Retrospective and prospective study of progressive retinal atrophy in dogs presented to the veterinary hospital of the Federal University of Paraná, Brazil

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Abstract

Background: Progressive retinal atrophy (PRA) is one of the main causes of blindness in dogs. Despite its clinical importance, there is limited epidemiological information available, particularly in South America.

Aim: The main objective of this study was to perform a retrospective, and prospective analysis of PRA in dogs admitted at the Veterinary Hospital of the Federal University of Paraná, Brazil.

Methods: Medical records of dogs admitted between 2014 and 2018 were selected through the archives of the Comparative Ophthalmology Laboratory. A total of 130 dogs with medical records indicating clinical signs suggestive of PRA, independent of the electroretinography confirmation, were selected. In order to investigate common characteristics, each patient's clinical history, ophthalmic examination, and visual status were reviewed (obstacle course, pupillary light reflex, dazzle reflex, visual tracking to a cotton ball, and menace responses). Additionally, a prospective study was performed, where flash electroretinography was performed on 30 animals with clinical signs suggestive of PRA, and 14 animals were selected for fundus photography. Data were assessed through descriptive and inferential statistics.

Results: A total of 2,055 dogs were evaluated between January 2014 and December 2018. Of those, 130 animals were presumptively diagnosed with PRA (6.33%), consisting of 18 different breeds and 27 dogs with a mixed pedigree. Poodles were the most prevalent breed (n = 26; 20.00%), followed by Cocker Spaniels (n = 19; 14.62%). In the reported caseload, Pomeranians showed a considerably higher odds ratio for PRA development (15.36%).

Conclusion: Pomeranians presented a high odds ratio, suggesting that further studies may be performed with breeds with a high potential for developing this disease.

Keywords: PRA, ERG, Retinal dystrophy.

Introduction

Retinal dystrophies are a group of inherited diseases that can cause blindness in humans and animals. Progressive retinal atrophy (PRA) is a group of inherited diseases characterized by progressive loss of photoreceptors and atrophy of the retina secondary to gene mutations (Magnusson, 1911; Petersen-Jones, 1998a), and it is one of the main types of retinal dystrophies affecting domestic animals. PRAs were first described in dogs in 1911 by Magnusson in Gordon Setters in Sweden but has also been reported in pigs (Li et al., 1998), sheep (Banin et al., 2015), horses (Winkler et al., 2020), non-human primates (Winkler et al., 2020), laboratory animals (Huang et al., 1993; Naash et al., 1993), and several breeds of cats (Rubin and Lipton, 1973, Giuliano and van der Woerd, 1999, Narfström et al., 2009). Since first reported in Gordon Setters, more than 100 breeds of dogs have been described with various PRAs (Petersen-Jones, 1998b; Petersen-Jones, 2005; Beltran, 2009). Histological and genetic analyzes have enabled PRAs to be further classified into different categories, most broadly based on whether the rods or cones are primarily affected and by the age of onset. Early-onset forms of PRAs are usually caused by a developmental abnormality of the photoreceptors (i.e., dysplasias) and are characterized by the onset of clinical signs prior to 1 year of age and rapid progression. Late-onset forms occur after normal maturation of the photoreceptors (i.e., degenerations) and affect middle-aged or older dogs, such as progressive rod-cone degeneration (PRCD) (Petersen-Jones, 2005).

Clinically, PRAs are characterized by the bilateral degeneration of the retina, causing progressive vision loss culminating in blindness, similar to retinitis pigmentosa, a homologous group of retinal degenerative disorders that are important causes of blindness in humans (Petersen-Jones, 1998a; Dias et al., 2018). Across breeds, nyctalopia (or night-blindness)
is the most common initial presenting clinical sign, although initial presentation with more advanced signs is not uncommon, such as mydriasis, bright eye-shine, secondary cataract formation, and loss of daytime vision and/or total blindness (Whitley et al., 1995; Petersen-Jones, 1998a; Petersen-Jones, 2005). Diagnosis is based primarily on history and clinical signs, with electrotetroretinography (ERG) being the definitive test. ERG usually changes precedes funduscopic changes. Progression of clinical signs varies with breeds and form of PRA, but PRAs culminate in blindness (Petersen-Jones, 1998a; Petersen-Jones, 2005).

Up until now, no therapeutic approach to limit progression or reverse vision loss associated with PRAs has been developed that is widely accepted, accessible, or feasible. However, gene therapy is showing promise in both veterinary and human ophthalmology (Petersen-Jones, 2012; Al-Saikhan, 2013; Apte, 2018; Takahashi et al., 2018; Arbabi et al., 2019). Studying domestic veterinary species with natural forms in PRAs has been pivotal in developing our understanding of the genetic basis of PRAs and new therapeutic strategies that may minimize the occurrence in both human and animal populations (Beltran, 2009; Petersen-Jones, 2012; Zeiss, 2013).

Previous studies have focused primarily on the clinical features and prevalence of PRAs in a specific breed or small group of breeds or have evaluated in detail the genetic aspects and molecular biology of a particular form of PRA. However, aspects including breed distribution have not been described. Considering the paucity of data information on PRAs in Brazil, the main objective of the present study was to evaluate the prevalence of this group of diseases and describe the nature of PRAs across the population of dogs presenting to the Laboratory of Comparative Ophthalmology (LABOCO) of the Veterinary Hospital of the Federal of Paraná (UFPR), as a means to identify new patterns of the disease and that may contribute to more objective studies in the future.

Materials and Methods
Data from 2,055 dogs presenting to the LABOCO with or without signs of retinal degeneration consistent with PRAs were collected by a clinic management system software (Vetus®, São Paulo, SP, Brazil) and through medical records of animals admitted between January 2014 and December 2018, regardless of diagnosis. Information about the breed, age, sex, presence of concomitant systemic and ocular diseases, presence of visual impairment or nyctalopia, ophthalmoscopy signs of retinal degeneration, presence of cataracts suspected to be secondary to retinal degeneration (due to clinical appearance, clinical history evidence, and ERG results), and personal data of the owners of the animals were collected and transcribed into an Excel spreadsheet (Microsoft®, Redmond, WA) for retrospective analysis. At each visit, a thorough ophthalmic examination was performed. Anterior segment exam was performed with a portable slit lamp (Hawk-Eye, Dioptrix, L’Union, France). Vision testing was assessed by the ability to track falling cotton balls and the ability to negotiate an obstacle course in both photopic and scotopic conditions. Neuroophthalmic parameters tested included the direct and consensual pupillary light reflex (PLR) and dazzle reflex using a 3.5 V Finoff halogen fiber optic transilluminator (Welch Allyn®, Skaneateles Falls, NY), as well as the menace response. Indirect ophthalmoscopy was performed after inducing mydriasis using 1% tropicamide (Mydriacyl®, Novartis®, São Paulo, SP, Brazil) with a 3.5 V Finoff halogen fiber optic transilluminator (Welch Allyn®, Skaneateles Falls, NY) and a 20D lens (OPTOMED OY LTD., Finland). In 14 dogs without advanced cataracts or nuclear sclerosis, fundus photography was performed using a ClearViewfundus camera (ClearView Optical Imaging System Optibrand, Ft Collins, CO). By nature of a retrospective analysis, not all animals had complete data recorded, including vision testing, in the medical record. Therefore, the resulting percentages reflect the number of eyes examined by each technique, and that was entered in the patient’s record.

ERG was performed on a total of 26 dogs within the study period and 4 dogs within the LABOCO archives, all of which had visual impairment with signs of retinal degeneration and/or cataract. Dogs with other causes of abnormal ERGs, such as posterior uveitis, retinal detachment, sudden acquired retinal degeneration syndrome (SARDS), and glaucoma, were excluded. The dogs were prepared for ERG with the instillation of a drop of Tropicamide 1% (Mydriacyl; AlconTM, São Paulo, SP, Brazil) combined with Phenylephrine 10% (Allergan, Guarulhos, SP, Brazil) in each eye every 5 minutes until complete mydriasis was achieved. Following mydriasis, dogs were dark-adapted for 20 minutes. A mini portable Ganzsfield (HMsERG model 1000, Ocuscience®, NV) was used to perform the flash electroretinogram. ERGs were performed using manual restraint without the use of anesthesia or sedation. Animals were placed in sternal recumbency with the active electrode (ERG-Jet, Fabrinal SA, La Chaux- de-Fonds, Switzerland) positioned on the cornea, and hypodermic platinum needles (Model E2, Grass Technologies, Warwick, NY) positioned as reference and ground electrodes. The reference electrode was positioned about 2 cm from the lateral canthus, and the ground electrode was positioned at the base of the neck. The protocol used was a short protocol, consisting of light flash intensities of 10, 3,000, and 10,000 mcd.s/m², based on the protocol described in Somma et al. (2016). A-wave and b-wave amplitudes, and implicit times (ITs), were measured by ERGVIEW 4.380V software (Ocuscience®, NV), and the data generated were analyzed using descriptive and inferential statistics. A Shapiro–Wilk test was used to assess data normality. For the sex-related data, the Chi-square test was used. The mean age of onset of the disease was compared...
in each breed by an unpaired Student’s \( t \)-test. Both analyses used a significance level of \( p < 0.05 \) and were performed using the GraphpadQuickCalcs statistical software (Graphpad Software Inc., La Jolla, CA). The analysis of non-normal data, such as ERG implicit time, was performed through the Mann–Whitney test \((p < 0.05)\). The odds ratio or the odds of a breed having PRA compared to the total population presenting to the LABOCO for PRA development in each breed was calculated by the MEDCALC® software (MedCalc Software bvba, Ostend, Belgium). The odds ratio values were considered significant when they did not present the value “1” in their confidence interval. Descriptive statistical data were calculated using the Excel software (Microsoft®, Redmond, WA).

**Ethical approval**
The project was registered in the Research Ethics Committee on Animal Use (Comissão de Ética no Uso de Animais—CEUA, in Portuguese) of UFPR, protocol number 045/2017.

**Results**

**Retrospective evaluation**
A total of 2,055 privately owned animals were admitted at LABOCO between 2014 and 2018. Females (1,107, 53.87%) outnumbered males (948, 46.13%). PRA was diagnosed in 130 dogs (6.33%), with no statistical difference (Chi Squared test) between sex (males 67, 51.53%; females 63, 48.47%; \( p < 0.05 \)). A total of 18 breeds were diagnosed with PRA, the most prevalent being the Miniature Poodles \((n = 26, 20.00\%)\), followed by English Cocker Spaniels \((n = 19, 14.62\%)\), and Lhasa Apsos \((n = 13, 10.00\%)\). When considering the prevalence of PRAs in relation to the entire population presenting to the LABOCO, the Pomeranian had a significantly higher odds ratio (Tables 1 and 2).

Presenting clinical signs reported by owners were nyctalopia (93, 71.54%) and general vision impairment (6, 4.62%). The mean age of onset of presenting clinical signs reported by the owners was 8.55 ± 3.92 years, and the mean age at the time of diagnosis of PRA was 9.06 ± 3.91 years (range 3 months to 16 years). No significant difference in the mean age of presentation or prevalence of PRAs was found across breeds when compared to the total population \((p <0.05)\) (Table 3).

**Clinical evaluation**
Among the 130 PRAs affected dogs, 208 eyes were evaluated through direct and consensual pupillary reflex, while 206 eyes were evaluated through dazzle reflex, menace response, and tracking (cotton wool

| Breed                        | Number of affected animals | Number of normal animals | % of affected | Total  |
|------------------------------|----------------------------|--------------------------|---------------|--------|
| Undefined breed (UB)         | 27                         | 471                      | 5.42          | 498    |
| Toy Poodle                   | 26                         | 194                      | 11.82         | 220    |
| Cocker Spaniel               | 19                         | 76                       | 20.00         | 95     |
| Lhasa Apso                   | 13                         | 224                      | 5.49          | 237    |
| Labrador Retriever           | 9                          | 34                       | 20.93         | 43     |
| Pomeranian                   | 5                          | 5                        | 50.00         | 10     |
| Shih Tzu                     | 4                          | 175                      | 2.23          | 179    |
| Yorkshire Terrier            | 4                          | 101                      | 3.81          | 105    |
| Maltese                      | 4                          | 38                       | 9.52          | 42     |
| Dachshund                    | 4                          | 26                       | 13.33         | 30     |
| Pinscher                     | 3                          | 90                       | 3.23          | 93     |
| American Pit Bull Terrier    | 3                          | 21                       | 12.50         | 24     |
| Bichon Frisé                 | 2                          | 6                        | 25.00         | 8      |
| Brazilian Terrier            | 2                          | 3                        | 40.00         | 5      |
| Pug                          | 1                          | 63                       | 1.56          | 64     |
| Beagle                       | 1                          | 38                       | 2.56          | 39     |
| Miniature Schnauzer          | 1                          | 33                       | 2.94          | 34     |
| Chow Chow                    | 1                          | 22                       | 4.35          | 23     |
| Whippet                      | 1                          | 3                        | 25.00         | 4      |
| Others                       | 0                          | 302                      | 0             | 302    |
| Total                        | 130                        | 1,925                    | 6.33          | 2,055  |
ball) response tests. In addition, 72 were submitted to the obstacle course test in photopic and scotopic conditions. Menace and tracking responses were the most commonly abnormal tests: 65.05% of the animals showed a negative response/absence of response. Moreover, 32.31% of the evaluated dogs presented at least scotopic impairment in the obstacle course, and 19.23% showed complete both night and day blindness (Tables 4 and 5).

Seventy-two animals (55.38%) were diagnosed with secondary cataracts. Of these, the highest rate was detected in poodles (21; 29.16%), followed by Cocker Spaniel (15; 20.83%) and mixed breed dogs (13; 18.05%). Equatorial and posterior cortical vacuoles were the most common initial presentation. All stages of cataracts maturity were found, with the incipient stage most common (48 eyes; 36.09%) (Table 6).

Fundus abnormalities consistent with PRAs, including early signs such as a granular tapetum, or more advanced sigs as tapetal hyperreflectivity, vascular attenuation, and optic disc atrophy, were found in 141 eyes (54.23%). The most prevalent fundus abnormality was retinal vascular attenuation, noted in 120 of the affected eyes (85.10%). The main funduscopic changes can be seen in Table 7.

### Table 2. Distribution of breeds diagnosed with PRAs relative to the total number of affected animals and their respective of odds ratio (2014–2018).

| Breed               | Number of diseased animals | % diseased dogs | Odds ratio | Confidence interval |
|---------------------|---------------------------|-----------------|------------|---------------------|
| UB                  | 27                        | 20.77           | 0.81       | 0.52–1.25           |
| Toy Poodle          | 26                        | 20.00           | 2.23       | 1.42–3.51           |
| Cocker Spaniel      | 19                        | 14.62           | 4.16       | 2.43–7.13           |
| Lhasa Apso          | 13                        | 10.00           | 0.84       | 0.47–1.52           |
| Labrador Retriever  | 9                         | 6.92            | 4.13       | 1.94–8.82           |
| Pomeranianian       | 5                         | 3.85            | 15.36      | 4.39–53.76          |
| Shih Tzu            | 4                         | 3.08            | 0.32       | 0.12 - 0.87         |
| Yorkshire Terrier   | 4                         | 3.08            | 0.57       | 0.21–1.58           |
| Dachshund           | 4                         | 3.08            | 2.32       | 0.80–6.75           |
| Maltese             | 4                         | 3.08            | 1.58       | 0.55–4.49           |
| Miniature Pinscher  | 3                         | 2.31            | 0.48       | 0.15–1.54           |
| American Pit Bull Terrier | 3 | 2.31 | 2.14 | 0.63–7.28 |
| Bichon Frisé        | 2                         | 1.54            | 5.00       | 1.00–25.01          |
| Brazilian Terrier   | 2                         | 1.54            | 10.01      | 1.66–60.45          |
| Beagle              | 1                         | 0.77            | 0.38       | 0.05–2.83           |
| Chow Chow           | 1                         | 0.77            | 0.67       | 0.09–5.01           |
| Pug                 | 1                         | 0.77            | 0.23       | 0.03–1.67           |
| Miniature Schnauzer | 1                         | 0.77            | 0.44       | 0.06–3.28           |
| Whippet             | 1                         | 0.77            | 4.97       | 0.51–48.08          |
| Total               | 130                       | 100             | -          | -                   |

### Discussion

Retrospective studies may provide a baseline for understanding the prevalence of disease in the specific
Table 3. Mean age of onset of clinical signs as described by owners and mean age at the time of diagnosis of PRA for different breeds of dogs.

| Breed         | Number of animals | Age of onset (mean/years) | Age at diagnosis (mean/years) |
|---------------|-------------------|---------------------------|-------------------------------|
| UB            | 27                | 8.85 ± 3.88               | 9.26 ± 3.79                   |
| Toy Poodle    | 26                | 9.05 ± 3.54               | 9.69 ± 4.17                   |
| Cocker Spaniel| 19                | 9.39 ± 3.68               | 9.79 ± 3.60                   |
| Lhasa Apso    | 13                | 9.08 ± 3.25               | 10.15 ± 3.36                  |
| Labrador Retriever | 9          | 6.78 ± 1.92               | 7.67 ± 1.58                   |
| Pomeranian    | 5                 | 3.15 ± 3.99               | 3.35 ± 4.30                   |
| Shih Tzu      | 4                 | 7.74 ± 2.06               | 8.00 ± 2.16                   |
| Yorkshire Terrier | 4            | 5.38 ± 4.39               | 7.25 ± 4.03                   |
| Dachshund     | 4                 | 9.75 ± 5.12               | 9.75 ± 5.12                   |
| Maltese       | 4                 | 8.25 ± 3.30               | 8.75 ± 4.11                   |
| Miniature Pinscher | 3         | 11.00 ± 2.65              | 11.00 ± 2.65                  |
| American Pit Bull Terrier | 3    | 2.75 ± 3.70               | 5.00 ± 3.46                   |
| Bichon Frisé  | 2                 | 9.50 ± 2.12               | 9.50 ± 2.12                   |
| Brazilian Terrier | 2         | 15.00 ± 1.41              | 16.00 ± 0.00                  |
| Beagle        | 1                 | 4                         | 4                             |
| Chow Chow     | 1                 | 13                        | 13                            |
| Pug           | 1                 | 7                         | 9                             |
| Miniature Schnauzer | 1   | 8                         | 8                             |
| Whippet       | 1                 | 4                         | 4                             |
| Total         | 130               | 8.55 ± 3.92               | 9.06 ± 3.91                   |

Table 4. The number of eyes with PRA resulting in abnormal clinical test results.

| Eye tests                      | Direct PLR | Indirect PLR | Dazzle reflex | Menace response | Tracking response (cotton wool ball) |
|--------------------------------|------------|--------------|---------------|-----------------|-------------------------------------|
| Present                        | 90 (43.27%)| 92 (44.23%)  | 80 (38.83%)   | 72 (34.95%)      | 72 (34.95%)                         |
| Absent/decreased               | 118 (56.73%)| 116 (55.77%)| 126 (61.17%)  | 134 (65.05%)     | 134 (65.05%)                        |
| Total                          | 208        | 208          | 206           | 206             | 206                                 |

Table 5. Obstacle course test results for dogs diagnosed with PRAs.

| Result—Obstacle course test | Number of animals |
|------------------------------|-------------------|
| Scotopic impairment          | 42 (32.31%)       |
| Scotopic impairment with discrete photopic impairment | 4 (3.07%) |
| Complete blindness           | 25 (19.23%)       |
| No deficit                   | 6 (4.62%)         |
| Unknown                      | 53 (40.77%)       |
| Total                        | 130               |

Table 6. Number of eyes diagnosed with cataracts secondary to PRAs and classified by stage of maturity in dogs with PRAs in LABOCO’s case series (2014–2018).

| Stage of maturity | Number of eyes |
|-------------------|----------------|
| Hypermature       | 24 (18.04%)    |
| Mature            | 22 (16.54%)    |
| Mature            | 39 (29.32%)    |
| Immature          | 48 (36.09%)    |
| Incipient         | 133            |
population and thus are essential steps in targeting areas in need of focused research. Considering limitations, the present investigation may have suffered an influence from a selection bias for the analysis, introduced by the selection of specific dogs that have been diagnosed with PRA in a Brazilian veterinary ophthalmology service (influenced by the most common local breeds). Thus, proper randomization was not achieved because the dogs were not randomly selected from the general population, and the general population's size was not determined. Other sources of bias should be considered as well; for example, Miniature Poodle owners might be more likely to have brought affected dogs to the facility for evaluation than owners of Miniature Schnauzers. Thus, extrapolation of results should be done with caution. Determining the number of cases among dogs presenting to LABOCO does not measure incidence within a breed or among all dogs in Brazil, but only among dogs that presented to the institution. Presumably, dogs presented to the LABOCO because they have health issues, and healthy dogs would likely be under-represented in this index population. Nevertheless, the present study provides the first report of the prevalence of PRAs in any canine population in South America. The prevalence of PRAs in the LABOCO was highest in Miniature Poodles (20.00%) and English Cocker Spaniels (14.62%), similar to what was previously reported in North America by Petersen-Jones (1998a). Lhasa Apso dogs were the third most prevalent; however, the odds ratio was not significant ($p = 0.05$). Considering all PRAs affected dogs, Pomeranians and the Brazilian Terrier had significantly higher odds ratio when compared to the total population, followed by Cocker Spaniel, Labrador retriever, and Toy Poodle. There is no published description of PRAs in Brazilian Terriers, an uncommonly presented breed at the Veterinary Hospital of the UFPR. The popularity of Pomeranians as a pet was increasing and were the

| Funduscopic evaluation | Hyperreflectivity | Vascular attenuation | Optic disc atrophy | Tapetal granulation | Total |
|------------------------|------------------|----------------------|--------------------|--------------------|-------|
|                        | 78 (55.31%)      | 120 (85.10%)         | 2 (1.41%)          | 2 (1.41%)          | 141   |

**Fig. 1.** Ocular fundus photographed by a ClearView camera and representative ERG tracings of dogs referred to LABOCO. A—Normal ocular fundus of a 5-month-old Pomeranian, showing a blue-green tapetal zone, adequate retinal vasculature emanating from the optic disc, dark spots representing the non-tapetal areas, and a well-myelinated optical disc. B—Normal 10,000 mcd.s/m² ERG tracing of the same dog as in (A), showing a well-defined a-wave (negative) and b-wave (positive). C—A 2-year-old Pomeranian with evidence of advanced PRA fundoscopically, showing pronounced vascular attenuation, tapetal hyperreflectivity, optic disc atrophy. D—ERG tracing of the same dog as in (C) at 10,000 mcd.s/m² showing an absent ERG waveform.
second most common breed recorded in the Brazilian Cynophilia Confederation (CBKC) in 2018, with a total of 18,828 registered dogs (CBKC, personal communication). Therefore, it is important to consider future studies on PRAs in Pomeranians to establish the features of these diseases and, in the future, make pedigree evaluations and molecular analysis studies to identify mutant genes and minimize the incidence and number of unaffected genetic carriers within the population.

The mean age of onset of clinical signs first noted by owners of the dogs affected with PRAs was 8.55 ± 3.92 years. However, the true age of onset of clinical signs could be earlier due to the owner’s difficulty recognizing the signs if subtle or if their pet was successfully adapting to visual impairment. American Pit Bull Terriers, Pomeranians, and Yorkshire Terriers have a lower mean age of onset. These findings are supported by Kijas et al. (2004) and Safatle et al. (2005), who identified a form of cone-rod dystrophy in young Pit Bulldogs with early-onset and rapid progression. However, a previous report described Yorkshire Terriers and Pomeranians with late-onset forms of PRAs (Downs et al., 2014).

Nystagmus is typically the first clinical sign noted due to initial rod photoreceptor degeneration, often described by owners as their pets having insecurity to navigate their environment in low light situations. Other behavioral changes, such as increased aggression or lethargy, have also been described (Barnett, 1965; Levin, 1998). Despite the ability to detect subtle changes of early PRAs on clinical examination, this opportunity does not frequently arise in practice. Due to the high level of adaptability that some dogs achieve as PRAs progress or due to the inability of an owner to detect subtle behavioral changes related to visual impairment, unfortunately, PRAs are often presented to a veterinarian for evaluation when in more advanced stages. Early stages of PRAs are most often detected incidentally on examination, often upon referral from a primary care veterinarian for secondary cataracts. In order to detect early PRA, younger close relatives of affected dogs should be evaluated (e.g., offspring of affected dogs).

Funduscopy is a useful tool in the early diagnosis of PRAs.常说 the first recognized on clinical examination. Early signs include light tapetal granularity and peripheral vascular taper, followed by tapetal hyperreflectivity and vascular attenuation coincident with progressive loss of photoreceptors and retinal thinning (Petersen-Jones, 1998a). The most prevalent lesions in the present study were vascular attenuation (85.10%) and tapetal hyperreflectivity (55.31%). Additionally, different types of PRAs have distinctive patterns of retinal degeneration. For example, PRCD in the Toy Poodle tends to present with early signs 5 years of age. Histologically, the photoreceptors at the posterior pole and central retina are affected before peripheral photoreceptors in this breed (Aguirre and Acland, 1988). PRCD in the Cocker Spaniel is typically slower, and the retina’s nasal and temporal corner aspects are spared until late, leaving healthy photoreceptor islands (Aguirre and Acland, 1988). In this study, no specific differences between the PRAs of different breeds were detected clinically. Other common abnormalities detected on ophthalmic exam, especially in dogs with advanced disease, include reduced or absent dazzle reflex and menace response. Direct and indirect PLC usually somewhat preserved, although diminished with longer to return to a resting state following light stimulation, due to the need for only a few functional photoreceptors to initiate the reflex pathway.

Secondary effects of PRAs are common, most notably being the development of cataracts (Petersen-Jones, 1998a; Park et al., 2009). In this study, 72 dogs (55.38%) were diagnosed with cataracts, of which 61 (84.72%) were bilateral. Cases of anterior cortical incipient cataracts were excluded because this is not the typical presentation for cataracts secondary to PRAs. This value is higher than those shown in a previous study, which reported that 44% of dogs with PRAs had cataracts (Kraijer-Huver et al., 2008). Poodles were the breed most commonly affected by cataracts, mostly of the incipient stage, consistent with previous reports (Park et al., 2009, Donzel et al., 2017). Between August 2016 and July 2018, 205 animals were diagnosed by the LABOCO with cataracts secondary to PRAs, and of these, 40 (19.51%) were Toy Poodles. Several owners reported the presence of a cloudy aspect in their dog’s eyes, and soon after, they lost their sight. The owner’s ability to detect cataracts is often limited to the late immature to mature stage, contributing significantly to visual impairment.

ERG can be useful to differentiate causes of visual impairment, such as SARDS, optic neuritis, and PRAs. It is also a helpful diagnostic tool when examining the posterior segment is limited due to alteration in the ocular media transparency, such as cataracts (Nafstrom and Petersen-Jones, 2013; Ekesten et al., 2013; Drazek et al., 2014). In this study, a flash ERG protocol was used in conscious animals under manual restraint, similar to that used preoperatively for cataract surgery staging (Ekesten et al., 2013; Somma et al., 2016). This technique does not allow the differentiation of cone and rod responses and is rarely used in studies of retinal dystrophies that require more complete protocols to evaluate with more detail the retinal electrical activity. Freeman et al. (2013) found that noise levels in non-sedated and non-anesthetized animals were not different than those with sedation or anesthesia, but that anesthesia and sedation both significantly reduced a- and b-wave amplitudes as well as ITs. Most generically, when using a short ERG protocol, PRAs cause a decrease in a-wave and b-wave amplitudes and an increase in their implicit
Our study corroborates these results; the a-wave and the b-wave of animals with PRAs were decreased at all light intensities or even absent in lower light intensities. In addition, both a-wave and b-wave ITs were significantly higher in the PRAs affected dogs, at light intensities of 3,000 and 10,000 med.s/m² ($p < 0.05$). Dogs with a clinical history of SARDs were excluded. However, some of the animals were selected for ERG even in the absence of a retinal abnormality on clinical examination, mainly because owners reported suspicious abnormalities in vision at home (e.g., especially in dim light). As a result, some normal ERG results were found.

Although the results only apply to the population of dogs presented at LABOCO and cannot be extrapolated to the entire breed, this study showed that Miniature Poodles and Cocker Spaniels were the breeds most commonly affected by PRAs presented at LABOCO. Further research on the prevalence of PRAs in populations of dogs throughout Brazil is recommended, which together may provide new avenues for research of inherited retinal diseases in domestic dogs. In addition, it is important to note that most dogs are highly adaptable to progressing visual impairment until a critical point is reached. Therefore, it is important to educate owners and raise awareness among the general population of signs to monitor for, particularly in high-risk breeds. Additional education is recommended for those owners with dogs diagnosed with PRAs, to help them successfully navigate the coming changes in their pet's lifestyle as visual impairment progresses. Pomeranians showed a high odds ratio within that population, and considering the increase in the popularity of this breed in recent years, monitoring the incidence of PRAs is recommended. Several commercially available tests for PRA-causing genetic variants for forms of PRA that occur in many breeds (e.g., see https://www.pawprintgenetics.com or https://breeder.wisdompanel.com). Blood samples were collected from all affected dogs in this investigation and will be screened for known PRA gene sequence variants in the near future.

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Conflict of interest

The authors declare that there are no conflicts of interest.

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Authors contribution

Henrique de Moura Freitas: Main executor of this project. Participated in the organization and preparation of all stages of this project including the schedule preparation, in the review of medical records and literature data, in the tabulation of data in tables, statistical analyses, in the registration of ophthalmoscopy and ERG images, writing, formatting and editing this work. Fabiano Montiani-Ferreira: Project advisor. Helped with the main idea and in the preparation and organization of all the stages of this project. Actively participated in the ophthalmoscopy and ERG and in the review of this paper. Also, participated in the edition of the images. André Tavares Sonna: Secondary author of this paper. Participated in the writing, review of the data, formatting and editing the figures and tables. Bret A. Moore: Secondary author of this paper. Participated in the writing, review of the data, formatting and editing the figures and tables.

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