

3382 Capital Circle NE Tallahassee, FL 32308

Genetic Testing Report Dolly

Generated on: 01/14/2025

Submitted By		Owned By
Cavapoo 3:16		Cavapoo 3:16
]	

Subject Dog	
Name: Dolly	Lab Reference #: 869011
Breed: Cavalier King Charles Spaniel	Sample Date: 12/04/2024
Phenotype: Blenheim	Research Date: 12/04/2024
Sex: Female	Microchip: 991003001661240
Birth: 09/14/2021	

Disorder Results(4	Disorder Results(4 of 14)				
CKCSID	n/n	Clear: Dog is negative for mutation associated with Curly Coat Dry Eye.			
DM	n/DM	Heterozygous: Dog carries one copy of the mutation associated with Degenerative Myelopathy. In some breeds, there is a low risk of the dog developing the disorder			
EFS	n/n	Clear: Dog is negative for mutation associated with Episodic Falling.			
PRA-prcd	n/n	Negative: Dog is negative for the mutation associated with prcd-PRA.			
Color Results(5 of 14)					
A-Locus	at/at	Dog has two copies of the gene causing tan points.			
B-Locus	B/B	Dog does not carry the mutation for most forms of chocolate coloration.			
D-Locus	D/D	Negative: Dog is negative for the mutation associated with a diluted coat color.			
E-Locus	e/e	Dog has two copies of cream/yellow.			
K-Locus	-Locus n/n Dog is negative for the KB allele, and the coat coloration will be based on the agouti genotype.				
Pattern Results(1	Pattern Results(1 of 14)				
S-Locus	S/S	Homozygous: Dog has two copies of S-Locus resulting in a nearly solid white, parti, or piebald coat color.			



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Genetic Testing Report Dolly

Trait Results(4 of 14) The dog is negative for the hair curl allele. The dog will have Curl 1&2 n/n non-curly hair, and will always pass on the allele responsible for non-curly hair to any offspring Furnishings n/n Non-Furnished: Dog is negative for the furnishings mutation. Two copies of the long-hair allele, dog will have longer than 1¹/1 Hair Length (1-5) average hair per the breed standard. Dog has no copies of the shedding allele. The dog will have Shedding n/n a low propensity towards shedding.



Certificate of Breed

OWNER'S NAME: North America Doodles

OWNER SUPPLIED BREED: Cavalier King Charles Spaniel DOG'S NAME: Cavapoos 3 :16 "s Dolly REGISTRATION ORGANIZATION: -- TEST DATE: September 6th, 2023 REGISTRATION NUMBER: --

This certifies the authenticity of **Cavapoos 3 :16 "s Dolly**'s canine genetic background as determined following careful analysis of more than 200,000 genetic markers. Welcome to the Embark family!

WOLFINESS 1.3% MEDIUM

MATERNAL **A224** HAPLOTYPE

CAVALIER KING CHARLES SPANIEL



100.0% Cavalier King Charles Spaniel

Adam Boyko, Ph.D.

Ryan Boyko CHIEF EXECUTIVE OFFICER





Test Date: September 6th, 2023

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FAMILY TREE







Test Date: September 6th, 2023

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BREED MIX

Cavalier King Charles Spaniel : 100.0%

GENETIC STATS

Wolfiness: 1.3 % **MEDIUM** Predicted adult weight: **21 lbs** Life stage: **Young adult** Based on your dog's date of birth provided.

TEST DETAILS

Kit number: EM-19080707 Swab number: 31210953206585







Fun Fact

The breed experienced two large bursts in popularity. The first is when Queen Victoria revived the dying breed. The second was when Charlotte, a popular character from the popular show Sex and the City adopted one on TV. Test Date: September 6th, 2023

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CAVALIER KING CHARLES SPANIEL

The Cavalier King Charles Spaniel is one of the most popular dog breeds in the United States, and with good reason. Their affectionate personalities combined with their need to be close to their humans make them a lovely breed of choice for families. They tend to get along well with children and peaceably with other dogs and animals in the home (though as the breed used to be used for hunting, caution around small animals should be exercised). The Cavalier has an interesting history -- their ancestors were dogs of the British monarchy, but over time, the breed began to die out as dogs with shorter muzzles were favored in the 1800s. They were crossed with Pugs and some other breeds to change their appearance. However, Roswell Eldridge sought out King Charles Spaniels that had longer muzzles, and recreated the Cavalier as it used to be from those dogs.



English Toy Spaniel Sibling breed



English Springer Spaniel Cousin breed



English Cocker Spaniel Cousin breed



Cocker Spaniel Cousin breed



Sussex Spaniel Cousin breed

Rembark

RELATED BREEDS





Test Date: September 6th, 2023

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MATERNAL LINE



Through Cavapoos 3 :16 "s Dolly's mitochondrial DNA we can trace her mother's ancestry back to where dogs and people first became friends. This map helps you visualize the routes that her ancestors took to your home. Their story is described below the map.

HAPLOGROUP: A1a

A1a is the most common maternal lineage among Western dogs. This lineage traveled from the site of dog domestication in Central Asia to Europe along with an early dog expansion perhaps 10,000 years ago. It hung around in European village dogs for many millennia. Then, about 300 years ago, some of the prized females in the line were chosen as the founding dogs for several dog breeds. That set in motion a huge expansion of this lineage. It's now the maternal lineage of the overwhelming majority of Mastiffs, Labrador Retrievers and Gordon Setters. About half of Boxers and less than half of Shar-Pei dogs descend from the A1a line. It is also common across the world among village dogs, a legacy of European colonialism.

HAPLOTYPE: A224

Part of the large A1a haplogroup, this haplotype is found in village dogs in Peru, Fiji, and Namibia. Among breeds, we see this haplotype most frequently in Cavalier King Charles Spaniels, Mastiffs, and Boston Terriers.





Test Date: September 6th, 2023

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TRAITS: BASE COAT COLOR

TRAIT

RESULT

Dark or Light Fur | E (Extension) Locus | Gene: Melanocortin Receptor 1 (MC1R) | Genetic Result: ee

This gene helps determine whether a dog can produce dark (black or brown) hairs or lighter yellow or red hairs. Any result except for **ee** means that the dog can produce dark hairs. An **ee** result means that the dog does not produce dark hairs at all, and will have lighter yellow or red hairs over their entire body.

Did You Know? If a dog has a **ee** result then the fur's actual shade can range from a deep copper to yellow/gold to cream - the exact color cannot be predicted solely from this result, and will depend on other genetic factors.

Dark brown pigment | Cocoa | Gene: HPS3 | Genetic Result: NN

Dogs with the **coco** genotype will produce dark brown pigment instead of black in both their hair and skin. Dogs with the **Nco** genotype will produce black pigment, but can pass the **co** variant on to their puppies. Dogs that have the **coco** genotype as well as the **bb** genotype at the B locus are generally a lighter brown than dogs that have the **Bb** or **BB** genotypes at the B locus.

Did You Know? The **co** variant and the dark brown "cocoa" coat color have only been documented in French Bulldogs. Dogs with the cocoa coat color are sometimes born with light brown coats that darken as they reach maturity.

Red Pigment Intensity LINKAGE | I (Intensity) Loci | Genetic Result: Intense Red Pigmentation

Intensity refers to the concentration of red pigment in the coat. Dogs with more densely concentrated (intense) pigment will be a deeper red, while dogs with less concentrated (dilute) pigment will be tan, yellow, cream, or white. Five locations in the dog genome explain approximately 70% of red pigmentation intensity variation across all dogs. Because the locations we test may not directly cause differences in red pigmentation intensity, we consider this to be a linkage test.

Did You Know? One of the genes that influences pigment intensity in dogs, TYR, is also responsible for intensity variation in domestic mice, cats, cattle, rabbits, and llamas. In dogs and humans, more genes are involved.

Light colored fur

(cream to red)

No impact on skin color

Any pigmented fur likely apricot or red





Test Date: September 6th, 2023

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TRAITS: BASE COAT COLOR (CONTINUED)

TRAIT

Brown or Black Pigment | B (Brown) Locus | Gene: Tyrosinase Related Protein 1 (TYRP1) | Genetic Result: BB

This gene helps determine whether a dog produces brown or black pigments. Dogs with a **bb** result produce brown pigment instead of black in both their hair and skin, while dogs with a **Bb** or **BB** result produce black pigment. Dogs that have **ee** at the E (Extension) Locus and **bb** at this B (Brown) Locus are likely to have red or cream coats and brown noses, eye rims, and footpads, which is sometimes referred to as "Dudley Nose" in Labrador Retrievers.

Did You Know? "Liver" or "chocolate" is the preferred color term for brown in most breeds; in the Doberman Pinscher it is referred to as "red".

Color Dilution | D (Dilute) Locus | Gene: Melanophilin (MLPH) | Genetic Result: DD

This gene helps determine whether a dog has lighter "diluted" pigment. A dog with a **Dd** or **DD** result will not be dilute. A dog with a **dd** result will have all their black or brown pigment lightened ("diluted") to gray or light brown, and may lighten red pigment to cream. This affects their fur, skin, and sometimes eye color. 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 one **d1** allele and one **d2** allele are typically dilute. 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.

Did You Know? There are many breed-specific names for these dilute colors, such as "blue", "charcoal", "fawn", "silver", and "Isabella". Dilute dogs, especially in certain breeds, have a higher incidence of Color Dilution Alopecia which causes hair loss in some patches.

Likely black colored

RESULT

Dark (non-dilute) skin





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TRAITS: COAT COLOR MODIFIERS

TRAIT

RESULT

Hidden Patterning | K (Dominant Black) Locus | Gene: Canine Beta-Defensin 103 (CBD103) | Genetic Result: k^yk^y

This gene helps determine whether the dog has a black coat. Dogs with a $k^y k^y$ result will show a coat color pattern based on the result they have at the A (Agouti) Locus. A $K^B K^B$ or $K^B k^y$ result means the dog is dominant black, which overrides the fur pattern that would otherwise be determined by the A (Agouti) Locus. These dogs will usually have solid black or brown coats, or if they have **ee** at the E (Extension) Locus then red/cream coats, regardless of their result at the A (Agouti) Locus. Dogs who test as $K^B k^y$ may be brindle rather than black or brown.

Did You Know? Even if a dog is "dominant black" several other genes could still impact the dog's fur and cause other patterns, such as white spotting.

Body Pattern | A (Agouti) Locus | Gene: Agouti Signalling Protein (ASIP) | Genetic Result: atat

This gene is responsible for causing different coat patterns. It only affects the fur of dogs that do not have **ee** at the E (Extension) Locus and do have **k**^y**k**^y at the K (Dominant Black) Locus. It controls switching between black and red pigment in hair cells, which means that it can cause a dog to have hairs that have sections of black and sections of red/cream, or hairs with different colors on different parts of the dog's body. Sable or Fawn dogs have a mostly or entirely red coat with some interspersed black hairs. Agouti or 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.

Did You Know? The ASIP gene causes interesting coat patterns in many other species of animals as well as dogs.

Facial Fur Pattern | E (Extension) Locus | Gene: Melanocortin Receptor 1 (MC1R) | Genetic Result: ee

In addition to determining if a dog can develop dark fur at all, this gene 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 \mathbf{E}^{m} in their result will have a mask, which is dark facial fur as seen in the German Shepherd and Pug. Dogs with no \mathbf{E}^{m} in their result but one or two copies of \mathbf{E}^{g} will instead have a "widow's peak", which is dark facial fur as seen in the German Shepherd and Pug. Dogs with no \mathbf{E}^{m} in their result but one or two copies of \mathbf{E}^{g} will instead have a "widow's peak", which is dark forehead fur.

Did You Know? The widow's peak is seen in the Afghan Hound and Borzoi, where it is called either "grizzle" or "domino".

No impact on coat color

No impact on coat pattern

No dark fur anywhere





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RESULT

TRAITS: COAT COLOR MODIFIERS (CONTINUED)

TRAIT

Saddle Tan | Gene: RALY | Genetic Result: NI

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.

Did You Know? The Saddle Tan pattern is characteristic of breeds like the Corgi, Beagle, and German Shepherd.

White Spotting | S (White Spotting) Locus | Gene: MITF | Genetic Result: spsp

This gene is responsible for most of the white spotting observed in dogs. Dogs with a result of **spsp** will have a nearly white coat or large patches of white in their coat. Dogs with a result of **Ssp** will have more limited white spotting that is breed-dependent. A result of **Ss** means that a dog likely has no white or minimal white in their coat. The S Locus does not explain all white spotting patterns in dogs and other causes are currently being researched. Some dogs may have small amounts of white on the paws, chest, face, or tail regardless of their result at this gene.

Did You Know? Any dog can have white spotting regardless of coat color. The colored sections of the coat will reflect the dog's other genetic coat color results.

Roan LINKAGE | R (Roan) Locus | Gene: USH2A | Genetic Result: rr

This gene, along with the S Locus, regulates whether a dog will have roaning. Dogs with at least one copy of **R** will likely have roaning on otherwise uniformly unpigmented white areas created by the S Locus. Roan may not be visible if white spotting is limited to small areas, such as the paws, chest, face, or tail. The extent of roaning varies from uniform roaning to non-uniform roaning, and patchy, non-uniform roaning may look similar to ticking. Roan does not appear in white areas created by other genes, such as a combination of the E Locus and I Locus (for example, Samoyeds). The roan pattern can appear with or without ticking.

Did You Know? Roan, tick, and Dalmatians' spots become visible a few weeks after birth. The R Locus is probably involved in the development of Dalmatians' spots.

No impact on coat pattern

Likely to have large white areas in coat

Likely no impact on coat pattern





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TRAITS: COAT COLOR MODIFIERS (CONTINUED)

TRAIT

Merle | M (Merle) Locus | Gene: PMEL | Genetic Result: mm

This gene is responsible for mottled or patchy coat color in some dogs. Dogs with an **M*m** result are likely to appear 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 have merle or double merle coat patterning. Dogs with an **mm** result are unlikely to have a merle coat pattern.

Did You Know? Merle coat patterning is common to several dog breeds including the Australian Shepherd, Catahoula Leopard Dog, and Shetland Sheepdog.

Harlequin | Gene: PSMB | Genetic Result: hh

This gene, along with the M Locus, determines whether a dog will have harlequin patterning. 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.

Did You Know? While many harlequin dogs are white with black patches, some dogs have grey, sable, or brindle patches of color, depending on their genotypes at other coat color genes.

No impact on coat color

RESULT

No impact on coat pattern





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TRAITS: OTHER COAT TRAITS

TRAIT

Furnishings LINKAGE | Gene: RSP02 | Genetic Result: II

This gene is responsible for "furnishings", which is another name for the mustache, beard, and eyebrows that are characteristic of breeds like the Schnauzer, Scottish Terrier, and Wire Haired Dachshund. A dog with an **FF** or **FI** result is likely to have furnishings. A dog with an **II** result will not have furnishings. We measure this result using a linkage test.

Did You Know? In breeds that are expected to have furnishings, dogs without furnishings are the exception - this is sometimes called an "improper coat".

Coat Length | Gene: FGF5 | Genetic Result: TT

This gene is known to affect hair/fur length in many different species, including cats, dogs, mice, and humans. In dogs, a **TT** result means the dog is likely to have a long, silky coat as seen in the Yorkshire Terrier and the Long Haired Whippet. A **GG** or **GT** result is likely to mean a shorter coat, like in the Boxer or the American Staffordshire Terrier.

Did You Know? In certain breeds, such as Corgi, the long coat is described as "fluff."

Shedding | Gene: MC5R | Genetic Result: TT

This gene affects how much a dog sheds. Dogs with furnishings or wire-haired coats tend to be low shedders regardless of their result for this gene. In other dogs, a **CC** or **CT** result indicates heavy or seasonal shedding, like many Labradors and German Shepherd Dogs. Dogs with a **TT** result tend to be lighter shedders, like Boxers, Shih Tzus and Chihuahuas.

Coat Texture | Gene: KRT71 | Genetic Result: CC

For dogs with long fur, dogs with a **TT** or **CT** result will likely have a wavy or curly coat like the coat of Poodles and Bichon Frises. Dogs with a **CC** result will likely have a straight coat—unless the dog has a "Likely Furnished" result for the Furnishings trait, since this can also make the coat more curly.

Did You Know? Dogs with short coats may have straight coats, whatever result they have for this gene.

Hairlessness (Xolo type) LINKAGE | Gene: FOXI3 | Genetic Result: NN

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Likely unfurnished (no mustache, beard, and/or eyebrows)

RESULT

Likely long coat

Likely light shedding

Likely straight coat





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TRAITS: OTHER COAT TRAITS (CONTINUED)

TRAIT

Hairlessness (Terrier type) | Gene: SGK3 | Genetic Result: NN

This gene is responsible for Hairlessness in the American Hairless Terrier. Dogs with the **DD** result are likely to be hairless. Dogs with the **ND** genotype will have a normal coat, but can pass the **D** variant on to their offspring.

Very unlikely to be hairless

Likely not albino

RESULT

Oculocutaneous Albinism Type 2 LINKAGE | Gene: SLC45A2 | Genetic Result: NN

This gene causes oculocutaneous albinism (OCA), also known as Doberman Z Factor Albinism. Dogs with a **DD** result will have OCA. Effects include severely reduced or absent pigment in the eyes, skin, and hair, and sometimes vision problems due to lack of eye pigment (which helps direct and absorb ambient light) and are prone to sunburn. Dogs with a **ND** result will not be affected, but can pass the mutation on to their offspring. We measure this result using a linkage test.

Did You Know? 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.





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TRAITS: OTHER BODY FEATURES

TRAIT

Muzzle Length | Gene: BMP3 | Genetic Result: CC

This gene affects muzzle length. A dog with a **AC** or **CC** result is likely to have a medium-length muzzle like a Staffordshire Terrier or Labrador, or a long muzzle like a Whippet or Collie. A dog with a **AA** result is likely to have a short muzzle, like an English Bulldog, Pug, or Pekingese.

Did You Know? At least five different genes affect snout length in dogs, with BMP3 being the only one with a known causal mutation. For example, the muzzle length of some breeds, including the long-snouted Scottish Terrier or the short-snouted Japanese Chin, appear to be caused by other genes. This means your dog may have a long or short snout due to other genetic factors. Embark is working to figure out what these might be.

Tail Length | Gene: T | Genetic Result: CC

This is one of the genes that can cause a short bobtail. Most dogs have a **CC** result and a long tail. Dogs with a **CG** result are likely to have a bobtail, which is an unusually short or absent tail. This can be seen in many "natural bobtail" 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 such a result do not survive to birth.

Did You Know? 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, it is not always caused by this gene. This suggests that other unknown genetic effects can also lead to a natural bobtail.

Hind Dew Claws | Gene: LMBR1 | Genetic Result: CC

This is one of the genes that can cause hind dew claws, which are extra, nonfunctional digits located midway between a dog's paw and hock. Dogs with a **CT** or **TT** result have about a 50% chance of having hind dewclaws. Hind dew claws can also be caused by other, still unknown, genes. Embark is working to figure those out.

Did You Know? Hind dew claws are commonly found in certain breeds such as the Saint Bernard.

Likely medium or long muzzle

RESULT

Likely normal-length tail

Unlikely to have hind dew claws

Kembark





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TRAITS: OTHER BODY FEATURES (CONTINUED)

TRAIT

Back Muscling & Bulk (Large Breed) | Gene: ACSL4 | Genetic Result: CC

This gene can cause heavy muscling along the back and trunk in characteristically "bulky" large-breed dogs including the Saint Bernard, Bernese Mountain Dog, Greater Swiss Mountain Dog, and Rottweiler. A dog with the **TT** result is likely to have heavy muscling. Leaner-shaped large breed dogs like the Great Dane, Irish Wolfhound, and Scottish Deerhound generally have a **CC** result. The **TC** result also indicates likely normal muscling.

Did You Know? This gene does not seem to affect muscling in small or even mid-sized dog breeds with lots of back muscling, including the American Staffordshire Terrier, Boston Terrier, and the English Bulldog.

Eye Color LINKAGE | Gene: ALX4 | Genetic Result: NN

This gene is associated with blue eyes in Arctic breeds like Siberian Husky as well as tri-colored (nonmerle) Australian Shepherds. Dogs with a **DupDup** or **NDup** result are more likely to have blue eyes, although some dogs may have only one blue eye or may not have blue eyes at all; nevertheless, they can still pass blue eyes to their offspring. Dogs with a **NN** result may have blue eyes due to other factors, such as merle or white spotting. We measure this result using a linkage test.

Did You Know? Embark researchers discovered this gene by studying data from dogs like yours. Who knows what we will be able to discover next? Answer the questions on our research surveys to contribute to future discoveries!

Likely normal muscling

RESULT

Less likely to have blue eyes





RESULT

DNA Test Report	Test Data: Sontomber Oil, agos	
	rest Date. September 6th, 2023	embk.me/cavapoos316sdolly
TRAITS: BODY SIZE		
TRAIT		
Body Size 1 Gene: IGF1 Genetic Result: II		RESULT
This is one of several genes that influence the size smaller body size. A result of NN is associated with	e of a dog. A result of II for this gene is associated with h larger body size.	ງ Smaller
Body Size 2 Gene: IGFR1 Genetic Result: GG		
This is one of several genes that influence the size smaller body size. A result of GG is associated with	e of a dog. A result of AA for this gene is associated with larger body size.	th Larger
Body Size 3 Gene: STC2 Genetic Result: AA		
This is one of several genes that influence the size smaller body size. A result of TT is associated with	of a dog. A result of AA for this gene is associated wit larger body size.	h Smaller
Body Size 4 Gene: GHR - E191K Genetic Result: AA	A	
This is one of several genes that influence the size smaller body size. A result of GG is associated with I	of a dog. A result of AA for this gene is associated with larger body size.	Smaller
Body Size 5 Gene: GHR - P177L Genetic Result: TT		
This is one of several genes that influence the size of smaller body size. A result of CC is associated with la	of a dog. A result of TT for this gene is associated with arger body size.	Smaller





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RESULT

TRAITS: PERFORMANCE

TRAIT

Altitude Adaptation | Gene: EPAS1 | Genetic Result: GG

This gene causes dogs to be especially tolerant of low oxygen environments, such as those found at high elevations. Dogs with a **AA** or **GA** result will be less susceptible to "altitude sickness."

Did You Know? This gene was originally identified in breeds from high altitude areas such as the Tibetan Mastiff.

Appetite LINKAGE | Gene: POMC | Genetic Result: NN

This gene influences eating behavior. An **ND** or **DD** result would predict higher food motivation compared to **NN** result, increasing the likelihood to eat excessively, have higher body fat percentage, and be more prone to obesity. Read more about the genetics of POMC, and learn how you can contribute to research, in our blog post (https://embarkvet.com/resources/blog/pomc-dogs/). We measure this result using a linkage test.

Did You Know? POMC is actually short for "proopiomelanocortin," and is a large protein that is broken up into several smaller proteins that have biological activity. The smaller proteins generated from POMC control, among other things, distribution of pigment to the hair and skin cells, appetite, and energy expenditure.

Normal altitude tolerance

Normal food motivation





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HEALTH REPORT

How to interpret Cavapoos 3 :16 "s Dolly's genetic health results:

If Cavapoos 3:16 "s Dolly 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 Cavapoos 3:16 "s Dolly 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.

Summary

Of the 255 genetic health risks we analyzed, we found 4 results that you should learn about.

Increased risk results (1)

Intervertebral Disc Disease (Type I)

On the second second

ALT Activity

Degenerative Myelopathy, DM

Proportionate Dwarfism

Clear results

Breed-relevant (3)

Other (248)





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BREED-RELEVANT RESULTS

Research studies indicate that these results are more relevant to dogs like Cavapoos 3 :16 "s Dolly, and may influence her chances of developing certain health conditions.

\odot	Intervertebral Disc Disease (Type I) (FGF4 retrogene - CFA12)	ncreased risk
\oslash	Dry Eye Curly Coat Syndrome (FAM83H Exon 5)	Clear
\oslash	Episodic Falling Syndrome (BCAN)	Clear
\oslash	Muscular Dystrophy (DMD, Cavalier King Charles Spaniel Variant 1)	Clear

CAVAPOOS 3 :16 "S DOLLY



DI	NA Test Report	Test Date: September 6th, 2023	embk.me/cavapoos316sdolly
0	THER RESULTS		
6	Canine Multifocal Retinopathy, cmr2 (BE	EST1 Exon 5, Coton de Tulear Variant)	Clear
6	Canine Multifocal Retinopathy, cmr3 (BE Lapponian Herder Variant)	EST1 Exon 10 Deletion, Finnish and Swedish Lapphund,	Clear
6	Canine Multiple System Degeneration (S	SERAC1 Exon 4, Chinese Crested Variant)	Clear
0	Canine Multiple System Degeneration (S	SERAC1 Exon 15, Kerry Blue Terrier Variant)	Clear
0	Cardiomyopathy and Juvenile Mortality (YARS2)	Clear
0	Centronuclear Myopathy, CNM (PTPLA)		Clear
\bigcirc) Cerebellar Hypoplasia (VLDLR, Eurasier V	(ariant)	Clear
\oslash) Chondrodystrophy (ITGA10, Norwegian El	khound and Karelian Bear Dog Variant)	Clear
\oslash) Cleft Lip and/or Cleft Palate (ADAMTS20,	Nova Scotia Duck Tolling Retriever Variant)	Clear
\oslash) Cleft Palate, CP1 (DLX6 intron 2, Nova Scc	otia Duck Tolling Retriever Variant)	Clear
\oslash) Cobalamin Malabsorption (CUBN Exon 8, I	Beagle Variant)	Clear
\oslash	Cobalamin Malabsorption (CUBN Exon 53	, Border Collie Variant)	Clear
\oslash	Collie Eye Anomaly (NHEJ1)		Clear
\oslash	Complement 3 Deficiency, C3 Deficiency (C3)	Clear
\oslash	Congenital Cornification Disorder (NSDHL,	Chihuahua Variant)	Clear
\oslash	Congenital Hypothyroidism (TPO, Rat, Toy,	Hairless Terrier Variant)	Clear
\oslash	Congenital Hypothyroidism (TPO, Tenterfie	ld Terrier Variant)	Clear
\oslash	Congenital Hypothyroidism with Goiter (TP	O Intron 13, French Bulldog Variant)	Clear





DN.	A Test Report	Test Date: September 6th, 2023	embk.me/cavapoos316sdolly
0	THER RESULTS		
\oslash) Congenital Hypothyroidism with Goiter (SLG	C5A5, Shih Tzu Variant)	Clear
\oslash) Congenital Macrothrombocytopenia (TUBB	1 Exon 1, Cairn and Norfolk Terrier Variant)	Clear
\oslash	Congenital Myasthenic Syndrome, CMS (CC	DLQ, Labrador Retriever Variant)	Clear
\oslash	Congenital Myasthenic Syndrome, CMS (CC	DLQ, Golden Retriever Variant)	Clear
\oslash	Congenital Myasthenic Syndrome, CMS (CH	IAT, Old Danish Pointing Dog Variant)	Clear
\oslash	Congenital Myasthenic Syndrome, CMS (CH	RNE, Jack Russell Terrier Variant)	Clear
\oslash	Congenital Stationary Night Blindness (LRIT	3, Beagle Variant)	Clear
\oslash	Congenital Stationary Night Blindness (RPE	65, Briard Variant)	Clear
\oslash	Craniomandibular Osteopathy, CMO (SLC37A	A2)	Clear
\oslash	Craniomandibular Osteopathy, CMO (SLC37A	2 Intron 16, Basset Hound Variant)	Clear
\oslash	Cystinuria Type I-A (SLC3A1, Newfoundland \	Variant)	Clear
\oslash	Cystinuria Type II-A (SLC3A1, Australian Cattl	e Dog Variant)	Clear
\oslash	Cystinuria Type II-B (SLC7A9, Miniature Pinsc	cher Variant)	Clear
\oslash	Day Blindness (CNGB3 Deletion, Alaskan Mala	amute Variant)	Clear
\oslash	Day Blindness (CNGA3 Exon 7, German Sheph	erd Variant)	Clear
\oslash	Day Blindness (CNGA3 Exon 7, Labrador Retrie	ever Variant)	Clear
\oslash	Day Blindness (CNGB3 Exon 6, German Shorth	naired Pointer Variant)	Clear
\oslash	Deafness and Vestibular Syndrome of Doberm	nans, DVDob, DINGS (MY07Δ)	Clear
			Clear

CAVAPOOS 3 :16 "S DOLLY



DNA	Test Report	Test Date: September 6th, 2023	embk.me/cavapoos316sdolly
ОТ	HER RESULTS		
\oslash	Demyelinating Polyneuropathy (SBF2/MTR	M13)	Clear
\oslash	Dental-Skeletal-Retinal Anomaly (MIA3, Ca	ne Corso Variant)	Clear
\oslash	Diffuse Cystic Renal Dysplasia and Hepatic	Fibrosis (INPP5E Intron 9, Norwich Terrier Variant)	Clear
\oslash	Dilated Cardiomyopathy, DCM (RBM20, Sch	nauzer Variant)	Clear
\oslash	Dilated Cardiomyopathy, DCM1 (PDK4, Dobe	erman Pinscher Variant 1)	Clear
\oslash	Dilated Cardiomyopathy, DCM2 (TTN, Dober	man Pinscher Variant 2)	Clear
\oslash	Disproportionate Dwarfism (PRKG2, Dogo A	rgentino Variant)	Clear
\oslash	Dystrophic Epidermolysis Bullosa (COL7A1,	Central Asian Shepherd Dog Variant)	Clear
\oslash	Dystrophic Epidermolysis Bullosa (COL7A1,	Golden Retriever Variant)	Clear
\oslash	Early Bilateral Deafness (LOXHD1 Exon 38, F	Rottweiler Variant)	Clear
\oslash	Early Onset Adult Deafness, EOAD (EPS8L2	Deletion, Rhodesian Ridgeback Variant)	Clear
\oslash	Early Onset Cerebellar Ataxia (SEL1L, Finnis	h Hound Variant)	Clear
\oslash	Ehlers Danlos (ADAMTS2, Doberman Pinsch	ner Variant)	Clear
\oslash	Enamel Hypoplasia (ENAM Deletion, Italian	Greyhound Variant)	Clear
\oslash	Enamel Hypoplasia (ENAM SNP, Parson Rus	sell Terrier Variant)	Clear
\oslash	Exercise-Induced Collapse, EIC (DNM1)		Clear
\oslash	Factor VII Deficiency (F7 Exon 5)		Clear
\oslash	Factor XI Deficiency (F11 Exon 7, Kerry Blue	Terrier Variant)	Clear

CAVAPOOS 3 :16 "S DOLLY



DNA	Test Report	Test Date: September 6th, 2023	embk.me/cavapoos316sdolly
от	HER RESULTS		
\oslash	Familial Nephropathy (COL4A4 Exon 3,	Cocker Spaniel Variant)	Clear
\oslash	Familial Nephropathy (COL4A4 Exon 30), English Springer Spaniel Variant)	Clear
\oslash	Fanconi Syndrome (FAN1, Basenji Varia	nt)	Clear
\oslash	Fetal-Onset Neonatal Neuroaxonal Dys	trophy (MFN2, Giant Schnauzer Variant)	Clear
\oslash	Glanzmann's Thrombasthenia Type I (IT	rGA2B Exon 13, Great Pyrenees Variant)	Clear
\oslash	Glanzmann's Thrombasthenia Type I (IT	「GA2B Exon 12, Otterhound Variant)	Clear
\oslash	Globoid Cell Leukodystrophy, Krabbe di	sease (GALC Exon 5, Terrier Variant)	Clear
\oslash	Glycogen Storage Disease Type IA, Von	Gierke Disease, GSD IA (G6PC, Maltese Variant)	Clear
\oslash	Glycogen Storage Disease Type IIIA, GS	D IIIA (AGL, Curly Coated Retriever Variant)	Clear
\oslash	Glycogen storage disease Type VII, Pho and English Springer Spaniel Variant)	osphofructokinase Deficiency, PFK Deficiency (PFKM,	Whippet Clear
\oslash	Glycogen storage disease Type VII, Pho Wachtelhund Variant)	osphofructokinase Deficiency, PFK Deficiency (PFKM,	Clear
\oslash	GM1 Gangliosidosis (GLB1 Exon 2, Portu	uguese Water Dog Variant)	Clear
\oslash	GM1 Gangliosidosis (GLB1 Exon 15, Shib	ba Inu Variant)	Clear
\oslash	GM1 Gangliosidosis (GLB1 Exon 15, Alas	skan Husky Variant)	Clear
\oslash	GM2 Gangliosidosis (HEXA, Japanese C	hin Variant)	Clear
\oslash	GM2 Gangliosidosis (HEXB, Poodle Varia	ant)	Clear
\oslash	Golden Retriever Progressive Retinal At	crophy 1, GR-PRA1 (SLC4A3)	Clear
\oslash	Golden Retriever Progressive Retinal At	crophy 2, GR-PRA2 (TTC8)	Clear





DNA Test Report	Test Date: September 6th, 2023	embk.me/cavapoos316sdolly
OTHER RESULTS		
⊘ Goniodysgenesis and Glaucoma	, Pectinate Ligament Dysplasia, PLD (OLFM3)	Clear
⊘ Hemophilia A (F8 Exon 11, Germa	an Shepherd Variant 1)	Clear
🔗 Hemophilia A (F8 Exon 1, Germar	n Shepherd Variant 2)	Clear
Hemophilia A (F8 Exon 10, Boxer	Variant)	Clear
Hemophilia B (F9 Exon 7, Terrier V	Variant)	Clear
🔗 Hemophilia B (F9 Exon 7, Rhodes	ian Ridgeback Variant)	Clear
🔗 Hereditary Ataxia, Cerebellar Deg	eneration (RAB24, Old English Sheepdog and Gordon Setter Vari	ant) Clear
Hereditary Cataracts (HSF4 Exon	9, Australian Shepherd Variant)	Clear
Hereditary Footpad Hyperkeratos	is (FAM83G, Terrier and Kromfohrlander Variant)	Clear
Hereditary Footpad Hyperkeratos	is (DSG1, Rottweiler Variant)	Clear
Hereditary Nasal Parakeratosis (S	UV39H2 Intron 4, Greyhound Variant)	Clear
Hereditary Nasal Parakeratosis, HI	NPK (SUV39H2)	Clear
Hereditary Vitamin D-Resistant Rid	ckets (VDR)	Clear
🔗 Hypocatalasia, Acatalasemia (CAT))	Clear
Hypomyelination and Tremors (FNI	IP2, Weimaraner Variant)	Clear
🖉 Hypophosphatasia (ALPL Exon 9, K	arelian Bear Dog Variant)	Clear
🖉 Ichthyosis (NIPAL4, American Bulld	log Variant)	Clear
⊘ Ichthyosis (ASPRV1 Exon 2, Germa	n Shepherd Variant)	Clear
		Clear

CAVAPOOS 3 :16 "S DOLLY



- Alexandre		
DNA Test Report	Test Date: September 6th, 2023	embk.me/cavapoos316sdolly
OTHER RESULTS		
🕗 Ichthyosis (SLC27A4, Great Da	ane Variant)	Clear
🔗 Ichthyosis, Epidermolytic Hyp	erkeratosis (KRT10, Terrier Variant)	Clear
🔗 Ichthyosis, ICH1 (PNPLA1, Gold	den Retriever Variant)	Clear
Inflammatory Myopathy (SLC2	5A12)	Clear
Inherited Myopathy of Great D	panes (BIN1)	Clear
Inherited Selected Cobalamin	Malabsorption with Proteinuria (CUBN, Komondor Variant)	Clear
Intestinal Lipid Malabsorption	(ACSL5, Australian Kelpie)	Clear
🧭 Junctional Epidermolysis Bullo	osa (LAMA3 Exon 66, Australian Cattle Dog Variant)	Clear
Junctional Epidermolysis Bull	osa (LAMB3 Exon 11, Australian Shepherd Variant)	Clear
Juvenile Epilepsy (LGI2)		Clear
🧭 Juvenile Laryngeal Paralysis a	nd Polyneuropathy (RAB3GAP1, Rottweiler Variant)	Clear
Juvenile Myoclonic Epilepsy (I	DIRAS1)	Clear
⊘ L-2-Hydroxyglutaricaciduria, L	2HGA (L2HGDH, Staffordshire Bull Terrier Variant)	Clear
⊘ Lagotto Storage Disease (ATG-	4D)	Clear
⊘ Laryngeal Paralysis (RAPGEF6	, Miniature Bull Terrier Variant)	Clear
Late Onset Spinocerebellar At	axia (CAPN1)	Clear
Late-Onset Neuronal Ceroid Li	pofuscinosis, NCL 12 (ATP13A2, Australian Cattle Dog Variant)	Clear
Leonberger Polyneuropathy 1	(LPN1, ARHGEF10)	Clear

Fembark





DNA	Test Report	Test Date: September 6th, 2023	embk.me/cavapoos316sdolly
от	HER RESULTS		
\oslash	Leonberger Polyneuropathy 2 (GJA9)		Clear
\oslash	Lethal Acrodermatitis, LAD (MKLN1)		Clear
\oslash	Leukodystrophy (TSEN54 Exon 5, Standard	d Schnauzer Variant)	Clear
\oslash	Ligneous Membranitis, LM (PLG)		Clear
\oslash	Limb Girdle Muscular Dystrophy (SGCD, Bo	oston Terrier Variant)	Clear
\oslash	Limb-Girdle Muscular Dystrophy 2D (SGCA	Exon 3, Miniature Dachshund Variant)	Clear
\oslash	Long QT Syndrome (KCNQ1)		Clear
\oslash	Lundehund Syndrome (LEPREL1)		Clear
\oslash	Macular Corneal Dystrophy, MCD (CHST6)		Clear
\oslash	Malignant Hyperthermia (RYR1)		Clear
\oslash	May-Hegglin Anomaly (MYH9)		Clear
\oslash	Methemoglobinemia (CYB5R3, Pit Bull Terr	rier Variant)	Clear
\oslash	Methemoglobinemia (CYB5R3)		Clear
\oslash	Microphthalmia (RBP4 Exon 2, Soft Coated	Wheaten Terrier Variant)	Clear
\oslash	Mucopolysaccharidosis IIIB, Sanfilippo Syr	ndrome Type B, MPS IIIB (NAGLU, Schipperke Variant)	Clear
\oslash	Mucopolysaccharidosis Type IIIA, Sanfilipp Variant)	o Syndrome Type A, MPS IIIA (SGSH Exon 6, Dachshun	d Clear
\oslash	Mucopolysaccharidosis Type IIIA, Sanfilipp Huntaway Variant)	o Syndrome Type A, MPS IIIA (SGSH Exon 6, New Zeala	ind Clear
\oslash	Mucopolysaccharidosis Type VI, Maroteaux Variant)	x-Lamy Syndrome, MPS VI (ARSB Exon 5, Miniature Pin	scher Clear





Clear

Clear

Clear

Clear

DNA Test Report	Test Date: September 6th, 2023	embk.me/cavapoos316sdolly
OTHER RESULTS		
O Mucopolysaccharidosis Type VII, Sl	y Syndrome, MPS VII (GUSB Exon 3, German Shepherd Varia	ant) Clear
Mucopolysaccharidosis Type VII, Sl	y Syndrome, MPS VII (GUSB Exon 5, Terrier Brasileiro Varian	t) Clear
Multiple Drug Sensitivity (ABCB1)		Clear
🔗 Muscular Dystrophy (DMD, Golden F	Retriever Variant)	Clear
Musladin-Lueke Syndrome, MLS (AI	DAMTSL2)	Clear
🧭 Myasthenia Gravis-Like Syndrome (CHRNE, Heideterrier Variant)	Clear
Myotonia Congenita (CLCN1 Exon 23	3, Australian Cattle Dog Variant)	Clear
🧭 Myotonia Congenita (CLCN1 Exon 7,	Miniature Schnauzer Variant)	Clear
Narcolepsy (HCRTR2 Exon 1, Dachs)	nund Variant)	Clear
Narcolepsy (HCRTR2 Intron 4, Dobe	rman Pinscher Variant)	Clear
Narcolepsy (HCRTR2 Intron 6, Labra	dor Retriever Variant)	Clear
Nemaline Myopathy (NEB, American	Bulldog Variant)	Clear
Neonatal Cerebellar Cortical Degen	eration (SPTBN2, Beagle Variant)	Clear
Neonatal Encephalopathy with Seiz	ures, NEWS (ATF2)	Clear
🔗 Neonatal Interstitial Lung Disease (I	LAMP3)	Clear

 \oslash Neuroaxonal Dystrophy, NAD (VPS11, Rottweiler Variant) Neuroaxonal Dystrophy, NAD (TECPR2, Spanish Water Dog Variant) \oslash Neuronal Ceroid Lipofuscinosis 1, NCL 1 (PPT1 Exon 8, Dachshund Variant 1)





DNA Test Report	Test Date: September 6th, 2023	embk.me/cavapoos316sdolly
OTHER RESULTS		
Neuronal Ceroid Lipofuscin	osis 10, NCL 10 (CTSD Exon 5, American Bulldog Variant)	Clear
Neuronal Ceroid Lipofuscin	osis 2, NCL 2 (TPP1 Exon 4, Dachshund Variant 2)	Clear
🔗 Neuronal Ceroid Lipofuscin	osis 5, NCL 5 (CLN5 Exon 4 SNP, Border Collie Variant)	Clear
🔗 Neuronal Ceroid Lipofuscin	osis 5, NCL 5 (CLN5 Exon 4 Deletion, Golden Retriever Variant)	Clear
🔗 Neuronal Ceroid Lipofuscin	osis 6, NCL 6 (CLN6 Exon 7, Australian Shepherd Variant)	Clear
🚫 Neuronal Ceroid Lipofuscin	osis 7, NCL 7 (MFSD8, Chihuahua and Chinese Crested Variant)	Clear
🔗 Neuronal Ceroid Lipofuscin	osis 8, NCL 8 (CLN8, Australian Shepherd Variant)	Clear
O Neuronal Ceroid Lipofuscin	osis 8, NCL 8 (CLN8 Exon 2, English Setter Variant)	Clear
O Neuronal Ceroid Lipofuscin	osis 8, NCL 8 (CLN8 Insertion, Saluki Variant)	Clear
Neuronal Ceroid Lipofuscing Variant)	osis, Cerebellar Ataxia, NCL4A (ARSG Exon 2, American Staffordsl	hire Terrier Clear
Oculocutaneous Albinism, C	DCA (SLC45A2 Exon 6, Bullmastiff Variant)	Clear
Oculocutaneous Albinism, C	DCA (SLC45A2, Small Breed Variant)	Clear
Oculoskeletal Dysplasia 2 (COL9A2, Samoyed Variant)	Clear
🔗 Osteochondrodysplasia (SL	C13A1, Poodle Variant)	Clear
Osteogenesis Imperfecta (0	COL1A2, Beagle Variant)	Clear
Osteogenesis Imperfecta (S	SERPINH1, Dachshund Variant)	Clear
Osteogenesis Imperfecta (C	COL1A1, Golden Retriever Variant)	Clear
P2Y12 Receptor Platelet Dis	sorder (P2Y12)	Clear





DNA	Test Report	Test Date: September 6th, 2023	embk.me/cavapoos316sdolly
от	HER RESULTS		
\oslash	Pachyonychia Congenita (KRT16, Dogue de	e Bordeaux Variant)	Clear
\oslash	Paroxysmal Dyskinesia, PxD (PIGN)		Clear
\oslash	Persistent Mullerian Duct Syndrome, PMDS	S (AMHR2)	Clear
\oslash	Pituitary Dwarfism (POU1F1 Intron 4, Karelia	an Bear Dog Variant)	Clear
\oslash	Platelet Factor X Receptor Deficiency, Scot	t Syndrome (TMEM16F)	Clear
\oslash	Polycystic Kidney Disease, PKD (PKD1)		Clear
\oslash	Pompe's Disease (GAA, Finnish and Swedis	sh Lapphund, Lapponian Herder Variant)	Clear
\oslash	Prekallikrein Deficiency (KLKB1 Exon 8)		Clear
\oslash	Primary Ciliary Dyskinesia, PCD (NME5, Ala	skan Malamute Variant)	Clear
\oslash	Primary Ciliary Dyskinesia, PCD (CCDC39 E	xon 3, Old English Sheepdog Variant)	Clear
\oslash	Primary Hyperoxaluria (AGXT)		Clear
\oslash	Primary Lens Luxation (ADAMTS17)		Clear
\oslash	Primary Open Angle Glaucoma (ADAMTS17	Exon 11, Basset Fauve de Bretagne Variant)	Clear
\oslash	Primary Open Angle Glaucoma (ADAMTS10	Exon 17, Beagle Variant)	Clear
\oslash	Primary Open Angle Glaucoma (ADAMTS10	Exon 9, Norwegian Elkhound Variant)	Clear
\oslash	Primary Open Angle Glaucoma and Primary Variant)	Lens Luxation (ADAMTS17 Exon 2, Chinese Shar-Pe	Clear
\oslash	Progressive Retinal Atrophy (SAG)		Clear
\oslash	Progressive Retinal Atrophy (IFT122 Exon 2	6, Lapponian Herder Variant)	Clear





DNA	Test Report	Test Date: September 6th, 2023	embk.me/cavapoos316sdolly
от	HER RESULTS		
\oslash	Progressive Retinal Atrophy, Bardet-Biedl S	Syndrome (BBS2 Exon 11, Shetland Sheepdog Variant)	Clear
\oslash	Progressive Retinal Atrophy, CNGA (CNGA1	Exon 9)	Clear
\oslash	Progressive Retinal Atrophy, crd1 (PDE6B, A	American Staffordshire Terrier Variant)	Clear
\oslash	Progressive Retinal Atrophy, crd4/cord1 (RI	PGRIP1)	Clear
\oslash	Progressive Retinal Atrophy, PRA1 (CNGB1)		Clear
\oslash	Progressive Retinal Atrophy, PRA3 (FAM161	A)	Clear
\oslash	Progressive Retinal Atrophy, prcd (PRCD Ex	ron 1)	Clear
\oslash	Progressive Retinal Atrophy, rcd1 (PDE6B E	xon 21, Irish Setter Variant)	Clear
\oslash	Progressive Retinal Atrophy, rcd3 (PDE6A)		Clear
\oslash	Protein Losing Nephropathy, PLN (NPHS1)		Clear
\oslash	Pyruvate Dehydrogenase Deficiency (PDP1,	Spaniel Variant)	Clear
\oslash	Pyruvate Kinase Deficiency (PKLR Exon 5, E	asenji Variant)	Clear
\oslash	Pyruvate Kinase Deficiency (PKLR Exon 7, B	eagle Variant)	Clear
\oslash	Pyruvate Kinase Deficiency (PKLR Exon 10,	Terrier Variant)	Clear
\oslash	Pyruvate Kinase Deficiency (PKLR Exon 7, La	abrador Retriever Variant)	Clear
\oslash	Pyruvate Kinase Deficiency (PKLR Exon 7, P	ug Variant)	Clear
\oslash	Raine Syndrome (FAM20C)		Clear
\oslash	Recurrent Inflammatory Pulmonary Disease	, RIPD (AKNA, Rough Collie Variant)	Clear

CAVAPOOS 3 :16 "S DOLLY



DNA	Test Report Test D	ate: September 6th, 2023	embk.me/cavapoos316sdolly
от	HER RESULTS		
\oslash	Renal Cystadenocarcinoma and Nodular Dermatofi	brosis (FLCN Exon 7)	Clear
\oslash	Retina Dysplasia and/or Optic Nerve Hypoplasia (S	IX6 Exon 1, Golden Retriever Variant)	Clear
\oslash	Sensory Neuropathy (FAM134B, Border Collie Varia	nt)	Clear
\oslash	Severe Combined Immunodeficiency, SCID (PRKDC	C, Terrier Variant)	Clear
\oslash	Severe Combined Immunodeficiency, SCID (RAG1,	Wetterhoun Variant)	Clear
\oslash	Shaking Puppy Syndrome (PLP1, English Springer S	Spaniel Variant)	Clear
\oslash	Shar-Pei Autoinflammatory Disease, SPAID, Shar-P	ei Fever (MTBP)	Clear
\oslash	Skeletal Dysplasia 2, SD2 (COL11A2, Labrador Retri	ever Variant)	Clear
\oslash	Skin Fragility Syndrome (PKP1, Chesapeake Bay Re	etriever Variant)	Clear
\oslash	Spinocerebellar Ataxia (SCN8A, Alpine Dachsbrack	e Variant)	Clear
\oslash	Spinocerebellar Ataxia with Myokymia and/or Seiz	ures (KCNJ10)	Clear
\oslash	Spongy Degeneration with Cerebellar Ataxia 1 (KCl	NJ10)	Clear
\oslash	Spongy Degeneration with Cerebellar Ataxia 2 (ATI	P1B2)	Clear
\oslash	Stargardt Disease (ABCA4 Exon 28, Labrador Retrie	ever Variant)	Clear
\oslash	Succinic Semialdehyde Dehydrogenase Deficiency	r (ALDH5A1 Exon 7, Saluki Variant)	Clear
\oslash	Thrombopathia (RASGRP1 Exon 5, American Eskim	o Dog Variant)	Clear
\oslash	Thrombopathia (RASGRP1 Exon 5, Basset Hound Va	ariant)	Clear
\oslash	Thrombopathia (RASGRP1 Exon 8, Landseer Varian	t)	Clear

CAVAPOOS 3 :16 "S DOLLY



DNA Te	st Report	Test Date: September 6th, 2023	embk.me/cavapoos316sdolly
отн	ER RESULTS		
⊘ ⊺	rapped Neutrophil Syndrome, TNS (VPS13)	В)	Clear
Θ υ	Ilrich-like Congenital Muscular Dystrophy	(COL6A3 Exon 10, Labrador Retriever Variant)	Clear
Ο υ	Ilrich-like Congenital Muscular Dystrophy	(COL6A1 Exon 3, Landseer Variant)	Clear
0 V	nilateral Deafness and Vestibular Syndrom	ne (PTPRQ Exon 39, Doberman Pinscher)	Clear
0 υ	rate Kidney & Bladder Stones (SLC2A9)		Clear
⊘ v	on Willebrand Disease Type I, Type I vWD ((VWF)	Clear
⊘ v	on Willebrand Disease Type II, Type II vWD	(VWF, Pointer Variant)	Clear
⊘ v	on Willebrand Disease Type III, Type III vW	D (VWF Exon 4, Terrier Variant)	Clear
⊘ v	on Willebrand Disease Type III, Type III vW	D (VWF Intron 16, Nederlandse Kooikerhondje Variar	nt) Clear
Ø ∨	on Willebrand Disease Type III, Type III vW	D (VWF Exon 7, Shetland Sheepdog Variant)	Clear
Ø X-	-Linked Hereditary Nephropathy, XLHN (CC	0L4A5 Exon 35, Samoyed Variant 2)	Clear
Ø X-	-Linked Myotubular Myopathy (MTM1, Labi	rador Retriever Variant)	Clear
⊘ x-	-Linked Progressive Retinal Atrophy 1, XL-F	PRA1 (RPGR)	Clear
Ø X-	-linked Severe Combined Immunodeficien	cy, X-SCID (IL2RG Exon 1, Basset Hound Variant)	Clear
⊘ x-	-linked Severe Combined Immunodeficien	cy, X-SCID (IL2RG, Corgi Variant)	Clear
⊘ Xa	anthine Urolithiasis (XDH, Mixed Breed Var	iant)	Clear
Øβ-	-Mannosidosis (MANBA Exon 16, Mixed-Br	eed Variant)	Clear





Test Date: September 6th, 2023

embk.me/cavapoos316sdolly

HEALTH REPORT

Increased risk result

Intervertebral Disc Disease (Type I)

Cavapoos 3 :16 "s Dolly inherited both copies of the variant we tested for Chondrodystrophy and Intervertebral Disc Disease, CDDY/IVDD, Type I IVDD Cavapoos 3 :16 "s Dolly is at increased risk for Type I IVDD

How to interpret this result

Cavapoos 3 :16 "s Dolly has two copies 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

Kembark





Test Date: September 6th, 2023

embk.me/cavapoos316sdolly

HEALTH REPORT

Ontable result

ALT Activity

Cavapoos 3:16 "s Dolly inherited one copy of the variant we tested for Alanine Aminotransferase Activity

Why is this important to your vet?

Cavapoos 3 :16 "s Dolly has one copy of a variant associated with reduced ALT activity as measured on veterinary blood chemistry panels. Please inform your veterinarian that Cavapoos 3 :16 "s Dolly has this genotype, as ALT is often used as an indicator of liver health and Cavapoos 3 :16 "s Dolly is likely to have a lower than average resting ALT activity. As such, an increase in Cavapoos 3 :16 "s Dolly'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: September 6th, 2023

embk.me/cavapoos316sdolly

HEALTH REPORT

Ontable result

Degenerative Myelopathy, DM

Cavapoos 3 :16 "s Dolly inherited one copy of the variant we tested for Degenerative Myelopathy, DM

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), Cavapoos 3 :16 "s Dolly is unlikely to develop this condition due to the variant. This result may be important if you decide to breed this dog - we recommend genetic testing potential mates for this condition.

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.





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HEALTH REPORT

Notable result

Proportionate Dwarfism

Cavapoos 3 :16 "s Dolly inherited both copies of the variant we tested for Proportionate Dwarfism Cavapoos 3 :16 "s Dolly is not known to be at increased risk for Proportionate Dwarfism

What does this result mean?

We do not know whether this increases the risk that Cavapoos 3:16 "s Dolly will develop this disease.

Scientific Basis

Research studies for this variant have been based on dogs of other breeds. Not enough dogs with the breeds in Cavapoos 3 :16 "s Dolly have been studied to know whether or not this variant will increase Cavapoos 3 :16 "s Dolly's risk of developing this disease.

What is Proportionate Dwarfism?

This variant in the GH1 gene may lead to a growth hormone (GH) abnormality and causes proportionately small stature, coat abnormalities and low blood sugar (hypoglycemia).

When signs & symptoms develop in affected dogs

Clinical signs may be visible as early as a few months of life, but coat and dental abnormalities and small size may not be clear until puppies are older.

How vets diagnose this condition

Clinical history, genetic testing, and laboratory testing can be used to diagnose this form of Proportionate Dwarfism.

How this condition is treated

Your veterinarian may recommend various treatments, including correcting blood sugar, as indicated."

Actions to take if your dog is affected

• Monitor for signs of hypoglycemia including not eating, lethargy, and inability to stand. Call your veterinarian immediately for advice if you notice these signs.





Test Date: September 6th, 2023

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INBREEDING AND DIVERSITY

CATEGORY

Inbreeding | Gene: n/a | Genetic Result: 37%

Inbreeding is a measure of how closely related this dog's parents were. The higher the number, the more closely related the parents. In general, greater inbreeding is associated with increased incidence of genetically inherited conditions.

Immune Response 1 | Gene: DRB1 | Genetic Result: No Diversity

Diversity in the Major Histocompatibility Complex (MHC) region of the genome has been found in some studies to be associated with the incidence of certain autoimmune diseases. Dogs that have less diversity in the MHC region—i.e. the Dog Leukocyte Antigen (DLA) inherited from the mother is similar to the DLA inherited from the father—are considered less immunologically diverse. A High Diversity result means the dog has two highly dissimilar haplotypes. A Low Diversity result means the dog has two similar but not identical haplotypes. A No Diversity result means the dog has inherited identical haplotypes from both parents. Some studies have shown associations between certain DRB1 haplotypes and autoimmune diseases such as Cushing's disease, but these findings have yet to be scientifically validated.

Immune Response 2 | Gene: DQA1 and DQB1 | Genetic Result: No Diversity

Diversity in the Major Histocompatibility Complex (MHC) region of the genome has been found in some studies to be associated with the incidence of certain autoimmune diseases. Dogs that have less diversity in the MHC region—i.e. the Dog Leukocyte Antigen (DLA) inherited from the mother is similar to the DLA inherited from the father—are considered less immunologically diverse. A High Diversity result means the dog has two highly dissimilar haplotypes. A Low Diversity result means the dog has two similar but not identical haplotypes. A No Diversity result means the dog has inherited identical haplotypes from both parents. A number of studies have shown correlations of DQA-DQB1 haplotypes and certain autoimmune diseases; however, these have not yet been scientifically validated.

RESULT

37%



No Diversity

How common is this amount of diversity in purebreds:



No Diversity

How common is this amount of diversity in purebreds:

