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# Characterization of Gene Expression Profiles of Normal Canine Retina and Brain Using a Retinal cDNA Microarray

## Abstract

**Purpose:** Construction of a canine retinal custom cDNA microarray for comprehensive retinal gene expression profiling and application for the identification of genes that are preferentially expressed in the retina and brain lobes using a brain pool reference tissue.

**Methods:** A cDNA microarray was constructed utilizing clones obtained from a normalized canine retinal expressed sequence tag library. Gene expression profiles were analyzed for normal retina, as well as the cortex of the frontal, occipital, and temporal brain regions. Each sample was studied against a reference sample of pooled brain RNA. Data from a quantified scanned image were normalized using the loess subgrid procedure. Retina-enriched genes were identified using the Significance Analysis of Microarrays (SAM) algorithm, and confirmed by northern blot analyses for selected genes. Differences between biological samples were displayed using principal component analysis (PCA).

**Results:** Expression profiles for each tissue set were analyzed against the common reference of pooled brain. Changes in expression between the sample and the reference were higher in the retina (27.9%) than the individual brain tissues (2-6.6%). Furthermore, all individual retinal samples were clearly separated from any of the hybridizations using brain tissue in the PCA. The accuracy of observed changes in expression has been confirmed by northern blot analysis using five randomly chosen genes that represented a wide range of different expression levels between retina and brain.

**Conclusions:** We have established an accurate and robust microarray system suitable for the investigation of expression patterns in the retina and brain. Characterization of the gene expression profiles in normal retina will facilitate the understanding of the processes that underline differences between normal and diseased retinas.

## Disciplines

Comparative and Laboratory Animal Medicine | Medicine and Health Sciences | Ophthalmology | Veterinary Medicine

## Comments

Paez, G. L., Sellers, K. F., Band, M., Acland, G. M., Zangerl, B., & Aguirre, G. D. (2006). Characterization of gene expression profiles of normal canine retina and brain using a retinal cDNA microarray. *Molecular Vision*, 12, 1048-1056. <http://www.molvis.org/molvis/v12/a118/>

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# Characterization of gene expression profiles of normal canine retina and brain using a retinal cDNA microarray

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**Purpose:** Construction of a canine retinal custom cDNA microarray for comprehensive retinal gene expression profiling and application for the identification of genes that are preferentially expressed in the retina and brain lobes using a brain pool reference tissue.

**Methods:** A cDNA microarray was constructed utilizing clones obtained from a normalized canine retinal expressed sequence tag library. Gene expression profiles were analyzed for normal retina, as well as the cortex of the frontal, occipital, and temporal brain regions. Each sample was studied against a reference sample of pooled brain RNA. Data from a quantified scanned image were normalized using the loess subgrid procedure. Retina-enriched genes were identified using the Significance Analysis of Microarrays (SAM) algorithm, and confirmed by northern blot analyses for selected genes. Differences between biological samples were displayed using principal component analysis (PCA).

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**Conclusions:** We have established an accurate and robust microarray system suitable for the investigation of expression patterns in the retina and brain. Characterization of the gene expression profiles in normal retina will facilitate the understanding of the processes that underline differences between normal and diseased retinas.

The domesticated dog, *Canis familiaris*, exhibits a diverse repertoire of morphological and behavioral characteristics that define the numerous specific breeds. This segregation, in effect the creation of genetic isolates, has resulted in the emergence of autosomal recessive inherited canine diseases that are often directly comparable to those observed in humans. Many of these affect the retina and result in a large group of genotypically distinct retinal disorders representing molecular and phenotypic counterparts in humans. By far the largest group of diseases in dog, termed progressive retinal atrophy (PRA), serve as disease homologs for retinitis pigmentosa (RP) in humans [1]. Although the final common pathway of photoreceptor degeneration in many retinal degenerative processes appears to be apoptosis [2-4], it is likely that differing mechanisms link the causative mutations to the cell death processes. Insights into these mechanisms are critical to develop therapeutic strategies, and these studies are dependent on animal models. One approach to examine the pathways involved in retinal degenerative diseases in animal models is gene expression profile analysis with the aim of identifying early alterations in gene expression that contribute to photoreceptor cell death [5]. The microarray technology shows

potential as a useful tool to identify genes and gene pathways involved in ocular disease progression and may therefore aid in the characterization and classification of these diseases, possibly identifying a common mechanism linking disorders into groups [6,7].

Microarrays have been useful in the simultaneous analysis of transcript levels of thousands of genes in different physiological states of an organism, tissue, or cell [8,9]. Construction of microarrays is most efficient when information is utilized from annotated genomes or expressed sequence tags (ESTs) and has led to new insights into animal development, cancer, infectious diseases, and aging [10-12]. A major limitation of the technique is the analysis of sequences represented on the array. This disadvantage is particularly problematic when analyzing highly specialized tissues such as the retina due to the repertoire of uniquely or preferentially expressed genes contributing to its structure and function. In the case of "orphan" species, commercial arrays are either not available or targeted to a broader research community and underrepresent the tissue of interest. The current available canine microarray is based on transcripts representing 11 nonretina tissues and gene predictions not supported by experimental data (Affymetrix Genechip® Canine Genome Array, Santa Clara, CA). To obtain insight into intrinsic processes of the canine and human retina, great efforts are directed toward the identification and characterization of transcripts with functional relevance to this tissue [13-15]. Despite these ad-

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vances, a remaining challenge is to obtain an expression map of the canine retina/retinal pigmented epithelium transcriptome, further facilitating the identification of retinal susceptibility genes but, most importantly, offering an invaluable resource for functional genomics studies.

To overcome these limitations and drive the development of the canine retina transcriptome, we selected about 4,500 genes from a normalized canine retinal EST database [16]. These were used to construct a canine retinal cDNA microarray for expression profiling of genes that are of interest in studies of normal development and function and pathological conditions of the retina. As the brain and retina have a common embryological origin, a brain pool was initially validated as a reference sample for comparing expression profiles in multiple samples from animals with different molecularly defined retinal diseases and at different stages of the disorders. As part of this study, we have compared gene expression profiles between normal retina and different regions of the brain cortex (frontal, occipital, temporal) to identify those genes preferentially expressed in either tissue. The overall aim of this study is the identification of retina specific gene expression patterns that will maximize the use in a variety of PRA dog models [1].

## METHODS

**Tissue collection and RNA preparation:** Retinas and brain samples were collected from three 16-week-old normal beagles (two males and one female) that were part of a specific-pathogen-free (SPF) colony at Cornell University. In addition, retinas and spleen samples were obtained from two 16-week-old crossbred dogs (one male and one female) that were part of NEI/NIH sponsored projects (R01EY06855, R01EY13132) and were kept at the Retinal Disease Studies Facility (RDSF) in Kennett Square, PA. All dogs were housed in an indoor kennel under 12 h cyclic light conditions, and eyes were collected at a single time period (noon) to avoid potential fluctuations in retinal gene expression with time of day [17-22]. Both eyes were enucleated from the light-adapted dogs following intravenous anesthesia with sodium pentobarbital, and the dogs were euthanized after enucleation with an overdose of the barbiturate. The retinas were collected within 1-2 min after enucleation, and brain and spleen samples obtained within 5-10 min after euthanasia; all tissues were flash frozen in liquid nitrogen and stored at -80 °C until use. The research was conducted in full compliance with the ARVO Resolution on the Use of Animals in Research.

**RNA isolation:** Total RNA was isolated using Trizol reagent (Invitrogen, Carlsbad, CA) and further purified by RNeasy mini kit (Qiagen, Valencia, CA). Purity and RNA quality were evaluated by absorbance at 260 nm and by denaturing formaldehyde agarose gel electrophoresis. High quality RNAs with  $A_{260}/A_{280}$  ratio over 1.8 and intact 28S and 18S RNA bands were used for microarray analysis. To generate an RNA reference sample for microarray hybridizations, we pooled equal amounts of total RNA from the cortex of the occipital, temporal, and frontal brain regions collected from three 16-week-old beagles to achieve a homogeneous pool of

transcripts. The pooled RNA was divided into aliquots (2 µg/µl) and stored at -80 °C until use. For the purpose of this work, four tissue groups have been established, containing five biological replicates for normal retina and three for each respective brain region. After initial validation of reproducibility, only one microarray experiment was used for each sample.

**Microarray construction:** For microarray production, 3,931 "UniGene" sequences are represented by individual clones from a canine retinal EST library [16]; another 120 unique sequences were added by two clones each to ensure complete coverage of cDNA. In addition, about 300 genes resulting from retinal subtraction libraries [16] or serving as controls were printed on the microarray. Each clone was picked from glycerol stocks manually, inoculated in 50 µl of LB Broth Base (Invitrogen) with 100 µg/ml ampicillin and grown overnight at 37 °C. PCR was performed in two 15 µl reactions for each clone by dipping the template into a master mix (1.5 µM MgCl<sub>2</sub>, 0.2 µM dNTP, 0.4 µM of each primer [T7: 5'-TAA TAC GAC TCA CTA TAG GG-3'; Sp6: 5'-ATT TAG GTG ACA CTA TAG-3'] and 1 U Taq polymerase) with a replicator, and amplified for 40 cycles at 94 °C for 10 s, 54 °C for 20 s and 72 °C for 1 min, after an initial denaturing step at 95 °C for 1 min, and followed by a final extension at 72 °C for 10 min. PCR products for each clone were pooled and purified using Millipore Montage PCR<sub>384</sub> Filter Plates (Millipore, Billerica, MA). PCR products were resuspended in 50 µl of distilled water, and 6 µl were run on a 1% agarose gel. Rhodopsin, (*RHO*), and β-Actin (*ACTB*) were used as positive controls, while microsatellite locus CUX20001 [23] was used as a genomic control spot. All PCR products plus Array Control™ PCR Spots (Ambion, Austin, TX; spots 1-8) were spotted on Telechem superamine glass slides (TeleChem International, Inc., Sunnyvale, CA) using a Cartesian Pysix 5000 printer.

**Microarray hybridization and scanning:** Ten µg of pooled canine brain total RNA was hybridized against 10 µg of the respective test sample. Tissues were fluorescently-labeled using an indirect labeling protocol (3 DNA Array Kit, Genesphere, Inc. Hatfield, PA) following the manufacturer's protocol. Hybridization signals were visualized with Cy5 and Cy3 fluorescent reporter molecules, and intensities were detected using a GenePix 4000B Scanner (Molecular Devices Corporation, Union City, CA). Scanned images were processed using GenePix Pro version 6.0 (Molecular Devices Corporation, Downingtown, PA). The main quantities of interest produced by the image analysis (segmentation and background correction) are the (R, G) fluorescence intensity pairs for each gene on each array (R=red for Cy5, and G=green for Cy3). Loess (locally weighted regression) normalization was performed to remove systematic variation that occurs in every microarray experiment. An MA plot was used to represent the (R, G) data, where  $M = \log_2(R/G)$  and  $A = \log_2(RG)$  [24]. For complete consideration surrounding the normalization process, loess normalization over the entire chip, and block-specific loess normalization followed by additional scaling via maximum likelihood estimation (MLE) or median absolute deviation (MAD) were also studied.

**Statistical analysis:** Each hybridization was annotated according to MIAME (Minimal Information About a Microarray Experiment) standards, and analyzed with GeneSpring version 7.2 (Silicon Genetics, Agilent Technologies, Palo Alto, CA; GPL3951, GPL3957, and GSE5208). Controls were used for quality control and normalization of the microarrays using GeneSpring, but ignored for the statistical data analysis. Significant changes in expression were identified with Significance Analysis of Microarrays (SAM, 1.15, Stanford University, Palo Alto, CA) [25] at a delta value resulting in a false discovery rate (FDR) of less than 10%. Only statistically significant changes will be reported here; however, the entire dataset is available from the authors on request.

Principal component analysis (PCA) was performed on a complete microarray dataset consisting of log-transformed expression values for each individual replicate. As a result, the dimensionality of the dataset was reduced and described in a series of axes that explain the variance within the data. In this analysis, 30.4%, 16.98%, and 14.1% of the variance within the data was accounted for by the X, Y, and Z axes, respectively. These were used to plot the data and give an impression of the variation between the individual datasets; distances between points on the graph are indicative of the differences between replicates.

**Northern analysis:** Northern analysis was used to confirm expression patterns observed in the microarray data. Total RNA (10 µg) from brain (equal representation of frontal, temporal, and occipital cortex regions), retina and spleen were separated on a 1% formaldehyde-agarose gel and transferred to a membrane (GeneScreen Plus Hybridization Transfer Membrane, Perkin Elmer Life Sciences, Inc. Boston, MA). The RNA was crosslinked to the membrane and hybridized to purified cDNA probes labeled with [ $\alpha^{32}$ P] dCTP using a DNA-labeling system (RadPrime DNA Labeling System, Invitrogen) at 68 °C overnight using ExpressHyb Solution (Clontech, Palo Alto, CA), and washed in 2X SSC, 0.05% SDS, and 0.1% SSC, 0.05% SDS for 40 min each.

Probes for northern analysis were obtained by RT-PCR with primers designed from the corresponding cDNA clone information [16] (Appendix 1). For RT-PCR, 0.5 µg of total RNA was incubated in a reaction mixture containing primer (50 µM oligo(dT)<sub>20</sub>, or 50 ng/µl random primer), 10 µM of dNTPs mix, 40 U/µl of RNase inhibitor, ThermoScript™ RT (15 U/µl; Invitrogen), 25 mM of MgCl<sub>2</sub>, 10X cDNA synthesis buffer in a final volume of 25 µl. The samples were incubated at 70 °C for 5 min to denature the RNA then put on ice and amplified as follows: 94 °C, 1 min; 50-63 °C, 1 min; 72 °C for 1 min, for 35 cycles. The RT-PCR product was fractionated by electrophoresis on a 1.4% agarose gel containing 1 µg/ml ethidium bromide in 1X TAE buffer. Each membrane was also hybridized with canine *ACTB* cDNA to normalize for RNA loading, and image intensities of the blots were quantified using a Bio-Imaging Analyzer-BAS 1000 (Fujifilm life Science, Tokyo, Japan) and its associated software.

## RESULTS

We produced a custom-made microarray chip featuring about 4,500 retina-specific genes to investigate expression profiles of the retina. First experiments evaluating reproducibility found negligible variance between technical replicates (data not shown). To allow cross-comparison between multiple conditions, the experiment was arranged as reference design thus allowing for an equal standardized error between each individual comparison and avoiding dye bias. Therefore, each biological sample was hybridized only once. Given the differences in test and reference sample, different normalization approaches were considered: block-specific loess normalization (subgrid normalization), general loess normalization on the entire array (global normalization), and subgrid loess normalizations followed by either a MLE or MAD scaling procedure (Appendix 1). This confirmed that the subgrid normalization, as performed by GeneSpring, did not introduce data bias, and was used for the presented results.

The canine retinal cDNA microarray was used to identify genes preferentially expressed in normal 16-week-old retina in comparison to a pooled brain reference sample. Quantitative differences in the level of gene expression are illustrated in a scatter plot (Figure 1) comparing the expression level for normal retina (X-axis) versus the expression level from the brain pool (Y-axis). High abundance of tissue-specific transcripts is observed in spots located outside the two-fold expression lines (Figure 1, lines 2 and 3). SAM was used with a FDR of 10%, to confirm that 13.2% of 3,070 valid genes strongly hybridized to retinal cDNA targets, implying that 405 genes on the microarray are preferentially expressed in the normal retina (Table 1, Appendix 1). Of the remainder, 2,213 transcripts showed equal expression between normal retina and the brain pool reference tissue, and 14.7%, 452 genes, were less abundant in the retina (Table 2). Note that we call the spotted DNA sequences “genes” whether they correspond to actual genes or ESTs.

To further characterize the genes found to be preferentially expressed in brain, we hybridized the same brain pool against cortex of frontal, occipital, and temporal brain regions of three the different animals. Three percent of 3,489 valid transcripts hybridized preferentially with the frontal region; this corresponds to 114 genes highly expressed in the frontal cortex. The occipital and temporal regions, respectively, had 22 and 20 genes (of 2,089 and 2,289 valid transcripts, respectively) that were more abundant. Most transcripts were unique to a single cortex area (Figure 2A, Table 3), and only two genes were upregulated in all three brain regions (DR010012A10F04, *TM4SF1*). Transcripts less abundant than in the pooled reference also demonstrated a low overlap between brain regions (Figure 2B, Table 4), with one gene downregulated in all three brain cortex areas (*FLJ10159*).

The obtained expression profiles were analyzed using PCA revealing independent clustering of experiments derived from retinal and brain samples (Figure 3). The PCA profile demonstrates that a specific normal retinal gene expression profile differs from the expression profiles of the frontal, occipital,

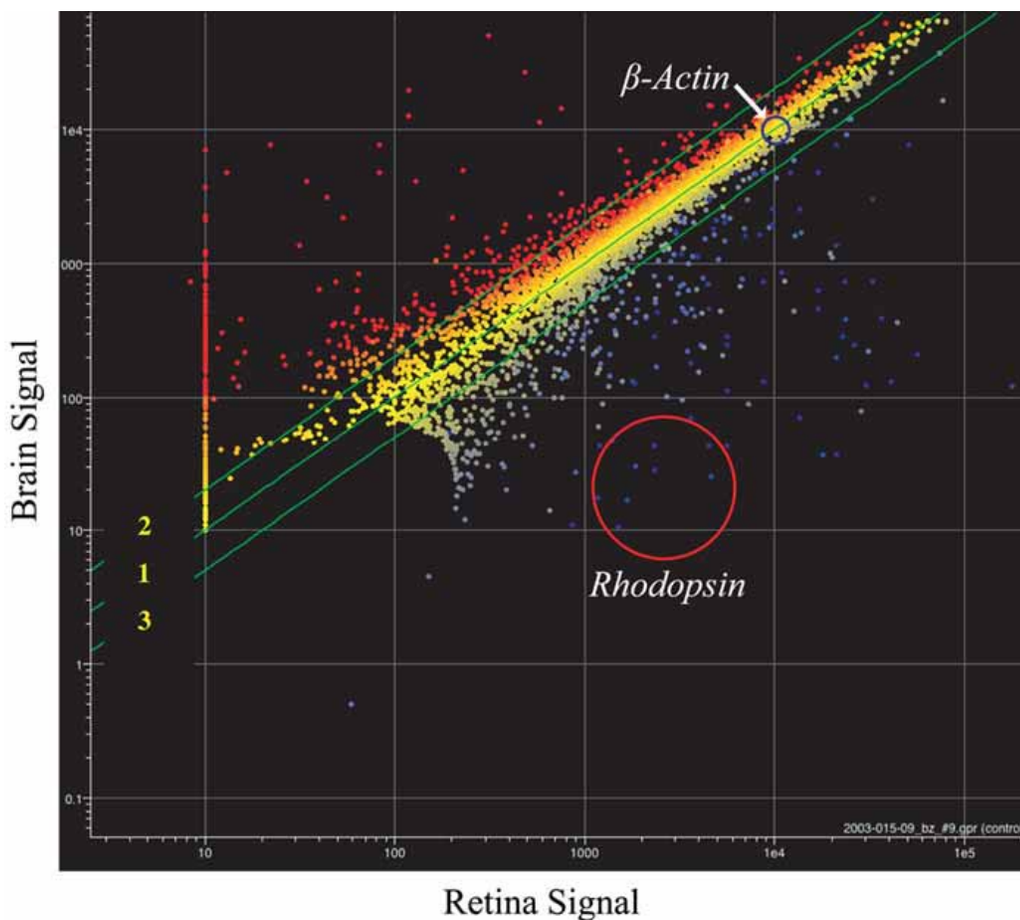


Figure 1. MA-plot comparing gene expression in retina and brain reference pool. Each point represents a single transcript/spot on the array, plotted as a function of its signal amplitude (expression level) for brain pool and retina. Line 1 denotes equivalent expression in retina and brain, whereas lines 2 and 3 indicate a two-fold difference in the signal strength. For illustration, spots representing *Rhodopsin*, much more abundant in retina than brain, are located within the red circle, while the uniformly expressed *β-Actin* spots lie within the blue circle.

TABLE 1. FUNCTIONAL ANNOTATION OF THE TOP 30 RETINA-ENRICHED GENES

Clone identifier	GenBank access number	Dog chromosome location	Gene symbol	Gene ID number	Human chromosome	Unigene	Retina/brain fold change	Function
DR010009B10H06	DT538190	25:47,884,218-47,884,349	SAG	6295	2q37.1	Hs.308 Cfa.7077	37	Rhodopsin-mediated signaling
DR010020A20B07	DT540322	14:21,958,280-21,958,543	GNAT1	2779	7q12.3	Hs.648519 Cfa.3191	36.5	Transport, visual system
DR010017A21C10	DT539700	7:22,113,973-22,114,385	PDC	5132	1q31.1	Hs.580 Cfa.1202	36	Signal, visual system
DR010018A20D11	DT539956	15:54,570,842-54,571,430	SFRP2	6423	4q31.3	Hs.481022	31.5	Unknown
DR010017A21D08	DT539707	20:8,475,328-8,475,483	RHO	6010	3q21-q24	Hs.247565 Cfa.7396	27.5	Signal, visual system
DR010024A10D06	DT541291	11:9,365,437-9,365,294				Hs.150406	26	EST
DR010013B10A12	DT538991	3:94,958,572-9,958,849	PDE6B	399653	10p14	Hs.59872 Cfa.3793	25	Signal, visual system
DR010015B10D11	DT539417	18:64,384,884-64,385,136	ROM1	6094	11q12.3	Hs.281564	24.5	Disk morphogenesis
DR010030A20H09	DT543034	4:35,677,194-35,677,735				10q23.1	23	EST
DR010008B20F12	DT538002	20:53,894,125-53,894,570	RDH8	50700	19p13.2	Hs.272405	21	Transport, visual system
DR010023B10C07	DT541135	17:37,457,058-37,457,430	DUSP2	1844	2q11.2	Hs.1183	20.5	Nucleic acid metabolism
DR010022A20E08	DT540832	12:51,422,323-51,422,562				Hs.520287	16	EST
DR010018B20H08	DT540046	2:87,515,835-87,515,994				1p36.22	11	EST
DR010008B20G12	DT538006	7:69,939,004-69,939,305	CLUL1	27098	18p11.32	Hs.274959 Cfa.3516	9.30	Unknown
DR010028B20H07	DT542573	27:4,960,724-4,961,087	NEUROD4	58158	12q13.2	Hs.131010	9.5	Metabolism
DR010028B10E10	DT542509	17:51,086,234-51,086,428	NLR	4901	14q11.1-q11.2	Hs.89606	9	DNA, visual system
DR010029A10E09	DT542612	6:19,524,381-19,524,521	KCTD13	253980	16p11.2	Hs.534590	9	DNA metabolism
DR010009B10A10	DT538119	28:32,984,199-32,984,621	INPP5F	22876	10q26.12	Hs.369755	6.5	Metabolism
DR010022B20B07	DT540940	8:35,335,208-35,335,594	OTX2	5015	14q21-q22	Hs.288655	6	Metabolism
DR010021B20E07	DT540724	20:38,082,009-38,082,300	LRTM1	57408	3p14.3	Hs.554867	6	Sensory organ development
DR010019B20C09	DT540148	14:58,294,157-58,294,685				7q31.2	5	EST
DR010014B20E10	DT539271	5:37,315,043-37,315,421				6p21.1	5	EST
DR010011A20D01	DT538558	18:9,132,499-9,132,941	ECOP	81552	7p11.2	Hs.488307	4.5	Signal transducer activity
DR010005B10H02	DT537179	5:59,339,081-59,339,539	GNB1	2782	1p36.33	Hs.430425	4.50	Signal transducer activity
DR010016B20C07	DT539561	2:74,241,431-74,241,808	HMG-17	403686	13p25.1	Hs.477784 Cfa.3549	4	Nucleic acid
DR010024B10C12	DT541427	22:7,990,538-7,990,876	HMGN2	3151	1p36.11	Hs.181163	4	Nucleic acid
DR010017B10E11	DT539798	23:35,612,629-35,613,006				11q12.3	4	EST
DR010027B20E09	DT542265	12:67,236,366-67,236,647	GLTP	51228	12q24.11	Hs.381256	4	Transport
DR010006B20G02	DT537497	2:78,953,814-78,954,066	DDOST	1650	1p36.12	Cfa.3884	3.5	Metabolism
DR010029B10D10	DT542720	26:11,674,727-11,675,099	PPP1CC	5501	12q24.11	Hs.79081 Cfa.451	3.5	Metabolism
DR010007A10G09	DT537586	33:26,792,981-26,793,197				3q13.33	3.5	EST

The top 30 genes with preferential retinal expression in a retina-brain pool comparisons are listed, with the corresponding identifiers, chromosomal location, expression ratio, and function. Hs represents *Homo sapiens*, Cfa represents *Canis familiaris*.

and temporal cortices of the brain, while these clusters are closely related. Within groups, experiments showed little variation, and the retinal profiles did not significantly differ with regard to litter or sex of the tested individuals. The brain lobe regions showed tissue-specific expression patterns, but one temporal sample was clearly separated. The result was confirmed by independent replication of the experiment and could represent an individual variation in this particular sample or an experimental artifact.

To confirm the microarray results, we selected five genes for northern analysis based on their microarray expression profiles. Two showed preferential expression in retina (*NEUROD4*, DR010030A20H09), two in brain (*SYN2*, *SPP1*),

and one was equally expressed in retina and brain (DR010024A20G01). Blots of total RNA from retina, brain, and spleen were hybridized with cDNA probes for these five genes (Appendix 1). We found that the northern blots showed the same expression patterns predicted by the microarray data with no detectable signal in spleen for any of the five genes (Figure 4A). Semiquantitative determination of expression in the northern blots correlated closely with the signal pattern obtained from the microarray experiments (Figure 4B).

**DISCUSSION**

We have undertaken the first step in the evaluation of the normal canine retinal expression profile to create a system that

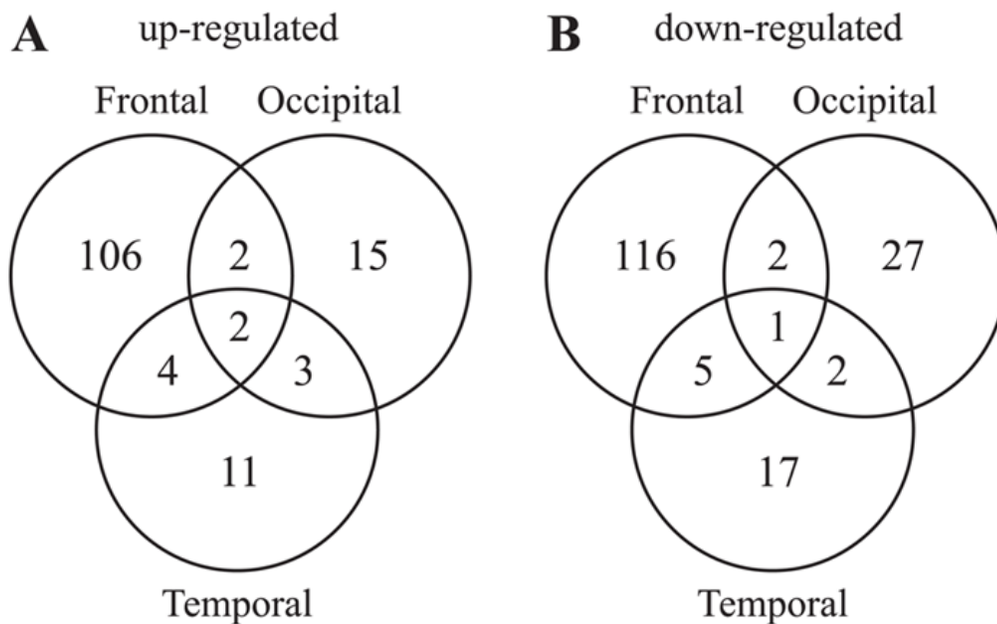


Figure 2. Region-dependent expression pattern for frontal, occipital, and temporal brain cortices. The diagrams show the number of genes up- (A) and downregulated (B) in the respective regions.

**TABLE 2. FUNCTIONAL ANNOTATION OF THE 30 TOP BRAIN ENRICHED GENES**

Clone identifier	GenBank access number	Dog chromosome location	Gene symbol	Gene ID number	Human chromosome	Unigene	Retina/brain fold change	Function
DR010023A20H05	DT541102	X:80,243,963-80,244,545	PLP1	5354	Xq22.2	Hs.1787 Cfa.3274	-25	Myelination
DR010027A10F10	DT542045	4:14,326,600-14,327,069	SPP1	6696	4q22.1	Hs.313	-20	Phosphoprotein
DR010005A20E04	DT537067	32:13,862,656-13,862,983	SPARCL1	8404	4q22.1	Hs.1424	-17	Extracellular matrix protein
DR010008A20A02	DT537874	20:40,129,412-40,129,709	NISCH	11188	3p21.1	Hs.435290	-16	Cytoskeleton and cellular components
DR010010B10C05	DT538398	6:62,422,039-62,422,316	LMO4	8543	1p22.3	Hs.436792	-11	Transcriptional regulator
DP010005000E06	DT536078	20:9,218,236-9,413,008	SYN2	6854	3p25	Hs.445503	-9	Neuronal phosphoproteins
DR010027E20A04	DT542218	16:55,443,443-55,443,919	GPM6A	2823	4q34.2	Hs.75819	-8	Myelin proteolipid
DR010014B20F03	DT539277	13:30,167,584-30,167,884	DDEF1	50807	8q24.21	Hs.106015 Cfa.10191	-6.5	Signal transduction
DR010015A20E10	DT539376	35:9,024,760-9,025,290	NRN1	51299	6p25.1	Hs.103291	-6.5	Neural activity
DR010019A20H03	DT540130	24:15,030,049-15,043,202	SNAP25	6616	2p12.2	Hs.167317	-6.5	Synaptic activity
DR010024A10E09	DT541302	26:31,587,239-31,587,700	SEPT5	5413	22q11.21	Hs.283743	-6	Nucleic acids activity
DR010024B10F05	DT541444	4:69,817,413-69,817,666			1q43		-6	EST
DR010026A20H08	DT541827	X:57,663,647-57,664,138	GDPD2	54857	Xq13.1	Hs.433812	-5	Cell differentiation
DR010013A10A11	DT538911	13:52,260,613-52,260,773	IGFBP7	3490	4q12	Hs.479808	-5	Insulin-like growth factor
DR010022A10B01	DT540751	24:15,694,097-15,694,559	C20orf103	24141	20p12.2	Hs.22920	-4.4	Protein precursor
DR010020A10B08	DT540252	17:62,843,281-62,843,815	MLLT11	10962	1q21.2	Hs.75823	-4.2	leukemogenesis
DR010025B10A02	DT541635	X:43,640,390-43,640,656			10q23.33		-4	Unknown
DR010026B20G05	DT541971	5:39,939,020-39,939,274			17p12		-4	Unknown
DR010025A21C11	DT541582	26:3,428,348-3,428,585	PXMP2	5827	12q24.33	Hs.430299	-3.6	Nucleic acitivity
DR010015A20E09	DT539375	15:5,881,332-5,881,851	PPT1	5538	1p34.2	Hs.3873 Cfa.10158	-3.6	Metabolism
DR010016B10H12	DT539539	20:48,103,437-48,103,712			19p13.11		-3	EST
DR010005B10E03	DT537156	2:79,096,079-79,096,475	CAMK2N1	55450	1p36.12	Hs.187922	-3	Metabolism
DR010012B20D10	DT539065	24:35,260,072-35,260,236	YWHAB	7529	20p13.12	Hs.279920	-2.5	Signal transduction
DR010020A20D02	DT540337	1:98,778,726-98,779,224	DIRAS2	54769	9q22.2	Hs.165636	-2.5	Metabolism
DR010024A20H05	DT541397	10:52,558,495-52,558,873	CALM2	805	2p21	Hs.468442	-2.5	Metabolism
DR010010A20G08	DT538371	1:111,515,746-111,516,031	CALM3	808	19q13.32	Hs.515487	-2.5	Metabolism
DR010013B10A11	DT538990	4:12,791,804-12,792,189			3p21.31		-2.5	EST
DR010013A10E12	DT539022	3:32,111,666-32,111,915			15q22.2		-2.5	EST
DR010028A20D02	DT542393	16:22,581,974-22,582,397			15q14		-2.5	EST
DR010012B20E05	DT538877	11,386,498-11,386,885	ATP2A2	488	12q24.11	Hs.506759	-2.5	Metabolism

The top 30 genes with preferential brain expression in a retina-brain pool comparison are listed, with the corresponding identifiers, chromosomal location, expression ratio, and function. Hs represents *Homo sapiens*, Cfa represents *Canis familiaris*.

allows comparison within and between different models of retinal degeneration. For this purpose, a reference design was chosen, which, ideally, would utilize a pool of normal retina as reference. It is not possible, however, to provide a retina pool that is large enough for the proposed scale of experiments, and, therefore, was substituted by a brain pool. The limitation of this reference is that genes not expressed in the brain cannot be validated against this common reference, and common data analysis procedures, such as normalization, might introduce experimental errors based on the tissue-specific differences. This study has validated the procedure, and identified potential problematic genes which subsequently can be investigated in alternate approaches such as quantitative RT-PCR or macroarrays [26] that selectively target for analysis those genes not expressed in the brain pool.

Microarray data are most commonly normalized using the loess algorithm, which assumes that only a small percentage of genes is differentially expressed, and maintains balance between the numbers of up- and downregulated expressed transcripts. Alternative approaches for normalization procedures propose the use of ANOVA models [27]. This method essentially performs only a global normalization, and does not correct for intensity or scale differences. Yang et al. [24] have found that the standard global median normalization can often be inadequate due to spatial and intensity dependent dye biases, and proposed loess subgrid normalization. We have applied different models to our data (Appendix 1), which further support these findings and suggests that we do not introduce experimental bias in the data by applying the loess subgrid normalization.

As an initial step in evaluating the microarray, we examined gene expression pattern differences between normal retina and the brain pool tissue reference sample. These comparisons allowed us to identify 405 out of 3,070 valid transcripts (13.2%) as preferentially expressed in retina (Table 1), and 452 genes with low abundance in the retina (Table 2). As expected, known photoreceptor genes were highly expressed in retina, demonstrating the specificity of the hybridization. Over a third of the differentially expressed clones represent potentially novel genes that are enriched or even specific to retina or brain [28]. Among the differential expressed genes were many important to neural activity, such as ion channels (transport class) and cytoskeletal proteins (structural class). Genes involved in more general processes, such as energy generation, also showed retina or brain preferred expression. Compared to the total number of genes with preferential expression, we found higher frequencies of proliferation and cell death genes expressed in the brain than retina, and nucleic acid processing genes in retina than brain. Among the transcripts preferentially expressed in the brain were many known and unknown genes not previously reported to have a preferential expression pattern in different regions of the brain [5,29]. We demonstrated also that expression patterns are quite restricted to individual cortex areas, as few differential expressed genes showed overlap between the examined brain regions (Figure 2).

It is clear that, despite the common embryonic origin of retina and brain tissues, the corresponding expression profiles differ. Even though a subset of the genes expressed in each of the three brain regions were unique to the tissue of origin, this

**TABLE 3. GENES UPREGULATED AND EXPRESSED IN TWO OR THREE BRAIN CORTICES**

Clone identifier	GenBank access number	Dog chromosome location	Gene symbol	Gene ID number	Human chromosome	Unigene	Overlapping regions	Function
DR010026A10E12	DT541032	1:75,209,657-75,209,810	Clorf63	57035	1p36.11	Hs.259412 Cfa.10157	F-O	Open reading frame
DR010027A10B02	DT541999	9:7,065,312-7,065,859			17q24.2		F-O	EST
DR010023B10A04	DT541111	22:64,149,260-64,149,794	UPF3A	65110	13q34	Hs.533855	F-T	Metabolism
DR010019B10C12	DT540150	12:7,009,358-7,009,914			6p21.31		F-T	EST
DR010017A10B01	DT539606	8:50,712,025-50,712,500	KIAA0317	9870	14q24.3	Hs.497417	F-T	Unknown
DR010018B10B12	DT539606	27:40,457,438-40,457,926	NECAP1	25977	12p13.31	Hs.555927	F-T	Involved in endocytosis
DR010023B10B01	DT541120	18:59,559,359-59,559,613			7q21.3		O-T	EST
DR010012B10B11	DT541128	10:14,366,170-14,366,540	CPSF6	11052	12q15	Hs.369606	O-T	Nucleic acids
DR010010B20H09	DT538497	7:12,476,293-12,476,485			14q23.1		O-T	EST
DR010012A10F04	DT538700	9:11,033,436-11,033,709			11q13.1		F-O-T	EST
DR010009B20B10	DT538214	23:47,202,203-47,202,727	TMSF1	4071	3q25.1	Hs.351316	F-O-T	Metabolism

Hs represents *Homo sapiens*, Cfa represents *Canis familiaris*, F represents Frontal, O represents Occipital, T represents Temporal.

**TABLE 4. GENES DOWNREGULATED AND EXPRESSED IN TWO OR THREE BRAIN CORTICES**

Clone identifier	Genbank access	Dog chromosome location	Gene symbol	Gene ID number	Human chromosome	Unigene	Overlapping regions	Function
DR010006A10H12	DT537289	32:35,442,842-35,442,964			4q25		F-O	EST
DR010027B10G05	DT542201	X:62,967,726-62,968,008	ATRX	546	Xq21.2	Hs.533523 Cfa.2632	F-O	Transcriptional regulator factor
DR010014A10D04	DT539133	24:28,803,524-28,803,789	RPN2	6185	20q11.23	Hs.370895	F-T	Metabolism
DR010009B10C03	DT538133	2:5,292,568-5,293,107			5q14.3		F-T	EST
DR010014B10D06	DT539212	2:81,584,176-81,584,587	RCC2	55920	1p36.13	Hs.380857	F-T	Cell division
DR010022A20D11	DT540746	24:28,877,230-28,877,628	MANBAL	63905	20q11.20	Hs.6126	F-T	Metabolism
DR010025A10A04	DT541537	25:22,008,506-22,008,977			12q13.11		F-T	EST
DR010012B10A09	DT538775	12:67,729,356-67,729,566	FLJ10159	55084	6q21	Hs.445244	F-O-T	Unknown
DR010007B20E10	DT537769	14:61,667,805-61,668,179	ING3	54556	7q31.33	Hs.489811	O-T	DNA binding
DR010012B10B11	DT538789	10:14,366,170-14,366,540	CPSF6	11052	12q15	Hs.369606	O-T	Nucleic acid binding

Hs represents *Homo sapiens*; Cfa represents *Canis familiaris*; F represents frontal; T represents temporal; O represents occipital.

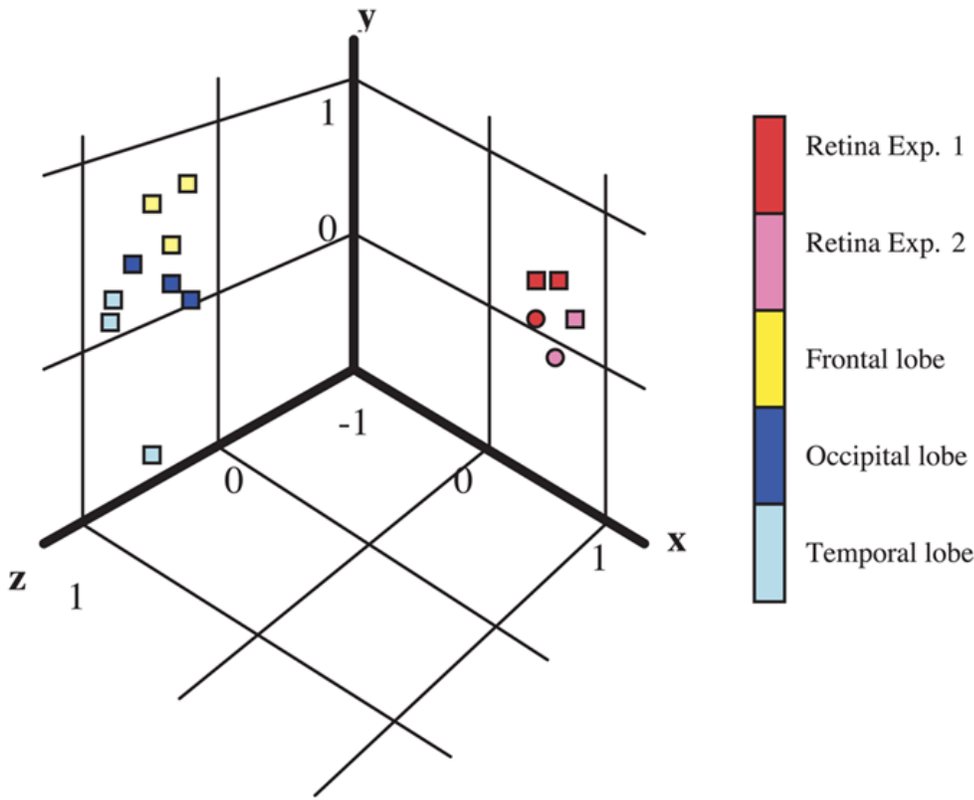


Figure 3. Principal component analysis illustrates differentiation of retinal gene expression profile from the three brain regions. No variance is found between individuals from different litters (experiment 1 [Exp.1] versus experiment 2 [Exp. 2]) or between males (squares) and females (circles).

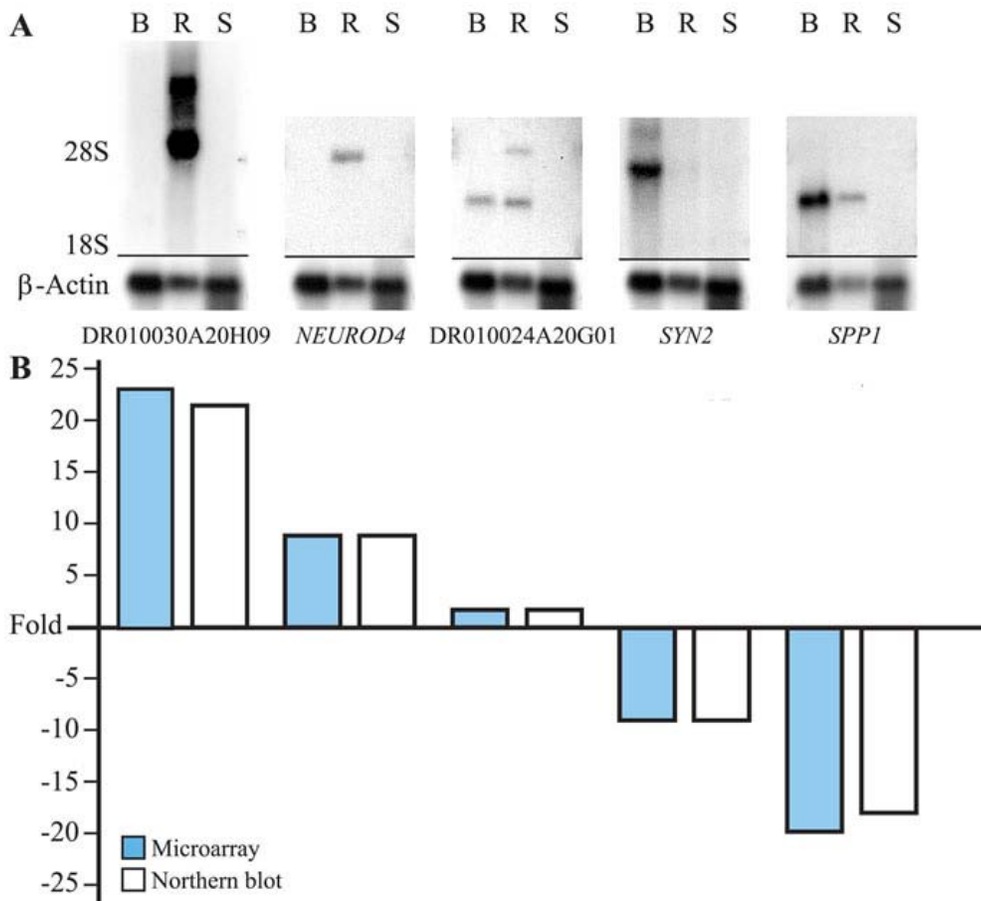


Figure 4. Northern blot validation of selected genes. **A**: Expression profile for selected genes was obtained from northern analysis using brain (B), retina (R), and spleen (S) through normalization against the canine  $\beta$ -actin loading control, and compared to microarray results (**B**). *NEUROD4*: neurogenic differentiation 4; *SYN2*: synapsin II; *SPP1*: secreted phosphoprotein 1.

was proportionately a small part of the total number of genes examined. Overall, the gene expression profiles of the three brain regions clustered closely, when compared to the different profile obtained from retina (Figure 3). The PCA also illustrates strong similarities in each of the replicates for a given tissue, showing the sensitivity and robustness of the custom cDNA microarray. Particularly for the retina, we did not observe significant differences in expression profiles between 16 week old dogs based on gender or litter as has previously been suggested from human studies [14].

Results of the microarray work were validated on northern analysis for individual transcripts (Figure 4) and confirmed to reflect the physiology of the analyzed tissues. For example, we found that *NEUROD4* had an increased expression level in retina compared to brain according with its function in the developing mammalian retina [30]. *CDKN1B* was equally expressed in normal retina and brain tissues; this gene encodes a protein that binds to and prevents the activation of cyclin E-CDK2 or cyclin D-CDK4 complexes, and thus controls the cell cycle progression at G1. The microarray results also identified ESTs representing potential novel genes that showed high preferential levels of expression in retina (e.g., DR010020A10D06, DR10030A20H09; Table 1). These novel genes of still unknown function are currently under further investigation as they may play critical roles in retinal development or maintenance, and are potential candidate genes for retinal disorders.

Recent expression studies conducted for human retina utilized the comparisons to different tissues, such as brain, liver, or kidney across SAGE, cDNA library, and microarray platforms [31,32]. In each case, the results yielding a few hundred retina-enriched genes can be classified into three different types: 1-genes already known to be retina enriched, such as guanine nucleotide binding protein (*GNAT1*) and arrestin (*ARR3*); 2-characterized genes previously not known to be retina enriched, such as WNT inhibitory factor 1 (*WIFI1*) and frizzled-related protein (*FRZB*), and 3-unknown genes, mostly EST clusters. Our data are not just comparable to those studies in terms of identified classes and the ratio of genes specific to the retina, but provide additional information for potential novel genes integrated in the original EST database that allows the subsequent characterization of these transcripts [16].

It has been suggested that genes involved in developmental regulation of the retina are among the top candidates for degenerative processes when mutated. Recently, a microarray profile between wild type and *Nrl*<sup>-/-</sup> mouse retinas produced a list of 1,000 genes with potential high influence in rod development [33]. Among those were several genes associated with human retinopathies (e.g., *GNAT1*, *RHO*, *SAG*, *ROM1*, *PDE6B*) that have also been identified through our analysis (Table 1, Appendix 1). Furthermore, this work presents comparable percentage of genes involved in metabolic pathways and unknown genes, suggesting that our microarray results are a suitable model system to address retina-specific biological questions. The results can now be extended to examine the gene expression profiles of different canine retinal degenera-

tion models aiming to differentiate those pathways that are unique and mutation-specific from those that are mutation-independent and common to several of the retinal disorders. This approach will provide new insights into the molecular mechanisms of these diseases, and help identify key molecules and pathways for therapeutic intervention.

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