The genetics of inherited retinal disorders in dogs: implications for diagnosis and management

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Abstract: Dogs are affected by many hereditary diseases just as humans are. One group of these diseases comprises of retinal disorders, which are a growing problem in canine breeding. These disorders are heterogeneous, with diverse causative mutations and modes of inheritance. Some affect only one breed, while others may affect many breeds; some breeds are affected by only one disease, while others can be affected by two or more. Dog breeders should take into account the presence of any deleterious alleles when choosing parents for the next generation.

Keywords: hereditary, retinal, disease, dog

Introduction

As of August 2015, a total of 653 disorders/traits were found in dogs, of which 263 were Mendelian traits, 193 Mendelian traits with a known key mutation, and 357 potentially usable models for human disease. Of these, 39 diseases affect the retina; some have a known genetic basis, while others are not yet known.

Canine Leber Congenital Amaurosis

Canine Leber congenital amaurosis (CLCA; formerly known as congenital stationary night blindness or CSNB) is one of the retinal diseases that recorded the largest shift. From finding the causative mutation through the introduction of genetic testing to gene therapy, very encouraging results were noted.

CLCA is a congenital disease (night blindness) with various degrees of visual impairment under photopic illumination (vision of affected dogs ranges from normal day vision to profound day blindness). The only breed affected by this disease is the Briard. Signs of disease occur early in the life of the affected dog (≈5–6 weeks). The causative mutation – deletion of four bases (AAGA) – of this disease was found in the \textit{RPE65} (retinal pigment epithelium-specific 65 kDa protein) gene. The result of the deletion is a frame shift and this leads to a premature stop codon. Some experimental replacement therapies to restore retinal function have been conducted with very encouraging results. Subretinal injections of recombinant adeno-associated virus (rAAV) were performed in affected dogs and improvements in visual behavior and electroretinographic (ERG) responses were observed. Visual improvement was long term; some improvement was found in younger and older dogs (≈30 months of age). In another study, a dog treated at the age of 30 months did not recover vision or retinal function. This suggests that there might be a preferred age for successful treatment with gene replacement therapy. The next question was whether visual
pathways leading from the defective retina to the visual cortex are intact. Therefore, RPE65-mutant dogs were studied using functional magnetic resonance imaging, and it was found that these dogs have markedly diminished retinal and subcortical responses to light.11 After treatment with rAAV, there was rapid and long-term restoration of cortical response. Human patients with RPE65 mutation (RPE65-LCA) have a preserved visual pathway and detectable cortical activation. Results from animal studies support the potential for human gene therapy to restore vision in congenitally blind patients with genetic retinal disease.11

**Central stationary night blindness**
True naturally occurring autosomal recessive canine central stationary night blindness (CSNB) was identified in Beagle dogs in Japan.12 Affected dogs had normal daylight vision and normal retinas but absent night vision, and they showed no detectable rod responses. The phenotype of this canine blindness is similar to the human Schubert-Bornschein form of complete CSNB. Most known candidate CSNB genes were excluded as being causative for this new form of canine blindness. The authors found reduced expression of three genes (GNAT1, CACNA2D4, and NYX) in both carriers and affected dogs; moreover, one gene (CACNA1F) was down-regulated only in the affected dogs. Ongoing studies aim to clarify the exact mechanism of night blindness, identify the causative gene, and understand the neuronal plasticity and retinal remodeling that occurs in the disease.12

**Progressive retinal atrophy**
Progressive retinal atrophy (PRA) is a group of canine retinal degenerations. This group is very heterogeneous because some of these diseases occur early while others occur late; some are recessive, others dominant, and others X-linked.13 But all these diseases show phenotypic similarities to retinitis pigmentosa (RP), a common cause of inherited blindness in humans.14 PRAs are rod–cone dystrophies wherein rod photoreceptors (PRs) are affected earlier than cone photoreceptors. The earliest clinical sign of the disease is night blindness.15

**Rod–cone dysplasia 1 (rcd1)**
The first retinal atrophy with mutation found was rod–cone dysplasia 1 (rcd1) in Irish Setter dogs. Rcd1 is a recessive trait in which PRs are rapidly lost. Farber et al16 found that affected dogs of this breed have elevated levels of retinal cyclic guanosine monophosphate (cGMP) resulting from deficient rod-specific cGMP phosphodiesterase (cGMP PDE) activity. Other researchers performed a study of affected and putative heterozygote Irish Setters and found that the affected PDEβ subunit mRNA contained a nonsense mutation on codon 807 (G to A transition converting TGG to TAG), which led to a premature stop codon. Polymerase chain reaction (PCR) studies with genomic DNA confirmed that normal, heterozygous, and affected dogs carry the expected alleles.17 The mutation was confirmed in Setters in the UK, and a fast PCR-based diagnostic test was developed.18 Finding treatment that would slow progression of the disease is a major goal of research on hereditary blindness. The calcium channel blocker D-cis-diltiazem, which in naturally occurring PDE6B murine model of RP alters the abnormal cellular mechanisms resulting from a mutant gene causing retinal degeneration, failed to repair PRs in affected Irish Setters.19

**Generalized PRA in Sloughi dogs**
Generalized PRA (gPRA or rcd1a) in Sloughis has an early onset similar to that in Irish Setters and is inherited in an autosomal recessive manner. As in the Irish Setters, in Sloughis too, the causative mutation for hereditary blindness was found in the PDE6B gene – an eight base pair (bp) insertion (TGAAGTCC) after codon 816, which causes a premature stop codon and truncates the PDE6B protein by 40 residues.20

**Rod–cone dysplasia type 2**
Rod–cone dysplasia type 2 (rcd2) is an early-onset autosomal recessive form of PRA and is phenotypically similar to LCA, an early-onset form of RP. This disease segregates naturally in the Collie breed of dogs. Kukekova et al15 mapped rcd2 to CFA7, an orthologue of human 1q32. Canine sequences corresponded to those of human CRB1 gene; a polymorphic microsatellite was identified in intron 5 of CRB1 and was used as a marker to test for cosegregation between CRB1 and rcd2, but based on linkage and RH (radiation hybrid mapping) data, this gene was excluded as a candidate for rcd2.15 The canine gene (named RD3) homologous to C1orf36 was identified, and a 22 bp insertion that causes a frame shift and continues the open reading frame beyond the normal stop codon was found.21

**PRA in the Cardigan Welsh Corgi (rcd3)**
Rcd3 in the Cardigan Welsh Corgi is inherited in an autosomal recessive manner and has early onset leading to blindness in young adult dogs similar to that from rcd1. After the earlier-mentioned causative transition in the PDE6B gene, the causative mutation (second in the dog) – the deletion
of 1 bp in codon 616 of PDE6A — was found in a Cardigan Welsh Corgi. The test for screening of the mentioned polymorphism was developed — the single-strand conformation polymorphism (SSCP) analysis. An allele-specific PCR-based test for rapid and inexpensive detection of the mutation was developed. The mutation in the PDE6A gene was excluded as causative for the gPRA in the following breeds: Chesapeake Bay Retriever, Entlebucher Sennenhund, Labrador Retriever, Tibetan Mastiff, Long- (LHD) and Wire-haired Dachshund (WHD), Tibetan Terrier, Miniature Poodle, Australian Cattle Dog, Cocker Spaniel, Saarloos/Wolfshund, Sloughi, and Collie. To avoid potential mistakes in genetic testing, in Welsh Corgis, the simple mismatch-PCR—restriction enzyme digestion test has been developed. The presence of digestion sites in the PCR product in both normal and mutant alleles provides a positive control for the activity of the restriction enzyme (Hinf1).

PRA in Gordon and Irish Setter breeds — rod–cone degeneration 4 (rcd4)
Late-onset PRA or rod–cone degeneration 4 (rcd4) was found in two Setter breeds. A frame-shift mutation was identified in the novel PRA locus, C2orf71. Because ∼10% of PRA cases in the Gordon Setter breed are not caused by the mentioned mutation, it seems that at least one other mutation causes PRA in this breed.

Generalized PRA in Schapendoes dog
In the Schapendoes dog, gPRA is characterized by initially affected rod PR vision causing night blindness, which is followed by progressive loss of cone PRs that leads to total blindness. The disease haplotype was mapped to CFA20 and, even though the causative mutation was not found, an indirect DNA test was developed for gPRA testing in Schapendoes dogs based on linkage analysis data. The causative mutation in the newly identified CCDC66 gene was found: a 1 bp insertion leads to a premature stop codon. The Schapendoes dogs and true null mouse provide valuable animal models to explore the precise role of the mentioned protein in retinal degeneration and the future development of a treatment for hereditary blindness.

PR dysplasia
In Miniature Schnauzers, the form of PRA known as PR dysplasia (pd) is an early-onset retinal disease that affects both rod and cone PRs. It is transmitted in an autosomal recessive manner. In pd-affected dogs, a missense mutation was found (codon 82, CGA to GGA) in the phosducin (PDC) gene. However, some pd-affected dogs were heterozygotes or homozygotes of the wild-type allele, and this indicates that another mutation may be present in this breed.

Early retinal degeneration
Early retinal degeneration (erd) was identified in Norwegian Elkhound dogs as an early-onset autosomal recessive disease corresponding to human LCA. In the STK38L/NDR2 (serine/threonine kinase 38-like protein – a protein kinase in the nuclear Dbf2-related family) gene, two mutations were found. One (a 4 bp deletion in exon 3) was, however, excluded as causative for erd. The second, the insertion of a short interspersed nuclear element (SINE) in exon 4, results in the absence of exon 4 from the mature STK38L mRNA, which causes dysfunction of the protein. This was the first time that the STK38L/NDR2 pathway was implicated in PR development and/or disease. This naturally occurring erd animal model will provide new insights into PR development and role of STK38L protein in other, especially neuronal, tissues.

PRA 1 in Golden Retriever (PRA1)
In the Golden Retriever breed, clinical signs of PRA were found, but one variant — progressive rod–cone degeneration (prcd) — only accounts for a small proportion of PRA cases. A novel PRA locus was identified in CFA37. In a solute carrier anion exchanger (SLC4A3) gene, a frame-shift mutation was identified. Because a large proportion of cases (∼44%) remains unexplained, it seems that three or more mutations cause PRA in the Golden Retriever breed.

PRA type 3 in Tibetan Spaniel and Tibetan Terriers (PRA3)
Most PRA cases in Tibetan Spaniels are clinically indistinguishable from other forms of PRA. The mode of inheritance seems to be autosomal recessive. The age of diagnosis is relatively late, at ∼5 years of age. Insertion of a SINE was identified in FAM161A gene, a ciliary gene associated with RP in humans. The mutation causes exon skipping and a shift in the reading frame, which results in a premature stop codon. The insertion is not present in all affected Tibetan Spaniels and Tibetan Terriers; this suggests that PRA in these breeds is genetically heterogeneous.

PRA in Papillon and Phalène dogs
Primary clinical signs that were found in Papillon and Phalène dogs include visual impairment in dim light. The causative mutation known to cause PRA was not found in
the mentioned breed (or breeds in Europe). Genome-wide association and linkage studies revealed a novel PRA locus in CFA2 in which the CNGBI (cyclic nucleotide gated channel beta 1) gene was found. In this gene, an insertion–deletion (indel) mutation (1 bp deletion followed by 6 bp insertion, causing a frame shift and hence premature stop codon) was found, and this indel was confirmed as causative for PRA in Papillon and Phalène dog breeds. The PRA and the causative mutation in the CNGBI gene in the Papillon breed was confirmed by a second group of authors.

**PRA in Basenji**

In the Basenji breed, the adult form of PRA was observed. First, night blindness presented, which progressed to total blindness. Many affected Basenjis retain adequate daylight vision for many years, sometimes for their entire life. This phenotype is similar to prcd disease, but the prcd mutation was excluded as causative in the Basenji breed. In the S-antigen (SAG) gene transition mutation (T-C), which changes a normal stop codon to a code for the amino acid arginine, which would result in a deduced addition of 25 amino acids, was identified.

**Progressive rod–cone degeneration**

Prcd is a phenotypically degenerative disorder in which, after normal postnatal development, rods and cones degenerate both structurally and functionally; for this reason, it is classified as a late-onset disorder. This retinal degeneration is inherited as an autosomal recessive trait and was originally characterized in a Miniature Poodle. The prcd locus was mapped to the centromeric end of canine chromosome 9 (CFA9); this region shows synteny conservation with HSA17qter – a region in which a form of human RP17 was mapped. On the basis of clinical similarities of human and canine diseases and chromosomal localization, prcd has been proposed as a locus homologue of RP17 in human. The prcd mutation for retinitis pigmentosa 3 (RP3) was found. One of these mutations, a five-nucleotide deletion (GAGAA), was found in the Siberian Husky dog. The same mutation (G4R), which replaces threonine with arginine (T4R), was confirmed as causal for dominant PRA in the English Mastiff. The same substitution was found in one affected Bull Mastiff, but another affected dog did not share this mutation; this suggests that in the Bull Mastiff there must be at least one other causative mutation that causes PRA.

**Dominant PRA**

PRA in the English Mastiff breed revealed a disease with ambiguous mode of inheritance. Examination of pedigrees showed that the majority of affected individuals had an affected parent, which indicates dominance. To confirm the dominant mode of inheritance, controlled outcross matings were performed. The appearance of affected offspring from the mating indicates the presence of a dominant allele. Analysis of a candidate gene, rhodopsin (RHO), identified two synonymous and one nonsynonymous substitution. The nonsynonymous substitution (C–G at position 11), which replaces threonine with arginine (T4R), was confirmed as causal for dominant PRA in the English Mastiff. The same substitution was found in one affected Bull Mastiff, but another affected dog did not share this mutation; this suggests that in the Bull Mastiff there must be at least one other causative mutation that causes PRA.

**X-linked PRA**

X-linked PRA (XLPRA) represents a spontaneous animal model of X-linked retinal degeneration in humans. It was first found in Siberian Husky dogs. In hemizygous males, the outer segments of rods are affected initially; then rod outer segments almost completely disappear and cone outer segment degeneration becomes apparent. Affected males begin to show clinical signs at the age of sexual maturity or young adulthood; female carriers demonstrate the rod specificity of disease. The disease locus was mapped to the short arm of the X–chromosome, which is homologous to human Xp21 in which the mutation for retinitis pigmentosa 3 (RP3) was found. In canids, three microdeletions in ORF15 (RPGR exon ORF15 (RP GTPase regulator exon open reading frame 15) were found. One of these mutations, a five-nucleotide deletion (GAGAA), was found in the Siberian Husky dog. The disease in this breed was renamed XLPRA1 to distinguish it from a second disease (XLPRA2) in mongrel dogs. The XLPRA1 mutation causes a frame shift and immediate premature stop; the same mutation was found in the Samoyed breed with a clinically similar X-linked disease. In XLPRA2, a two-nucleotide deletion (GA) was found, which results in
a frame shift that significantly changes the deduced peptide sequence. The XLPRA2 causes a very severe phenotype. The third mutation (a three-nucleotide deletion – eliminates a glutamic acid) in red wolves does not alter the remainder of the protein and does not cause disease.\textsuperscript{50} Because gene-based therapy has limitations (eg, it requires identification of the mutated gene), the use of neuroprotective agents that can rescue PRs regardless of the genetic and/or environmental causes was developed. In several rodent and large animal models, ciliary neurotrophic factor (CNTF) was successfully used for rescue of PRs. In a group of 16 affected XLPRA2 dogs, CNTF treatment was used but no significant neuroprotective effect in the XLPRA2 retina was found. On the other hand, in the rcd1 control group, a statistically significant rescue of PRs with CNTF was observed. This suggests that one agent cannot treat all retinal degenerations but can be successfully used in some of these deleterious diseases.\textsuperscript{51} Gene therapy based on AAV 2/5 vector-mediated transfer was performed, with some positive results in XLPRA1- and XLPRA2-affected dogs. The full-length human RPGRorf15 complementary DNA driven by the human G-protein-coupled receptor protein kinase 1 (hGRK1) was used. In XLPRA1, treatment was initiated before PR loss (28 weeks of age); in XLPRA2, the injections were performed at 5 weeks of age. In XLPRA1, treatment before disease onset prevented disease development; in XLPRA2, treatment arrested disease progression and the morphology of the remaining PRs was restored to normal.\textsuperscript{52}

In working Border Collies in France, PRA with X-linked mode of transmission was found. The exclusion of mutations in the RPGR gene, which causes XLPRA1 and XLPRA2, suggests that this PRA in Border Collies might correspond to a new form of X-linked PRA. This new form was named XLPRA3.\textsuperscript{53}

Cone–rod dystrophies
In cone–rod dystrophies, the cone system is more affected; severe loss of central vision, color vision, and photophobia are seen in affected individuals. Onset may occur in childhood or early adolescence.\textsuperscript{54}

Cone–rod dystrophy 1 (crd1 or crd4)
Cone–rod dystrophy 1 (crd1 or crd4) in Miniature Long-haired Dachshunds (MLHDs) was primarily described as rod–cone dystrophy. First ophthalmoscopically observable signs were found in affected puppies at the age of 25 weeks (bilateral patchy and mild hyperreflectivity in tapetal fundus). Over the next few months, the intensity of hyperreflectivity increased and was accompanied by attenuation of the retinal blood vessels.\textsuperscript{54} A 44 bp insertion (A\textsubscript{29}GGAAGCAACAG-GATG) in RP GTPase regulator-interacting protein 1 (RPGRIP1) gene was found in affected MLHDs from the research group; and 5.3% (two individuals) of dogs from the pet population were homozygous for the mutation but without clinical signs of disease.\textsuperscript{55}To confirm or exclude the insertion as causal for crd1, dogs from a research colony expanded from the original study were used (recessive homozygotes with insertion, as well as heterozygotes) and clinical findings, ERG, objective vision testing, histology, and other tests were performed; the 44 bp insertion was consequently excluded from direct causal association with crd1.\textsuperscript{56} Then, an additional single locus on canine chromosome 15 was found in which the majority of early-onset cases were homozygous for a 1.49 Mb interval. It seems that homozygosity at both loci must be present for early-onset retinal degeneration to develop.\textsuperscript{57}

Cone–rod dystrophy in Standard Wire-haired Dachshund
First behavioral changes in affected puppies of Standard Wire-haired Dachshund (SWHD) breed were seen at the age of 6 weeks, but age of onset and progression of retinal degeneration showed marked diversity. Initial onset of changes was observed between 10 months and 3 years, and changes were always bilateral and symmetrical. A complete retinal atrophy was evident at the age of 5–6 years. Analysis of the ERG recordings showed severely reduced cone single-flash a- and b-wave amplitudes.\textsuperscript{58} The mode of inheritance of cone–rod dystrophy (crd) in the SWHD is autosomal recessive. The causative mutation, deletion of 180 bp, was found in the nephroptosis 4 (NPHP4, also known as nephroretinin) gene. The affected protein lacks the domain interacting with RPGRIP1 in retina.\textsuperscript{59}

Cone–rod dystrophy 1 and 2 (crd1 and crd2)
In American Staffordshire Terrier dogs and American Pit Bull Terrier dogs, two early-onset autosomal recessive retinal degenerations were identified.\textsuperscript{60} In both breeds, the disease affects young dogs (<1 year of age). Dogs are affected with severe photopic and scotopic visual impairment, which progresses to more severe blindness in early adulthood. The disorders are similar in both breeds and so were termed crd1 and crd2, respectively. For the same reason, and because the breeds share a common ancestor and are physically similar, a cross-breeding complementation test was undertaken to
prove that the two degenerations were nonallelic. This was confirmed, and in crd1-affected dogs, mutation in the PDE6B gene was found (three-base deletion in exon 21, which causes deletion of one amino acid – asparagine – at position 802 of the protein). In crd2-affected dogs, a 1 bp insertion in exon 10 of the IQCB1 (IQ motif-containing protein B1 or nephrocystin 5 – NPHe5) gene was identified, which causes a frame-shift mutation and premature stop codon.\(^\text{61}\)

### Cone–rod dystrophy 3 (crd3) or gPRA

In the Irish Glen of Imaal Terrier (GIT) breed of dogs, the disorder termed cone-rod dystrophy was identified. It is an adult-onset disease, with the first signs presenting in affected dogs at 3 years of age and progressing to end-stage retinal degeneration over several years. This disease is orthologous to human cone–rod dystrophy 9 (CORD9), which is caused by a mutation in the A Disintegrin and Metalloprotease domain, family member 9 (ADAM9) gene. In the same gene, in dogs, a large genomic deletion was found (it removes exons 15 and 16 from the ADAM9 transcript, introduces a premature stop codon, and would remove critical domains from the encoded protein).\(^\text{62}\) The deletion was confirmed in another group of affected GIT dogs with gPRA, which indicates that crd3 and gPRA in GIT is the same disease with the same causative deletion in the ADAM9 gene.\(^\text{63}\)

### Achromatopsia

Achromatopsia or rod monochromacy and day blindness is a rare autosomal hereditary disease that results in complete loss of cone PR function, while the rods remain intact. The disease is characterized by decreased visual acuity, photophobia, nyctalagus, and complete color blindness.\(^\text{64}\) Cone degeneration (cd) in Alaskan Malamute and German Shorthaired Pointer breeds is inherited in an autosomal recessive manner and is homologous to human achromatopsia. Mutations in the canine homologue of the cyclic nucleotide-gated channel β-subunit gene (CNGB3), responsible for human ACHM3 disease phenotype, were found. In Alaskan Malamute-derived dogs, the deletion that removes all exons of the mentioned gene was found. In the German Shorthaired Pointer, a missense mutation in exon 6 (D262N) within a conserved region of CNGB3 gene was found. Affected dogs from both breeds provide a natural model for the study of disease mechanisms and development of potential therapeutic methods.\(^\text{65}\) The same deletion as in the Alaskan Malamute was found in some other breeds. Two of them were Siberian Husky and Alaskan Sled dog, which are from a subgroup of distance runners and share a common ancestor, but one other, the Miniature Australian Shepherd, is not genetically related to the Alaskan Malamute; this suggests that other breeds may potentially carry the same cd allele and be affected by achromatopsia.\(^\text{66}\)

Gene replacement therapy using rAAV serotype 5 (rAAV5) was successfully used for restoration of cone function and associated photopic vision in both canine achromatopsia models. Results of this study hold promise for future clinical trials in human patients with CNGB3 achromatopsia.\(^\text{67}\)

### Canine multifocal retinopathy

Best macular dystrophy (BMD) is a human autosomal dominant disease of the retina caused by mutations in the bestrophin (BEST1) gene. BMD typically presents in childhood.\(^\text{68}\) Even though dogs lack the foveomacular region affected in BMD, lesions observed in a disease termed canine multifocal retinopathy (cmr) closely resemble the vitelliform lesions of BMD. This hereditary retinal abnormality was seen in several dog breeds. Autosomal recessive mode of inheritance was supposed after pedigree analysis and prospective matings. On the basis of similarities between BMD and cmr, the BEST1/ VMD2 (canine) gene was selected for phenotype-directed candidate gene analysis. Two mutations in the coding sequence were found. The first, which causes cmr1 in Great Pyrenees dogs, English Mastiff, and Bull Mastiff breeds, was a nonsense mutation in exon 2 at codon 25 (C – T), which replaces an arginine residue with a stop codon. The second, which causes cmr2 in the Coton de Tulear breed, was a missense mutation (G – A) in exon 5.\(^\text{69}\) In dogs from the Lapponian Herder breed, a retinal disease similar to cmr was found, but mutations for cmr1 and cmr2 were excluded as causative for this variant of the disease. The novel form of retinopathy was named cmr3, and two mutations in the coding sequence of BEST1 were identified: a deletion at nucleotide position 1,388 (results in a frame shift and introduces a premature stop codon) and a substitution at nucleotide position 1,466 (leads to change in the amino acid sequence – glycine to valine, and in combination with the first-mentioned mutation, causes an additional stop codon). Mutations were found in complete linkage disequilibrium. In dogs from the Bernese Mountain dog breed, two other potentially deleterious mutations were found, but only in the heterozygous state, and no homozygous affected dog has been identified. This indicates that there may be another variant of cmr. Causative mutation for cmr1 was identified in the new breeds Dogue de Bordeaux and Italian Cane Corso,\(^\text{70}\) Australian Shepherd,\(^\text{71}\) and Boerboel.\(^\text{72}\) The affected dogs with cmr1 and cmr2 were used in a detailed study to characterize BEST disease in this naturally occurring large animal model.\(^\text{73}\)
Retinal and skeletal dysplasia or oculoskeletal dysplasia

Retinal and skeletal dysplasia or oculoskeletal dysplasia (OSD) is a hereditary disease, which presents in the Labrador Retriever and Samoyed dog breeds. In this disease, normally organized retinal cells are disorganized and can form folds. In severe forms of disease, the retina can be unattached and other abnormalities can be seen in the eye (cataract). Depending on the severity of the disease, dogs may have reduced eyesight or can be totally blind. Skeletal problems include joint abnormalities, the breakage of pieces of affected bones, and stunted growth of bones in legs. Cross-breeding of an OSD-affected Labrador Retriever to an OSD-affected Samoyed resulted in a nondwarf progeny; this established that these two disorders are nonallelic. OSD in the Labrador Retriever breed has been termed drd1 (dwarfism with retinal dysplasia type 1) and OSD in the Samoyed breed has been termed drd2 (dwarfism with retinal dysplasia type 2). In two types of collagen, causative mutations for both forms of disease were found, with drd1 cosegregating with an insertional mutation (an insertion of a guanine in exon 1) in COL9A3 gene and drd2 cosegregating with a deletion mutation (1,267 bp deletion in the 5′-end of the gene) in COL9A2. Both mutations are predicted to truncate the respective protein product. These naturally occurring canine models can be used to further evaluate the relationships between the COL9 alpha chains, their regulatory mechanisms, as well as future treatment possibilities.

Collie eye anomaly

Collie eye anomaly (CEA) is a hereditary visual impairment with heterogeneous signs. Primarily, the retina is not affected, but in severe forms of the disease, retinal detachment can be present. CEA affects many dog breeds that are members of the herding group or those breeds that were used for their creation (Collie Rough and Smooth, Border Collie, Australian Shepherd, Shetland Sheepdog – Sheltie, Nova Scotia Duck Tolling Retriever, Lancashire Heeler, Longhaired Whippet, Boykin Spaniel, and the Hokkaido dog breed). Some other breeds may be affected by CEA, but the disease has not been identified in others to date. The causative mutation for CEA was identified in the nonhomologous end joining factor 1 (NHEJ1) gene as a large deletion in intron 4, which contains several highly conserved elements (reviewed by Palanova).

Retinal diseases with unidentified genetic basis

The retinopathy observed in the Swedish Vallhund – based on its inheritance patterns and the progressive nature of its vision loss – appears to be a form of PRA. The phenotype of this trait is rather different from most forms of PRA (multifocal, rather than diffuse, distribution of retinal degeneration). Age of disease onset and disease progression can vary considerably. This progressive retinopathy affects both rod and cone PRs, as well as the RPE. Affected dogs do not share common alleles, and so six known canine retinal disease genes (BEST1, PDE6A, PDE6B, RPE65, NPHP4, and CNGB3) were excluded as causative for novel PRA in Swedish Vallhunds. Because of the individual variability in disease progression, even among related dogs, it seems that genetic disease modifiers are present in this breed.

Retinal degeneration II (central PRA or pigment epithelial dystrophy) is a disease of the PR cells and retinal blood vessels. The first sign of this disease is central vision loss. Later on in the course of the disease, cataracts may develop. Two genes have been associated with this disease (GNB3 [encoding guanine nucleotide binding protein or G protein], beta polypeptide 3, and PRPH2 [encoding peripherin 2]), but no causative mutations were found. But the majority of central PRA has been considered nutritional rather than genetic, eg, central PRA in the Walker Hounds and Beagles fed with a vitamin E-deficient diet.

Retinal detachment was observed, eg, in the Shi-Tzu breed. Retinal and vitreous degeneration were also frequently observed in affected dogs from the mentioned breed.

Retinal dysplasia affects many breeds, is due to the incorrect formation of the retina, and results in complete vision loss.

Retinal dysplasia and persistent hyperplastic primary vitreous were found in miniature Schnauzer dogs. It is a congenital hereditary trait with autosomal recessive mode of inheritance, and so far, the causative gene has not been found.

Retinal dystrophy or choroideremia was observed in the Great Dane breed. It exhibits hereditary progressive PR degeneration. The CHM gene was screened in 22 breeds with various forms of retinal dystrophies but variants were found to be nonpathogenic.

Retinoschisis is not simple to detect because it is often confused with retinal degeneration. The most common form is secondary retinoschisis subsequent to other disorders. Rod dysplasia is a hereditary retinal disease inherited as a recessive trait in the Norwegian Elkhound.

Rod–cone dysplasia is supposed to be associated with GNGT2 (guanine nucleotide binding protein [G protein], gamma transducing activity polypeptide 2) gene, but altered...
localization of GNGT2 protein in affected retinas was found only in cd-affected dogs.\textsuperscript{85}

**New findings in hereditary retinal diseases**

A number of genetic mutations for different forms of retinal degenerations were found, but key components and the molecular events that link specific mutations to PR degeneration remain poorly characterized. Multiple pathways (proapoptotic and prosurvival) are associated with PR degeneration.\textsuperscript{86} Epigenetic mechanisms (eg, microRNA [miRNA] regulation) also play an essential role in the control of the complex visual processes during eye development and disease.\textsuperscript{87} The work of Genini et al\textsuperscript{88} focused on expression of miRNAs in healthy and XLPR\textsuperscript{4} (as well as rcd1, erd, and prcd)-affected retinas brings forth some interesting findings. The expression of miRNAs was monitored in three phases of retinal development after birth (at 3 weeks – induction phase; 7 weeks – execution phase; and 16 weeks – chronic cell death phase in affected tissue). Results demonstrated that miRNA expression differences were minimal between 3 weeks and 7 weeks, but a substantial number of altered miRNAs were identified at 16 weeks of age in XLPR\textsuperscript{4}-affected retinal tissues compared to normal healthy tissues. This suggests that miRNAs may be effectors or may arise in response to retinal disease progression, but they are not initiators of the PR degeneration process. Although different mutations trigger retinal diseases, the authors\textsuperscript{88} observed commonalities in the miRNA expression pattern, which appear to be associated with the kinetics of PR cell death. This suggests that the use of miRNAs as a target for future therapeutic design might be effective for treating retinal degenerative diseases regardless of the causative mutation.\textsuperscript{88}

**Genetic testing of hereditary retinal diseases**

Many genetic tests for screening for causative mutations of hereditary retinal diseases were developed,\textsuperscript{3,17,22,23,89} and some of them were further improved\textsuperscript{26,90,91} for faster, simple, and accurate identification of affected dogs and carriers of the mutations. For a detailed review of genetic testing of hereditary diseases of dogs, refer the work of Mellersh.\textsuperscript{92} The advantages of DNA tests are that, unlike clinical diagnostic methods, they can detect unaffected carriers and subclinical cases prior to the onset of disease. If there are other forms of retinal diseases or any other hereditary disease with unidentified mutation in the breed, currently available DNA tests are not able to exclude all inherited diseases in the breed.\textsuperscript{93}

**Implications for diagnosis and management of hereditary retinal diseases**

Because hereditary diseases represent a growing problem in many dog breeds all over the world, it is necessary to fight them. Screening for clinical signs of retinal diseases and genetic testing of known causative mutations should be standard in breeding practice.

But not all retinal diseases and other hereditary traits have causative mutations identified. These traits should also be taken into account to assess the overall health and genetic potential of the dog.\textsuperscript{93}

It is important to preserve genetic variability within and across dog breeds. One of the common problems in dog breeding is common sire phenomenon, but some breeder clubs have solved it successfully (for a detailed review focused on dog breeding problems, refer the work of Leroy\textsuperscript{94}).

In some dog breeds, only one hereditary retinal disease presents, so it is possible to eliminate the disease allele successfully by simple elimination of all carriers of this allele (especially if the frequency of disease allele is low). In some extraordinary cases, when the affected dog (or carrier of causative mutation) is an excellent individual of the breed, breeding can be permitted; all of the progeny should be tested for the presence of the disease allele. Other breeds carry more than one hereditary retinal disease and, sometimes, other hereditary diseases with known or unknown genetic bases. This complicates the situation for breeders and breeder clubs. Generally, it is not possible to choose completely genetically healthy parents for the next generation of dogs (we are speaking of known genetically transmitted traits only, because these are what we can control). Breeders must take into account not only the genetic health of the dog, but also its good character, which is frequently neglected, and the working utilization of the dog for which the breed was created. First, the frequently occurring disease/trait should be solved, and after the reduction of incidence of this disease allele, another allele should be chosen to fight it with. The dog, man’s best friend, represents not only our assistant in many facets of our lives, it is our pet, friend, and a suitable animal model of many diseases too. Dogs are dependent on our care and protection, and that is why we must take care of their health and welfare.
Disclosure
The author reports no conflicts of interest in this work.

References


