

Sample ID: 2309-W-56206  
Report Date: 29/09/2023  
Our Ref: 9115

**Veterinary Surgeon**

Kintra Vet  
Malcolmwood Farm  
Blantyre  
G72 9UL  
United Kingdom  
  
info@kintravet.com

**Owner**

Ms Bozena Bienkowska  
Foreside Farm  
Neilston  
Glasgow  
G78 3AQ  
United Kingdom

**Animal Details**

<b>Animal:</b>	Cat (Feline)	<b>D.O.B</b>	11/06/2020
<b>Name</b>	Barnycooniecats Peanut (Lady Grey)	<b>Microchip No.</b>	9001333000456229
<b>Breed</b>	Maine Coon	<b>Registration</b>	SBT 110620035
<b>Sex</b>	Female	<b>Tattoo No.</b>	

**Sample**

<b>Sample Material</b>	EDTA-Blood	<b>Sample received:</b>	21/09/2023
<b>Sample Date:</b>	31/07/2023		

**Test**

<b>Test Name:</b>	HCM (Hypertrophic cardiomyopathy )
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**Result**

<b>Genotype:</b>	N/N (Genetically Clear)
<b>Interpretation:</b>	The examined animal is homozygous for the wild type allele. It does not carry the causative mutation for Hypertrophic Cardiomyopathy in the MYBPC3-gene (A31P).  Trait of inheritance: autosomal-dominant  Scientific studies found correlation between the mutation and symptoms of the disease in the following breeds: Maine Coon and related breeds  The test result is only valid for the submitted sample.

\*\*\*\*\* END OF RESULT \*\*\*\*\*

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<b>Sex</b>	Female	<b>Tattoo No.</b>	

**Sample**

<b>Sample Material</b>	EDTA-Blood	<b>Sample received:</b>	21/09/2023
<b>Sample Date:</b>	31/07/2023		

**Test**

**Test Name:** 8015CGD PK Deficiency (Pyruvate Kinase Deficiency)

**Result**

**Genotype:** N/ N (Genetically Clear)  
**Interpretation:** The examined animal is homozygous for the wild type allele. It does not carry the causative mutation for Pyruvate Kinase Deficiency in the PKLR-gene.  
  
Trait of inheritance: autosomal-recessive.  
  
The result is only valid for the submitted sample.

\*\*\*\*\* END OF RESULT \*\*\*\*\*

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UK Office: LABOKLIN (UK) Unit 20, Wheel Forge Way, Trafford Park, Manchester, M17 1EH

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<b>Sex</b>	Female	<b>Tattoo No.</b>	

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**Test**

<b>Test Name:</b>	8729 Factor XI Deficiency
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**Result**

<b>Genotype:</b>	N / N (Genetically Clear)
<b>Interpretation:</b>	The examined animal is homozygous for the wildtype-allele. It does not carry the associated variant for Factor XI deficiency in the FXI-gene.  Trait of inheritance: autosomal recessive  A correlation between the mutation and symptoms of the disease was found in the following breed: Maine Coon  The current result is only valid for the sample submitted to our laboratory.

\*\*\*\*\* END OF RESULT \*\*\*\*\*

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<b>Sex</b>	Female	<b>Tattoo No.</b>	

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<b>Sample Material</b>	EDTA-Blood	<b>Sample received:</b>	21/09/2023
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**Test**

**Test Name:** 8900C Genetic Blood Groups in cats

**Result**

**Genotype:** N / N

**Interpretation:** The examined animal is homozygous for the N allele. It does not carry the causative genetic variant found in correlation with the serologic blood group B and AB (C) so far.

The test detects three genetic variants for the 'b' allele (268T>A, 179G>T, 1322delT) and one variant for the 'c' allele (364C>T).

Allelic series: N>c>b

The result is only guaranteed for the submitted sample.

\*\*\*\*\* END OF RESULT \*\*\*\*\*

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**Test**

**Test Name:** SMA (Spinal Muscular Atrophy) in Maine Coon

**Result**

**Genotype:** N / N (Genetically Clear)  
**Interpretation:** The analyzed cat is a noncarrier of the mutation, that is suggested to cause Spinal Muscular Atrophy (SMA).  
  
Since it cannot pass the mutation onto its offspring it can be mated to any other cat.  
  
The currently known mutation has been analysed.  
  
This result is only valid for the submitted sample material and for the Maine Coon breed.

\*\*\*\*\* END OF RESULT \*\*\*\*\*

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Your cat's DNA will be stored for 5 years and will be available for further testing. If you wish to order further tests on this cat, you don't have to submit a new sample; simply write the above sample ID on the order form, or mark which test(s) you require from the list below and return together with the order form. You can also order by calling us on 0161 282 3066 and quoting the sample ID.

The following tests are available for *Maine Coon*:

- 8604 Cystinuria (Feline Cystinuria) (CY)
- 8729 Factor XI Deficiency (F11)
- 8900CGD Genetic Blood groups in cats
- 8080 HCM 1 (Hypertrophic cardiomyopathy ) Mutation Meurs (G-- > C) A31P
- 8776 MDR1 Gene Defect
- 8473 Mucopolysaccharidosis type VII (MPS VII / MPS7)
- 8461 Myotonia Congenita (Fainting Goat)
- 8015CGD PK Deficiency (Pyruvate Kinase Deficiency)
- 8118CGD SMA (Spinal Muscular Atrophy )
- 618 Serological Evaluation of blood Groups (**EDTA Blood ONLY**)
- 8350 **Special Offer:** HCM, HCR, GSD4, PKD, PRA, PK-Def., SMA, Blood Groups
- 8718 **Maine Coon DNA bundle:** HCM1 + SMA + PK-Def

**Coat Colours**

- 8242 Coat Colour Albino
- 8099Cats Coat Colour Dilution
- 8257 Coat Colour: Chocolate and Cinnamon
- 8147Cats Coat Length
- 8090 Coat colour Variant Agouti (Tabby)
- 8258 Colourpoint Siamese and Burmese ( Siamese , Burmese and Mink )
- 8591 Dominant White / White Spotting
- 8778 Tabby (Mackerel , Blotched ) Taqpep
- 8777 Ticked cat (Ticking)
- 8280 **Feline Coat Colours OFFER:** Agouti + Burma + Chocolate + Cinnamon + Dilution + Siam

\*\*\*\*\* END OF RESULT \*\*\*\*\*

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