


Sloughi	
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Ocular disorders known or presumed to be inherited (published)

	Diagnosis	Description and comments specific to the breed	Inheritance	Gene/ marker test	References
A	Entropion	Lower lateral	Unknown	NO	1
B	Progressive Retinal Atrophy (PRA) (rcd1a)	> 2 y.o. and progressively deteriorates > 4 y.o.	Autosomal recessive	PDE6B	2
C	Multifocal retinopathy	Dogs > 2 y.o	Unknown	NO	1

The ECVO's advice relating to hereditary eye disease control

Please see ECVO Manual chapter 8: VET Advice

Recommendations regarding age and frequency for eye examinations

Please see ECVO Manual chapter 7: ECVO Age and Frequency recommendations

Other ocular disorders (reported)

	Diagnosis	Source
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References

1. Chaudieu G, Chahory S. 2013. Affections oculaires héréditaires ou à prédisposition héréditaire chez le chien. Rueil-Malmaison : Ed. du Point Vétérinaire, Wolters Kluwer France.
2. Dekomien G et al (2000) Generalized progressive retinal atrophy of Sloughi is due to a 8 bp insertion in exon 21 of the PDE6B gene. Cytogenet Cell Genet 90 : 261-267.