



# Radiation hybrid mapping of cataract genes in the dog

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**Purpose:** To facilitate the molecular characterization of naturally occurring cataracts in dogs by providing the radiation hybrid location of 21 cataract-associated genes along with their closely associated polymorphic markers. These can be used for segregation testing of the candidate genes in canine cataract pedigrees.

**Methods:** Twenty-one genes with known mutations causing hereditary cataracts in man and/or mouse were selected and mapped to canine chromosomes using a canine:hamster radiation hybrid RH5000 panel. Each cataract gene ortholog was mapped in relation to over 3,000 markers including microsatellites, ESTs, genes, and BAC clones. The resulting independently determined RH-map locations were compared with the corresponding gene locations from the draft sequence of the canine genome.

**Results:** Twenty-one cataract orthologs were mapped to canine chromosomes. The genetic locations and nearest polymorphic markers were determined for 20 of these orthologs. In addition, the resulting cataract gene locations, as determined experimentally by this study, were compared with those determined by the canine genome project. All genes mapped within or near chromosomal locations with previously established homology to the corresponding human gene locations based on canine:human chromosomal synteny.

**Conclusions:** The location of selected cataract gene orthologs in the dog, along with their nearest polymorphic markers, serves as a resource for association and linkage testing in canine pedigrees segregating inherited cataracts. The recent development of canine genomic resources make canine models a practical and valuable resource for the study of human hereditary cataracts. Canine models can serve as large animal models intermediate between mouse and man for both gene discovery and the development of novel cataract therapies.

As the leading cause of worldwide blindness and low vision, cataracts affect millions of individuals, and billions of dollars are spent annually on cataract medical and surgical expenses [1]. The etiology of cataracts varies; they can be hereditary, traumatic, or secondary to ocular inflammatory or systemic diseases, or as a sequelae to taking certain pharmaceuticals/medications. The onset of cataract also varies. Congenital cataracts occur in over two out of 10,000 live births [2] and are the leading cause of treatable visual impairment in children worldwide [3,4]. Approximately 1/3 of all congenital cataracts are inherited [5]. Age-related cataracts, however, are the most common type of cataract. Approximately 26.6 million Americans older than 40 years have cataract or have had cataract surgery, and this number is projected to increase to 39.6 million by the year 2020 [6]. The risk factors associated with the development of age-related cataract include ultraviolet light exposure [7], cigarette smoking [8], alcohol consumption [9], and nutrition [10]. In addition, the role of genetics in age-related cataract has been reviewed and sug-

gests an interaction between genetic background and environmental risk factors [11].

Most of our knowledge of cataractogenesis comes from genetic studies of hereditary cataracts in man and mice. Mutations causing nonsyndromic inherited cataract have been identified in at least 15 genes in humans [12], and at least 18 genes in the mouse (Mouse Genome Informatics). Here, we propose the development of the dog as a complementary large animal model for molecular studies of cataracts to facilitate gene discovery and the development of novel and alternative cataract therapies. There are over 300 breeds of dog, and of these, 20 breeds are known to have hereditary cataracts while another 125 breeds are suspected of having hereditary cataract [13]. It is likely that some hereditary cataracts in dogs will be caused by mutations in genes already identified in humans or mice. However, to date no mutations causing cataract have been identified in any canine pedigrees. Because of the recent progress in canine genomics [14-19], and the success in identifying canine models for retinal diseases [20-22], it is likely that similar progress will be possible with molecular studies of canine cataracts.

To begin the molecular identification of cataract phenotypes in the dog we have selected 21 genes known to cause cataracts in man and/or mouse. These include structural proteins (6), crystallin genes (6), and transcription factors (9). We have determined the canine chromosomal locations of each

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gene using a radiation hybrid (RH) 5000 rad panel [23] and compared our experimentally derived chromosomal locations for each gene with that of the draft sequence of the canine

genome. RH-mapping and genome sequencing are complementary procedures which identify gene locations, order, and distance, along with regions of conserved synteny across spe-

**TABLE 1. PCR PRIMER SEQUENCES USED TO PLACE CATARACT ORTHOLOGS ON CANINE CHROMOSOMES USING AN RH5000 PANEL**

Gene	GenBank accession number	Primer sequences	T <sub>m</sub> (°C)	Product size (bp)
<b>Structural protein genes</b>				
BFSP2	NM_003571	F: CCTCTTCATAGTTCCTTGAC R: CCCTACAGTTCTATGCTTCCAG	50.7 55.8	419
CX46	NM_021954	F: CCAGAGAATGCTAATTTGTCCC R: CAGCACATGCACTTACACAGG	54.9 57.6	365
CX50	NM_005267	F: GGGATAACCAGAGGGTAGCAC R: GGATCTCGAAGTTCTCCTACC	57.8 54.9	386
LIM2	NM_030657	F: GGCAACAAGTGCTACCTGC R: CCCAGTTTCCAACCTAGTTC	56.9 53.0	353
MIP	NM_012064	F: GGGCACGAGTACGAGCACTG R: GCCTGCACGCTTCGCAAGC	61.0 63.5	308
SPARC	NM_003118	F: CTGATCATGTGGCCTTGAGC R: GGCCTAATCTGATCTGCTAAG	56.2 52.5	343
<b>Crystallin genes</b>				
CRYBB2	NM_000496	F: GGAGAGCAGTTTGTGTTGAG R: GTGCAATGTGGCAACCCCTTAG	55.9 56.9	356
CRYGA	NM_014617	F: CCTACGAAGGAATCCCTTTTG R: GAGTGGAGCTCATTGAGCCG	53.2 58.1	349
CRYGB	NM_005210	F: GCTCCTGGCATCCAGTGGG R: CTGCTCCTGCTGCTCTACTC	61.0 57.3	404
CRYGC	NM_020989	F: CTCTAACCAGACTCTGAACC R: GGACCCAGCAGCCCTCCAG	54.3 63.3	373
CRYGD	NM_006891	F: GACAGCGGCTGCTGGATGC R: CCAGGACAGGACCTATTGCTG	62.1 57.4	381
CRYGS	NM_017541	F: CAGCTCATCCCAAGACTGAATG R: GACACCATCCAGCACCTTACTG	55.7 57.7	390
<b>Transcription factors</b>				
CHX10	NM_182894	F: CGTAAGTTAAGTACCAGAGGG R: CTAAACCTGTGATTTCTGTGCC	52.1 54.0	390
EYA1	NM_172060	F: CTACTGCCAAACTCATCAC R: CCCACAGAAAGAATGTAGATG	53.2 50.7	386
FOXE3	NM_012186	F: GCAAGTCACTGCAGGGACTG R: GTTCTTTAGTGGTGCTGGAGG	58.6 55.5	342
HSF4	NM_001538	F: GATTCTGAGCTCTACCATAG R: GCCAGGGTCTGGTTGAAGC	51.7 58.9	463
MAF	NM_005360	F: GTTAACAGCACGGAGGTTACAC R: CCCTTCCACTGCATCTGAGTC	55.9 57.6	340
PAX6	NM_001604	F: GCCACATCTTCAGTACAAAG R: TAGTTCAGGCATTGACTGATG	49.6 50.3	300
PITX3	NM_005029	F: CCTCCCTGAACAGGGTGATAG R: CCGCACCATTGCACACCACG	57.5 63.0	304
SIX5	NM_175875	F: GACCCGCAGCTTCTCAAGC R: CCTTCCTACTGCAAAGTGAGC	59.0 55.8	371
SOX1	NM_005986	F: CTCAAGAATGATATCCACTGCTTC R: GTACATTTTCAGAGTCAATGTGGC	53.5 54.1	299

The column labeled "T<sub>m</sub>" indicates primer melting temperature. PCR product size and corresponding NCBI reference sequences are shown.

cies. Combining data from RH-maps with that from genomic sequences maximizes the information obtained from both methodologies [24] and helps to resolve discrepancies that might arise with the draft sequence information currently available. The identification of cataract gene locations along with their nearest polymorphic markers provides an integrated resource essential for segregation analysis of cataract phenotypes in canine cataract pedigrees. This resource will aid in the development of the dog as a complementary large animal model for the molecular study of cataracts and will help to advance our knowledge of naturally occurring cataracts in mammalian species. This may one day contribute to the development of new therapies to prevent, delay, or even reverse cataracts.

## METHODS

*Selection of cataract gene orthologs:* Twenty-one gene orthologs were selected as cataract candidate genes in dogs based on their ability to cause cataracts in man and/or mouse. These candidate genes include structural proteins: beaded filament structural protein 2 (*BFSP2*), connexin 46 (*CX46*), connexin 50 (*CX50*), lens intrinsic membrane protein 2 (*LIM2*), major intrinsic protein of lens fiber (*MIP*), and secreted protein acidic cysteine-rich (*SPARC*); crystallins:  $\beta$ B2-crystallin (*CRYBB2*),  $\gamma$ A-crystallin (*CRYGA*),  $\gamma$ B-crystallin (*CRYGB*),  $\gamma$ C-crystallin (*CRYGC*),  $\gamma$ D-crystallin (*CRYGD*), and  $\gamma$ S-crystallin (*CRYGS*); transcription factors: CEH10 homeodomain-containing homolog (*CHX10*), homolog of *Drosophila* eyes absent 1 (*EYA1*), forkhead Box E3 (*FOXE3*), heat shock transcription factor 4 (*HSF4*), avian musculoaponeurotic fibrosarcoma oncogene homolog (*MAF*), paired-box 6 transcription factor (*PAX6*), paired-like homeodomain transcription factor 3 (*PITX3*), homolog of *Drosophila* sine oculis homeobox (*SIX5*), and sex-determining region-Y (SRY)-box 1 (*SOX1*).

*Canine orthologous sequences:* Human gene cDNAs obtained from the National Center for Biotechnology Information (NCBI) database were analyzed against drafts of the canine genome to obtain orthologous sequences. The corresponding accession numbers for the human cDNA sequences are listed in Table 1. Human gene cDNAs for *BFSP2*, *CX46*, *CX50*, *MIP*, *PAX6*, and *PITX3* were analyzed against the original 1.5X (poodle) canine genome sequence through collaboration with The Institute for Genomic Research (TIGR). Human gene cDNAs for *CHX10*, *CRYBB2*, *CRYGA*, *CRYGB*, *EYA1*, *FOXE3*, *HSF4*, *MAF*, *SIX5*, *SOX1*, and *SPARC* were analyzed against the publicly available 1.5X (poodle) canine genome using the BLAST function, and cDNAs for *CRYGC*, *CRYGD*, *CRYGS*, and *LIM2* were analyzed against the 7.5X (boxer) canine whole genome shotgun sequence using the BLAT function. Primers were designed from putative intronic or untranslated regions of canine orthologs of each cataract candidate gene using Amplify software (Bill Engels, University of Wisconsin, Madison, WI). Primers were designed to amplify canine-specific sequences suitable for RH mapping and are summarized in Table 1. The average length of primers was 25 nucleotides with an average melting temperature of 56.2 °C, and the average PCR product size was 363 bp. All PCRs

were performed on an MJ Research Tetrad 2 thermocycler (Global Medical Instrumentation Inc., Ramsey, MN). To ensure canine specificity, initial PCRs were carried out at 25  $\mu$ l volume on control canine, hamster, and canine:hamster (1:5 ratio) DNA. PCR conditions were as follows: 2 min 94 °C, followed by 30 cycles of 30 s at 94 °C, 30 s at 56 °C, 30 s at 72 °C, and a final extension of 5 min at 72 °C. The PCR product sizes ranged from 144-463 bp. PCRs were optimized by adjusting the annealing temperature where needed.

*RH5000 panel:* Canine cataract candidate genes were positioned on a canine radiation hybrid map using an RH5000 panel consisting of 118 cell lines made by fusing 5,000 rad-irradiated canine fibroblast cells with TK-HTK3 hamster cells [23]. Information on the RH5000 panel may be obtained from the Canine Genetics Research Projects web site. The RH5000 panel has a retention frequency of 22% and a resolution limit of 600 kb [17]. The PCR reactions on the RH5000 panel were performed with 15  $\mu$ l volumes, using the same PCR conditions already described, and run in duplicate or triplicate. PCR products were analyzed on 2% agarose gels containing 0.05% ethidium bromide at 50-65 volts. Gel bands were visualized using a UV transilluminator, and images were taken of each gel. Gels were then scored for presence, absence, or ambiguity of the band of interest in each of the 118 cell lines.

*Radiation hybrid map construction:* Radiation hybrid data collected from the 21 cataract gene orthologs was merged with previous data from 3,270 markers [17], and analyzed using Multimap® software [25]. Two-point linkage analysis was performed for each canine ortholog to determine its nearest markers and chromosomal location. A subset of the linked markers was selected and used as a framework map based on their map locations as previously provided [17]. The Multimap® multipoint mapping algorithm was then used to place the cataract gene orthologs on canine chromosomes in relation to the markers within the framework map.

The May 2005 7.5X canine draft sequence predicted locations of each cataract ortholog were compared with the corresponding locations of each PCR amplicon used for RH-mapping. The resulting genome locations for each ortholog and corresponding sequence-tagged site (STS) are listed in Table 2.

## RESULTS & DISCUSSION

*Two-point linkage analysis:* Twenty-one cataract gene orthologs were positioned on canine chromosomes using an RH5000 panel of 118 cell lines. Orthologs were located on 16 of the 39 canine chromosomes with more than one ortholog represented on three chromosomes (*CFA1*, 5, and 37; Table 2). The nearest chromosomal markers were determined for each cataract gene ortholog using Multimap® two-point linkage analysis [25]. Logarithm of odds (LOD) scores for the two closest linked markers ranged from 8.0 (*CRYGD*:BAC\_374\_E1) to 27.7 (*CRYGS*:BAC\_374\_K23), with an average LOD score of 18.9 (Table 2). Theta ( $\theta$ ), a measure of the frequency of breakage between two markers, varies from 0 (no breakage between markers) to 1 (complete breakage between markers), and represents the distance be-

tween markers. Theta ranged from 0 (*CRYGC*:BAC\_381\_J22, *SIX5*:BAC\_375\_N13, and *SIX5*:FH2598) to 0.354 (*CRYGD*:BAC\_374\_E1) with an average  $\theta$  of 0.095 for the two closest linked markers (Table 2).

Two-point linkage analysis placed the  $\gamma$ -crystallin genes (*CRYGA*, *CRYGB*, *CRYGC*, and *CRYGD*) on CFA37. There are estimated to be seven  $\gamma$ -crystallin genes in humans; these include *CRYGA*, *CRYGB*, *CRYGC*, *CRYGD*, and *CRYGS*, and two pseudogenes, *CRYGE* and *CRYGF*. There is sequence homology between *CRYGA*, *CRYGB*, *CRYGC*, *CRYGD*, *CRYGE*, and *CRYGF* and these genes form a cluster at HSA 2q33-35, and in mouse they form a cluster on chromosome 1 (MMU1). In humans, *CRYGS* maps to 3q25-qter, and in mouse

to MMU16. Similarly, canine *CRYGS* was not part of the  $\gamma$ -crystallin cluster on CFA37, but mapped to CFA34 near BAC\_374\_K23 and marker REN266K05. Mutations causing cataracts have previously been identified in murine *Cryga* [26,27], *Crygb* [26], *Crygc* [27], *Crygd* [27,28], and *Crygs* [29,30] as well as human *CRYGC* [31], *CRYGD* [31], and *CRYGS* [32]. As a result, there is a high likelihood that the  $\gamma$ -crystallins will also be involved in canine cataracts.

LOD scores for the two closest linked markers to  $\gamma$ -crystallin genes A, B, C, and D ranged from 8.0 (*CRYGD*:BAC\_374\_E1) to 24.95 (*CRYGC*:BAC\_381\_J22) and  $\theta$  ranged from 0 (*CRYGC*:BAC\_381\_J22) to 0.354 (*CRYGD*:BAC\_374\_E1; Table 2). The two closest linked

TABLE 2. TWENTY-ONE CATARACT GENES MAPPED ONTO THE CANINE GENOME

Gene	Chromosome	Closest markers	LOD	$\theta$	STS location	Canine ortholog location
BFSP2	CFA23	REN156G20	22.448	0.086	33427673-33428089	33388029-33450927
		REN210D03	17.827	0.169		
CX46	CFA25	EST18C6	9.688	0.276	20988791-20989147	20986634-20987933
		BAC_372_A23	9.617	0.261		
CX50	CFA17	CFOR12C10	20.47	0.056	61506499-61506884	61498923-61505209
		PEZ8	19.834	0.058		
LIM2	CFA1	KLK2	18.201	0.035	108473188-108473537	108472780-08479290
		REN06N11	17.642	0.066		
MIP	CFA10	FH2537	25.039	0.09	3684287-3684594	3684852-3689232
		ATP5B	21.428	0.132		
SPARC	CFA4	FH4018	14.097	0.048	60879945-60880285	60873070-60887224
		BAC_374_E21	12.897	0.093		
CRYBB2	CFA26	CRYBB1	25.249	0.043	22335395-22335748	22331283-22338858
		REN131L06	24.11	0.064		
CRYGA	CFA37	AHT133	20.482	0.092	19453604-19453950	19453388-19455353
		REN105M20	20.482	0.092		
CRYGB	CFA37	BAC_381_J22	15.426	0.123	20124535-20124930	19440507-19442393
		FH2708	13.738	0.157		
CRYGC	CFA37	BAC_381_J22	24.954	0	19431676-19432046	19431504-19433284
		EST22D1	23.468	0.026		
CRYGD	CFA37	CRYGB	15.027	0.045	19425779-19426157	19424588-19426264
		BAC_374_E1	8.03	0.354		
CRYGS	CFA34	BAC_374_K23	27.708	0.021	22165843-22166230	22165569-22166681
		REN266K05	27.283	0.022		
CHX10	CFA8	EST10D4	23.799	0.025	50510969-50511356	50492693-50511026
		EST27E3	21.735	0.051		
EYA1	CFA29	STS246A16	19.01	0.119	23177126-23177509	23176722-23342854
		C29_188	18.473	0.138		
FOXE3	CFA15	FH3888	23.211	0.046	16294587-16294926	16295972-16298254
		FH3886	22.806	0.047		
HSF4	CFA5	AHTH201REN	13.127	0.052	85205320-85205780	85204186-85208678
		REN122J03	11.927	0.1		
MAF	CFA5	FH3450	9.517	0.244	74976764-74977099	74970434-74976948
		FH3113	9.427	0.224		
PAX6	CFA18	WT1	17.245	0.148	38687699-38688008	38671041-38693640
		FH3824	17.15	0.148		
PITX3	CFA28	EST3H1	19.137	0.083	17763487-17763792	17761261-17772191
		COL17A1	18.606	0.085		
SIX5	CFA1	BAC_375_N13	21.594	0	112730879-112731247	112726865-12731063
		FH2598	21.594	0		
SOX1	CFA22	EST3G2	26.711	0.022	62864830-62865127	62861615-62864949
		BAC_376_O1	24.774	0.045		

PCR primers (Table 1) from 21 cataract genes were used for radiation hybrid-mapping using an RH5000 panel. Multimap® two-point linkage analysis was used to identify the chromosome location and two closest markers [17], along with the corresponding LOD score, and  $\theta$  value (distance). To confirm these locations, PCR amplicons were analyzed against the 7.5X canine draft sequence to obtain sequence-tagged sites (STS) locations, which were compared with the corresponding canine ortholog locations within the draft sequence.

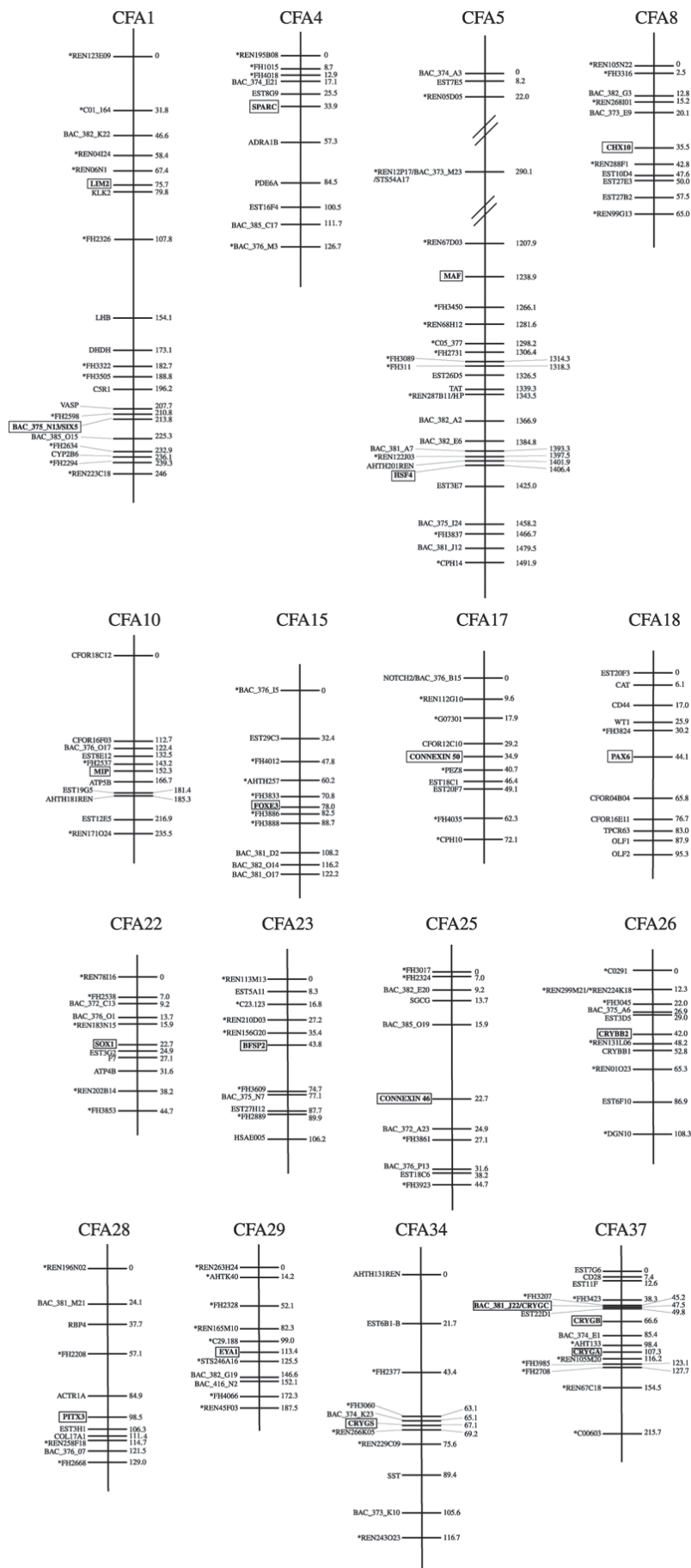


Figure 1. Radiation hybrid maps locating 20 cataract genes to canine chromosomes. In each *Canis familiaris* (CFA) chromosome representation, unboxed markers on left represent framework markers [17]. Markers with an asterisk are polymorphic. Boxed markers are location of canine orthologs of cataract associated genes. Numbers on right are distances in centiRays from the top framework marker.

markers to *CRYGD* were *CRYGB* from a previous study [17] (LOD=15.027,  $\theta=0.045$ ) and BAC\_374\_E1 (LOD=8.03,  $\theta=0.354$ ). Although in the previous study, the *CRYGB* marker was “unlinked” to any other marker, the corresponding 171 bp amplicon was located on CFA37 (19,425,908-19,426,078). Also, in the previous study, markers were incorporated into a 1,500 marker map using Multimap® pairwise calculation at a threshold of LOD>8 [15]. The *CRYGB* marker in that study may have linked to other markers in the data set at a lower threshold. Decreased marker density in the vicinity of a gene’s map location can also result in unlinked gene markers. However, marker density in the vicinity of *CRYGB* should have been sufficient in the previous study since they mapped the same markers, which linked to the  $\gamma$ -crystallins in this study.

The next closest linked marker to *CRYGD* (BAC\_374\_E1: LOD=8.03,  $\theta=0.354$ ) was also on CFA37. The decrease in LOD score from 15.027 (*CRYGD:CRYGB*) to 8.03 supports linkage but the increase in  $\theta$  from 0.045 (*CRYGD:CRYGB*) to 0.354 indicates an increase in the distance between these two markers. All of the markers that linked to *CRYGD* with LOD>4 were located on CFA37 (data not shown). In addition, the *CRYGD* PCR amplicon was located on CFA37 (19,425,779-19,426,157) within the predicted location for *CRYGD* based on reference sequence NM006891 (19,424,588-19,426,264). Thus, based on two-point linkage analysis results and the location of the *CRYGD* PCR amplicon, we place *CRYGD* on CFA37.

$\alpha$ A- and  $\alpha$ B-crystallin genes were previously mapped to CFA31 (*CRYAA*) and CFA5 (*CRYAB*) [33], and  $\beta$ A1- and  $\beta$ B1-crystallin to CFA9 (*CRYBA1*) [17] and CFA26 (*CRYBB1*) [17,33]. Here we placed *CRYBB2* on CFA26 near *CRYBB1* and *REN131L06*. Similarly, *CRYBB1* and *CRYBB2* both map to 22q11.2-12.2 in man, and chromosome 5 in mouse (MMU5). *MIP* was mapped to CFA10 near *FH2537* and *ATP5B*, in agreement with previously reported data [33]. Both *LIM2* and *SIX5* mapped to CFA1, and *HSF4* and *MAF* to CFA5. All other orthologs (excluding the  $\gamma$ -crystallins) mapped to individual chromosomes as summarized in Table 2.

**Sequence-tagged sites:** Sequence-tagged sites (STS) for each cataract ortholog were obtained by analyzing the corresponding PCR amplicons against the May 2005 7.5X canine draft sequence. With the exception of *CRYGB*, all of the resulting STS were located inside or within 1,300 bp of the predicted gene location based on human reference sequences (Table 2). The *CRYGB* STS (20,124,535-20,124,930) was not within the predicted location for canine *CRYGB* (19,440,507-19,442,393) based on the human reference sequence (NM\_005210). Primers used to amplify *CRYGB* were designed from the 1.5X (poodle) whole genome shotgun reference sequence AACN010184836. This sequence was selected from 1.5X canine sequences having homology to human *CRYGB* mRNA (NM\_005210). When analyzed against the 7.5X (boxer) canine genomic sequence, AACN010184836 had sequence homology to four unique sites on CFA37 (19,417,160-19,418,711; 19,426,006-19,426,485; 19,432,821-19,455,395; 20,124,527-20,126,527). These included the human reference sequence (NM\_005210) *CRYGB* predicted site (19,440,507-

19,442,393). However, the actual AACN010184836 sequence was located on CFA37 from 20,124,527-20,126,527, and the 396 bp portion of this sequence which corresponded to the *CRYGB* amplicon only had homology to CFA37 at 20,124,535-20,124,930 (which is the *CRYGB* amplicon location). Therefore, even though the *CRYGB* primers were designed from the AACN010184836 sequence which had homology to human *CRYGB*, this sequence and the PCR amplicon derived from it were >682 kb downstream from the expected *CRYGB* site.

Furthermore, there are four *CRYGB* predicted sites on CFA37 based on human protein homology (19,415,025-19,433,068; 19,424,652-19,455,135; 19,440,575-19,442,393; and 20,126,034-20,128,506). Human protein sequence homology is determined by analyzing human protein sequences against the human genome to find exonic sequences corresponding to amino acid sequences. The amino acid sequences are then analyzed against the canine genomic sequence using tBLASTn (protein query vs. translated database) to find predicted canine exons. Three of the four *CRYGB* predicted sites based on protein homology overlap and cover a 40 kb region (19,415,026-19,455,135). This region is 670 kb upstream of the fourth predicted *CRYGB* site (20,126,033-20,128,569). The *CRYGB* STS amplified here was located 1,104 bp upstream to this fourth site. Because there are multiple predicted sites for *CRYGB* on CFA37 in the dog genome, and these sites share sequence homology, it is difficult to design primers specific for any single *CRYGB* predicted site. The multiple sites of *CRYGB* sequence homology make the actual location of the canine *CRYGB* gene unclear. As a result, our *CRYGB* STS should not be used to predict the exact location of canine *CRYGB*.

**Radiation hybrid-mapping:** Markers linked to cataract orthologs, as determined by two-point linkage analysis, were used to create a framework map with previously established marker positions [17]. This framework map was then used to determine the map locations of the cataract orthologs (Figure 1). The orthologs were positioned between the two closest markers using the multipoint mapping algorithm of Multimap® software. In two cases, the cataract orthologs mapped to the same exact position as a BAC end marker in the framework map ( $\theta=0$ ). For example, *SIX5* mapped to CFA1 at the same location as BAC\_375\_N13, and *CRYGC* mapped to CFA37 at the same location as BAC\_381\_J22. Since BACs represent large (155 kb) DNA fragments [34], the mapping of these two genes to the same location as the two BACs suggests that these genes may be located within the BACS.

The RH maps generated by the Multimap® multipoint mapping algorithm were generally consistent with the results of two-point linkage analysis, and the orthologs mapped on top of, between, or near the closest linked markers as determined by two-point linkage. An exception was *CRYGD* which was placed on CFA37 based on two-point linkage results and the location of the *CRYGD* STS (Table 2) but could not be accurately localized within the framework map. The framework map used to localize the  $\gamma$ -crystallin genes on CFA37 included the  $\gamma$ -crystallin gene vectors (*CRYGA*, *CRYGB*, *CRYGC*, *CRYGD*) along with framework markers whose lo-

cations were previously determined [17]. As a result, the framework map did not include the unlinked *CRYGB* marker from the previous study [17], which was the closest linked marker to *CRYGD* based on two-point linkage analysis. Instead, the closest linked marker to *CRYGD* in the framework map was BAC\_374\_E1 which linked to *CRYGD* with LOD=8.03, and  $\theta=0.354$ . All of the other markers in the framework map which linked to *CRYGD* had a LOD score <8.03 and a  $\theta$  value >0.354. These  $\theta$  values indicated an increased frequency of breakage (increased distance) between *CRYGD* and the markers on the framework map and decreased the likelihood of accurately localizing *CRYGD* on the map. As a result, although we placed *CRYGD* on CFA37 based on two-point linkage analysis and the location of the *CRYGD* amplicon, we were unable to accurately localize this gene on the framework map using the Multimap® multipoint mapping algorithm due to the decreasing LOD scores and increasing  $\theta$  values of the markers linked to *CRYGD*. The lack of markers linked to *CRYGD* with high LOD scores and low  $\theta$  values may indicate low marker density in the vicinity of *CRYGD*. Even though all of the  $\gamma$ -crystallins orthologs (*CRYGA*, *CRYGB*, *CRYGC*, *CRYGD*) were located on CFA37 and were mapped against each other, this did not resolve the problem of marker density in the vicinity of *CRYGD*. To resolve this problem *CRYGD* should be mapped on the new RH9000 panel [24].

$\gamma$ A-,  $\gamma$ B-, and  $\gamma$ C-crystallin genes mapped to CFA37 as expected based on results of two-point-linkage analysis. The 5'→3' RH-map order on CFA37 (*CRYGC*, *CRYGB*, *CRYGA*; Figure 1) matched the order of the predicted gene locations within the 7.5X draft sequence (Table 2, Column G). However, this RH-map order did not correspond with the order of the STS locations within the 7.5X draft sequence (*CRYGC*, *CRYGA*, *CRYGB*; Table 2, Column F). Two-point linkage analysis linked the *CRYGB* ortholog to multiple markers on CFA37. The two closest linked markers were BAC\_381\_J22 (LOD=15.426,  $\theta=0.123$ ), and FH2708 (LOD=13.738,  $\theta=0.157$ ). The Multimap® multipoint mapping algorithm placed the *CRYGB* ortholog on CFA37 between *EST22D1* and BAC\_374\_E1, which are also between BAC\_381\_J22 and *FH2708*. Thus, two-point linkage analysis and radiation hybrid mapping results are consistent and place *CRYGB* on CFA37 in the same region, however this location does not correspond with the *CRYGB* STS location.

RH maps are physical maps based on the statistical analysis of the pattern of marker retention in a panel of RH-cell lines. As a result, the order and location of markers within RH-maps do not always correspond exactly with the order and location of markers within genomic sequences. Marker order and position on the canine RH5000 map [17] for example, are not equivalent with their order and position in the 7.5X canine genomic sequence in all instances. Thus, it is not surprising that the *CRYGB* marker in this study mapped to a location on the RH-map that does not exactly correspond to the physical location of this marker within the canine genome. Nevertheless, RH-maps are valuable in localizing markers and genes to specific regions and providing statistically close markers for association and linkage testing.

Furthermore, the discrepancy between the order of *CRYGA*, *CRYGB*, and *CRYGC* on the RH-map and their STS in the draft sequence may also reflect the limited power for fine resolution of the RH5000 panel, which has a resolution limit of 600 kb. The distance between *CRYGA* and *CRYGB* STS is about 670 kb, and the distance between *CRYGB* and *CRYGC* STS is about 692 kb. These distances are close to the resolution limit for the RH5000 panel. Further, the distance between *CRYGA* and *CRYGC* STS is only 21,558 bp and is outside the resolution limit for the panel. Therefore, the inconsistency between the RH-map order and the STS order for *CRYGA*, *CRYGB*, and *CRYGC* may be due to the resolution limit of the RH5000 panel. The new RH9000 panel has a resolution of 200 kb [24] and can be used to resolve mapping issues for the  $\gamma$ -crystallin genes in the dog.

The canine chromosomal locations of the 21 cataract orthologs were consistent with human and canine established chromosomal synteny [35-37]. All of the cataract orthologs were mapped to canine chromosomal locations which were syntenic with the corresponding human gene locations, except for four orthologs (*SPARC*, *HSF4*, *CX46*, and *PITX3*) that mapped to locations directly adjacent to the corresponding syntenic region of the human gene location. Although *CRYGD* was not localized on the RH-map, the results of the two-point linkage analysis did place this gene on CFA37 in a region with homology to *HSA2q33-35* where the gene is located in humans.

To facilitate the molecular characterization of naturally occurring cataracts in the dog we have provided the RH location of 20 cataract-associated genes along with their associated polymorphic markers (Figure 1). Primers for selected polymorphic markers near cataract-associated genes can be obtained from the RH5000 website and serve as a resource for association and linkage testing in canine pedigrees segregating inherited cataracts. Although *CRYGD* is not represented in Figure 1, two-point linkage analysis did link *CRYGD* to BAC\_374\_E1 which localizes it on CFA37 to some degree. Also, the *CRYGD* STS location (19,425,779-19,426,157) can be used to identify proximal markers in the canine genome sequence. Though the RH5000 panel was used to map cataract-associated genes in this study, these results have not been incorporated into the RH5000 map. Nevertheless, the RH5000 map can be used to locate the markers referred to in this study (Table 2 and Figure 1). In addition, while the RH5000 and RH9000 maps will not be merged, a subset of 545 BACs has been mapped on both RH-panels and is part of both maps allowing navigation between the RH5000 and RH9000 maps.

Animal models provide a tremendous tool in the study of cataract, and molecular studies in mice have been instrumental in advancing our knowledge of the mechanisms of cataract formation. The dog has recently emerged as a promising model organism for gene discovery and genomic studies, and is a complementary species to the mouse for molecular genetic studies. There are over 300 known breeds of dogs, and most function as genetic isolates due to restricted mating. As a result, there is a decrease in genetic diversity within breeds and an increase in diversity between breeds such that genetic dis-

eases within one breed tend to be caused by the same mutations (founder effect). This facilitates the mapping of disease loci. In addition, developments in canine genomic resources that include integrated RH maps [14-17,24], the availability of the canine genomic sequence [18,19], and comparative chromosome maps of the human and dog genomes which identify homologous segments and conserved synteny between canine and human karyotypes [35-37] make it practical to develop and use canine models.

Although mutations causing human cataracts have been identified in 15 genes there are many other genes with mutations yet to be identified. The study of naturally occurring cataracts in the dog could reveal novel genes or mutations not yet identified in either man or the mouse. The dog represents a valuable yet largely untapped resource to study the molecular genetics of cataracts. This study takes some initial steps toward developing the tools needed to begin studies on the canine cataract model. We have identified the chromosomal locations of 20 canine orthologs of cataract-associated genes and their nearest polymorphic markers (Figure 1) that can be used for association and linkage testing in cataract pedigrees. This resource, along with the recent advances in canine genomics, will help facilitate the development of such models for the study of naturally occurring inherited cataracts. Once developed, canine cataract models should help to identify new cataract genes and disease-causing mutations, assist in the development of new therapies, and provide an intermediate large animal model between mouse and human for preclinical trials of new therapies.

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