permit hot-start PCR. Thermocycling consisted of an initial

denaturation step of 93°C for 3 min, and a final extension step of 72° for 20 min. Marker amplifications were for 7 cycles of 93° for 20 sec, 65° for 30 sec, 72° for 2 min; 5 cycles of 93° for 20 sec, 58° for 30 sec, 72° for 2 min; and 25 cycles of 93° for 20 sec, 55° for 30 sec, 72° for 2 min.

An aliquot of PCR product $(0.4 \,\mu\text{ls})$ was combined with $0.5 \,\mu\text{ls}$ of GeneScan 500 LIZ (Applied Biosystems) and $10 \,\mu\text{ls}$ Hi-Di formamide, and denatured for 3 min at 95°. An aliquot $(1 \,\mu\text{l})$ was separated by electrophoresis and detected by fluorescence using ABI 3730 capillary instruments. The command line option in *GeneMapper4.0* was used to automatically pre-process ABI 3730 files remotely. Genotypes were scored with allele assignments that were consistent across families. All genotypes were manually curated.

SNP genotyping: Pedigree members of the F₂ intercross family were genotyped with a commercially available *Infinium CanineSNP20 BeadChip* according to the manufacturer's instructions. The canine BeadChip assays 22,362 unique SNPs distributed across the genome with ~110 kb spacing. Data were collected using an *Illumina BeadStation* scanner and dedicated data collection software. Genotypes were generated with *BeadStudio*.

FISH: BAC clones corresponding to genomic regions were obtained from CHORI (Oakland, CA). BAC-derived DNA was labeled with Spectrum Green or Spectrum Orange using nick translation. Paired sets of chromosome-specific probes (300 ng each) were differentially labeled and co-hybridized to canine metaphase spreads prepared from cultured canine lymphocytes. FISH preparations were washed (0.4x SSC, 0.3% NP-40) and counter-stained with DAPI. Slides were analyzed and images were captured using a Genus Cytogenetics digital microscopy station.

Data integrity: Microsatellite and SNP data were inspected for Mendelian inheritance using *PedCheck* (O'Connell and Weeks, 1998). When a single non-Mendelian inheritance was detected for a marker within a family, all potentially errant genotypes among the siblings or parent (*i.e.*, *PedCheck* output) were removed from the dataset. When multiple genotypes for a locus were inconsistent with inheritance in a family, all of the genotypes for the marker for that family were removed. If the number of errant genotypes detected by inheritance exceeded 10% for a given locus across families, the locus was discarded.

Map assembly algorithm: Markers were positioned according to their respective sequence coordinates in the *canFam2* reference genome (www.ncbi.nim.nih.gov). Correct chromosome assignment was assessed by testing all pairwise marker combinations for linkage using the *twopoint* option of *CRIMAP* (Green *et al.*, 1990). Markers showing stronger linkage to at least two loci on a different chromosome were reassigned to that chromosome. The *all* option of *CRIMAP* was used to scrutinize the order of markers along each chromosome. Every marker was iteratively removed and reassigned based solely on genetic data. Markers that could be placed elsewhere on the chromosome with equivalent statistical support (within 1 LOD) were deemed either error-prone, poorly informative, or incorrectly positioned on the sequence build. These markers were not included in the framework map. The *flips* option of *CRIMAP* was used to further test local marker order. Maximum likelihoods were calculated for each permuted order in a three-marker sliding window along the chromosome. The order inferred from physical coordinates was maintained only if it was within 1 LOD unit of the highest likelihood obtained.

The effects of positive interference render tight double recombinants biologically unlikely. Double crossover events (DCO), which usually involve a single marker out of phase with the surrounding markers, are most likely indicative of genotyping errors, mutations, gene conversions, sequence mis-assemblies, or segregating chromosome polymorphisms. The *chrompic* option of *CRIMAP* was used to identify multiple crossovers over short genetic distances. The output was graphed with a custom tool that visually emphasized interruptions in the parental origin of phased chromosomes (*Kodachrompic*, A.K. Wong, unpublished). A double crossover event isolated to a single offspring was interpreted as a genotyping error of that individual. DCOs shared among siblings suggested a genotype error in a parent. The distribution of alleles in the family was scrutinized to identify the most probable errant genotype, which was removed. Terminal markers, which could not be evaluated in the same way, were scored twice by independent readers to improve the reliability of the genotype data.

Map expansion caused by undetected errors was estimated by iteratively (1) removing a marker, (2) rebuilding the local map, (3) re-calculating inter-marker recombination fractions, and (4) comparing the original and new recombination fractions. Markers that led to map inflation (> 2 cM) were excluded. Sex-averaged, male-specific, and female-specific autosomal genetic distances, as well as the female-specific distances of the *X* chromosome, were calculated with the *fixed* option of *CRIMAP* using the Kosambi map function.

Quantifying Inter-individual Variation: The number of recombination events in parental meioses was inferred from offspring genotypes using the *chrompic* option in *CRIMAP*.

Recombination events localized to chromosome termini (first and last 3 markers) were discounted; distal exchanges are challenging to distinguish from genotyping errors. The mean

recombination rate in each mother (or father) was estimated as the average number of recombination events in the haploid gametes transmitted to her (or his) progeny. One-way ANOVA was applied to test for significant differences among mothers and fathers. The recombination count in a given meiotic product was used as the response variable, and parental identity was treated as the factor. The significance of resultant F-statistics was empirically determined using the quantile position of the realized statistic along the distribution of $10^4 F$ -values derived by permutation.

Sequence Correlates: Markers on the framework map were binned into non-overlapping 5 and 10 Mb windows across the genome. For each window harboring a sufficient number of markers $(n \ge 2)$, the regional recombination rate was estimated by the slope of a simple linear regression of genetic map position in cM units on physical genome position in Mb. Imposing a harsher criterion of $n \ge 5$ makers per window had no significant impact on our findings, but did vastly diminish the number of regions that could be included in the analysis, particularly at the 5 Mb scale. Results reported in the main text derive from the more lax criterion.

The number of repetitive elements (LINES, SINES, LTRs, poly-A repeats, poly-C repeats, poly-G repeats, poly-T repeats, simple repeats, AT-rich, CA-rich, satellites, DNA repeats, and low complexity repeats), base composition measures (GC%, number of CpG dinucleotides, number of CpG islands), incidence of the CCNCCNTNNCCNC putatively recombinogenic motif (Myers et al. 2008), the number of genes (from RefSeq and the N-SCAN gene prediction track implemented in the UCSC Genome Browser), and the distance to the telomere as a fraction of total chromosome length were computed for each window using the most recent genome

assembly (*canFam2*). Spearman rank correlation tests between recombination rate and individual sequence variables were performed in the *R* environment for statistical computing.

RESULTS

Assembling markers and meioses: Perfect CA-repeat microsatellites were computationally selected from the reference sequence (Lindblad-Toh *et al.*, 2005) to achieve sub-megabase and possibly sub-centiMorgan spacing on a linkage map. Because markers were newly developed and untested, the collection (n = 3,349 loci) was preliminarily screened against a panel of 36 purebred dogs (Supplemental Table 1). Results indicated that the markers were amenable to high-throughput genotyping and were sufficiently informative for map assembly (HET = 0.46 ± 0.16 ; Supplemental Table 3 and Supplemental File 1).

The markers were typed on familial resources comprising 450 mostly phase-known meioses (Table 1). The families represented an efficient meiotic mapping resource, with 281 dogs from three pedigrees totaling 24 sibships and an average of 8 offspring per cross.

Genotyping was performed in six phases, with each phase including markers to span the genome in ~4 Mb increments. This approach allowed a more frequent quality control of genotyping, as well as a new map assembly following each phase. Intermediate map builds were posted online during the course of the study (http://www.vgl.ucdavis.edu/dogmap/). Completion of the phases yielded a million genotypes for assembly of a final map. The meiotic contributions of families are summarized in Table 1. Summary statistics are given in Table Supplemental 3.

Clarifying marker order: For map assembly, markers were ordered along chromosomes according to their physical coordinates in *canFam2*. Chromosome assignments and relative order were tested against the genetic data. Several loci were discrepant (n = 17), showing significantly stronger linkage to two or more loci on a different chromosome than the one to which they had been assigned in *canFam2*. Genetic data were given preference and discrepant markers were re-assigned (Supplemental Table 4). The vast majority of marker positions were concordant for both types of positional information (99.4%), indicating the *canFam2* build was of high quality. This implied there was considerable value in using the physical scaffold to guide and clarify marker order. It should be noted that the power to detect macro-assembly errors was presumably greater than the power to discern micro-assembly errors. Fine-scale errors in the sequence build may have been undetected in our analyses.

Estimating genetic distances: Given an established marker order, genetic distances for intermarker intervals were estimated for male-specific, female-specific, and sex-averaged maps. Sexaveraged distances were tested for map inflation, which can result from cryptic errors producing artifactual crossovers. Each locus was iteratively removed from the map, and relevant intermarker distances were re-calculated. Approximately 1% of markers (n = 46) were suspected of having undetected genotyping errors, as evidenced by a significant reduction in map length (> 2 cM) upon their iterative removal. These potentially errant loci were excluded from further assembly.

Autosomal maps: Of autosomal markers genotyped (n = 3,549), 1,469 qualified as anchor loci based on an ability to 'sample' a large number of meioses (*i.e.*, > 100 informative meioses) with

concomitant statistical support for linkage (*i.e.*, pairwise LOD > 11). These markers formed the basis of a framework map. An additional 1,606 markers were considered 'map validators' (*i.e.*, a pairwise LOD > 3); these loci contributed modestly to estimates of recombination rate, but supported the mapping content of anchor loci (*e.g.*, by substantiating phase transitions of observed crossovers). Table 2 lists summary statistics for the framework and comprehensive maps.

A SNP-based map: In addition to microsatellite markers, ~22,000 SNPs distributed across the genome (every $104 \text{ kb} \pm 25 \text{ kb}$) were genotyped on the F_2 intercross family (Table 1). The quality of the SNP data was high, as reflected by call rate (99.8%), inheritance (0.09% non-Mendelian errors), and informativeness (15,375 polymorphic loci; average MAF \geq 0.20). The resulting SNP-based map detected most of the same crossovers observed in the microsatellite-based map (718/836). Undetected crossovers were mostly attributable to uncertain phase of diallelic SNPs. The SNP-based map exhibited 64 crossovers at the ends of autosomes that had not been captured by microsatellite markers. This suggested the sex-averaged length of the canine autosomal genome might be somewhat greater than 21 Morgans. Overall, the two maps were in strong agreement, consistent with the microsatellite-based map being of high integrity.

An X map and the pseudo-autosomal region (PAR): Markers were analyzed separately for the X chromosome to accommodate female-specific recombination. Of 140 markers analyzed, 82 served as anchor loci and 55 served as validating loci. Markers were ordered according to physical coordinates, and genetic distances were computed from informative maternal meioses.

The results suggested a genetic length for the *X* chromosome of 111 cM (Table 3 and Supplemental Figure 1).

The vast majority of X-linked loci showed a single allele in males, as expected for hemizygosity. However, five markers exhibited significant heterozygosity in males (avg. HET = 0.80). None of these markers exhibited fixed alleles, arguing against duplicated regions. The five loci clustered physically (Supplemental Figure 1), and their allelic variation segregated with autosomal inheritance, consistent with their belonging to the canine pseudo-autosomal region (PAR). The genotype data for 565 SNPs along the X chromosome were checked for a similar pattern. Fifty SNPs showed heterozygosity in males (avg. HET = 0.56), and these were from the same physical region as the five microsatellites (Supplemental Figure 1). This suggested the PAR on the metacentric X localized to the telomeric end (Supplemental Figure 1). This corresponded to a female-specific length of 7 cM, and a male-specific length of 28 cM. Reference sequence data was available from female dogs only (Kirkness $et\ al.$, 2003; Lindblad-Toh $et\ al.$, 2005), precluding a characterization of the physical arrangement of the PAR with Y synteny.

Observed variation in genome-wide recombination: Although the map showed pronounced regional differences in recombination rates, several systematic patterns were evident. Nearly every autosome was characterized by low recombination rate at the centromeric end and high rate at the telomeric end (Figure 1). Two chromosomes (*CFA27* and *CFA32*) exhibited a reversed pattern (*i.e.*, high rates at the centromere and low rates at the telomere; Figure 1). Given the consistency of the recombinational profiles across the other 36 autosomes and the generality of

this pattern in mammals (Kong *et al.*, 2002; Shifman *et al.*, 2006), the simplest interpretation was that the orientation of the chromosomes was mis-specified in *canFam2*.

We tested this prediction directly using fluorescence in situ hybridization (FISH) analysis. Differentially labeled probes were generated from BAC clones, which were anchored to the ends of each autosomal linkage map by markers. The probes were localized by FISH with canine metaphase spreads. The results strongly supported the chromosome orientation inferred from the linkage map rather than the one listed in *canFam2* (Supplemental Figure 3; Supplemental Table 5).

Recombination rate was inversely associated with the physical length of chromosomes (Figure 2). The smallest chromosome (*CFA38*) exhibited a three-fold greater rate than the largest chromosome (*CFA1*) (Table 3). These results were generally consistent with the expectation of at least one crossover per chromosome, a nearly universal requirement of meiosis from yeast to man (Kaback *et al.*, 1989; Kong *et al.*, 2002).

The rate of recombination also systematically varied by sex. Female meioses exhibited a 1.2-fold greater average rate than males (Table 2). The influence of sex was not uniformly distributed across the genome (Supplemental Figure 2), with differences in sex ratio being most striking at autosome ends. Female meioses exhibited a four-fold greater rate near centromeres, whereas male meioses exhibited a four-fold greater rate near telomeres. Similar observations of sex differences have been made in human (Broman *et al.*, 1998) and mouse (Paigen *et al.*, 2008).

Inter-individual variation in recombination rate was also observed (Supplemental Figure 4). The female with the highest recombination rate exhibited $\sim\!20\%$ more crossovers per meiosis than the female with the lowest rate, though these differences could be chance variation (one-

way ANOVA: F = 1.35; P = 0.16). There was substantially more variation in genome-wide rates (nearly two-fold) among male dogs, and these differences were statistically significant (F = 1.77; P = 0.042). This finding complements observations of individual differences in recombination in a variety of species, including human (Broman *et al.*, 1998; Kong *et al.*, 2004), *Drosophila* (Brooks and Marks, 1986; Chinnici, 1971; Kidwell, 1972), Tribolium (Dewees, 1975) and laboratory strains of mice (Reeves *et al.*, 1990; Koehler *et al.*, 2002; Paigen *et al.*, 2008). Interindividual variation may therefore be a pervasive characteristic of meiotic recombination in sexually reproducing species.

Sequence correlates of recombination rate: A sequence-explicit framework motivated investigation of the relationship between sequence variables and regional heterogeneity in recombination rate. Twenty sequence features were tested for rank correlation with recombination rate (Table 4). Non-overlapping 5 and 10 Mb windows were tested across the genome. The strongest predictor of sex-averaged recombination rates for both window sizes was proximity to the telomere. In addition, several classes of repetitive elements, the number of CpG dinucleotide sites, and the incidence of a recombinogenic sequence motif (Myers *et al.*, 2005; Myers *et al.*, 2008) were also predictive of local recombination rates, similar to what has been observed in other mammals (Kong *et al.*, 2002; Shifman *et al.*, 2006; Jensen-Seaman *et al.*, 2004; Myers *et al.*, 2005).

We examined these sequence correlates in the context of sex-specific influences. This analysis revealed distinct differences in males and females. Several GC-related measures (GC-rich repeats, CpG dinucleotide sites, and CpG islands) were positively correlated with male recombination rates, but weakly or negatively correlated with female recombination rates (Table

4). These findings were intriguing in light of the fact that males disproportionately drive the evolution of GC content in the human genome (Dreszer *et al.*, 2007; Duret and Arndt, 2008), which may stem from a male-intensified, biased gene conversion toward G/C alleles during recombinational repair (Marais, 2003). The putative recombinogenic motif (CCNCCNTNNCCNC) was also positively correlated with male recombination rates, but uncorrelated with female rates. This sex-specificity in the dog was similar to that observed in the house mouse (Shifman *et al.*, 2006), but different from the motif's recombinogenic properties in both human sexes (Myers *et al.*, 2008).

Sex-dependent correlations were also found for several classes of repetitive elements.

AT-rich repeats, satellites, low-complexity repeats, polyT repeats, polyA repeats, and long terminal repeats were positively correlated with female recombination rates, but non-correlated with male rates. Together, these findings suggest that regional recombination is mediated differently in the two sexes by features of the sequence landscape.

Interpolated maps for SNPs and scanning set loci: The resolution and integrity of the canine map afforded an opportunity to place additional markers on the map through linear interpolation (Kong *et al.*, 2002). Positions for 2.4 million publicly available SNPs were interpolated against the sex-averaged map. Results have been made available here (Supplemental File 2), and are also available upon request as a custom track for a publicly available browser (www.genome.ucsc.edu). Microsatellite markers from the latest minimal scanning set (MMS3, 507 markers; (Sargan *et al.*, 2007)) were similarly positioned by interpolation to afford multipoint analyses (Supplemental File 3). An electronic file with inferred inter-marker genetic distances recombination fractions has also been made available (Supplemental File 4).

DISCUSSION

We have addressed a principal unmet need in our field by assembling a linkage map that enables navigating the dog genome in a structured and systematic way. The map we have created (i) informs on the basic biology of meiosis and the evolution of recombination rate; (ii) provides important clues on the genomic determinants of regional heterogeneity and sex-specificity of recombination rates; (iii) facilitates unbiased genetic access to natural variation and phenotypic diversity in canids; and (iv) lays out an integrated wet bench and bioinformatics approach for developing a *de novo* map for any species for which a draft sequence becomes available.

Basic biology of meiosis: Although classical genetics in model organisms has dissected the mechanisms governing meiosis, observations from linkage maps have uniquely informed on natural variation in underlying processes (Dumont and Payseur, 2008; reviewed in Coop and Przeworski, 2007). The canine genome represents a natural experiment in meiosis, with 38 pairs of unusually short, acrocentric autosomes. Despite the unique challenges such a karyotype might present to the meiotic apparatus, patterns of recombination in the genetic map suggest this karyotype has been accommodated by conventional means. An up-regulation of recombination rate among physically smaller chromosomes, for instance, is a conserved feature of meiosis from yeast to man (Kaback *et al.*, 1989; Kong *et al.*, 2002). This non-random distribution ensures the obligatory crossover per chromosome arm that helps homologs attain bipolar orientation during Meiosis I.

Despite the unique karyotypic features of the dog, the sex-averaged rate of recombination across the genome (.97 cM/Mb) is within the range of other characterized

mammals (0.5 cM/Mb – 1.1 cM/Mb). Genetic map-based estimates of recombination rate are available for two additional carnivore species, house cat (~1.1 cM/Mb; Menotti-Raymond *et al.*, 1999) and silver fox (~0.6 cM/Mb; Kukekova *et al.*, 2007). While large differences in map quality and coverage preclude detailed comparisons, the rate of recombination in dog is clearly not an outlier among carnivores. Although fine-scale recombination rates evolve on short evolutionary time-scales (Winckler *et al.*, 2005; Ptak *et al.*, 2005) the dog genetic map adds to evidence for more rigid evolutionary constraints on broader scale recombination rates (Myers et. al 2005; Dumont and Payseur 2008).

The elevated recombination rate in female meioses, a salient feature of eutherian maps, was also evident in the dog, though the sex ratio was not as great as it is in man (1.2-fold in dog versus 1.7-fold in humans (Broman *et al.*, 1998; Kong *et al.*, 2002)). In part, this may be attributable to karyotype—mouse and rat also have mostly acrocentric autosomes, and similarly exhibit a muted sex-difference relative to human (Shifman *et al.*, 2006). Karyotypic organization accounts for only some of the differences between sexes (Hunt and Hassold, 2002). For reasons not yet clear, sex differences are modest in cow (Ihara *et al.*, 2004) and sheep (Maddox *et al.*, 2001), for instance, and are reversed in the two metatherian mammals (marsupials) studied to date (Zenger *et al.*, 2002; Samollow *et al.*, 2004).

As in other placental mammals, sex differences in the dog were not uniformly distributed across the genome. Female meioses exhibited a greater recombination rate near centromeres whereas male meioses showed a greater rate near telomeres. In general, it appears female meioses are at greater risk for non-disjunction in man (Antonarakis, 1991), and this risk is exacerbated among chromosomes with distal crossover events (Lamb *et al.*, 1996). Distal crossovers decrease fidelity in yeast as well, suggesting crossover placement fundamentally

influences chromosome orientation on the meiotic spindle. Interestingly, segregation in yeast can be rescued with an experimental tether, but only if the tether is located at the centromere (Lacefield and Murray, 2007). This implies that physical ties near the centromere (*e.g.*, crossovers) intrinsically promote bipolar orientation. If so, the greater recombination rate near the centromere in females could be compensatory—an adaptation that offsets the greater sensitivity to non-disjunction in oogenesis.

Evolution of patterned recombination: The canine linkage map may offer an unprecedented opportunity to address the evolution of recombination rate. By generating favorable allelic combinations and breaking down negative linkage disequilibrium, recombination can facilitate the response to selection (Fisher, 1930; Felsenstein, 1965). This has led to the hypothesis that species under frequent and intense selection evolve toward increased recombination rates (Burt and Bell, 1987). Recent work has shown that the purebred domesticated chicken exhibits higher recombination rates than Red Jungle fowl, its wild ancestral counterpart (Groenen *et al.*, 2009). In our case, describing the effects of domestication requires a genetic map assembled for the wolf, *C. lupus*. There are logistical constraints to this, but a broad assessment of recombination patterns could be inferred from an analysis of LD in wolf populations, as has been done in man (McVean *et al.*, 2004).

The dog also presents an opportunity to address microevolutionary changes in recombination rate. Mouse strains show systematic differences in recombination (Shiroishi *et al.*, 1991; Paigen *et al.*, 2008), and inter-individual variation, from Drosophila to man, has been tied to causative genes (Kong *et al.*, 2008; Coop *et al.*, 2008; Chinnici, 1971; Kidwell, 1972; Brooks and Marks, 1986; Shiroishi *et al.*, 1991; Koehler *et al.*, 2002). Consistent with these

observations, we documented significant variation among male dogs. If this variation is heritable, we might also expect differences in recombination rate among breeds. However, our use of admixed pedigrees for map assembly complicated the detection of such differences. The relatively short timescale separating dog breeds and the artificial selection that has shaped breed phenotypes should motivate the construction of breed-specific maps to understand the effects of recent isolation and strong selection on the evolution of recombination rate.

Genomic control of recombination rate: Sequence correlates provide mechanistic insights into recombination rate variation. Several sequence correlates found in human (Yu et al., 2001; Kong et al., 2002), mouse (Jensen-Seaman et al., 2004) and rat (Jensen-Seaman et al., 2004) were also found in the dog. The number of CpG dinucleotides, for instance, is positively correlated with recombination rate in each species (Kong et al., 2002; Jensen-Seaman et al., 2004), including the dog. Cross-species comparisons have also revealed differences. Although recombination is correlated with LINES in human, mouse, and rat (Kong et al., 2002; Jensen-Seaman et al., 2004), no such correlation was observed in the dog. Distance from the centromere is the best predictor of recombination in man and dog, whereas sequence features remain the best predictors in mouse and rat (Jensen-Seaman et al., 2004).

Many sequence-based correlations were significant, but most were relatively weak. In this respect, the dog joins a growing list of mammals for which the majority of genomic variation cannot be explained by sequence characteristics (Kong *et al.*, 2002; Myers *et al.*, 2005; Jensen-Seaman *et al.*, 2004; Shifman *et al.*, 2006). Alternative sources of variation, for which there is mounting evidence, are epigenetics and chromatin state (Winckler *et al.*, 2005; Neumann and Jeffreys, 2006; Sandovici *et al.*, 2006; Sigurdsson *et al.*, 2009). CpG islands, the principal

targets of methylation, are correlated with recombination rate in human, mouse, rat, and dog.

Interestingly, CpG association in the dog differed in sign between the sexes, suggesting opposing processes in male and female meioses.

The canine genetic map provides important clues to the origins of sex differences in recombination. This dimorphism has long been a puzzle (Haldane, 1922; Morgan, 1912; Dunn, 1920). The most striking result to date is a sequence variant in humans that increases recombination rates in males, but decreases recombination rates in females (Kong *et al.*, 2008). Thus, causative genes and cis-sequence motifs may combine to account for sex-specific differences in recombination rate, possibly mediated by chromatin state (Petkov *et al.*, 2007).

Application to mapping phenotypes: The *de novo* map makes known the full genetic landscape of the dog genome, and thus allows assessing empirical coverage provided by linkage-scanning sets. The genetic positions of markers in commonly used sets were interpolated so that researchers could derive these benefits. Known positions and inter-marker distances are also essential for multipoint analyses to maximize the power and resolution of fine-mapping and quantitative trait loci (QTL) mapping. The genetic map may also inform on the pattern and extent of LD in the purebred dog. LD is 20- to 100-fold more extensive in breed isolates than in human populations (Sutter *et al.*, 2004; Lindblad-Toh *et al.*, 2005), a consequence of introgression and historical bottlenecks. Recombination is one of several forces that shape LD; knowledge of local recombination rates may therefore be useful for discerning the contributions of other factors (*e.g.*, selection).

Map-building extensibility: An available reference sequence presented us with an uncommon opportunity to assemble a map efficiently. The genome sequence afforded an abundance of putative markers that could be computationally mined from a physical framework and selected for even spacing and complete coverage. These markers performed well, and their standardized design facilitated wet bench genotyping. Systematic marker ascertainment affords other opportunities, such as studying mutation rates in the context of the "slippery genome" hypothesis of dog domestication (Fondon and Garner, 2004). Positional information for markers was valuable in guiding and clarifying locus order. This resulted in a map that was more inclusive of typed markers. These strategies for building a map *de novo* are applicable to any natural species with a draft sequence. This exemplifies how genomics is continuing to enhance access to natural variation in a broader array of non-model organisms.

Online maps and ancillary data: We have provided map figures (Supplemental Figure 5) and electronic files with marker and map content. We have also posted detailed map builds online with hyperlinks to ancillary data. The genetic maps and associated tabular data are available at: http://www.vgl.ucdavis.edu/dogmap/.

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Supplemental material to be made available online

Supplemental Figure 1. An X chromosome linkage map with denoted PAR (PDF 488 KB).

Supplemental Figure 2. Sex-specific recombination rates across the autosomes (PDF 317 KB).

Supplemental Figure 3. FISH analysis of suspected mis-specification of autosome orientation in *canFam2* (PDF 460 KB).

Supplemental Figure 4. Inter-individual variation in recombination rate (PDF 84 KB).

Supplemental Figure 5. Chromosome-specific linkage maps (*CFA1-19*) (PDF 6 MB).

Supplemental Figure 6. Chromosome-specific linkage maps (*CFA20-38*) (PDF 4 MB).

Supplemental Table 1. Panel of 36 purebred dogs for assessing marker performance (PDF 42 KB).

Supplemental Table 2. Design parameters for markers (PDF 33 KB).

Supplemental Table 3. Marker information (PDF 1.3 MB).

Supplemental Table 4. Loci reflecting discrepancies between the map and reference Sequence (PDF 53 KB).

Supplemental Table 5. Testing autosome orientation against observed chromosomal patterns of recombination rate variation (PDF 50 KB).

Supplemental File 1. Marker spreadsheet (MSExcel 777 KB).

Supplemental File 2. SNP interpolation (MSExcel 1 MB).

Supplemental File 3. MSS interpolation (MSExcel 65 KB).

Supplemental File 4. Inter-marker recombination fractions for multipoint analysis with MSS3 (MSExcel 452 KB).

Figure Legends

Figure 1. Variation in recombination rate across the autosomal genome. Recombination rate along each of the 38 autosomes is shown from centromeric to telomeric end (left to right). *x*-axis, physical length, scaled to accommodate different chromosome sizes. *y*-axis, sex-averaged recombination rate. Sliding windows of 5 Mb were used to calculate cM/Mb along the chromosome. The pattern of recombination rate for two autosomes (*CFA27 and CFA32*) suggested their orientation may be incorrectly specified in the *canFam2* build.

Figure 2. Correlation of recombination with physical chromosome size. (*A*) Genetic length of autosomes plotted against physical length. (*B*) Mean recombination rate across autosomes plotted against physical length. Results are shown for canine (black), human (blue), and mouse (red) autosomes. Data describing physical sizes and sex-averaged recombination rates were obtained from (Kong *et al.*, 2002) and (Shifman *et al.*, 2006) for human and mouse, respectively. Physical chromosome sizes in the dog were calculated from the *canFam2* coordinates of the first and last markers for each autosome. The average recombination rate for canine chromosomes was calculated from the autosomal linkage maps.

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FIGURE 1

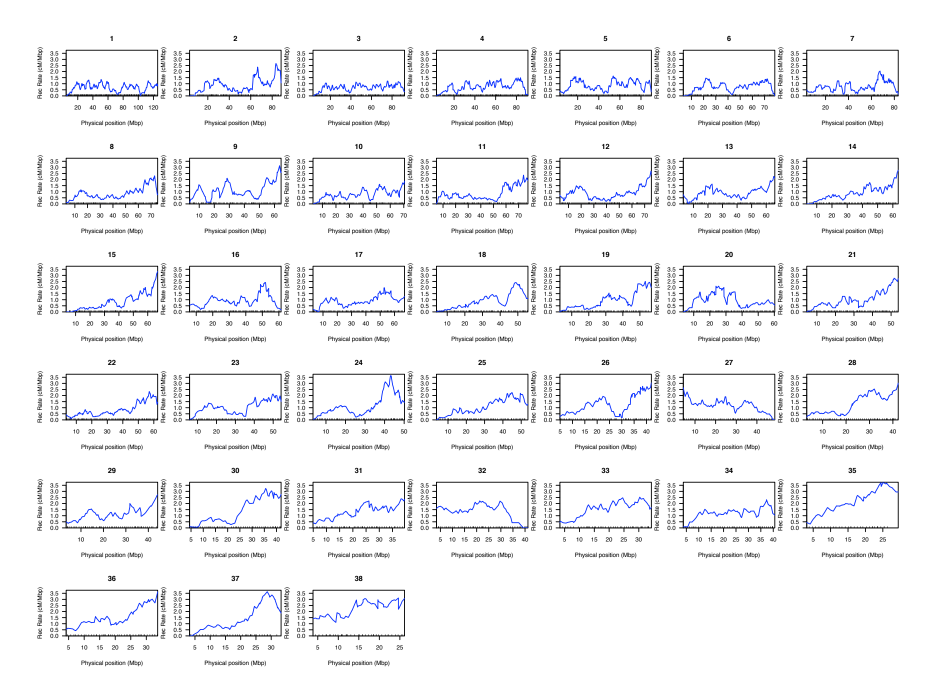
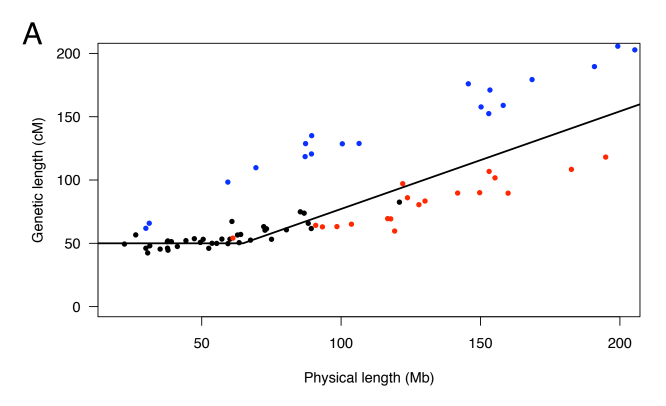


FIGURE 2



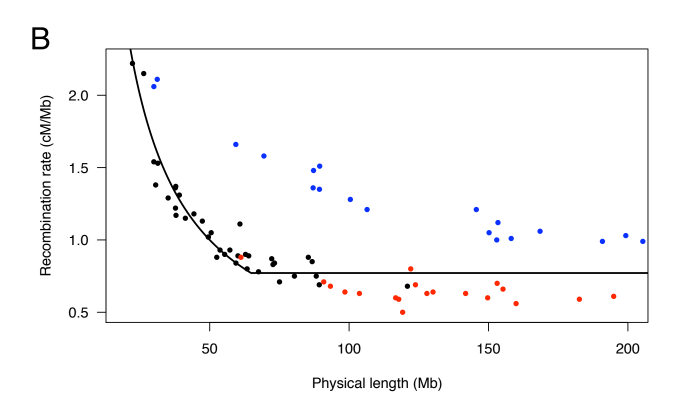


TABLE 1

Meiotic mapping resources used in this study

	No. of	Max.	Ava	Avg.	
Familial Resource	Dogs	Meioses	Avg. Parental HET	Informative Meioses	
CF Reference ^a	213	333	0.54 ± 0.08	149 ± 75	
F₂ Intercross ^ь	28	52	0.62 ± 0.01	26 ± 18	
Silken Windhound	40	65	0.50 ± 0.07	27 ± 19	
Total Resources	281	450	0.61 ± 0.09	201 ± 91	

^aCF, Cornell Families, described in (Neff et al. 1999).

^bParental types were Border Collie (paternal) and Newfoundland (maternal).

TABLE 2
Summary statistics of linkage maps^a

	Framework	Combined
No. of Markers	1496	3075
Genetic Length	1937 cM	2085 cM
Genetic Spacing	1.3 ± 1.7 cM	$0.7 \pm 0.9 \text{ cM}$
Physical Spacing	1.4 ± 1.0 Mb	0.7 ± 0.4 Mb
Gaps < 5 cM	94%	98%
Recombination Rate	0.92 cM/Mb	0.97 cM/Mb
Genetic Female/Male	1.28	1.19
Avg. Inf. Meioses	266 ± 57	201 ± 91

^aNot inclusive of the X chromosome.

TABLE 3Physical and genetic lengths of dog chromosomes

	Physical	Cov Aver (old)	Mala (aM)	Famala (aM)	A	No. of Mandage	
CFA (Mb)ª		Sex-Avg. (cM)	Male (cM)	Female (cM)	Avg. cM/Mb	No. of Markers	
1	120.9	82.5	69.3	96.5	0.68	174	
2	85.4	74.9	70.3	77.6	0.88	120	
3	89.3	61.7	50.2	74.0	0.69	121	
4	88.2	65.8	59.5	72.1	0.75	129	
5	86.8	73.8	69.8	76.6	0.85	123	
6	75.0	53.2	42.0	65.2	0.71	100	
7	80.4	60.7	51.6	69.8	0.75	108	
8	72.7	60.4	53.0	68.9	0.83	101	
9	60.8	67.3	69.1	65.5	1.11	85	
10	67.5	52.5	45.9	60.1	0.78	99	
11	73.3	61.5	56.5	65.9	0.84	94	
12	72.2	63.2	60.5	66.1	0.87	107	
13	62.9	56.6	47.3	67.8	0.90	87	
14	60.1	53.3	49.3	57.2	0.89	83	
15	63.4	50.5	53.0	47.8	0.80	92	
16	55.4	49.9	43.2	59.4	0.90	80	
17	64.0	57.0	62.7	55.0	0.89	87	
18	52.5	46.1	34.5	58.0	0.88	80	
19	53.7	50.0	49.5	52.1	0.93	76	
20	57.2	53.4	50.5	57.6	0.93	83	
21	49.5	50.7	45.8	54.4	1.02	74	
22	59.4	49.7	47.3	53.5	0.84	91	

Total	2274.4	2085.1	1910.0	2387.7	NA	3212
Χ	124.9	NA	NA	111.1	0.89	137
38	22.3	49.4	48.4	52.6	2.22	42
37	30.6	42.4	42.6	44.3	1.38	42
36	30.0	46.1	49.2	43.0	1.54	51
35	26.3	56.7	49.5	63.5	2.15	34
34	37.9	44.6	33.8	54.2	1.17	53
33	31.4	48.0	49.5	46.8	1.53	47
32	37.8	51.9	54.3	49.8	1.37	52
31	35.1	45.4	38.1	51.9	1.29	51
30	37.7	51.3	38.6	61.8	1.36	53
29	41.3	47.5	49.4	46.1	1.15	58
28	39.1	51.3	45.1	57.3	1.31	58
27	44.3	52.1	42.8	62.5	1.18	62
26	37.7	46.2	48.9	43.2	1.22	57
25	50.5	53.2	45.9	60.6	1.05	70
24	47.4	53.6	45.2	63.8	1.13	80
23	49.5	50.7	47.9	54.1	1.02	71

^aFrom the first and last marker coordinates in *canFam2*.

NA. not applicable.

TABLE 4 Sequence correlates of genome-wide variation in recombination rate.^a

	5Mb			10Mb		
Sequence Feature	Female	Male	Sex-Avg	Female	Male	Sex-Avg
GC%	-0.20	0.10	-0.04	-0.23 ⁻	-0.01	-0.14
No. CpG Sites	-0.05	0.32	0.16 ⁻	-0.08	0.30"	0.14
No. CpG Islands	-0.19	0.25	0.09	-0.22 ⁻	0.24 ⁻	0.07
Distance to Telomere	-0.23	-0.46***	-0.47***	-0.17 [·]	-0.60***	-0.56
Motif Count	-0.02	0.29	0.15 ⁻	-0.01	0.30"	0.16
No. RefSeq Genes	-0.04	0.16 ⁻	0.05	0.11	0.17 ⁻	0.10
No. NScan Genes	-0.06	0.22	0.08	0.05	0.12	0.02
SINES	0.19"	0.14 ⁻	0.12 ⁻	0.33***	80.0	0.09
LINES	-0.08	-0.05	-0.08	0.31***	-0.04	-0.01
LTRs	0.28	0.01	0.11	0.48***	-0.01	0.12
polyA	0.30	0.15 ⁻	0.23***	0.49***	0.12	0.24 [.]
polyC	-0.02	0.15 ⁻	0.06	0.09	0.06	0.02
polyG	-0.04	0.17 [.]	0.09	0.07	0.07	0.02
polyT	0.28	0.20	0.24	0.47***	0.12	0.23 [.]
Simple Repeats	0.32	0.06	0.18 ⁻	0.54***	0.06	0.20 [.]
AT-rich Repeats	0.24	-0.07	0.04	0.41***	-0.11	0.05
GC-rich Repeats	-0.08	0.25	0.12	-0.04	0.16	0.04
Satellites	-0.16 ⁻	-0.02	-0.09	-0.17 [·]	-0.08	-0.15
Low Complexity Repeats	0.24	-0.04	0.06	0.45	-0.07	0.09
DNA Repeats	0.30	0.20"	0.23	0.43	0.09	0.16

^a Rank correlation coefficients were calculated in 5 Mb and 10 Mb sliding windows.

P < 0.01 P < 10⁴ P < 10⁴