Embark Veterinary Veterinary Practice



Swab code: zoe100100_swab Swab activated on 7/29/2022 Results completed on 7/29/2022 Report accessed on 7/29/2022 Ordered by Lorenz Connelly

vetsupport@embarkvet.com 1-855-203-8271

Patient Information

Client Information

Breed Information

Zoe

4 yrs 2 mths - F

Genetic Age: 40 human years Predicted Adult Weight: 75 lbs **Dean Knowles**

contact@example.com

555-555-4222

100.0% Rhodesian Ridgeback



1 Notable Result

Early Onset Adult Deafness, EOAD

Page 2



219 Clear Results

Zoe is not at increased risk for 219 of the genetic health variants that Embark tests.

Page 4



Glossary

Page 20

100.0% Rhodesian Ridgeback Results completed on: 7/29/2022



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1 Notable Result

Early Onset Adult Deafness, EOAD

Rhodesian Ridgeback Variant

How to interpret this result

Zoe has one copy of this variant in the EPS8L2 gene. Because this variant is inherited in an autosomal recessive manner (meaning dogs need two copies of the variant to develop the disease), Zoe is unlikely to develop Early Onset Adult Deafness, EOAD due to this variant.

Additionally, at this time, the clinical impact of this variant for dogs of Zoe's breeds is unknown. While she can pass this variant on to the next generation, the influence of this result in breeding decisions with dogs of the same breed(s) is undetermined. You can email vetsupport@embarkvet.com to discuss this further with a genetic counselor.

What is Early Onset Adult Deafness, EOAD?

Hearing loss can be categorized into three groups.

- Age-related hearing loss, referred to as presbycusis, is a slow but progressive loss of hearing commonly occurring in older animals and is a complex sensory disorder with a mixture of genetic and environmental components.
- Congenital deafness (hearing loss present at birth) may be attributed to environmental factors such as infections or genetic factors.
- Hearing loss can also take place neonatally or during puppy or early adulthood.

Similar to congenital deafness, the etiology of early-onset hearing loss is diverse, with a complex interaction of genetic and environmental components. Different etiological patterns (such as the age of onset and progression of the hearing condition) appear to be correlated with specific genes.

Variant Info

FPS8I 2 Deletion Exon 12 Recessive inheritance 1 copy of the variant



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This form of EOAD is characterized by bilateral deafness before two years of age and is not a pigmentation-related deafness.

Any questions?

You can email vetsupport@embarkvet.com or call 1-855-203-8271 should you desire to speak with a genetic counselor for more information.

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All Conditions Tested

To view COI and traits information, log into your account.

Auditory (2)

| | Gene | Copies | Results |
|--|-------------------------|--------|---------|
| Deafness and Vestibular Syndrome of Dobermans, DVDob, DINGS | МҮО7А | 0 | Clear |
| Early Onset Adult Deafness, EOAD - Rhodesian Ridgeback Variant | EPS8L2 Deletion Exon 12 | 1 | Notable |

Cardiac (4)

| Dilate | d Cardiomyopathy | Gene | Copies | Results |
|----------|--|---------------|-------------|----------------------|
| Ø | Dilated Cardiomyopathy, DCM1 - Doberman Pinscher Variant 1 | PDK4 | 0 | Clear |
| Ø | Dilated Cardiomyopathy, DCM2 - Doberman Pinscher Variant 2 | TTN | 0 | Clear |
| | | | | |
| Other | | Gene | Copies | Results |
| Other | Cardiomyopathy and Juvenile Mortality - Belgian Shepherd Variant | Gene YARS2 | Copies 0 | Results Clear |

Endocrine (3)

| Hypothyroidism | Gene | Copies | Results |
|--|------------|--------|---------|
| Congenital Dyshormonogenic Hypothyroidism with Goiter - Shih Tzu Variant | SLC5A5 | 0 | Clear |
| Congenital Hypothyroidism - Rat, Toy Fox, and Hairless Terrier Variant | TPO Exon 3 | 0 | Clear |
| Congenital Hypothyroidism - Tenterfield Terrier Variant | TPO Exon 9 | 0 | Clear |

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Gastrointestinal (4)

| Gastroenteropathy | Gene | Copies | Results |
|---|--------------|--------|---------|
| ✓ Lundehund Syndrome | LEPREL1 | 0 | Clear |
| Malabsorptive Disorder | Gene | Copies | Results |
| Imerslund-Grasbeck Syndrome, Selective Cobalamin Malabsorption - Beagle Variant | CUBN Exon 8 | 0 | Clear |
| Imerslund-Grasbeck Syndrome, Selective Cobalamin Malabsorption - Border Collie Variant | CUBN Exon 53 | 0 | Clear |
| Inherited Selected Cobalamin Malabsorption with Proteinuria - Komondor Variant | CUBN | 0 | Clear |

Hematologic (32)

| Coagulopathy | Gene | Copies | Results |
|--|----------------|--------|---------|
| Bernard-Soulier Syndrome, BSS - Cocker Spaniel Variant | GP9 | 0 | Clear |
| Congenital Macrothrombocytopenia - Cairn and Norfolk Terrier Variant | TUBB1 Exon 1 | 0 | Clear |
| Factor IX Deficiency, Hemophilia B - Rhodesian Ridgeback Variant | F9 Exon 7 | 0 | Clear |
| Factor IX Deficiency, Hemophilia B - Terrier Variant | F9 Exon 7 | 0 | Clear |
| Factor VII Deficiency | F7 Exon 5 | 0 | Clear |
| Factor VIII Deficiency, Hemophilia A - Boxer Variant | F8 Exon 10 | 0 | Clear |
| Factor VIII Deficiency, Hemophilia A - German Shepherd Variant 1 | F8 Exon 11 | 0 | Clear |
| Factor VIII Deficiency, Hemophilia A - German Shepherd Variant 2 | F8 Exon 1 | 0 | Clear |
| Glanzmann's Thrombasthenia Type I - Great Pyrenees Variant | ITGA2B Exon 13 | 0 | Clear |
| Glanzmann's Thrombasthenia Type I - Otterhound Variant | ITGA2B Exon 12 | 0 | Clear |
| May-Hegglin Anomaly - Pug Variant | MYH9 | 0 | Clear |

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| | Gene | Copies | Results |
|---|-----------------|--------|---------|
| P2Y12 Receptor Platelet Disorder - Greater Swiss Mountain Dog Variant | P2Y12 | 0 | Clear |
| Platelet Factor X Receptor Deficiency, Scott Syndrome - German Shepherd Dog | Variant TMEM16F | 0 | Clear |
| Prekallikrein Deficiency - Shih Tzu Variant | KLKB1 Exon 8 | 0 | Clear |
| Thrombopathia - American Eskimo Dog Variant | RASGRP1 Exon 5 | 0 | Clear |
| Thrombopathia - Basset Hound Variant | RASGRP1 Exon 5 | 0 | Clear |
| ✓ Thrombopathia - Landseer Variant | RASGRP1 Exon 8 | 0 | Clear |
| ✓ Von Willebrand Disease Type I, Type I vWD | VWF | 0 | Clear |
| ✓ Von Willebrand Disease Type II, Type II vWD - Pointer Variant | VWF | 0 | Clear |
| ✓ Von Willebrand Disease Type III, Type III vWD - Shetland Sheepdog Variant | VWF Exon 7 | 0 | Clear |
| ✓ Von Willebrand Disease Type III, Type III vWD - Terrier Variant | VWF Exon 4 | 0 | Clear |
| Red Blood Cell Abnormality | Gene | Copies | Results |
| Canine Elliptocytosis - Labrador Retriever Variant | SPTB Exon 30 | 0 | Clear |
| Methemoglobinemia - Pomeranian Variant | CYB5R3 | 0 | Clear |
| Pyruvate Kinase Deficiency - Basenji Variant | PKLR Exon 5 | 0 | Clear |
| Pyruvate Kinase Deficiency - Beagle Variant | PKLR Exon 7 | 0 | Clear |
| Pyruvate Kinase Deficiency - Labrador Retriever Variant | PKLR Exon 7 | 0 | Clear |
| Pyruvate Kinase Deficiency - Pug Variant | PKLR Exon 7 | 0 | Clear |
| Pyruvate Kinase Deficiency - Terrier Variant | PKLR Exon 10 | 0 | Clear |

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| White | Blood Cell Abnormality | Gene | Copies | Results |
|----------|---|----------------|--------|---------|
| | Canine Leukocyte Adhesion Deficiency Type I, CLAD I - Setter Variant | ITGB2 Exon 3 | 0 | Clear |
| ⊘ | Canine Leukocyte Adhesion Deficiency Type III, CLAD III - German Shepherd Variant | FERMT3 | 0 | Clear |
| ⊘ | Trapped Neutrophil Syndrome, TNS | VPS13B Exon 19 | 0 | Clear |
| Other | | Gene | Copies | Results |
| Ø | Ligneous Membranitis, LM - Scottish Terrier Variant | PLG | 0 | Clear |

Immunologic (6)

| | Gene | Copies | Results |
|--|--------------|--------|---------|
| Complement 3 Deficiency, C3 Deficiency - Brittany Variant | C3 | 0 | Clear |
| Severe Combined Immunodeficiency, SCID - Terrier Variant | PRKDC | 0 | Clear |
| Severe Combined Immunodeficiency, SCID - Wetterhoun Variant | RAG1 | 0 | Clear |
| Shar-Pei Autoinflammatory Disease, SPAID, Shar-Pei Fever | MTBP | 0 | Clear |
| X-linked Severe Combined Immunodeficiency, X-SCID - Basset Hound Variant | IL2RG Exon 1 | 0 | Clear |
| X-linked Severe Combined Immunodeficiency, X-SCID - Corgi Variant | IL2RG | 0 | Clear |

Integument (18)

| Collagen Abnormality | Gene | Copies | Results |
|---|----------------|--------|---------|
| Dystrophic Epidermolysis Bullosa - Central Asian Shepherd Dog Variant | COL7A1 | 0 | Clear |
| Dystrophic Epidermolysis Bullosa - Golden Retriever Variant | COL7A1 Exon 68 | 0 | Clear |



| | | Gene | Copies | Results |
|----------|---|-----------------|--------|---------|
| | Ehlers Danlos - Doberman Pinscher Variant | ADAMTS2 | 0 | Clear |
| ⊘ | Musladin-Lueke Syndrome, MLS - Beagle Variant | ADAMTSL2 Exon 7 | 0 | Clear |
| Kerati | n Abnormality | Gene | Copies | Results |
| ⊘ | Congenital Keratoconjunctivitis Sicca and Ichthyosiform Dermatosis, Dry Eye Curly 6 Syndrome, CKCSID - Cavalier King Charles Spaniel Variant | Coat FAM83H | 0 | Clear |
| ⊘ | Focal Non-Epidermolytic Palmoplantar Keratoderma, Pachyonychia Congenita - Dogue de Bordeaux Variant | KRT16 Exon 6 | 0 | Clear |
| | Hereditary Footpad Hyperkeratosis - Rottweiler Variant | DSG1 | 0 | Clear |
| | Hereditary Footpad Hyperkeratosis - Terrier and Kromfohrlander Variant | FAM83G | 0 | Clear |
| | Hereditary Nasal Parakeratosis, HNPK - Labrador Retriever Variant | SUV39H2 | 0 | Clear |
| Ø | Ichthyosis, Epidermolytic Hyperkeratosis - Terrier Variant | KRT10 Intron 5 | 0 | Clear |
| ② | Ichthyosis, ICH1 - Golden Retriever Variant | PNPLA1 Exon 8 | 0 | Clear |
| Ø | Ichthyosis - American Bulldog Variant | NIPAL4 Exon 6 | 0 | Clear |
| Ø | Ichthyosis - Great Dane Variant | SLC27A4 | 0 | Clear |
| Other | | Gene | Copies | Results |
| Ø | Bald Thigh Syndrome - Greyhound Variant | IGFBP5 | 0 | Clear |
| | Ectodermal Dysplasia, Skin Fragility Syndrome - Chesapeake Bay Retriever Variant | PKP1 Intron 1 | 0 | Clear |
| ② | Lethal Acrodermatitis, LAD - Bull Terrier Variant | MKLN1 | 0 | Clear |

| | Gene | Copies | Results |
|---|---------|--------|---------|
| Oculocutaneous Albinism, OCA - Small Breed Variant | SLC45A2 | 0 | Clear |
| X-linked Ectodermal Dysplasia, Anhidrotic Ectodermal Dysplasia, XHED - German Shepherd D Variant | og EDA | 0 | Clear |

| Metabolic (33) | | | |
|---|--------------|--------|---------|
| Enzyme Deficiency | Gene | Copies | Results |
| Hypocatalasia, Acatalasemia - Beagle Variant | CAT | 0 | Clear |
| L-2-Hydroxyglutaricaciduria, L2HGA - Staffordshire Bull Terrier Variant | L2HGDH | 0 | Clear |
| Pyruvate Dehydrogenase Deficiency - Spaniel Variant | PDP1 | 0 | Clear |
| Storage Disease | Gene | Copies | Results |
| Canine Fucosidosis - English Springer Spaniel Variant | FUCA1 | 0 | Clear |
| | GLB1 Exon 15 | 0 | Clear |
| GM1 Gangliosidosis - Portuguese Water Dog Variant | GLB1 Exon 2 | 0 | Clear |
| GM1 Gangliosidosis - Shiba Inu Variant | GLB1 Exon 15 | 0 | Clear |
| GM2 Gangliosidosis - Japanese Chin Variant | HEXA | 0 | Clear |
| GM2 Gangliosidosis - Poodle Variant | HEXB Exon 3 | 0 | Clear |
| Globoid Cell Leukodystrophy, Krabbe Disease - Terrier Variant | GALC Exon 5 | 0 | Clear |
| Glycogen Storage Disease Type IA, Von Gierke Disease, GSD IA - Maltese Variant | G6PC | 0 | Clear |
| Glycogen Storage Disease Type II, Pompe's Disease, GSD II - Finnish and Swedish Lapphund, Lapponian Herder Variant | GAA Exon 15 | 0 | Clear |
| Glycogen Storage Disease Type IIIA, GSD IIIA - Curly Coated Retriever Variant | AGL GDE | 0 | Clear |



| | | Gene | Copies | Results |
|------------|---|---------------|--------|---------|
| | Glycogen storage disease Type VII, Phosphofructokinase Deficiency, PFK Deficiency - Nachtelhund Variant | PFKM Exon 8 | 0 | Clear |
| | Glycogen storage disease Type VII, Phosphofructokinase Deficiency, PFK Deficiency - Whippet and English Springer Spaniel Variant | PFKM Exon 21 | 0 | Clear |
| ⊘ L | Lagotto Storage Disease | ATG4D Exon 10 | 0 | Clear |
| ⊘ L | ate-Onset Neuronal Ceroid Lipofuscinosis, NCL12 - Australian Cattle Dog Variant | ATP13A2 | 0 | Clear |
| ⊘ N | Mucopolysaccharidosis IIIB, Sanfilippo Syndrome Type B, MPS IIIB - Schipperke Variant | NAGLU | 0 | Clear |
| ⊘ N | Mucopolysaccharidosis Type IIIA, Sanfilippo Syndrome Type A, MPS IIIA - Dachshund Va | iriant SGSH | 0 | Clear |
| | Mucopolysaccharidosis Type IIIA, Sanfilippo Syndrome Type A, MPS IIIA - New Zealand Huntaway Variant | SGSH | 0 | Clear |
| ⊘ N | Mucopolysaccharidosis Type VII, Sly Syndrome, MPS VII - German Shepherd Variant | GUSB | 0 | Clear |
| ⊘ N | Mucopolysaccharidosis Type VII, Sly Syndrome, MPS VII - Terrier Brasileiro Variant | GUSB | 0 | Clear |
| ⊘ N | Neuronal Ceroid Lipofuscinosis 1, NCL1 - Dachshund Variant | PPT1 Exon 8 | 0 | Clear |
| ⊘ N | Neuronal Ceroid Lipofuscinosis 10, NCL10 - American Bulldog Variant | CTSD Exon 5 | 0 | Clear |
| ⊘ N | Neuronal Ceroid Lipofuscinosis 2, NCL2 - Dachshund Variant | TPP1 Exon 4 | 0 | Clear |
| | Neuronal Ceroid Lipofuscinosis 5, NCL5 - Border Collie and Australian Cattle Dog Variant | CLN5 Exon 4 | 0 | Clear |
| ⊘ N | Neuronal Ceroid Lipofuscinosis 5, NCL5 - Golden Retriever Variant | CLN5 Exon 4 | 0 | Clear |
| ⊘ N | Neuronal Ceroid Lipofuscinosis 6, NCL6 - Australian Shepherd Variant | CLN6 Exon 7 | 0 | Clear |
| ⊘ N | Neuronal Ceroid Lipofuscinosis 7, NCL7 - Chihuahua and Chinese Crested Variant | MFSD8 | 0 | Clear |
| | Neuronal Ceroid Lipofuscinosis 8, NCL8 - Australian Shepherd and German Shorthaired Pointer Variant | CLN8 | 0 | Clear |

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| | Gene | Copies | Results |
|---|-------------|--------|---------|
| Neuronal Ceroid Lipofuscinosis 8, NCL8 - English Setter Variant | CLN8 Exon 2 | 0 | Clear |
| Neuronal Ceroid Lipofuscinosis 8, NCL8 - Saluki Variant | CLN8 | 0 | Clear |
| Neuronal Ceroid Lipofuscinosis, Cerebellar Ataxia, NCL4A - American Staffordshire Terrier Variant | ARSG Exon 2 | 0 | Clear |

Muscular (13)

| Musculai (13) | | | |
|---|----------------|--------|---------|
| Movement Disorder | Gene | Copies | Results |
| Myotonia Congenita - Australian Cattle Dog Variant | CLCN1 Exon 23 | 0 | Clear |
| Myotonia Congenita - Miniature Schnauzer Variant | CLCN1 Exon 7 | 0 | Clear |
| Muscular Dystrophy | Gene | Copies | Results |
| Limb Girdle Muscular Dystrophy - Boston Terrier Variant | SGCD | 0 | Clear |
| Muscular Dystrophy - Cavalier King Charles Spaniel Variant 1 | DMD | 0 | Clear |
| Muscular Dystrophy - Golden Retriever Variant | DMD | 0 | Clear |
| Ullrich-like Congenital Muscular Dystrophy - Labrador Retriever Variant 1 | COL6A3 Exon 10 | 0 | Clear |
| Myopathy | Gene | Copies | Results |
| Centronuclear Myopathy, CNM - Labrador Retriever Variant | PTPLA | 0 | Clear |
| Exercise-Induced Collapse, EIC | DNM1 | 0 | Clear |
| Inflammatory Myopathy - Dutch Shepherd Variant | SLC25A12 | 0 | Clear |
| Inherited Myopathy of Great Danes | BIN1 | 0 | Clear |

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| | | Gene | Copies | Results |
|----------|--|-------------|--------|---------|
| ⊘ | Myotubular Myopathy 1, X-linked Myotubular Myopathy, XL-MTM - Labrador Retriever Variant | MTM1 Exon 7 | 0 | Clear |
| Ø | Nemaline Myopathy - American Bulldog Variant | NEB | 0 | Clear |
| Other | | Gene | Copies | Results |
| Ø | Myostatin Deficiency, Bully Whippet Syndrome | MSTN | 0 | Clear |

Neurologic (32)

| Brain or Seizure Disorder | Gene | Copies | Results |
|--|---------------|--------|---------|
| Alaskan Husky Encephalopathy, Subacute Necrotizing Encephalomyelopathy | LC19A3 Exon 2 | 0 | Clear |
| Alexander Disease - Labrador Retriever Variant | GFAP Exon 4 | 0 | Clear |
| Benign Familial Juvenile Epilepsy, Remitting Focal Epilepsy - Lagotto Romagnolo Variant | LGI2 Exon 8 | 0 | Clear |
| Cerebellar Abiotrophy, Neonatal Cerebellar Cortical Degeneration, NCCD - Beagle Variant | t SPTBN2 | 0 | Clear |
| Cerebellar Hypoplasia - Eurasier Variant | VLDLR | 0 | Clear |
| Hereditary Ataxia, Cerebellar Degeneration - Old English Sheepdog and Gordon Setter Variant | RAB24 Exon 1 | 0 | Clear |
| Neonatal Encephalopathy with Seizures, NEWS - Poodle Variant | ATF2 | 0 | Clear |
| Progressive Early-Onset Cerebellar Ataxia - Finnish Hound Variant | SEL1L | 0 | Clear |
| Spinocerebellar Ataxia with Myokymia and/or Seizures - Terrier Variant 2 | KCNJ10 | 0 | Clear |
| Spinocerebellar Ataxia, Late-Onset Ataxia, LoSCA - Terrier Variant 1 | CAPN1 | 0 | Clear |



| | | Gene | Copies | Results |
|----------|--|--------|--------|---------|
| ⊘ | Spongy Degeneration with Cerebellar Ataxia 1, SDCA1, SeSAME/EAST Syndrome - Shepherd Variant 1 | KCNJ10 | 0 | Clear |
| Ø | Spongy Degeneration with Cerebellar Ataxia 2, SDCA2 - Shepherd Variant 2 | ATP1B2 | 0 | Clear |
| oven | nent Disorder | Gene | Copies | Results |
| | Degenerative Myelopathy, DM | SOD1A | 0 | Clear |
| ⊘ | Hypomyelination and Tremors - Weimaraner Variant | FNIP2 | 0 | Clear |
| Ø | Juvenile Myoclonic Epilepsy - Rhodesian Ridgeback Variant | DIRAS1 | 0 | Clear |
| | Progressive Neuronal Abiotrophy, Canine Multiple System Degeneration, CMSD - Chinese Crested Variant | SERAC1 | 0 | Clear |
| ⊘ | Progressive Neuronal Abiotrophy, Canine Multiple System Degeneration, CMSD - Kerry Blue Terrier Variant | SERAC1 | 0 | Clear |
| ⊘ | Shaking Puppy Syndrome, X-linked Generalized Tremor Syndrome - English Springer Spaniel Variant | PLP1 | 0 | Clear |
| arcol | epsy | Gene | Copies | Results |
| Ø | Narcolepsy - Dachshund Variant | HCRTR2 | 0 | Clear |
| | Narcolepsy - Doberman Pinscher Variant | HCRTR2 | 0 | Clear |
| ⊘ | Narcolepsy - Labrador Retriever Variant | HCRTR2 | 0 | Clear |
| euro | degenerative Disorder | Gene | Copies | Results |
| ⊘ | Fetal-Onset Neonatal Neuroaxonal Dystrophy - Giant Schnauzer Variant | MFN2 | 0 | Clear |
| Ø | Neuroaxonal Dystrophy, NAD - Rottweiler Variant | VPS11 | 0 | Clear |
| ⊘ | Neuroaxonal Dystrophy, NAD - Spanish Water Dog Variant | TECPR2 | 0 | Clear |

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| Neuropathy | Gene | Copies | Results |
|--|------------------|--------|---------|
| Alaskan Malamute Polyneuropathy, AMPN | NDRG1 | 0 | Clear |
| Demyelinating Polyneuropathy - Miniature Schnauzer Variant | SBF2/MTRM13 | 0 | Clear |
| Juvenile Laryngeal Paralysis and Polyneuropathy, Polyneuropathy with Ocular Abnormalities and Neuronal Vacuolation, POANV - Rottweiler Variant | RAB3GAP1 | 0 | Clear |
| Juvenile-Onset Polyneuropathy, Leonberger Polyneuropathy 1, LPN1 | ARHGEF10 Exon 17 | 0 | Clear |
| Juvenile-Onset Polyneuropathy, Leonberger Polyneuropathy 2, LPN2 | GJA9 | 0 | Clear |
| Laryngeal Paralysis - Miniature Bull Terrier Variant | RAPGEF6 | 0 | Clear |
| Sensory Neuropathy | Gene | Copies | Results |
| Hereditary Sensory Autonomic Neuropathy, Acral Mutilation Syndrome, AMS - Spaniel Pointer Variant | and GDNF-AS | 0 | Clear |
| Sensory Neuropathy - Border Collie Variant | FAM134B | 0 | Clear |
| Neuromuscular (7) | | | |
| Junctionopathy | Gene | Copies | Results |

| Junctionopathy | Gene | Copies | Results |
|---|--------------|--------|---------|
| Congenital Myasthenic Syndrome, CMS - Golden Retriever Variant | COLQ Exon 13 | 0 | Clear |
| Congenital Myasthenic Syndrome, CMS - Heideterrier Variant | CHRNE | 0 | Clear |
| Congenital Myasthenic Syndrome, CMS - Jack Russell Terrier Variant | CHRNE Exon 7 | 0 | Clear |
| Congenital Myasthenic Syndrome, CMS - Labrador Retriever Variant | COLQ Exon 14 | 0 | Clear |
| Congenital Myasthenic Syndrome, CMS - Old Danish Pointing Dog Variant | CHAT Exon 6 | 0 | Clear |

| Movement Disorder | Gene | Copies | Results |
|---|---------------|--------|---------|
| Episodic Falling Syndrome - Cavalier King Charles Spaniel Variant B | CAN Exons 1-4 | 0 | Clear |
| Paroxysmal Dyskinesia, PxD - Soft Coated Wheaten Terrier Variant | PIGN | 0 | Clear |
| Ophthalmologic (31) | | | |
| Glaucoma | Gene | Copies | Results |
| Goniodysgenesis and Glaucoma, Pectinate Ligament Dysplasia, PLD - Border Collie Varia | nt OLFML3 | 0 | Clear |
| Primary Open Angle Glaucoma and Primary Lens Luxation - Chinese Shar-Pei Variant | ADAMTS17 | 0 | Clear |
| Primary Open Angle Glaucoma - Basset Fauve de Bretagne Variant | ADAMTS17 | 0 | Clear |
| Primary Open Angle Glaucoma - Beagle Variant | ADAMTS10 | 0 | Clear |
| Primary Open Angle Glaucoma - Norwegian Elkhound Variant | ADAMTS10 | 0 | Clear |
| Iris or Lens | Gene | Copies | Results |
| Hereditary Cataracts, Early-Onset Cataracts, Juvenile Cataracts - Australian Shepherd Va | nriant HSF4 | 0 | Clear |
| Primary Lens Luxation | ADAMTS17 | 0 | Clear |
| Retinopathy | Gene | Copies | Results |
| Achromatopsia - German Shepherd Variant | CNGA3 Exon 7 | 0 | Clear |
| Achromatopsia - Labrador Retriever Variant | CNGA3 Exon 7 | 0 | Clear |
| Autosomal Dominant Progressive Retinal Atrophy - English Mastiff and Bullmastiff Variant | RHO Exon 1 | 0 | Clear |
| Canine Multifocal Retinopathy, cmr1 BEST1 | I/VMD2 Exon 2 | 0 | Clear |



| | Gene | Copies | Results |
|---|--------------------|--------|---------|
| Canine Multifocal Retinopathy, cmr2 - Coton de Tulear Variant | BEST1/VMD2 Exon 5 | 0 | Clear |
| Canine Multifocal Retinopathy, cmr3 - Finnish and Swedish Lapphund, Lapponian Herder Variant | BEST1/VMD2 Exon 10 | 0 | Clear |
| Collie Eye Anomaly, Choroidal Hypoplasia, CEA | NHEJ1 Intron 4 | 0 | Clear |
| Congenital Stationary Night Blindness - Beagle Variant | LRIT3 | 0 | Clear |
| Congenital Stationary Night Blindness - Briard Variant | RPE65 | 0 | Clear |
| Oay Blindness, Cone Degeneration, Achromatopsia - Alaskan Malamute Variant | CNGB3 Deletion | 0 | Clear |
| Day Blindness, Cone Degeneration, Achromatopsia - German Shorthaired Pointe Variant | er CNGB3 Exon 6 | 0 | Clear |
| Golden Retriever Progressive Retinal Atrophy 1, GR-PRA1 | SLC4A3 Exon 16 | 0 | Clear |
| Golden Retriever Progressive Retinal Atrophy 2, GR-PRA2 | TTC8 Exon 8 | 0 | Clear |
| Macular Corneal Dystrophy, MCD - Labrador Retriever Variant | CHST6 | 0 | Clear |
| Progressive Retinal Atrophy, CNGA - Shetland Sheepdog Variant | CNGA1 Exon 9 | 0 | Clear |
| Progressive Retinal Atrophy, Cone-Rod Dystrophy 1, crd1 - American Staffordshill Variant | re Terrier PDE6B | 0 | Clear |
| Progressive Retinal Atrophy, Cone-Rod Dystrophy 4, crd4/cord1 | RPGRIP1 Exon 2 | 0 | Clear |
| Progressive Retinal Atrophy, PRA1 - Papillon Variant | CNGB1 | 0 | Clear |
| Progressive Retinal Atrophy, PRA3 - Tibetan Spaniel and Terrier Variant | FAM161A | 0 | Clear |
| Progressive Retinal Atrophy, Progressive Rod-Cone Degeneration, prcd | PRCD Exon 1 | 0 | Clear |
| Progressive Retinal Atrophy, Rod-Cone Dysplasia 1, rcd1 - Irish Setter Variant | PDE6B Exon 21 | 0 | Clear |
| Progressive Retinal Atrophy, Rod-Cone Dysplasia 3, rcd3 - Corgi Variant | PDE6A | 0 | Clear |

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| | Gene | Copies | Results |
|--|--------------|--------|---------|
| Progressive Retinal Atrophy - Basenji Variant | SAG | 0 | Clear |
| X-Linked Progressive Retinal Atrophy 1, XL-PRA1 - Samoyed and Husky Variant | RPGR Exon 15 | 0 | Clear |
| Oral Cavity (4) | | | |
| Developmental Disorder | Gene | Copies | Results |
| Cleft Lip and/or Cleft Palate - Nova Scotia Duck Tolling Retriever Variant | ADAMTS20 | 0 | Clear |
| Tooth Structure Defect | Gene | Copies | Results |
| Autosomal Recessive Amelogenesis Imperfecta, Familial Enamel Hypoplasia - Italian Greyhound Variant | ENAM | 0 | Clear |
| Autosomal Recessive Amelogenesis Imperfecta, Familial Enamel Hypoplasia - Parson Ru Terrier Variant | ssell ENAM | 0 | Clear |
| Raine Syndrome, Canine Dental Hypomineralization Syndrome - Border Collie Variant | FAM20C | 0 | Clear |
| Personalized Medicine (3) | | | |
| | Gene | Copies | Results |
| Alanine Aminotransferase Activity | GPT | 0 | Clear |
| MDR1 Drug Sensitivity | ABCB1 | 0 | Clear |
| Malignant Hyperthermia | RYR1 | 0 | Clear |
| Pulmonary (4) | | | |

Neonatal Interstitial Lung Disease - Airedale Terrier Variant

Results

Clear

Copies

0

Gene

LAMP3



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| | Gene | Copies | Results |
|---|--------|--------|---------|
| Primary Ciliary Dyskinesia, PCD - Alaskan Malamute Variant | NME5 | 0 | Clear |
| Primary Ciliary Dyskinesia, PCD - Old English Sheepdog Variant | CCDC39 | 0 | Clear |
| Recurrent Inflammatory Pulmonary Disease, RIPD - Rough Collie Variant | AKNA | 0 | Clear |

| Skeletal (10) | | | | | | | |
|-------------------------------|--|-----------------|--------|---------|--|--|--|
| Chondrodystrophy | | Gene | Copies | Results | | | |
| Chondrodystrophy Retrogene | and Intervertebral Disc Disease, CDDY/IVDD, Type I IVDD - | FGF4 - chr12 | 0 | Clear | | | |
| Chondrodystrophy | r - Norwegian Elkhound and Karelian Bear Dog Variant | ITGA10 | 0 | Clear | | | |
| Oculoskeletal Dysp Variant | olasia 2, Dwarfism-Retinal Dysplasia 2, drd2, OSD2 - Samoyed | COL9A2 5' UTR | 0 | Clear | | | |
| Osteochondrodysp | olasia, Skeletal Dwarfism - Miniature Poodle Variant | SLC13A1 | 0 | Clear | | | |
| Skeletal Dysplasia | 2, SD2 - Labrador Retriever Variant | COL11A2 | 0 | Clear | | | |
| Decreased Bone Strength | | Gene | Copies | Results | | | |
| Hereditary Vitamin | D-Resistant Rickets - Pomeranian Variant | VDR Exon 4 | 0 | Clear | | | |
| Osteogenesis Imp | erfecta, Brittle Bone Disease - Beagle Variant | COL1A2 | 0 | Clear | | | |
| Osteogenesis Imp | erfecta, Brittle Bone Disease - Dachshund Variant | SERPINH1 Exon 5 | 0 | Clear | | | |
| Osteogenesis Imp | erfecta, Brittle Bone Disease - Golden Retriever Variant | COL1A1 Exon 18 | 0 | Clear | | | |
| Other | | Gene | Copies | Results | | | |

Craniomandibular Osteopathy, CMO - Terrier and Australian Shepherd Variant

SLC37A2 Exon 15

Clear

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Urogenital (14)

| Nephro | ppathy | Gene | Copies | Results |
|----------|--|----------------|--------|---------|
| Ø | Autosomal Recessive Hereditary Nephropathy, Familial Nephropathy, ARHN - Cocker Spaniel Variant | COL4A4 Exon 3 | 0 | Clear |
| | Autosomal Recessive Hereditary Nephropathy, Familial Nephropathy, ARHN - English Springer Spaniel Variant | COL4A4 Exon 30 | 0 | Clear |
| | Fanconi Syndrome - Basenji Variant | FAN1 | 0 | Clear |
| Ø | Polycystic Kidney Disease, PKD - Bull Terrier Variant | PKD1 Exon 29 | 0 | Clear |
| Ø | Protein Losing Nephropathy, PLN - Soft Coated Wheaten and Airedale Terrier Variant | NPHS1 | 0 | Clear |
| ⊘ | X-Linked Hereditary Nephropathy, XLHN - Samoyed Variant 2 | COL4A5 Exon 35 | 0 | Clear |
| Urolith | iasis | Gene | Copies | Results |
| Ø | 2,8-Dihydroxyadenine Urolithiasis, 2,8-DHA Urolithiasis - American Indian Dog Variant | APRT Exon 3 | 0 | Clear |
| | Cystinuria Type I-A - Newfoundland Variant | SLC3A1 Exon 2 | 0 | Clear |
| Ø | Cystinuria Type II-A - Australian Cattle Dog Variant | SLC3A1 Exon 6 | 0 | Clear |
| | Cystinuria Type II-B - Miniature Pinscher Variant | SLC7A9 Exon 9 | 0 | Clear |
| Ø | Hyperuricosuria and Hyperuricemia or Urolithiasis, HUU | SLC2A9 Exon 5 | 0 | Clear |
| ⊘ | Primary Hyperoxaluria - Coton de Tulear Variant | AGXT Exon 2 | 0 | Clear |
| Other | | Gene | Copies | Results |
| | Persistent Mullerian Duct Syndrome, PMDS - Miniature and Standard Schnauzer Varia | nt AMHR2 | 0 | Clear |
| Ø | Renal Cystadenocarcinoma and Nodular Dermatofibrosis, RCND - German Shepherd D Variant | og FLCN | 0 | Clear |

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Glossary

Key Terms

Increased Risk Result

The dog is at risk for showing clinical signs (phenotype) of a given condition. For recessive conditions, this means a dog has inherited two copies of an associated variant. For dominant, codominant, and additive conditions, this means a dog has inherited at least one copy of the variant. X-linked conditions will vary based on sex of the dog.

A dog's breed(s) and genetic background are also considered in this assessment. Genetic testing is an assessment of risk and not a clinical diagnosis, and not all dogs in this category will develop clinical signs.

Notable Result

A result may be notable for several reasons. The variant may not induce a disease state but rather inform patient care (this may include the tests listed under Personalized Medicine). The dog may have only one copy of a variant with a recessive mode of inheritance (meaning the dog is a carrier and is not expected to show the phenotype associated with the variant). The impact of the variant may also be influenced by a dog's breed(s). Based on the available research within the breed or related breeds, you will see more specific text within the results.

Clear Result

A dog with two healthy copies of a gene sequence is not at risk for developing the associated disease due to that variant. Many diseases can manifest as a result of other unknown genetic variants and/or environmental factors.

Variant

An alteration in the DNA with the potential to cause a change in phenotype (i.e. disease). A report may state that the dog has zero, one, or two copies of the variant for which we test. The term "variant" may be used interchangeably with "mutation."

Genotype

The genetic code related to the variant being present or absent in the dog's DNA.

Phenotype

The physical impact or appearance directed by the genotype. The phenotype is often described as an expression of the genotype.

Complex Phenotype

The condition, appearance, or other physical expression of the genotype controlled by both genetic and environmental factors.

Penetrance

Proportion of dogs with a particular genotype that expresses the associated phenotype. There are two types of penetrance.

- 1. Incomplete penetrance means that not all dogs with the genotype will develop the clinical signs of the phenotype.
- 2. Complete penetrance means that all dogs with the genotype will develop the clinical signs of the phenotype.

Carrier

This term has traditionally been used to describe a dog that has one copy of the variant but is not expected to show the phenotype associated with the variant (this is applicable to variants with a recessive mode of inheritance (MOI) as described below). If used in a breeding pair, a carrier may pass the variant to its litter.

At-risk

This indicates that the dog may manifest the disease and generally is used when a dog has two copies of the variant (but this depends on the MOI).

Embark uses the term "at-risk" and not "affected" because genetic testing is an assessment of risk and not a clinical diagnosis.

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Linkage Disequilibrium Test

When a causal variant cannot be identified or when the variant is incompatible with the genotyping platform constraints, allelic association or linkage disequilibrium (LD) tests can be utilized. This is typically done to assist dog breeders in selectively breeding out a deleterious condition. LD tests are based on a statistical association between two loci that are physically very close in the DNA. The coupling of the chosen proxy marker to the causal variant is known mathematically for the most relevant populations.

LD-based tests have a slightly increased incidence of false positives and false negatives, which are test-specific and known. Embark offers limited numbers of these tests. Embark continuously works to refine LD-based tests by assaying the direct variant in a subset of dogs using alternative methods. These inputs help to refine the tests over time.

Provisional Result

Embark combines random sampling and sequencing with the use of blinded controls to confirm that each test is performing to standard at >99% genotyping accuracy and reproducibility. Our standard health tests have been validated using known heterozygous and homozygous samples to ensure design accuracy and use multiple probes per condition to ensure reproducibility. Provisional tests are for rare disorders for which DNA samples from carrier and/or at-risk individuals are not available for calculating test reliability, or for structural variants where more testing is needed to ensure the same level of accuracy.

If you have access to DNA from carrier or at-risk individuals and are interested in helping us validate a test, please contact us at vetsupport@embarkvet.com

Modes of Inheritance

Recessive

A dog is thought to need two copies of a variant to be considered at-risk for the clinical disease or to have the visible phenotype for traits. This may apply to autosomal or X-linked variants, however. Read below for additional details regarding X-linked variants.

Dominant

A dog is thought to need only one copy of the variant to be considered at-risk for the clinical disease or to have the visible phenotype for traits.

Codominant/Additive

In general, these terms are used to describe variants in which dogs with one copy of the variant have a different phenotype compared to dogs with zero or two copies of the variant (although there is a slight difference between the two terms).

X-linked

The variant resides on the X chromosome, and male dogs need just one copy of the variant to be considered at-risk. For recessively inherited X-linked conditions, female dogs typically require two copies of the variant to be considered at-risk. Female dogs who have one copy of a recessively inherited X-linked variant are often referred to as carriers, but they can exhibit signs of disease that range from clinically asymptomatic to fully affected. This is due to a normal phenomenon known as X-chromosome inactivation, where one X chromosome is silenced in each cell.

Weight

The Embark DNA test provides a genetic size based not just on breed ancestry but on over a dozen genes known to influence a dog's weight, as well as sex and breed-specific modifiers.

Our algorithm explains over 85% of the variance in healthy adult weight. However, due to a few as-yet-undiscovered genes and genetic interactions that affect size, this algorithm sometimes under or over-predicts weight.

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Genetic Age

Dogs age at very different rates due to a number of genetic and environmental factors. Embark's genetic age calculates how old a dog would be if he or she were aging at an average human rate (using humans in the USA as the baseline). This measure is more personalized than "one dog year = seven human years".

View the patient's profile see the personalized genetic age table for this dog.

We start by asking the dog's approximate calendar age. We then calculate genetic age by factoring a dog's breed composition along with information from genes that affect size, sex, and the dog's inbreeding coefficient (COI).

Impact of Breed

When determining whether or not a variant is expected to have a clinical impact for a breed, we have taken into account research either published, internal, or otherwise presented by a subject matter authority as our primary criteria. So, while a dog may have the variant associated with a disease (one or two copies for dominant variants and two copies for autosomal recessive variants), he or she may not be known to be at significant clinical risk from that variant.

Based on the available research within the breed or highly related breeds, you may see text similar to the following options:

- 1. This genetic variant is not likely to significantly increase the risk that this dog will develop the clinical disease.
- 2. This genetic variant is associated with an increased risk that this dog will develop the clinical disease.
- 3. We do not know whether this variant increases the risk that this dog will develop the clinical disease.

Embark is continuing to explore the relationship of genotype to phenotype, and risk assessment may be updated as more data is reviewed. You can contact vetsupport@embarkvet.com or call 1-855-203-8271 to report any clinical diagnoses.