ORTHOPEDIC	FOUNDATION FOR	ANIMALS, INC.			
TIMBERVIEW TYLER		AB1C390B01 gistration no.			
POODLE, MINIATURE	М				
film/test/lab #	dat	8/17/2021 te of birth	OFA		
tattoo/microchip/DNA profile		} e at evaluation in months	A Not-For-Profit Organization		
2362424 application number	P	O-BCA2642/13M/P-N0			
05/23/2022 date of report	Th	F.A. NUMBER is number issued with the right voke by the Orthopedic Founda	to correct or		
RESULTS:		, ,			
Normal cardiovascular examination via auscultation - No evidence of congenital or acquired heart disease was noted. Since acquired heart disease may develop later, these evaluation results remain valid for one year, and annual examinations are recommended to continue to monitor cardiac health.					
		ORMAL/CLEAR - PRA	ACTITIONER		
DUANE OTTO	OFA eCert	AAK	eller DIM		
ổ 351 E CR 100 N ARCOLA IL 61910			D.V.M., M.S., DACVR ERINARY SERVICES		
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	www.ofa.org				

This electronic OFA certificate was generated on: 05/23/2022

This certification can be verified on the OFA website by entering the dog's registration number into the orange search box located at the top of the page or by scanning the QR code above.

If there are any errors on this certificate, please email CORRECTIONS@OFFA.ORG to request a correction.

Orthopedic Foundation for Animals, Inc. 2300 E. Nifong Blvd. Columbia, MO 65201-3806

OFA website: www.ofa.org E-mail address: ofa@offa.org Phone number: 573-442-0418 Fax number: 573-875-5073

ORTHOPEDIC FOUNDATION FOR ANIMALS, INC.

TIMBERVIEW TYLER registered name

POODLE, MINIATURE *sex/breed*

film/test/lab #

tattoo/microchip/DNA profile

2362424 application number

05/23/2022 date of report

RESULTS:

owner

The results of the examination submitted to OFA indicate that no evidence of patellar luxation was recognized.

NORMAL - PRACTITIONER

UAB1C390B01

registration no.

03/17/2021

age at evaluation in months

date of birth

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G.G.KELLER. D.V.M., M.S., DACVR CHIEF OF VETERINARY SERVICES

DUANE OTTO 351 E CR 100 N

ARCOLA IL 61910



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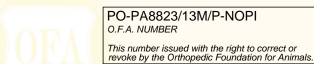
If there are any errors on this certificate, please email CORRECTIONS@OFFA.ORG to request a correction.

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A Not-For-Profit Organization









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BREED ANCESTRY

Poodle (Small) : 100.0%

GENETIC STATS

Predicted adult weight: **21 lbs** Genetic age: n/a (Date of birth unknown)

TEST DETAILS

Kit number: EM-38581636 Swab number: 31211051900912





Test Date: June 13th, 2022

Rembark

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POODLE (SMALL)

Miniature and toy poodles are varieties of the poodle breed which originated in Germany in the 15th century. Unlike the larger standard poodle (>15 inches tall), these small poodles were not developed for hunting---except for truffles!---and were generally used as lap dogs and companions. Small poodles are frequently used to create designer dogs like Schnoodles and Maltipoos with low-shedding, hypoallergenic coats. All poodles are highly intelligent and energetic, and need daily exercise and stimulation. They are overall healthy dogs, although heritable eye disease, epilepsy and allergies are relatively common, and toy poodles also have a heightened risk of accidents/trauma due to their small size.

Alternative Names Toy Poodle, Miniature Poodle

Fun Fact

Although Toy Poodles are the most popular dog breed in Japan, Poodles as a group are the eight most popular breed in the US, with miniature poodles being the most common variety.







Test Date: June 13th, 2022

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MATERNAL LINE



Through Tyler's mitochondrial DNA we can trace his mother's ancestry back to where dogs and people first became friends. This map helps you visualize the routes that his ancestors took to your home. Their story is described below the map.

HAPLOGROUP: B1

B1 is the second most common maternal lineage in breeds of European or American origin. It is the female line of the majority of Golden Retrievers, Basset Hounds, and Shih Tzus, and about half of Beagles, Pekingese and Toy Poodles. This lineage is also somewhat common among village dogs that carry distinct ancestry from these breeds. We know this is a result of B1 dogs being common amongst the European dogs that their conquering owners brought around the world, because nowhere on earth is it a very common lineage in village dogs. It even enables us to trace the path of (human) colonization: Because most Bichons are B1 and Bichons are popular in Spanish culture, B1 is now fairly common among village dogs in Latin America.

HAPLOTYPE: B74

Part of the large B1 haplogroup, this haplotype occurs most frequently in mixed breed dogs.







Test Date: June 13th, 2022

embk.me/tyler143

PATERNAL LINE



Through Tyler's Y chromosome we can trace his father's ancestry back to where dogs and people first became friends. This map helps you visualize the routes that his ancestors took to your home. Their story is described below the map.

HAPLOGROUP: A1b

For most of dog history, this haplogroup was probably quite rare. However, a couple hundred years ago it seems to have found its way into a prized male guard dog in Europe who had many offspring, including the ancestors of many European guard breeds such as Doberman Pinchers, St. Bernards, and Great Danes. Despite being rare, many of the most imposing dogs on Earth have it; strangely, so do many Pomeranians! Perhaps this explains why some Poms are so tough, acting like they're ten times their actual size! This lineage is most commonly found in working dogs, in particular guard dogs. With origins in Europe, it spread widely across other regions as Europeans took their dogs across the world.

HAPLOTYPE: Ha.7

Part of the A1b haplogroup, this haplotype is found in village dogs from Lebanon and Indonesia. Among breeds, it is also found in Miniature Schnauzer and Toy Poodle.







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RESULT

TRAITS: COAT COLOR

TRAIT

E Locus (MC1R)

The E Locus determines if and where a dog can produce dark (black or brown) hair. Dogs with two copies of the recessive **e** allele do not produce dark hairs at all, and will be "red" over their entire body. The shade of red, which can range from a deep copper to yellow/gold to cream, is dependent on other genetic factors including the Intensity loci. In addition to determining if a dog can develop dark hairs at all, the E Locus can give a dog a black "mask" or "widow's peak," unless the dog has overriding coat color genetic factors. Dogs with one or two copies of the **Em** allele usually have a melanistic mask (dark facial hair as commonly seen in the German Shepherd and Pug). Dogs with no copies of **Em** but one or two copies of the **Eg** allele usually have a melanistic "widow's peak" (dark forehead hair as commonly seen in the Afghan Hound and Borzoi, where it is called either "grizzle" or "domino").

No dark hairs anywhere (ee)

K Locus (CBD103)

The K Locus **K^B** allele "overrides" the A Locus, meaning that it prevents the A Locus genotype from affecting coat color. For this reason, the **K^B** allele is referred to as the "dominant black" allele. As a result, dogs with at least one **K^B** allele will usually have solid black or brown coats (or red/cream coats if they are **ee** at the E Locus) regardless of their genotype at the A Locus, although several other genes could impact the dog's coat and cause other patterns, such as white spotting. Dogs with the **k**^y**k**^y genotype will show a coat color pattern based on the genotype they have at the A Locus. Dogs who test as **K^Bk**^y may be brindle rather than black or brown.

Not expressed (k^yk^y)







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RESULT

TRAITS: COAT COLOR (CONTINUED)

TRAIT

Intensity Loci LINKAGE

Areas of a dog's coat where dark (black or brown) pigment is not expressed either contain red/yellow pigment, or no pigment at all. Five locations across five chromosomes explain approximately 70% of red pigmentation "intensity" variation across all dogs. Dogs with a result of **Intense Red Pigmentation** will likely have deep red hair like an Irish Setter or "apricot" hair like some Poodles, dogs with a result of **Intermediate Red Pigmentation** will likely have tan or yellow hair like a Soft-Coated Wheaten Terrier, and dogs with **Dilute Red Pigmentation** will likely have cream or white hair like a Samoyed. Because the mutations we test may not directly cause differences in red pigmentation intensity, we consider this to be a linkage test.

Any pigmented hair likely apricot or red (Intense Red Pigmentation)

A Locus (ASIP)

The A Locus controls switching between black and red pigment in hair cells, but it will only be expressed in dogs that are not **ee** at the E Locus and are $\mathbf{k}^{\mathbf{y}}\mathbf{k}^{\mathbf{y}}$ at the K Locus. Sable (also called "Fawn") dogs have a mostly or entirely red coat with some interspersed black hairs. Agouti (also called "Wolf Sable") dogs have red hairs with black tips, mostly on their head and back. Black and tan dogs are mostly black or brown with lighter patches on their cheeks, eyebrows, chest, and legs. Recessive black dogs have solid-colored black or brown coats.

D Locus (MLPH)

The D locus result that we report is determined by two different genetic variants that can work together to cause diluted pigmentation. These are the common **d** allele, also known as "**d1**", and a less common allele known as "**d2**". Dogs with two **d** alleles, regardless of which variant, will have all black pigment lightened ("diluted") to gray, or brown pigment lightened to lighter brown in their hair, skin, and sometimes eyes. There are many breed-specific names for these dilute colors, such as "blue", "charcoal", "fawn", "silver", and "Isabella". Note that in certain breeds, dilute dogs have a higher incidence of Color Dilution Alopecia. Dogs with one **d** allele will not be dilute, but can pass the **d** allele on to their puppies. To view your dog's **d1** and **d2** test results, click the "SEE DETAILS" link in the upper right hand corner of the "Base Coat Color" section of the Traits page, and then click the "VIEW SUBLOCUS RESULTS" link at the bottom of the page.

Not expressed (a^ya^t)

Not expressed (DD)





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RESULT

TRAITS: COAT COLOR (CONTINUED)

TRAIT

Cocoa (HPS3)

Dogs with the coco genotype will produce dark brown pigment instead of black in both their hair and skin.No co alleles, notDogs with the Nco genotype will produce black pigment, but can pass the co allele on to their puppies.expressed (NN)Dogs that have the coco genotype as well as the bb genotype at the B locus are generally a lighter brownthan dogs that have the Bb or BB genotypes at the B locus.

B Locus (TYRP1)

Dogs with two copies of the **b** allele produce brown pigment instead of black in both their hair and skin. Dogs with one copy of the **b** allele will produce black pigment, but can pass the **b** allele on to their puppies. E Locus **ee** dogs that carry two **b** alleles will have red or cream coats, but have brown noses, eye rims, and footpads (sometimes referred to as "Dudley Nose" in Labrador Retrievers). "Liver" or "chocolate" is the preferred color term for brown in most breeds; in the Doberman Pinscher it is referred to as "red".

Likely black colored nose/feet (BB)

Saddle Tan (RALY)

The "Saddle Tan" pattern causes the black hairs to recede into a "saddle" shape on the back, leaving a tan face, legs, and belly, as a dog ages. The Saddle Tan pattern is characteristic of breeds like the Corgi, Beagle, and German Shepherd. Dogs that have the **II** genotype at this locus are more likely to be mostly black with tan points on the eyebrows, muzzle, and legs as commonly seen in the Doberman Pinscher and the Rottweiler. This gene modifies the A Locus **a**^t allele, so dogs that do not express **a**^t are not influenced by this gene.

•

Not expressed (NI)

S Locus (MITF)

The S Locus determines white spotting and pigment distribution. MITF controls where pigment is produced, and an insertion in the MITF gene causes a loss of pigment in the coat and skin, resulting in white hair and/or pink skin. Dogs with two copies of this variant will likely have breed-dependent white patterning, with a nearly white, parti, or piebald coat. Dogs with one copy of this variant will have more limited white spotting and may be considered flash, parti or piebald. This MITF variant does not explain all white spotting patterns in dogs and other variants are currently being researched. Some dogs may have small amounts of white on the paws, chest, face, or tail regardless of their S Locus genotype.

Likely to have little to no white in coat (SS)







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RESULT

TRAITS: COAT COLOR (CONTINUED)

TRAIT

M Locus (PMEL)

Merle coat patterning is common to several dog breeds including the Australian Shepherd, Catahoula Leopard Dog, and Shetland Sheepdog, among many others. Merle arises from an unstable SINE insertion (which we term the "M*" allele) that disrupts activity of the pigmentary gene PMEL, leading to mottled or patchy coat color. Dogs with an **M*m** result are likely to be phenotypically merle or could be "non-expressing" merle, meaning that the merle pattern is very subtle or not at all evident in their coat. Dogs with an **M*M*** result are likely to be phenotypically merle. Dogs with an **mm** result have no merle alleles and are unlikely to have a merle coat pattern.

Note that Embark does not currently distinguish between the recently described cryptic, atypical, atypical+, classic, and harlequin merle alleles. Our merle test only detects the presence, but not the length of the SINE insertion. We do not recommend making breeding decisions on this result alone. Please pursue further testing for allelic distinction prior to breeding decisions.

R Locus (USH2A) LINKAGE

The R Locus regulates the presence or absence of the roan coat color pattern. Partial duplication of the USH2A gene is strongly associated with this coat pattern. Dogs with at least one **R** allele will likely have roaning on otherwise uniformly unpigmented white areas. Roan appears in white areas controlled by the S Locus but not in other white or cream areas created by other loci, such as the E Locus with **ee** along with Dilute Red Pigmentation by I Locus (for example, in Samoyeds). Mechanisms for controlling the extent of roaning are currently unknown, and roaning can appear in a uniform or non-uniform pattern. Further, non-uniform roaning may appear as ticked, and not obviously roan. The roan pattern can appear with or without ticking.

Likely no impact on coat pattern (rr)

No merle alleles (mm)

H Locus (Harlequin)

This pattern is recognized in Great Danes and causes dogs to have a white coat with patches of darker pigment. A dog with an **Hh** result will be harlequin if they are also **M*m** or **M*M*** at the M Locus and are not **ee** at the E locus. Dogs with a result of **hh** will not be harlequin. This trait is thought to be homozygous lethal; a living dog with an **HH** genotype has never been found.

No harlequin alleles (hh)







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RESULT

TRAITS: OTHER COAT TRAITS

TRAIT

Furnishings (RSPO2) LINKAGE

Dogs with one or two copies of the F allele have "furnishings": the mustache, beard, and eyebrows Likely furnished characteristic of breeds like the Schnauzer, Scottish Terrier, and Wire Haired Dachshund. A dog with two I (mustache, beard, alleles will not have furnishings, which is sometimes called an "improper coat" in breeds where and/or eyebrows) (FF) furnishings are part of the breed standard. The mutation is a genetic insertion which we measure indirectly using a linkage test highly correlated with the insertion.

Coat Length (FGF5)

The FGF5 gene is known to affect hair length in many different species, including cats, dogs, mice, and humans. In dogs, the T allele confers a long, silky haircoat as observed in the Yorkshire Terrier and the Long Haired Whippet. The ancestral G allele causes a shorter coat as seen in the Boxer or the American Staffordshire Terrier. In certain breeds (such as Corgi), the long haircoat is described as "fluff."

Likely long coat (TT)

Shedding (MC5R)

Dogs with at least one copy of the ancestral C allele, like many Labradors and German Shepherd Dogs, are heavy or seasonal shedders, while those with two copies of the T allele, including many Boxers, Shih Tzus (CC) and Chihuahuas, tend to be lighter shedders. Dogs with furnished/wire-haired coats caused by RSPO2 (the furnishings gene) tend to be low shedders regardless of their genotype at this gene.

Hairlessness (FOXI3) LINKAGE

A duplication in the FOXI3 gene causes hairlessness over most of the body as well as changes in tooth shape and number. This mutation occurs in Peruvian Inca Orchid, Xoloitzcuintli (Mexican Hairless), and Chinese Crested (other hairless breeds have different mutations). Dogs with the NDup genotype are likely to be hairless while dogs with the NN genotype are likely to have a normal coat. The DupDup genotype has never been observed, suggesting that dogs with that genotype cannot survive to birth. Please note that this is a linkage test, so it may not be as predictive as direct tests of the mutation in some lines.

Hairlessness (SGK3)

Very unlikely to be Hairlessness in the American Hairless Terrier arises from a mutation in the SGK3 gene. Dogs with the DD hairless (NN) result are likely to be hairless. Dogs with the ND genotype will have a normal coat, but can pass the D

Likely light shedding

Very unlikely to be hairless (NN)

Kembark





RESULT

TRAITS: OTHER COAT TRAITS (CONTINUED)

TRAIT

Oculocutaneous Albinism Type 2 (SLC45A2) LINKAGE

Dogs with two copies **DD** of this deletion in the SLC45A2 gene have oculocutaneous albinism (OCA), also known as Doberman Z Factor Albinism, a recessive condition characterized by severely reduced or absent pigment in the eyes, skin, and hair. Affected dogs sometimes suffer from vision problems due to lack of eye pigment (which helps direct and absorb ambient light) and are prone to sunburn. Dogs with a single copy of the deletion **ND** will not be affected but can pass the mutation on to their offspring. This particular mutation can be traced back to a single white Doberman Pinscher born in 1976, and it has only been observed in dogs descended from this individual. Please note that this is a linkage test, so it may not be as predictive as direct tests of the mutation in some lines.

Coat Texture (KRT71)

Dogs with a long coat and at least one copy of the **T** allele have a wavy or curly coat characteristic of Poodles and Bichon Frises. Dogs with two copies of the ancestral **C** allele are likely to have a straight coat, but there are other factors that can cause a curly coat, for example if they at least one **F** allele for the Furnishings (RSPO2) gene then they are likely to have a curly coat. Dogs with short coats may carry one or two copies of the **T** allele but still have straight coats.





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RESULT

TRAITS: OTHER BODY FEATURES

TRAIT

Muzzle Length (BMP3)

Dogs in medium-length muzzle (mesocephalic) breeds like Staffordshire Terriers and Labradors, and long muzzle (dolichocephalic) breeds like Whippet and Collie have one, or more commonly two, copies of the ancestral **C** allele. Dogs in many short-length muzzle (brachycephalic) breeds such as the English Bulldog, Pug, and Pekingese have two copies of the derived **A** allele. At least five different genes affect muzzle length in dogs, with BMP3 being the only one with a known causal mutation. For example, the skull shape of some breeds, including the dolichocephalic Scottish Terrier or the brachycephalic Japanese Chin, appear to be caused by other genes. Thus, dogs may have short or long muzzles due to other genetic factors that are not yet known to science.

Likely medium or long muzzle (CC)

Tail Length (T)

Whereas most dogs have two **C** alleles and a long tail, dogs with one **G** allele are likely to have a bobtail, which is an unusually short or absent tail. This mutation causes natural bobtail in many breeds including the Pembroke Welsh Corgi, the Australian Shepherd, and the Brittany Spaniel. Dogs with **GG** genotypes have not been observed, suggesting that dogs with the **GG** genotype do not survive to birth. Please note that this mutation does not explain every natural bobtail! While certain lineages of Boston Terrier, English Bulldog, Rottweiler, Miniature Schnauzer, Cavalier King Charles Spaniel, and Parson Russell Terrier, and Dobermans are born with a natural bobtail, these breeds do not have this mutation. This suggests that other unknown genetic mutations can also lead to a natural bobtail.

Hind Dewclaws (LMBR1)

Common in certain breeds such as the Saint Bernard, hind dewclaws are extra, nonfunctional digits located midway between a dog's paw and hock. Dogs with at least one copy of the **T** allele have about a 50% chance of having hind dewclaws. Note that other (currently unknown to science) mutations can also cause hind dewclaws, so some **CC** or **TC** dogs will have hind dewclaws. Likely normal-length tail (CC)

Unlikely to have hind dew claws (CC)

Fembark





TRAITS: OTHER BODY FEATURES (CONTINUED)

TRAIT

Blue Eye Color (ALX4) LINKAGE

Embark researchers discovered this large duplication associated with blue eyes in Arctic breeds like Siberian Husky as well as tri-colored (non-merle) Australian Shepherds. Dogs with at least one copy of the duplication (**Dup**) are more likely to have at least one blue eye. Some dogs with the duplication may have only one blue eye (complete heterochromia) or may not have blue eyes at all; nevertheless, they can still pass the duplication and the trait to their offspring. **NN** dogs do not carry this duplication, but may have blue eyes due to other factors, such as merle. Please note that this is a linkage test, so it may not be as predictive as direct tests of the mutation in some lines.

Back Muscling & Bulk, Large Breed (ACSL4)

The **T** allele is associated with heavy muscling along the back and trunk in characteristically "bulky" largebreed dogs including the Saint Bernard, Bernese Mountain Dog, Greater Swiss Mountain Dog, and Rottweiler. The "bulky" **T** allele is absent from leaner shaped large breed dogs like the Great Dane, Irish Wolfhound, and Scottish Deerhound, which are fixed for the ancestral **C** allele. Note that this mutation does not seem to affect muscling in small or even mid-sized dog breeds with notable back muscling, including the American Staffordshire Terrier, Boston Terrier, and the English Bulldog.

Likely normal muscling (CC)

Less likely to have blue

eyes (NN)







DNA Test Report	Test Date: June 13th, 2022	embk.me/tyler143	
TRAITS: BODY SIZE			
TRAIT		RESULT	
Body Size (IGF1)		Smaller (II)	
The I allele is associated with smaller bod	y size.		
Body Size (IGFR1)		Larger (GG)	
The A allele is associated with smaller boo	dy size.		
Body Size (STC2)		Intermediate (TA)	
The A allele is associated with smaller boo	dy size.	internediate (IA)	
Body Size (GHR - E191K)		Smollor (AA)	
The A allele is associated with smaller boc	dy size.	Smaller (AA)	
Body Size (GHR - P177L)		Larger (CC)	
The T allele is associated with smaller bod	ly size.		





Test Date: June 13th, 2022



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RESULT

TRAITS: PERFORMANCE

TRAIT

Altitude Adaptation (EPAS1)

This mutation causes dogs to be especially tolerant of low oxygen environments (hypoxia), such as those found at high elevations. Dogs with at least one **A** allele are less susceptible to "altitude sickness." This mutation was originally identified in breeds from high altitude areas such as the Tibetan Mastiff.

Appetite (POMC) LINKAGE

This mutation in the POMC gene is found primarily in Labrador and Flat Coated Retrievers. Compared toNormal fooddogs with no copies of the mutation (NN), dogs with one (ND) or two (DD) copies of the mutation are moreNormal foodlikely to have high food motivation, which can cause them to eat excessively, have higher body fatmotivation (NN)percentage, and be more prone to obesity. Read more about the genetics of POMC, and learn how you cancontribute to research, in our blog post (https://embarkvet.com/resources/blog/pomc-dogs/). Wemeasure this result using a linkage test.motivation (NI)

TYLER



DNA Test Report

Test Date: June 13th, 2022

CLINICAL TOOLS

These clinical genetic tools can inform clinical decisions and diagnoses. These tools do not predict increased risk for disease.

Alanine Aminotransferase Activity (GPT)

Tyler's baseline ALT level may be Low Normal

Why is this important to your vet?

Tyler has one copy of a variant associated with reduced ALT activity as measured on veterinary blood chemistry panels. Please inform your veterinarian that Tyler has this genotype, as ALT is often used as an indicator of liver health and Tyler is likely to have a lower than average resting ALT activity. As such, an increase in Tyler's ALT activity could be evidence of liver damage, even if it is within normal limits by standard ALT reference ranges.

What is Alanine Aminotransferase Activity?

Alanine aminotransferase (ALT) is a clinical tool that can be used by veterinarians to better monitor liver health. This result is not associated with liver disease. ALT is one of several values veterinarians measure on routine blood work to evaluate the liver. It is a naturally occurring enzyme located in liver cells that helps break down protein. When the liver is damaged or inflamed, ALT is released into the bloodstream.

How vets diagnose this condition

Genetic testing is the only way to provide your veterinarian with this clinical tool.

How this condition is treated

Veterinarians may recommend blood work to establish a baseline ALT value for healthy dogs with one or two copies of this variant.







Test Date: June 13th, 2022

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HEALTH REPORT

How to interpret Tyler's genetic health results:

If Tyler inherited any of the variants that we tested, they will be listed at the top of the Health Report section, along with a description of how to interpret this result. We also include all of the variants that we tested Tyler for that we did not detect the risk variant for.

A genetic test is not a diagnosis

This genetic test does not diagnose a disease. Please talk to your vet about your dog's genetic results, or if you think that your pet may have a health condition or disease.

Tyler is at increased risk for one genetic health condition.

And inherited three variants that you should learn more about.

Chondrodystrophy and Intervertebral Disc Disease, CDDY/IVDD, Type I IVDD		٥
Methemoglobinemia		0
Degenerative Myelopathy, DM		0
Dilated Cardiomyopathy, DCM1		0
Breed-Relevant Genetic Conditions	5 variants not detected	<
Additional Genetic Conditions	210 variants not detected	<





Test Date: June 13th, 2022



embk.me/tyler143

HEALTH REPORT

Chondrodystrophy and Intervertebral Disc Disease, CDDY/IVDD, Type I IVDD (FGF4 retrogene - CFA12)

Tyler inherited one copy of the variant we tested

Tyler is at increased risk for Type I IVDD

How to interpret this result

Tyler has one copy of an FGF4 retrogene on chromosome 12. In some breeds such as Beagles, Cocker Spaniels, and Dachshunds (among others) this variant is found in nearly all dogs. While those breeds are known to have an elevated risk of IVDD, many dogs in those breeds never develop IVDD. For mixed breed dogs and purebreds of other breeds where this variant is not as common, risk for Type I IVDD is greater for individuals with this variant than for similar dogs.

What is Chondrodystrophy and Intervertebral Disc Disease, CDDY/IVDD, Type I IVDD?

Type I Intervertebral Disc Disease (IVDD) is a back/spine issue that refers to a health condition affecting the discs that act as cushions between vertebrae. With Type I IVDD, affected dogs can have a disc event where it ruptures or herniates towards the spinal cord. This pressure on the spinal cord causes neurologic signs which can range from a wobbly gait to impairment of movement. Chondrodystrophy (CDDY) refers to the relative proportion between a dog's legs and body, wherein the legs are shorter and the body longer. There are multiple different variants that can cause a markedly chondrodystrophic appearance as observed in Dachshunds and Corgis. However, this particular variant is the only one known to also increase the risk for IVDD.

When signs & symptoms develop in affected dogs

Signs of CDDY are recognized in puppies as it affects body shape. IVDD is usually first recognized in adult dogs, with breed specific differences in age of onset.

Signs & symptoms

Research indicates that dogs with one or two copies of this variant have a similar risk of developing IVDD. However, there are some breeds (e.g. Beagles and Cocker Spaniels, among others) where this variant has been passed down to nearly all dogs of the breed and most do not show overt clinical signs of the disorder. This suggests that there are other genetic and environmental factors (such as weight, mobility, and family history) that contribute to an individual dog's risk of developing clinical IVDD. Signs of IVDD include neck or back pain, a change in your dog's walking pattern (including dragging of the hind limbs), and paralysis. These signs can be mild to severe, and if your dog starts exhibiting these signs, you should schedule an appointment with your veterinarian for a diagnosis.

How vets diagnose this condition

For CDDY, dogs with one copy of this variant may have mild proportional differences in their leg length. Dogs with two copies of this variant will often have visually longer bodies and shorter legs. For IVDD, a neurological exam will be performed on any dog showing suspicious signs. Based on the result of this exam, radiographs to detect the presence of calcified discs or advanced imaging (MRI/CT) to detect a disc rupture may be recommended.

How this condition is treated

IVDD is treated differently based on the severity of the disease. Mild cases often respond to medical management which includes cage rest and pain management, while severe cases are often treated with surgical intervention. Both conservative and surgical treatment should be followed up with rehabilitation and physical therapy.







Test Date: June 13th, 2022

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HEALTH REPORT

Methemoglobinemia (CYB5R3)

Tyler inherited one copy of the variant we tested

What does this result mean?

Because this variant is inherited in an autosomal recessive manner (meaning dogs need two copies of the variant to develop the disease), Tyler is unlikely to develop this condition due to the variant.

Impact on Breeding

Your dog carries this variant and will pass it on to ~50% of his offspring. You can email breeders@embarkvet.com to discuss with a genetic counselor how the genotype results should be applied to a breeding program.

What is Methemoglobinemia?

Oxygen is carried in the blood by hemoglobin. Methemoglobin forms when hemoglobin iron is oxidized, and it cannot carry oxygen in the blood. Methemoglobinemia is a disease where too much methemoglobin is present and the body no longer has the oxygen supply it needs to function. This disease was first described in a mixed breed dog.

When signs & symptoms develop in affected dogs

Signs often first appear with a concurrent disease, such as a respiratory infection, that causes affected dogs to decompensate.

How vets diagnose this condition

Genetic and laboratory testing can be used to diagnose this condition. Please note that there are also toxins that can cause this condition.

How this condition is treated

Methylene blue can be administered to control the clinical signs, however, this is not a cure and is a long term therapy. Treatment of concurrent infections or inflammation is also recommended.

Actions to take if your dog is affected

• Please see your veterinarian as soon as possible if you suspect a respiratory infection or any other breathing difficulties as these can become life threatening if not addressed.







Test Date: June 13th, 2022

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HEALTH REPORT

Degenerative Myelopathy, DM (SOD1A)

Tyler inherited one copy of the variant we tested

What does this result mean?

Because this variant is inherited in an autosomal recessive manner (meaning dogs need two copies of the variant to develop the disease), Tyler is unlikely to develop this condition due to the variant.

Impact on Breeding

Your dog carries this variant and will pass it on to ~50% of his offspring. You can email breeders@embarkvet.com to discuss with a genetic counselor how the genotype results should be applied to a breeding program.

What is Degenerative Myelopathy, DM?

The dog equivalent of Amyotrophic Lateral Sclerosis, or Lou Gehrig's disease, DM is a progressive degenerative disorder of the spinal cord. Because the nerves that control the hind limbs are the first to degenerate, the most common clinical signs are back muscle wasting and gait abnormalities.

When signs & symptoms develop in affected dogs

Affected dogs do not usually show signs of DM until they are at least 8 years old.

How vets diagnose this condition

Definitive diagnosis requires microscopic analysis of the spinal cord after death. However, veterinarians use clues such as genetic testing, breed, age, and other diagnostics to determine if DM is the most likely cause of your dog's clinical signs.

How this condition is treated

As dogs are seniors at the time of onset, the treatment for DM is aimed towards increasing their comfort through a combination of lifestyle changes, medication, and physical therapy.

Actions to take if your dog is affected

• Giving your dog the best quality of life for as long as possible is all you can do after receiving this diagnosis.







Test Date: June 13th, 2022

HEALTH REPORT

Dilated Cardiomyopathy, DCM1 (PDK4, Doberman Pinscher Variant 1)

Tyler inherited one copy of the variant we tested

Tyler is not likely to be at increased risk for DCM1

What does this result mean?

Research indicates that this genetic variant is not likely to increase the risk that Tyler will develop this condition.

Scientific Basis

Dogs with Tyler's breed have been included in research studies or have had follow-up by our experts that indicate that this genetic variant is not likely to increase the risk of Tyler developing clinical disease.

Impact on Breeding

This genetic result should not be the primary factor in your breeding decisions.

What is Dilated Cardiomyopathy, DCM1?

DCM is the most common acquired heart disease of adult dogs. The heart has two heavily muscled ventricles that pump blood away from the heart. This disease causes progressive weakening of the ventricles by reducing the muscle mass, which causes the ventricles to dilate. Dilated ventricles do not contract and circulate oxygenated blood well, which eventually leads to heart failure.

When signs & symptoms develop in affected dogs

This disease can rarely be seen in puppies and young adults. It is typically seen in middle aged to older dogs.

How vets diagnose this condition

The earlier a diagnosis can be reached, the better the outcome. If you are concerned about your dog's heart, discuss it with your veterinarian who can run basic preliminary tests. They may recommend a visit to a veterinary cardiologist for a complete evaluation, including an ultrasound of the heart (echocardiogram).

How this condition is treated

Treatment is completely dependent on how advanced the disease is at the time of diagnosis. It can range from monitoring the patient periodically to intensive hospitalization at specialty veterinary practices.

Actions to take if your dog is affected

- The cause of this disease is multifactorial and not completely understood. Genetics, nutrition, infections and environmental exposures can all play a role in the development of DCM. In fact, DCM has recently been featured extensively in the news due to suspected nutritional deficiencies in some grain free diets.
- Annual echocardiograms by a board certified cardiologist and annual Holter monitoring are the best ways to diagnose DCM early.







Test Date: June 13th, 2022

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BREED-RELEVANT CONDITIONS TESTED



Tyler did not have the variants that we tested for, that are relevant to his breed:

- Von Willebrand Disease Type I, Type I vWD (VWF)
- Progressive Retinal Atrophy, prcd (PRCD Exon 1)
- GM2 Gangliosidosis (HEXB, Poodle Variant)
- Neonatal Encephalopathy with Seizures, NEWS (ATF2)
- 💽 Osteochondrodysplasia, Skeletal Dwarfism (SLC13A1, Poodle Variant)





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ADDITIONAL CONDITIONS TESTED

Tyler did not have the variants that we tested for, in the following conditions that the potential effect on dogs with Tyler's breed may not yet be known.

- 🛃 MDR1 Drug Sensitivity (ABCB1)
- P2Y12 Receptor Platelet Disorder (P2Y12)
- 🔀 Factor IX Deficiency, Hemophilia B (F9 Exon 7, Terrier Variant)
- 🌄 Factor IX Deficiency, Hemophilia B (F9 Exon 7, Rhodesian Ridgeback Variant)
- Factor VII Deficiency (F7 Exon 5)
- 🔀 Factor VIII Deficiency, Hemophilia A (F8 Exon 10, Boxer Variant)
- 🌄 Factor VIII Deficiency, Hemophilia A (F8 Exon 11, German Shepherd Variant 1)
- Sactor VIII Deficiency, Hemophilia A (F8 Exon 1, German Shepherd Variant 2)
- Thrombopathia (RASGRP1 Exon 5, Basset Hound Variant)
- Thrombopathia (RASGRP1 Exon 8, Landseer Variant)
- 😴 Thrombopathia (RASGRP1 Exon 5, American Eskimo Dog Variant)
- 🚫 Von Willebrand Disease Type III, Type III vWD (VWF Exon 4, Terrier Variant)
- 😴 Von Willebrand Disease Type III, Type III vWD (VWF Exon 7, Shetland Sheepdog Variant)
- 🗸 Von Willebrand Disease Type II, Type II vWD (VWF, Pointer Variant)
- 😴 Canine Leukocyte Adhesion Deficiency Type I, CLAD I (ITGB2, Setter Variant)
- 😴 Canine Leukocyte Adhesion Deficiency Type III, CLAD III (FERMT3, German Shepherd Variant)
- Congenital Macrothrombocytopenia (TUBB1 Exon 1, Cairn and Norfolk Terrier Variant)
- Canine Elliptocytosis (SPTB Exon 30)
- 😴 Glanzmann's Thrombasthenia Type I (ITGA2B Exon 13, Great Pyrenees Variant)
- 😴 Glanzmann's Thrombasthenia Type I (ITGA2B Exon 12, Otterhound Variant)
- 🔀 May-Hegglin Anomaly (MYH9)
- 💎 Prekallikrein Deficiency (KLKB1 Exon 8)
- 🔽 Pyruvate Kinase Deficiency (PKLR Exon 5, Basenji Variant)

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DNA Test Report

Test Date: June 13th, 2022

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ADDITIONAL CONDITIONS TESTED

- 💎 Pyruvate Kinase Deficiency (PKLR Exon 7, Beagle Variant)
- Pyruvate Kinase Deficiency (PKLR Exon 10, Terrier Variant)
- Trapped Neutrophil Syndrome, TNS (VPS13B)
- 🌄 Ligneous Membranitis, LM (PLG)
- 🔀 Platelet Factor X Receptor Deficiency, Scott Syndrome (TMEM16F)
- 장 Bernard-Soulier Syndrome, BSS (GP9, Cocker Spaniel Variant)
- Congenital Hypothyroidism (TPO, Tenterfield Terrier Variant)
- 🔇 Congenital Hypothyroidism (TPO, Rat, Toy, Hairless Terrier Variant)
- 😴 Congenital Dyshormonogenic Hypothyroidism with Goiter (SLC5A5, Shih Tzu Variant)
- Complement 3 Deficiency, C3 Deficiency (C3)
- 😴 Severe Combined Immunodeficiency, SCID (PRKDC, Terrier Variant)
- 😴 Severe Combined Immunodeficiency, SCID (RAG1, Wetterhoun Variant)
- 🔀 X-linked Severe Combined Immunodeficiency, X-SCID (IL2RG Exon 1, Basset Hound Variant)
- 🔀 X-linked Severe Combined Immunodeficiency, X-SCID (IL2RG, Corgi Variant)
- 💽 Progressive Retinal Atrophy, rcd1 (PDE6B Exon 21, Irish Setter Variant)
- Progressive Retinal Atrophy, rcd3 (PDE6A)
- Progressive Retinal Atrophy, CNGA (CNGA1 Exon 9)
- Progressive Retinal Atrophy, PRA1 (CNGB1)
- 🗸 Progressive Retinal Atrophy (SAG)
- 😴 Golden Retriever Progressive Retinal Atrophy 1, GR-PRA1 (SLC4A3)
- 😴 Golden Retriever Progressive Retinal Atrophy 2, GR-PRA2 (TTC8)
- 📀 Progressive Retinal Atrophy, crd1 (PDE6B, American Staffordshire Terrier Variant)
- Progressive Retinal Atrophy, crd4/cord1 (RPGRIP1)
- 🔇 X-Linked Progressive Retinal Atrophy 1, XL-PRA1 (RPGR)
- 📀 Progressive Retinal Atrophy, PRA3 (FAM161A)





Test Date: June 13th, 2022

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ADDITIONAL CONDITIONS TESTED

- 🔀 Collie Eye Anomaly, Choroidal Hypoplasia, CEA (NHEJ1)
- 🌄 Day Blindness, Cone Degeneration, Achromatopsia (CNGB3 Deletion, Alaskan Malamute Variant)
- 🛃 Day Blindness, Cone Degeneration, Achromatopsia (CNGB3 Exon 6, German Shorthaired Pointer Variant)
- 🚫 Achromatopsia (CNGA3 Exon 7, German Shepherd Variant)
- 🚫 Achromatopsia (CNGA3 Exon 7, Labrador Retriever Variant)
- 💽 Autosomal Dominant Progressive Retinal Atrophy (RHO)
- 💽 Canine Multifocal Retinopathy, cmr1 (BEST1 Exon 2)
- 🔇 Canine Multifocal Retinopathy, cmr2 (BEST1 Exon 5, Coton de Tulear Variant)
- 😴 Canine Multifocal Retinopathy, cmr3 (BEST1 Exon 10 Deletion, Finnish and Swedish Lapphund, Lapponian Herder Variant)
- 😴 Primary Open Angle Glaucoma (ADAMTS10 Exon 9, Norwegian Elkhound Variant)
- 💽 Primary Open Angle Glaucoma (ADAMTS10 Exon 17, Beagle Variant)
- 😴 Primary Open Angle Glaucoma (ADAMTS17 Exon 11, Basset Fauve de Bretagne Variant)
- 🌄 Primary Open Angle Glaucoma and Primary Lens Luxation (ADAMTS17 Exon 2, Chinese Shar-Pei Variant)
- 🔇 Goniodysgenesis and Glaucoma, Pectinate Ligament Dysplasia, PLD (OLFM3)
- 🛃 Hereditary Cataracts, Early-Onset Cataracts, Juvenile Cataracts (HSF4 Exon 9, Australian Shepherd Variant)
- Primary Lens Luxation (ADAMTS17)
- 🔇 Congenital Stationary Night Blindness (RPE65, Briard Variant)
- 💽 Congenital Stationary Night Blindness (LRIT3, Beagle Variant)
- 🔀 Macular Corneal Dystrophy, MCD (CHST6)
- 🔀 2,8-Dihydroxyadenine Urolithiasis, 2,8-DHA Urolithiasis (APRT)
- Cystinuria Type I-A (SLC3A1, Newfoundland Variant)
- Cystinuria Type II-A (SLC3A1, Australian Cattle Dog Variant)
- Cystinuria Type II-B (SLC7A9, Miniature Pinscher Variant)
- 🔀 Hyperuricosuria and Hyperuricemia or Urolithiasis, HUU (SLC2A9)
- 🔽 Polycystic Kidney Disease, PKD (PKD1)





ADDITIONAL CONDITIONS TESTED

- 💎 Primary Hyperoxaluria (AGXT)
- 🌄 Protein Losing Nephropathy, PLN (NPHS1)
- 🔀 X-Linked Hereditary Nephropathy, XLHN (COL4A5 Exon 35, Samoyed Variant 2)
- 🌄 Autosomal Recessive Hereditary Nephropathy, Familial Nephropathy, ARHN (COL4A4 Exon 30, English Springer Spaniel Variant)
- 🏷 🛛 Autosomal Recessive Hereditary Nephropathy, Familial Nephropathy, ARHN (COL4A4 Exon 3, Cocker Spaniel Variant)
- 💎 Fanconi Syndrome (FAN1, Basenji Variant)
- Primary Ciliary Dyskinesia, PCD (CCDC39 Exon 3, Old English Sheepdog Variant)
- 💽 Primary Ciliary Dyskinesia, PCD (NME5, Alaskan Malamute Variant)
- Congenital Keratoconjunctivitis Sicca and Ichthyosiform Dermatosis, Dry Eye Curly Coat Syndrome, CKCSID (FAM83H Exon 5)
- 🌄 X-linked Ectodermal Dysplasia, Anhidrotic Ectodermal Dysplasia, XHED (EDA Intron 8)
- 🔀 Renal Cystadenocarcinoma and Nodular Dermatofibrosis, RCND (FLCN Exon 7)
- 🔀 Canine Fucosidosis (FUCA1)
- 🌄 Glycogen Storage Disease Type II, Pompe's Disease, GSD II (GAA, Finnish and Swedish Lapphund, Lapponian Herder Variant)
- 🔇 Glycogen Storage Disease Type IA, Von Gierke Disease, GSD IA (G6PC, Maltese Variant)
- 😴 Glycogen Storage Disease Type IIIA, GSD IIIA (AGL, Curly Coated Retriever Variant)
- 😴 Mucopolysaccharidosis Type IIIA, Sanfilippo Syndrome Type A, MPS IIIA (SGSH Exon 6, Dachshund Variant)
- 😴 Mucopolysaccharidosis Type IIIA, Sanfilippo Syndrome Type A, MPS IIIA (SGSH Exon 6, New Zealand Huntaway Variant)
- 🛃 Mucopolysaccharidosis Type VII, Sly Syndrome, MPS VII (GUSB Exon 5, Terrier Brasileiro Variant)
- 🔀 Mucopolysaccharidosis Type VII, Sly Syndrome, MPS VII (GUSB Exon 3, German Shepherd Variant)
- Glycogen storage disease Type VII, Phosphofructokinase Deficiency, PFK Deficiency (PFKM, Whippet and English Springer Spaniel Variant)
- 😴 Glycogen storage disease Type VII, Phosphofructokinase Deficiency, PFK Deficiency (PFKM, Wachtelhund Variant)
- Lagotto Storage Disease (ATG4D)
- 🔀 Neuronal Ceroid Lipofuscinosis 1, NCL 1 (PPT1 Exon 8, Dachshund Variant 1)
- 💽 Neuronal Ceroid Lipofuscinosis 2, NCL 2 (TPP1 Exon 4, Dachshund Variant 2)
- 🔀 Neuronal Ceroid Lipofuscinosis, Cerebellar Ataxia, NCL4A (ARSG Exon 2, American Staffordshire Terrier Variant)

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ADDITIONAL CONDITIONS TESTED

- Neuronal Ceroid Lipofuscinosis 5, NCL 5 (CLN5 Exon 4 SNP, Border Collie Variant)
- 🌄 Neuronal Ceroid Lipofuscinosis 6, NCL 6 (CLN6 Exon 7, Australian Shepherd Variant)
- 🚫 Neuronal Ceroid Lipofuscinosis 8, NCL 8 (CLN8 Exon 2, English Setter Variant)
- 🏷 Neuronal Ceroid Lipofuscinosis 7, NCL 7 (MFSD8, Chihuahua and Chinese Crested Variant)
- 🚫 Neuronal Ceroid Lipofuscinosis 8, NCL 8 (CLN8, Australian Shepherd Variant)
- 🚫 Neuronal Ceroid Lipofuscinosis 10, NCL 10 (CTSD Exon 5, American Bulldog Variant)
- 🏷 Neuronal Ceroid Lipofuscinosis 5, NCL 5 (CLN5 Exon 4 Deletion, Golden Retriever Variant)
- 🚫 Neuronal Ceroid Lipofuscinosis 8, NCL 8 (CLN8 Insertion, Saluki Variant)
- 🍼 Late-Onset Neuronal Ceroid Lipofuscinosis, NCL 12 (ATP13A2, Australian Cattle Dog Variant)
- 💽 GM1 Gangliosidosis (GLB1 Exon 15, Shiba Inu Variant)
- 💽 GM1 Gangliosidosis (GLB1 Exon 15, Alaskan Husky Variant)
- 🔀 GM1 Gangliosidosis (GLB1 Exon 2, Portuguese Water Dog Variant)
- 💽 GM2 Gangliosidosis (HEXA, Japanese Chin Variant)
- 🔀 Globoid Cell Leukodystrophy, Krabbe disease (GALC Exon 5, Terrier Variant)
- 🌄 Autosomal Recessive Amelogenesis Imperfecta, Familial Enamel Hypoplasia (ENAM Deletion, Italian Greyhound Variant)
- 🔇 Autosomal Recessive Amelogenesis Imperfecta, Familial Enamel Hypoplasia (ENAM SNP, Parson Russell Terrier Variant)
- Persistent Mullerian Duct Syndrome, PMDS (AMHR2)
- 💽 Deafness and Vestibular Syndrome of Dobermans, DVDob, DINGS (MYO7A)
- 🚫 Shar-Pei Autoinflammatory Disease, SPAID, Shar-Pei Fever (MTBP)
- 🚫 Neonatal Interstitial Lung Disease (LAMP3)
- C Recurrent Inflammatory Pulmonary Disease, RIPD (AKNA, Rough Collie Variant)
- 💙 Alaskan Husky Encephalopathy, Subacute Necrotizing Encephalomyelopathy (SLC19A3)
- Alexander Disease (GFAP)
- 😴 Cerebellar Abiotrophy, Neonatal Cerebellar Cortical Degeneration, NCCD (SPTBN2, Beagle Variant)
- 🍼 Cerebellar Ataxia, Progressive Early-Onset Cerebellar Ataxia (SEL1L, Finnish Hound Variant)

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DNA Test Report

ADDITIONAL CONDITIONS TESTED

- Cerebellar Hypoplasia (VLDLR, Eurasier Variant)
- 🔀 Spinocerebellar Ataxia, Late-Onset Ataxia, LoSCA (CAPN1)
- 😴 Spinocerebellar Ataxia with Myokymia and/or Seizures (KCNJ10)
- 🌄 Hereditary Ataxia, Cerebellar Degeneration (RAB24, Old English Sheepdog and Gordon Setter Variant)
- Senign Familial Juvenile Epilepsy, Remitting Focal Epilepsy (LGI2)
- 😴 Fetal-Onset Neonatal Neuroaxonal Dystrophy (MFN2, Giant Schnauzer Variant)
- 🔀 Hypomyelination and Tremors (FNIP2, Weimaraner Variant)
- 😴 Shaking Puppy Syndrome, X-linked Generalized Tremor Syndrome (PLP1, English Springer Spaniel Variant)
- Neuroaxonal Dystrophy, NAD (TECPR2, Spanish Water Dog Variant)
- Neuroaxonal Dystrophy, NAD (VPS11, Rottweiler Variant)
- C L-2-Hydroxyglutaricaciduria, L2HGA (L2HGDH, Staffordshire Bull Terrier Variant)
- 💽 Alaskan Malamute Polyneuropathy, AMPN (NDRG1 SNP)
- 🔀 Narcolepsy (HCRTR2 Intron 4, Doberman Pinscher Variant)
- 🚫 Narcolepsy (HCRTR2 Intron 6, Labrador Retriever Variant)
- Narcolepsy (HCRTR2 Exon 1, Dachshund Variant)
- 🌄 Progressive Neuronal Abiotrophy, Canine Multiple System Degeneration, CMSD (SERAC1 Exon 15, Kerry Blue Terrier Variant)
- 😴 Progressive Neuronal Abiotrophy, Canine Multiple System Degeneration, CMSD (SERAC1 Exon 4, Chinese Crested Variant)
- Suvenile Laryngeal Paralysis and Polyneuropathy, Polyneuropathy with Ocular Abnormalities and Neuronal Vacuolation, POANV (RAB3GAP1, Rottweiler Variant)
- 🌄 Hereditary Sensory Autonomic Neuropathy, Acral Mutilation Syndrome, AMS (GDNF-AS, Spaniel and Pointer Variant)
- 🔀 Sensory Neuropathy (FAM134B, Border Collie Variant)
- 😴 Juvenile-Onset Polyneuropathy, Leonberger Polyneuropathy 1, LPN1 (LPN1, ARHGEF10)
- Juvenile Myoclonic Epilepsy (DIRAS1)
- 😴 Juvenile-Onset Polyneuropathy, Leonberger Polyneuropathy 2, LPN2 (GJA9)
- Spongy Degeneration with Cerebellar Ataxia 1, SDCA1, SeSAME/EAST Syndrome (KCNJ10)
- 🔀 Spongy Degeneration with Cerebellar Ataxia 2, SDCA2 (ATP1B2)





ADDITIONAL CONDITIONS TESTED

- Dilated Cardiomyopathy, DCM2 (TTN, Doberman Pinscher Variant 2)
- Long QT Syndrome (KCNQ1)
- 💽 Cardiomyopathy and Juvenile Mortality (YARS2)
- 🚫 Muscular Dystrophy (DMD, Cavalier King Charles Spaniel Variant 1)
- 🚫 Muscular Dystrophy (DMD, Golden Retriever Variant)
- 🔀 Limb Girdle Muscular Dystrophy (SGCD, Boston Terrier Variant)
- 🍼 Ullrich-like Congenital Muscular Dystrophy (COL6A3 Exon 10, Labrador Retriever Variant)
- Centronuclear Myopathy, CNM (PTPLA)
- Exercise-Induced Collapse, EIC (DNM1)
- Inherited Myopathy of Great Danes (BIN1)
- Myostatin Deficiency, Bully Whippet Syndrome (MSTN)
- 🚫 Myotonia Congenita (CLCN1 Exon 7, Miniature Schnauzer Variant)
- 💽 Myotonia Congenita (CLCN1 Exon 23, Australian Cattle Dog Variant)
- 💽 Nemaline Myopathy (NEB, American Bulldog Variant)
- 🍼 Myotubular Myopathy 1, X-linked Myotubular Myopathy, XL-MTM (MTM1, Labrador Retriever Variant)
- 🚫 Inflammatory Myopathy (SLC25A12)
- 🚫 Hypocatalasia, Acatalasemia (CAT)
- 💽 Pyruvate Dehydrogenase Deficiency (PDP1, Spaniel Variant)
- 🚫 Malignant Hyperthermia (RYR1)
- 🌄 Imerslund-Grasbeck Syndrome, Selective Cobalamin Malabsorption (CUBN Exon 53, Border Collie Variant)
- 🌄 Imerslund-Grasbeck Syndrome, Selective Cobalamin Malabsorption (CUBN Exon 8, Beagle Variant)
- 🔇 Inherited Selected Cobalamin Malabsorption with Proteinuria (CUBN, Komondor Variant)
- C Lundehund Syndrome (LEPREL1)
- 🔇 Congenital Myasthenic Syndrome, CMS (CHAT, Old Danish Pointing Dog Variant)
- 🌄 Congenital Myasthenic Syndrome, CMS (COLQ, Labrador Retriever Variant)





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ADDITIONAL CONDITIONS TESTED

- Congenital Myasthenic Syndrome, CMS (CHRNE, Jack Russell Terrier Variant)
- Congenital Myasthenic Syndrome, CMS (COLQ, Golden Retriever Variant)
- 🔀 Myasthenia Gravis-Like Syndrome (CHRNE, Heideterrier Variant)
- C Episodic Falling Syndrome (BCAN)
- 💽 Paroxysmal Dyskinesia, PxD (PIGN)
- C Demyelinating Polyneuropathy (SBF2/MTRM13)
- 🔀 Laryngeal Paralysis (RAPGEF6, Miniature Bull Terrier Variant)
- 🔇 Dystrophic Epidermolysis Bullosa (COL7A1, Golden Retriever Variant)
- 😴 Dystrophic Epidermolysis Bullosa (COL7A1, Central Asian Shepherd Dog Variant)
- 😴 Ectodermal Dysplasia, Skin Fragility Syndrome (PKP1, Chesapeake Bay Retriever Variant)
- 💽 Ichthyosis, Epidermolytic Hyperkeratosis (KRT10, Terrier Variant)
- 💽 Ichthyosis, ICH1 (PNPLA1, Golden Retriever Variant)
- 🛃 Ichthyosis (SLC27A4, Great Dane Variant)
- 💽 Ichthyosis (NIPAL4, American Bulldog Variant)
- 😴 Focal Non-Epidermolytic Palmoplantar Keratoderma, Pachyonychia Congenita (KRT16, Dogue de Bordeaux Variant)
- 😴 Hereditary Footpad Hyperkeratosis (FAM83G, Terrier and Kromfohrlander Variant)
- 🔇 Hereditary Footpad Hyperkeratosis (DSG1, Rottweiler Variant)
- Hereditary Nasal Parakeratosis, HNPK (SUV39H2)
- 🔀 Musladin-Lueke Syndrome, MLS (ADAMTSL2)
- 🗸 Oculocutaneous Albinism, OCA (SLC45A2, Small Breed Variant)
- Bald Thigh Syndrome (IGFBP5)
- 🔀 Lethal Acrodermatitis, LAD (MKLN1)
- 💽 Ehlers Danlos (ADAMTS2, Doberman Pinscher Variant)
- 🔀 Cleft Lip and/or Cleft Palate (ADAMTS20, Nova Scotia Duck Tolling Retriever Variant)
- 💽 Hereditary Vitamin D-Resistant Rickets (VDR)





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ADDITIONAL CONDITIONS TESTED

- 😴 Oculoskeletal Dysplasia 2, Dwarfism-Retinal Dysplasia 2, drd2, OSD2 (COL9A2, Samoyed Variant)
- 🜄 Osteogenesis Imperfecta, Brittle Bone Disease (COL1A2, Beagle Variant)
- 🔇 Osteogenesis Imperfecta, Brittle Bone Disease (SERPINH1, Dachshund Variant)
- 🔇 Osteogenesis Imperfecta, Brittle Bone Disease (COL1A1, Golden Retriever Variant)
- 🔀 Skeletal Dysplasia 2, SD2 (COL11A2, Labrador Retriever Variant)
- 🔽 Craniomandibular Osteopathy, CMO (SLC37A2)
- 🛃 Raine Syndrome, Canine Dental Hypomineralization Syndrome (FAM20C)
- 🔀 Chondrodystrophy (ITGA10, Norwegian Elkhound and Karelian Bear Dog Variant)
- 🏷 Mucopolysaccharidosis IIIB, Sanfilippo Syndrome Type B, MPS IIIB (NAGLU, Schipperke Variant)
- 🔇 Early Onset Adult Deafness, EOAD (EPS8L2 Deletion, Rhodesian Ridgeback Variant)





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INBREEDING AND DIVERSITY

CATEGORY

Coefficient Of Inbreeding

Our genetic COI measures the proportion of your dog's genome where the genes on the mother's side are identical by descent to those on the father's side.

MHC Class II - DLA DRB1

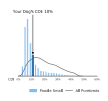
A Dog Leukocyte Antigen (DLA) gene, DRB1 encodes a major histocompatibility complex (MHC) protein involved in the immune response. Some studies have shown associations between certain DRB1 haplotypes and autoimmune diseases such as Addison's disease (hypoadrenocorticism) in certain dog breeds, but these findings have yet to be scientifically validated.

MHC Class II - DLA DQA1 and DQB1

DQA1 and DQB1 are two tightly linked DLA genes that code for MHC proteins involved in the immune response. A number of studies have shown correlations of DQA-DQB1 haplotypes and certain autoimmune diseases; however, these have not yet been scientifically validated.

RESULT

10%



High Diversity

How common is this amount of diversity in purebreds:



High Diversity

How common is this amount of diversity in purebreds:

