Collie Eye Anomaly (CEA) / Choroidal Hypoplasia (CH)

**Affected breeds:**
- Australian Shepherd
- Bearded Collie
- Border Collie
- Lancashire Heeler
- Nova Scotia Duck Tolling Retriever
- Rough Collie
- Shetland Sheepdog
- Smooth Collie
- Whippet

Collies share Collie Eye Anomaly (CEA) with several other breeds – it’s not just a problem for collies. CEA is more technically known as Choroidal Hypoplasia (CH). It is a recessively inherited eye disorder that causes abnormal development of the choroid - an important layer of tissue under the retina of the eye. This disease is seen most frequently in U.S. collies, but also worldwide in Rough and Smooth Collies, Border Collies, Australian Shepherds, Lancashire Heelers, and Shetland Sheepdogs. Since the choroid layer does not develop normally from the start, the primary abnormality can be diagnosed at a very young age. Regrettably, there is no treatment or cure for CEA.

CEA is caused by a recessive genetic mutation. This means that dogs which carry the mutation ("CARRIERS") are normal but will pass the mutation on to an average of 50% of their offspring. Puppies which inherit two copies of the mutation will develop CEA ("AFFECTED").

**This test is particularly useful for breeders:**
- To identify carriers among their breeding stock so that they can avoid CARRIER X CARRIER mating combinations which would risk AFFECTED puppies.
- To conclusively confirm Progressive Retinal Atrophy caused by CEA

**This test will be reported as:**
- CLEAR : no evidence of the CEA mutation
- CARRIER : carries one copy of the defect, which will be passed to 50% of offspring
- AFFECTED : carries two copies of the defect, and will have CEA

The genetic status of dogs can be used to predict breeding outcomes when different combinations are mated:
- CLEAR X CLEAR = 100% CLEAR
- CARRIER X CLEAR = 50% CARRIER, 50% CLEAR
- CARRIER X CARRIER = 25% AFFECTED, 50% CARRIER, 25% CLEAR

**References**