


Maine coon HCM (hypertrophic cardiomyopathy) test

		Vizi Barbara	Test number: 11092019-Viz2
Cattery HU*Diamond Star	DOB 14.04.2018	Electronic chip	Report date: 13.09.2019
Breed Maine Coon	Cat Sina's Little Giants Crispy, black silver classic tabby highwhite	Sex ♂	

Result

N/N

Result Codes:

N/N – Cat is homozygous for a normal type (i.e. carry two normal copies of the *MYBPC3* gene).

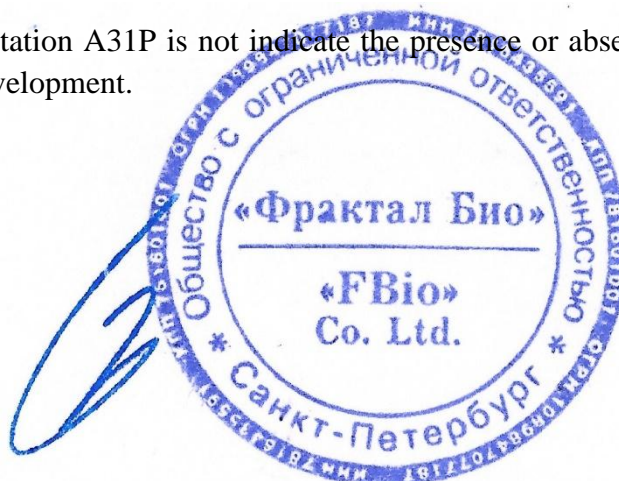
N/HCMmc – Cat is heterozygous for the mutant allele (one copy of the *MYBPC3* gene carry the A31P mutation).

HCMmc/HCMmc – Cat is homozygous for the A31P mutation (two copies of the *MYBPC3* gene carry the A31P mutation).

This test only detects the A31P mutation associated with HCM in Maine Coon cats and outcrosses as described by Meurset *al.*, 2005.

The presence or absence of the mutation A31P is not indicate the presence or absence of the disease, but only describes the risk of its development.

Chief of molecular
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Bagmanova S. Elena






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Feline ADPKD test (autosomal dominant polycystic kidney disease)

		Vizi Barbara		Test number: 11092019-Viz2
Cattery HU*Diamond Star	DOB 14.04.2018	Electronic chip		Report date: 13.09.2019
Breed Maine Coon	Cat Sina's Little Giants Crispy, black silver classic tabby highwhite	Sex ♂		

Result

N/N

Result Codes:

N/N – Cat is homozygous for a normal type (i.e. carry two normal copies of the *PKD1* gene).

N/pkd1 – Cat is heterozygous for the mutant allele (one copy of the *PKD1* gene carry the 10063C>A mutation).

pkd1/pkd1 – Cat is homozygous for the 10063C>A mutation (two copies of the *PKD1* gene carry the 10063C>A mutation). Homozygote form could be lethal in utero or at a very early age.


This test detects the 10063C>A mutation associated with autosomal dominant polycystic kidney disease in cats as described by Lyons *et al.*, 2004.

The presence of the only one copy of the *PKD1* gene containing the 10063C>A mutation causes ADPKD.

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Feline PK Deficiency test (Pyruvate Kinase Deficiency in Felines)

		Vizi Barbara	Test number: 11092019-Viz2
Cattery HU*Diamond Star	DOB 14.04.2018	Electronic chip	Report date: 13.09.2019
Breed Maine Coon	Cat Sina's Little Giants Crispy, black silver classic tabby highwhite	Sex ♂	

Result

N/N

Result Codes:

N/N – Cat is homozygous for a normal type (i.e. carry two normal copies of the *PKLR* gene).

N/k – Cat is heterozygous for the mutant allele (one copy of the *PKLR* gene carry the mutation).

k/k – Cat is homozygous for the mutant allele (two copies of the *PKLR* gene carry the mutation).


This test detects the *c.693+304G>A* mutation associated with pyruvate kinase deficiency in Felines as described by Grahn *et al.*, 2012.

The PKdef disease affects cats with two mutant copies of the *PKLR* gene only. Cats with only one mutant copy of the *PKLR* gene are clinically without any symptoms but are the carriers.

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Feline SMA test (autosomal recessive spinal muscular atrophy)

		Vizi Barbara	Test number: 11092019-Viz2
Cattery HU*Diamond Star	DOB 14.04.2018	Electronic chip	Report date: 13.09.2019
Breed Maine Coon	Cat Sina's Little Giants Crispy, black silver classic tabby highwhite	Sex ♂	

Result

N/N

Result Codes:

N/N – Cat is homozygous for a normal type (i.e. carry two normal copies of the *LIX1-LNPEP* locus).

N/lix1-lnpep – Cat is heterozygous for the mutant alleles (one copy of the *LIX1-LNPEP* locus carry the deletion).

lix1-lnpep/lix1-lnpep – Cat is homozygous for the mutant alleles (two copies of the *LIX1-LNPEP* locus carry the deletion).


This test detects the 140-kb deletion in the *LIX1-LNPEP* locus associated with autosomal recessive spinal muscular atrophy in domestic cats as described by Fyfe *et al.*, 2006.

The SMA disease affects cats with two mutant copies of the *LIX1-LNPEP* locus only. Cats with only one mutant copy of the *LIX1-LNPEP* locus are clinically without any symptoms.

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AB Blood Group DNA-test in Felines

		Vizi Barbara		Test number: 11092019-Viz2
Cattery HU*Diamond Star	DOB 14.04.2018	Electronic chip		Report date: 08.10.2019
Breed Maine Coon	Cat Sina's Little Giants Crispy, black silver classic tabby highwhite	Sex ♂		

Result

N/N

Result Codes:

N/N – Cat is type A or type AB.

N/b – Cat is a carrier of B factor; serotype could be Type A or Type AB.

b/b – Cat is type B.

The DNA blood group test identifies cats that have the B serotype (two copies of b allele) and also that are B carriers (one copy of the b allele, A or AB serotypes). Because the DNA test cannot distinguish between a Type A versus a Type AB cat, the non-b allele is reported as N. The cat could have a Type A or Type AB serotype.

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