Identification of a variant in NDP associated with X-linked retinal dysplasia in the English cocker spaniel dog

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Abstract

Purpose
Three related male English Cocker Spaniels (ECS) were reported to be congenitally blind. Examination of one of these revealed complete retinal detachment. A presumptive diagnosis of retinal dysplasia (RD) was provided and pedigree analysis was suggestive of an X-linked mode of inheritance. We sought to investigate the genetic basis of RD in this family of ECS.

Methods
Following whole genome sequencing (WGS) of the one remaining male RD-affected ECS, two distinct investigative approaches were employed: a candidate gene approach and a whole genome approach. In the candidate gene approach, COL9A2, COL9A3, NHEJ1, RS1 and NDP genes were investigated based on their known associations with RD and retinal detachment in dogs and humans. In the whole genome approach, affected WGS was compared with 814 unaffected canids to identify candidate variants, which were filtered based on appropriate segregation and predicted pathogenic effects followed by subsequent investigation of gene function. Candidate variants were tested for appropriate segregation in the ECS family and association with disease was assessed using samples from a total of 180 ECS.

Results
The same variant in NDP (c.653_654insC, p.Met114Hisfs*16) that was predicted to result in 15 aberrant amino acids before a premature stop in norrin protein, was identified independently by both approaches and was shown to segregate appropriately within the ECS family. Association of this variant with X-linked RD was significant (P = 0.0056).
Conclusions
For the first time, we report a variant associated with canine X-linked RD. *NDP* variants are already known to cause X-linked RD, along with other abnormalities, in human Norrie disease. Thus, the dog may serve as a useful large animal model for research.

Introduction
Retinal dysplasia (RD) is a developmental disorder that is defined as abnormal differentiation of the retina with proliferation of one or more of its components [1]. It has been described in multiple dog breeds and can be subdivided into focal, multifocal, geographic and total retinal dysplasia types [2]. RD can present as a nonsyndromic form such as that described in the Bedlington and Sealyham Terriers or it can present as a syndromic RD such as in oculoskeletal dysplasia described in the Labrador Retriever, Samoyed and Northern Inuit dog [3–7]. Genetic variants associated with oculoskeletal dysplasia have been identified in all of these breeds [7, 8]. Total RD presents clinically as total retinal detachment. This can be associated with non-attachment due to failure of contact between the neurosensory retina and retinal pigment epithelium during embryogenesis or due to complete detachment of the retina. It may also be seen in conjunction with microphthalmos, vitreal dysplasia, leukocoria and a rotary-searching nystagmus [2].

Retinal detachment has also been reported as a clinical feature in Collie Eye Anomaly (CEA) in multiple breeds [9]. CEA is characterized by regional hypoplasia of the choroid. In severe cases and especially those with colobomas, affected dogs can develop retinal detachments, intraocular haemorrhage and subsequent blindness [9, 10]. The genetic variant for CEA has been identified in non-homologous end joining factor 1 (*NHEJ1*) [11]. The mode of inheritance of total RD and CEA has been reported to be autosomal recessive in all breeds [3, 4, 7, 8, 10]. Although other retinal pathologies have been found to be X-linked in dogs, such as XLPRA in the Siberian Husky as well as other breeds, to the author’s knowledge an X-linked RD has not been reported before [12].

Many genetic disorders that affect humans have an equivalent disease that is recognised in the dog [13]. Retinal diseases that can result in blindness in people that are known to be X-linked include retinoschisis and *NDP* (norrin cystine knot growth factor)-related retinopathies [14, 15]. Retinoschisis can present in infants as young as two months of age but more commonly presents in males at five years and upwards with macular dysfunction. In more severe forms of the disease retinal detachment can occur with a variable reported incidence of 16.6–30% [14, 16, 17]. Variants in *XLRS1* (X-linked (juvenile) retinoschisis 1) have previously been associated with retinoschisis [18–20].

*NDP*-related retinopathies are characterised by a spectrum of fibrous and vascular changes of the retina at birth that progress and cause varying degrees of visual impairment in males. The most severe phenotype described is Norrie disease that presents clinically with a grey-yellow fibrovascular mass (pseudoglioma) secondary to retinal vascular dysgenesis and retinal detachment with congenital blindness [21]. Cognitive impairment and behavioural disturbances occur in approximately 33% and 45% of people with the disease respectively. Progressive sensorineural hearing loss usually develops, with one study revealing 85–90% of patients experiencing the onset of hearing loss by their mid-20s and cryptorchidism has also been rarely reported in cases of Norrie disease [22–24]. Associated genetic variants have been identified within *NDP*, which encodes a small, presumably secreted protein (norrin), with a
cysteine-knot motif [25]. Greater than 100 different disease-causing NDP variants have been identified in humans to date [26]. In mice, norrin has been shown to play important roles in angiogenesis, not only in development of the eye but also the ear, brain, and female reproductive system [27].

Here we describe a novel form of RD associated with a variant in NDP in the English Cocker Spaniel (ECS), a small dog of the gundog breed, first recognised as a separate variety in 1893 [28]. In a family of ECS where three of the males were reported to be congenitally blind and pedigree analysis was suggestive of an X-linked mode of inheritance, we sought to investigate the genetic basis of the RD. Using two distinctive investigative approaches; a candidate gene approach and a whole genome sequencing (WGS) approach, we identified a provocative candidate variant. To the best of our knowledge, this is the first description of canine X-linked RD and the first time a variant in NDP has been associated with any disease in the dog. These findings also reveal the dog as a potential large animal model for studying NDP-related retinopathies in humans.

Materials and methods

Ethics statement

All dogs used for this study were privately owned pet dogs that were examined at the owners’ request following informed and written owner consent. Buccal mucosal swab sampling was performed following written owner consent and no in vivo experiments were undertaken. All clinical examinations were conducted as part of routine ophthalmic examination and not specifically for research purposes. All sample collection was approved by the Animal Health Trust Ethics Committee (04–2018).

Animals used

In this study a total of 1260 samples of 178 breeds were used. This consisted of DNA samples from one affected ECS, nine related ECS, 170 unaffected ECS, 264 unaffected dogs, WGS data from an additional 814 canids and clinical information only from two more affected ECS. Affected dogs were defined as those with RD in the form of retinal detachment. Twelve related ECS from two litters (three affected and nine unaffected) were included in the study. DNA was collected from ten of the twelve dogs, including one affected. The remaining two affected were deceased and were therefore unavailable for DNA collection.

DNA samples from an additional 170 unaffected ECS and from 264 unaffected dogs of 93 other breeds that were previously collected for unrelated research projects were used in the candidate variant genotyping.

WGS data from 814 canids were used as ‘controls’ in the whole genome approach. These comprised two distinct groups: WGS data from 198 unaffected dogs (comprising of 98 breeds and multiple crossbreeds) previously sequenced by the authors for other studies, and WGS from 616 canids of unknown phenotypes (comprising 122 dog breeds, multiple crossbreeds, and eight wolves, of unknown clinical status) derived from the Dog Biomedical Variant Database Consortium (DBVDC) [29].

Clinical investigation

Ocular examinations were performed on all twelve related ECS, between 2010–2018, at the Animal Health Trust (AHT) veterinary hospital’s ophthalmology department, by veterinary ophthalmologists, due to owners’ concerns of visual impairment in three of the males. The original two affected males were examined at 6 weeks old in 2010, whilst the affected case was examined
at 6 weeks, 12 weeks and 12 months old in 2017–18. Five generation pedigrees were obtained for all related ECS (Fig 1). Ocular ultrasonography was performed in the three affected males. The sire of the second litter had been examined for hereditary eye diseases as part of the British Veterinary Associated/Kennel Club/International Sheep Dog Society Eye Scheme.

**Ocular examination**

All ophthalmic examinations performed in the study included: vision assessment via maze testing and visual responses; rebound tonometry (TonoVetTM iCare, FI-01510 Vantaa, Finland); slit-lamp biomicroscopy (Kowa SL-17, Torrance, California, USA); and direct and indirect ophthalmoscopy (Welch Allyn Ltd, Ashby de la Zouch, Leicestershire, UK and Keeler Professional, Windsor, Surrey, UK) after dilation with 1% tropicamide (Minims, Bausch & Lomb, Kingston-upon-Thames, Surrey, UK). B-mode ocular ultrasound was performed on the three affected ECS (Easoate MyLabONE, 10-18MHz 30mm linear probe, Imotek, Somersham, Cambs, UK). Topical anaesthetic 0.5% proxymetacaine (Minims, Bausch & Lomb, Kingston-upon-Thames, Surrey, UK) was instilled in the eyes prior to ocular ultrasonography. Carbomer gel (Viscotears, Bausch & Lomb, Kingston-upon-Thames, Surrey, UK) was used as a coupling medium for ocular ultrasonography, and the probe was placed in a horizontal position in contact with the central cornea and axial eye length was measured from the corneal endothelium to the internal sclera. Fundus photography was performed in four ECS (RetCam Shuttle, Clarity, Macclesfield, Cheshire, UK).

**DNA extraction**

DNA was extracted from the buccal mucosal swabs using the QIAamp DNA Blood Mini Kit (Qiagen, Manchester, UK) according to the manufacturer’s instructions. DNA concentration and purity were determined by using the NanoDrop 1000 spectrophotometer (Thermo Fisher Scientific, Waltham, Massachusetts, USA).
Scientific, Loughborough, UK) and/or the Qubit fluorometer with the Qubit dsDNA broad range (BR) Assay Kit (Invitrogen, Loughborough, UK). Initial primers for the candidate gene approach were designed using Primer3 and purchased from Integrated DNA Technologies (IDT, Leuven, Belgium) [30].

Whole genome sequencing

WGS of DNA from one RD-affected ECS was outsourced to Edinburgh Genomics, UK. Illumina sequencing of a TruSeq Nano library on a HiSeq X sequencing platform and paired-end sequencing generated with read lengths of 150 base pairs and approximately 30x coverage of the dog genome. Reads were then aligned to the canine reference genome (CanFam3.1) using the Burrows-Wheeler Aligner and variant calls were made using GATK (HaplotypeCaller) [31, 32]. The WGS was also visualised and manually interrogated using Integrative Genomics Viewer (IGV) [33]. WGS data from this dog is available from the European Nucleotide Archive (ENA, Accession number: PRJEB39869).

Candidate gene approach

Five candidate genes that have been associated with retinal detachment in dogs or X-linked retinal detachments in humans were selected (NHEJ1, COL9A3, COL9A2, RS1 and NDP) [7, 8, 11, 18, 34]. Using IGV, the exons and introns of each candidate gene and 5000bp of flanking DNA were examined for single nucleotide variants (SNVs), insertions and deletions using WGS from an unaffected American cocker spaniel and the CanFam3.1 for reference and comparison. The potential variants were filtered by consideration of their genomic context (i.e. if they are within coding or exonic sequence, or are splice site variants), their presence in the control genomes, their location with respect to repetitive regions and their mapping quality score. Ensembl genome browser, PolyPhen and Provean were used to further analyse potential causal variants to predict pathogenicity [35–37]. PredictProtein and Swiss-Model were used to assess structural differences between the normal and altered NDP protein [38, 39].

Whole genome approach

WGS from 198 dogs were processed through the same analysis pipeline as the ECS case. Genomic Variant Call Format (VCF) files from 199 canine WGS (198 “controls” and one ECS case) were combined using HaplotypeCaller into a multi-sample VCF file. For cross-genome analysis, Variant Effect Predictor (VEP) was run on the multi-sample file. Two greyhounds were affected with a similar phenotype and were therefore omitted from the multi-sample file. Variants predicted to result in premature start/stop codons, splicing variants, nonsense variants, coding insertions or deletions, or missense variants predicted by Polyphen and Provean to be pathogenic were retained for further investigation [35, 37]. The remaining variants were then filtered based on appropriate segregation (homozygous for the risk-variant in the case and homozygous wild type or heterozygous for the risk variant in the controls). The DBVDC genome dataset containing 616 additional canids was used for additional variant filtering based on appropriate segregation. Pubmed, OMIM, Retnet and Varelect were used to investigate gene function, expression and phenotype of remaining variants [40–42].

Candidate variant genotyping

Candidate variants that remained after filtering were investigated further by Sanger sequencing of the 10 related ECS (1 affected and 9 unaffected ECS) and appropriate segregation for an autosomal recessive or X-linked mode of inheritance was assessed (S1 Appendix).
The remaining candidate variant in NDP was genotyped in 170 ECS and 264 dogs of 93 breeds, using allele specific probes in an allelic discrimination assay. Custom primers designed to be 135 base pairs in length (forward: 5’-GGAGAGGATGTACCGGTAGGT; reverse: ACTT CGGCTGCGCTGTT) and allele-specific probes (variant: FAM-CCTCATGCCCCCCG; reference: VIC-AGCCTCATGCCCCCCG) (Thermo Fisher Scientific, Loughborough, UK) were used. Reactions were carried out in 8 μL volumes including 4 μL LUNA Universal qPCR Master Mix (New England Biolabs), 1.4 μL MQ, 0.4 μL DMSO, 0.2 μL of primers and 2 μL genomic DNA. Cycling parameters were 25˚C for 30 seconds, 95˚C for 3 minutes, 40 cycles of 95˚C for 3 seconds and 60˚C for 10 seconds, and finally 25˚C for 30 seconds.

Statistics
Using Microsoft Excel, a Fisher’s exact test using a 2 x 2 contingency table was used to assess significance of association of the candidate variant with RD using the one affected and 170 unaffected ECS. Statistical significance was set at P ≤ 0.05.

Results
Clinical findings
The related ECS were all purebred dogs that resided in the UK. A pedigree was established of those related ECS (Fig 1). In the original litter examined (seven puppies), two males were found to have behavioural signs suggestive of severe visual deficits (determined by observing their movements in unfamiliar surroundings), absent pupillary light reflexes, wandering nystagmus, darkened irises, and extensive posterior segment haemorrhage. Ocular ultrasonography revealed total retinal detachment and increased echogenicity within posterior segment in all eyes. The globe diameters (anterior to posterior) were comparable with their normal litter mates at 16mm in all dogs except one affected dog, whose globes were 15mm OU. Both males were euthanised, one dog when less than one year old due to behavioural abnormalities and blindness, the other at six years old due to suspected glaucoma and aggression. One male was found to be bilaterally cryptorchid on clinical examination.

The sire of the second litter had no abnormalities detected on ocular examination. In the second litter examined (four puppies), one male displayed behavioural signs suggestive of severe visual deficits (determined by observing his movements in unfamiliar surroundings), and was found to have absent dazzle and pupillary light reflexes, a wandering nystagmus, corneal endothelial opacities ventrally in the right eye, darkened irises, hyphema of the right eye and extensive posterior segment haemorrhage in both eyes, with the retina in the left eye appearing detached and as a fibrovascular retrolental mass (Figs 2 and 3). Ocular ultrasonography revealed total retinal detachment and increased echogenicity within the posterior segment in both eyes. The left globe appeared microphthalmic with an anterior-posterior diameter of 16.42mm versus 21.88mm in the right eye. The affected male was subsequently re-examined, the left eye appeared relatively unchanged whereas the corneal pathology had advanced to diffuse corneal degeneration with neovascularisation and dense crystalline stromal deposits in the right eye (Fig 4). Microphakia was evident along with incipient nuclear cataract being present (Fig 5). On clinical examination the dog was found to be cryptorchid at 10 months old.

Identification of candidate variants by a candidate gene approach
Manual visual inspection of the DNA sequence of the five candidate genes, comparing the affected ECS with 1 unaffected dog and the CanFam3.1 reference using IGV, identified 196 variants that were initially narrowed down to 12 potential variants by the manual interrogation.
described above. This was then further narrowed down on the basis of predicted pathogenicity (using Provean and Polyphen-2) to two potential variants. The first was a missense variant in exon seven in \textit{COL9A3} (c.1240G$>$C, p.Asp414His), predicted to be “probably damaging” by PolyPhen and “neutral” by Provean. The second was a frameshift variant in \textit{NDP} (c.338_339insC), that was predicted to result in 15 aberrant amino acids before a premature stop in norrin protein (p.Met114Hisfs’16), resulting in a loss of the last five amino acids. Each of the aberrant amino acids was independently assessed for pathogenicity using PolyPhen-2 and Provean (Table): four changes were predicted to be “Probably Damaging” by PolyPhen-2 and nine changes were predicted to be “Deleterious” by Provean (Table 1). The NDP protein is made up of 133 amino acids; therefore, 15.0% of the protein is predicted to be either changed or lost. In addition, significant structural differences between the wildtype and altered proteins are predicted (Fig 6).

\textbf{Identification of candidate variants by a whole genome sequencing approach}

WGS data from the RD-affected ECS was compared to those from 198 unaffected control dogs of 98 other breeds. There was a total of 27,874,423 variants in at least one of these genomes. These variants were first filtered based on predicted effect on a protein using the Ensembl Variant Effect Predictor (VEP) [43, 44], resulting in 530 variants. These variants were further
filtered based on appropriate segregation for an autosomal recessive mode of inheritance which resulted in 23 remaining variants (Table 2).

Of the 23 variants only 14 were within known genes that had human orthologues. Five of these variants were found in the additional 648 canid genomes within the DBVDC and were excluded from further investigation. This resulted in 9 variants in 9 individual genes. Gene function was investigated using Pubmed, OMIM, Retnet and Varelect. Only two of these genes were known to be involved in retinal development: *NDP* and *FJX1* (four jointed box kinase 1). The variant in *NDP* was identical to that discovered by the candidate gene approach (c.338_339insC). The variant in *FJX1* was a frameshift variant (c.199_260del).

**Genotyping the candidate variants**

Sanger sequencing was used to genotype the nine related ECS and the one RD-affected dog for which DNA was available. The RD-affected dog was homozygous for two of the variant alleles (*NDP*, *COL9A3*) but was homozygous for the wildtype *FJX1* allele with sanger sequencing. Manual inspection of the *FJX1* variant region in the affected WGS revealed highly repetitive sequence and reads with poor alignment with CanFam3.1, and the calling of this as a variant is therefore, a false positive. Three unaffected ECS were also homozygous for the *COL9A3* variant, whereas all nine unaffected dogs were either heterozygous or homozygous for the wildtype *NDP* allele and all nine unaffected dogs were homozygous wild type for *FJX1*. Therefore, only the *NDP* variant segregated appropriately for an autosomal recessive or X-linked mode of inheritance among the nine related ECS (Fig 6). An allelic discrimination assay was then performed in the additional 170 unaffected ECS and 264 unaffected dogs of 93 other breeds. All
these dogs were homozygous for the wild type allele. Association of the \textit{NDP} variant with RD was significant ($P = 0.0056$).

**Discussion**

In this study, two different approaches, candidate-gene and whole genome, were used to identify a single exonic insertion variant in \textit{NDP} which was shown to be associated with X-linked RD in the ECS. This variant is predicted to result in a 15 residue shift in the reading frame, a premature stop codon and norrin protein truncated by five residues, out of a normal length of 133 amino acids. While structural variants near exons would have been detected in the candidate gene approach, our current whole genome analysis pipeline does not detect structural variants. It is, therefore, worth noting that the presence of structural variation causing disease in this study cannot be fully excluded. In addition, the canine genome is not well annotated with regulatory regions and lncRNAs, and these were therefore not investigated and cannot be excluded. Indeed, a lncRNA (NDP-AS1) on the reverse strand is annotated on the human genome build GRCh38.p13, but it is unclear if the canine genome shares this feature, and if it does whether the variant described here affects the NDP protein, the lncRNA or both. It is also worth noting linkage disequilibrium (LD) was not investigated due the small number of WGS available for ECS in this study (four in total including the case), which was considered too few...
Fig 5. Ocular defects associated with affected ECS. Showing corneal and lenticular abnormalities.

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| Amino acid change | PolyPhen Result   | PolyPhen Score | Provean Result | Provean Score (Threshold < -2.5) |
|-------------------|-------------------|----------------|----------------|---------------------------------|
| M114H             | Possibly Damaging | 0.566          | Neutral        | -0.102                          |
| R115E             | Possibly Damaging | 0.462          | Neutral        | -0.647                          |
| L116A             | Possibly Damaging | 0.816          | Deleterious    | -3.109                          |
| T117H             | Probably Damaging | 0.974          | Deleterious    | -3.353                          |
| A118R             | Probably Damaging | 0.958          | Neutral        | -1.118                          |
| T119H             | Probably Damaging | 0.974          | Neutral        | -1.647                          |
| Y120L             | Possibly Damaging | 0.816          | Deleterious    | -6.235                          |
| R121P             | Possibly Damaging | 0.827          | Deleterious    | -4.941                          |
| Y122V             | Possibly Damaging | 0.816          | Deleterious    | -6.882                          |
| I123H             | Possibly Damaging | 0.944          | Deleterious    | -1.588                          |
| L124P             | Probably Damaging | 0.974          | Neutral        | -1.588                          |
| S125L             | Possibly Damaging | 0.633          | Deleterious    | -3.176                          |
| C126L             | Possibly Damaging | 0.462          | Deleterious    | -10                              |
| H127S             | Possibly Damaging | 0.462          | Neutral        | 2.255                            |
| C128L             | Possibly Damaging | 0.462          | Deleterious    | -7.118                          |

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Fig 6. Comparison of the wildtype (A) and altered protein (B). Using PredictProtein reveal that the two proteins are expected to differ throughout with regards to secondary structure, protein binding and conservation. In addition, modelling of both proteins using Swiss-Model (C) reveal that in the altered molecule, the 15 amino acids at the C-terminus, involved in the shifted reading frame, do not align with the N-terminus (white and red in C) of the wildtype protein at all, and also do not align with any other known protein structures.

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Table 2. 23 remaining variants.

| Chromosome | Genomic coordinates | Reference allele | Alternate allele | Consequence | Gene Symbol | Gene Full Name | Human Orthologue | Present in DBVDC | Base change | Amino acid change | CDS Position | Protein Position |
|------------|---------------------|------------------|-----------------|-------------|-------------|----------------|------------------|----------------|-------------|-----------------|------------|------------------|
| 8          | 3597623             | A                | C               | missense_variant | DACT1       | Dishevelled Binding Antagonist of Beta Catenin 1 | DACT1 | Yes | gAG/gGg       | 1700       | 567              |
| 25         | 35076169            | C                | G               | missense_variant | BMP1        | Bone Morphogenic Protein 1 | BMP1 | Yes | gGo/gGc        | 11        | 38               |
| 26         | 947613              | TA               | TTTCTCTGTGTTAAA, TTTCTCTGTTAA | frameshift_variant | ENSCAFG00000000764 | None | Yes | An/Ac | I/X | 1549 | 517 |
| 31         | 2139695             | G                | GP              | frameshift_variant | KLHDC9      | kelch domain containing 9 | KLHDC9 | No | -/A           | 75–76     | 305–316          |
| 38         | 5947278             | G                | A               | missense_variant | COL6A2      | Collagen Type IX Alpha 2 Chain | COL6A2 | Yes | aGa/aAt | 2495 | 832          |
| 17         | 2001410             | G                | OC              | frameshift_variant | GAREM2      | Grb2-associated regulator of Erk/MAPK1 | GAREM2 | No | -/-X         | 735–736   | 245–246         |
| 18         | 32393127            | C                | frameshift_variant | FJX1 | Four Jointed Box Kinase 1 | FJX1 | No | CTGGAGGAGCGGGTGCC | CCAGATACGAGCAGATcc/aCCA | 67–87 |
| 6          | 8011462             | A Pituitary CTG  | A,AGCTCTCTCTCTCTG | missense_variant | ENSCAFG00000000764 | None | Yes | aCCAGCACTCTAACAGCATCTCCAGAaTACGAGCAGACcc | FF/LSPF | 68–75 |
| 12         | 520701              | GAGAGAAAGAGAGAAAA | frameshift_variant | FJX1 | Four Jointed Box Kinase 1 | FJX1 | No | CTGGAGGAGCGGGTGCC | CCAGATACGAGCAGATcc/aCCA | 67–87 |
| 16         | 1239695             | A                | G               | missense_variant | ENSCAFG00000000764 | None | Yes | aGo/aGa | 422 | 141 |
| 11         | 5402599             | G                | GAAGAAAGAAAGAAAGA | frameshift_variant | ENSCAFG00000000764 | None | Yes | aGo/aGa | 77+77 | 20+26 |
| 21         | 6799663             | T                | G               | missense_variant | CCDC82      | Coiled-coil Domain Containing 82 | CCDC82 | Yes | goT/goG | 171 | 57 |

(Continued)
| Chromosome | Genomic coordinates | Reference allele | Alternate allele | Consequence | Gene Symbol | Gene Full Name | Human Orthologue | Present in DBVDC | Base change | Amino acid change | CDS Position | Protein Position |
|------------|-------------------|----------------|----------------|-------------|-------------|----------------|-----------------|----------------|-------------|----------------|--------------|----------------|
| 25         | 44173062          | G              | A              | missense_variant | CHRND       | Cholinergic Receptor Nicotinic Delta Subunit | CHRND | No            | cGc/cAc     | R/H          | 1400          | 407           |
| 34         | 19179708          | G              | A              | missense_variant | TBCCD1      | TBCC Domain Containing 1 | TBCCD1 | No            | aG/aTg      | T/M          | 167           | 56            |
| 39         | 1615231           | G              | A              | missense_variant| ARSF        | Arylsulfatase F | ARSF | No            | aG/aA      | R/K          | 161           | 34            |
| 39         | 3790069           | G              | GC             | frameshift_variant | NDP        | Norrin Cystine Knot Growth Factor | NDP | No            | ggc/ggGc    | G/GX         | 338–339       | 113           |
| 6          | 5860792           | C              | G              | missense_variant | CCDC18      | Coiled-Coil Domain Containing 18 | CCDC18 | Yes           | aG/aC      | M/I          | 1230          | 410           |
| 7          | 9830091           | C              | A              | missense_variant | TRAF5       | TNF Receptor Associated Factor 5 | TRAF5 | No            | G/aA      | Q/K          | 1003          | 335           |
| 9          | 59422620          | G              | A              | missense_variant | DENND1A     | DENN Domain Containing 1A | DENND1A | No            | cGc/cAc    | R/H          | 2279          | 760           |

Two remaining variants FJX1 and NDP involved in retinal development highlighted in blue.

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samples for an accurate estimate of LD in the breed in general and especially in the vicinity of the variant of interest by the authors.

This is the first time a variant in \textit{NDP} has been described in the dog and the first time X-linked RD has been reported in the dog. The results suggest that this variant is likely to be private to this family of ECS as it was not identified in the additional 170 ECS and the 1078 other non-ECS canids. The pedigree analysis was helpful in determining it was most likely a recessive condition as only a small proportion of the progeny was affected and, because it was only observed in male progeny, a X-linked mode of inheritance was also hypothesized. This narrowed down the potential candidate genes to be investigated. Genomic material was only available from one affected as the other two affected were deceased. The low number of affected animals is a limiting factor for this study. Ocular tissue from RD-affected ECS would also be helpful for assessing changes in the mRNA and protein expression within the retina of these dogs in comparison to age matched control dog retinas. Further work is required to prove this is the causal variant of RD in the ECS, although the predicted effect on the protein is supportive of this hypothesis.

WGS is a very useful and cost-effective method of identifying variants, when faced with a small number of affected cases. Unlike genome wide association studies, a small number, or even a single case may be sufficient to identify underlying variant as in this study. If successful, WGS can help identify risk variants early on in the disease emergence process, hence eliminating it before the disease becomes widespread within the population, which is especially important in dogs due to high levels of inbreeding, popular sire effects and relatively fast turnover of progeny [45, 46]. A limitation to using WGS as with other sequencing techniques, is the inherent difficulty in the detection of structural variants, for example, transposons, inversions, or large insertions and deletions which our current WGS pipeline cannot identify. Genome-wide association studies can be useful in identifying the approximate location of the disease variant even when there is a large structural variant, so its use in combination with WGS could help mitigate this limitation. Unfortunately, due to our low case numbers, this was not possible in this study. It is therefore worth noting that the presence of structural variation causing disease cannot be fully excluded.

The candidate gene approach was considered appropriate for this study because similar phenotypes had been reported in humans and dogs which have been shown to be associated with a limited number of genes. This approach is useful when trying to identify the genetic basis of a disease thought to be of simple inheritance using only a small number of cases [47]. It requires existing knowledge about the known or presumed biology of the phenotype under investigation, either in other breeds of dog or in other species, which was available in this instance [48]. This method is however biased and limited, as it only includes genes known to be associated with the relevant phenotype, meaning variants in novel genes not previously associated with the phenotype will go undetected [49]. A whole genome approach was also used, which does not require prior knowledge of gene candidacy and thus is considered a less biased approach compared to candidate gene analysis [50].

\textit{NDP} encodes for norrin, a cysteine knot growth factor, that is required for angiogenesis in the eye, ear, brain and female reproductive system [47]. Norrin is an atypical Wnt (wingless-related integration site) ligand, that has binding sites for both \textit{Fz4} (Frizzled 4) and \textit{LRP5/6} (low-density lipoprotein receptor-related protein) which activates β-catenin signalling, that plays a central role in retinal vascularisation [47, 51]. Its structure is a semi-circular shaped homodimer linked through three intermolecular disulphide bonds. Disease-causing variants will affect the protein folding and stabilisation of the norrin structure and hence affect protein function [52].
Over 100 disease-causing NDP variants have been reported in humans [26]. Analysis of gnomAD data within the equivalent human C-terminal region affected by the frameshift and truncation (X:43809045–43809107, based on genome build GRCh37/hg19) revealed eight variants. Of these, three are missense, five are synonymous and none are truncating. However, analysis of the ClinVar data in gnomAD revealed five additional “likely pathogenic” variants, including three missense and two nonsense/stop gained variants [53]. These variants can result in inherited retinopathies including Norrie disease, X-linked familial exudative vitreoretinopathy, retinopathy of prematurity and Coat’s disease [54]. Retinal detachment has been reported in all of these conditions to varying degrees at different clinical stages [55–57]. Retinal detachment in Norrie disease has even been reported in utero during the third trimester in a known carrier of the condition [58]. The affected ECS males were examined at 6 weeks old, although blindness was suspected earlier than this. Puppies open their eyes at approximately two weeks of age [59]. In the affected ECS, retinal detachment was suspected to be congenital, but this cannot be confirmed.

Histopathological characterisation is usually not possible in humans with NDP-related retinopathies. Therefore Ndph (Norrie disease homologue)-knockout mice have been used as a small animal model to study early histopathological changes [60]. In mice with targeted inactivation of Ndp, histological data suggest retrolental structures and a disorganized ganglion cell layer as primary pathogenic events [61]. Histopathology of the retina and posterior segment of an RD-affected ECS would be useful to better identify the phenotypic characteristics of this type of RD in the dog. No ocular tissue was available from the affected dog in this study, so the pathogenesis can only be hypothesised based on the clinical findings observed. Although there were subtle ophthalmic differences in the three affected dogs, the overall ocular changes were similar, as they all shared the congenital blindness, extensive posterior segment haemorrhage and the retinal detachment. The congenital retinal detachments in the form of fibrovascular retrolental masses and extensive vitreal haemorrhage, is more similar to that described in clinical cases of human Norrie disease than the other inherited retinopathies associated with NDP variants [62, 63].

Cognitive impairment, behavioural disturbances and progressive sensorineural hearing loss have been described in humans with Norrie disease. The two affected males of the initial litter of ECS were euthanised early in life due to behavioural abnormalities that were thought to be related to their lack of vision. The affected male in the second litter was not reported to show behavioural abnormalities. No males were reported to have any hearing impairment, although brainstem auditory evoked response testing was not performed to confirm this, in the affected ECS to confirm this. Hearing loss is reported to develop in humans in their early twenties so it is possible that the affected males had not yet reached an appropriate age for this to have developed and the one surviving affected male could develop hearing loss as he continues to mature [22].

This is the first time a variant in NDP has been associated with a retinal disease in the dog and also the first X-linked RD to be reported in the dog. Therefore, this study also suggests the dog may serve as a useful model for understanding human NDP-related retinopathies and the development of gene and other therapies.

Supporting information

S1 Appendix. Reaction mixes and thermal cycling parameters: Sanger sequencing of candidate variants.

(DOCX)
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