



ГЕНЕТИЧЕСКАЯ ЭКСПЕРТИЗА № НЦ0002492

Дата регистрации заказа:	03.11.2019	Дата взятия материала:	02.11.2019
Ф.И.О. владельца:	Ванина	Дата выполнения:	15.11.2019
Вид животного:	Кошка	Микрочип №:	643094100609413
Порода:	Бенгальская	Дата сверки чипа:	
Возраст (лет, мес.):	0,9	Пол:	М
Лечащий врач:	Пономарева	Кличка:	Victoria Grand Sherlock
		Заказчик:	Монморанси
Материал:	буккальный эпителий	Метод:	Генетические исследования

Результаты исследования:

Прогрессирующая атрофия сетчатки rdAc:

NN, Не несет аллель заболевания. Заболевание, ассоциированное с исследованной мутацией, не будет развиваться. Животное не передаст аллель заболевания потомству.

Врач КЛД, эксперт: **Руковицина Вера Михайловна**
Дата выдачи: **15.11.2019**

Лаборатория сертифицирована
ГОСТ Р ИСО 9001-2008
РОСС RU.ИК76.K00071



Лаборатория является участником Федеральной системы внешней оценки качества лабораторных исследований МЗ РФ ФСВОК (код участника 10705)

Претензии по результатам анализа принимаются в течение месяца с даты готовности.
Архивное хранение результата гарантировано в течение 1 года.



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		Заказчик:	Монморанси
Материал:	буккальный эпителий	Метод:	Генетические исследования

Результаты исследования:

Дефицит эритроцитарной пируваткиназы (PK deficiency):

NN, Не несет аллель заболевания. Заболевание, ассоциированное с исследованной мутацией, не будет развиваться. Животное не передаст аллель заболевания потомству.

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Архивное хранение результата гарантировано в течение 1 года.



VETERINARY GENETICS LABORATORY
SCHOOL OF VETERINARY MEDICINE
ONE SHIELDS AVENUE
DAVIS, CALIFORNIA 95616-8744

TELEPHONE: (530) 752-2211
FAX: (530) 752-3556

BENGAL PRA TEST REPORT

TATIANA LOPATINA STR. DEKABRISTOV, 28-2-42 MOSCOW, 127273 RUSSIAN FEDERATION		Case: CAT87661 Date Received: 11-Aug-2016 Print Date: 12-Aug-2016 Report ID: 6253-7714-9862-4180 Verify report at www.vgl.ucdavis.edu/myvgl/verify.html
Cat: QUICK HUNTERS VIVIEN DOB: 06/01/2015 Sex: Female Breed: Bengal Color: Brown (black) spotted tabby		Reg: RU 0220.010615.18304.LO.BEN
Sire: DESTINYBENGALS BLACK JAGUAR Dam: VICTORIA GRAND FELISIA		Reg: Reg:

BENGAL PRA RESULT

N/N

Result Codes:

- N/N Normal - no copies of the PRA-b mutation.
- N/PRA Carrier - 1 copy of the PRA-b mutation; vision will be normal.
- PRA/PRA Affected - 2 copies of the PRA-b mutation; cat will develop clinical signs of Bengal PRA.

For more detailed information on Bengal PRA results, please go to:
www.vgl.ucdavis.edu/services/cat/BengalPRA.php

BENGAL PROGRESSIVE RETINAL ATROPHY

The Veterinary Genetics Laboratory at the University of California – Davis and the Lyons Feline Genetics and Comparative Medicine Laboratory at the University of Missouri announce the release of the **new genetic test for progressive retinal atrophy (PRA)**, which causes an autosomal recessive blindness in Bengal cats. The disease causes the destruction of the cells that register light (photoreceptors) in the back of the eye (the retina). The loss of the cells begins around 7 weeks of age and slowly progresses until the cat has very compromised vision by approximately 2 years of age¹. However, blindness develops at different rates in different cats. We have examples of cats over 2 years of age chasing a laser pointer; however vision testing by an ophthalmologist indicated that the cats should be blind. Blind cats tend to have more difficulty at night, sometimes becoming more vocal and more attached to their owners. The pupils are usually more dilated for affected cats than for cats with normal vision in the same lighting conditions. Affected cats also tend to carry their whiskers in a more forward position. Once affected cats know their surroundings, they are very mobile and active. Our thanks to the breeders who came forward and helped us establish a colony so that we could define the condition and find the gene responsible for this defect.

The mutant DNA variant appears to be novel to the Bengal breed and occurred early in a popular lineage of the Bengals. We expect Bengal cats worldwide to have the condition and we have had reports of affected cats in the United Kingdom, Europe and the USA. Bengal PRA is autosomal recessive, thus two copies of the mutant DNA variant are required for the cats to be blind. The blindness can be detected either by the DNA test or by an eye exam prior breeding age. Carriers, cats with one copy of the mutation, can only be detected by the DNA test.

Because the mutation is recent in evolutionary time, an extended DNA region and several genes around the casual mutation had to be examined as many DNA variants appeared to be perfectly linked with the blindness. This is a common and well known phenomenon, called linkage disequilibrium. Other examples of linkage disequilibrium have been published for cat DNA variants that cause *Brown* coloration² and cat AB blood group.³

Several DNA variants within the region needed to be evaluated and after working with European breeders, many non-causal variants were eliminated. Initially, a gene mutation in a known vision gene was implicated, but after screening hundreds of cats, a second mutation in a different and novel vision gene was identified as being perfectly

linked with the disease. Only by continued cooperation and participation of breeders was this discovery possible!

To the best of our ability and knowledge, the new test is correct and perfectly linked with the disease. However, we are limited by the cat samples obtained for research and we will continue studies to demonstrate the function of this gene in cats. We will be publishing this research in a scientific journal, however, we are releasing the new test ahead of the publication because breeders are continuing to use carrier cats and even blind cats in their breeding programs, and thus Bengal PRA blindness is rapidly spreading.

All cats previously tested for Bengal PRA by the VGL have been retested for the new mutation and new reports will be issued to clients.

1: Ofri R, Reilly CM, Maggs DJ, Fitzgerald PG, Shilo-Benjamini Y, Good KL, Grahn RA, Splawski DD, Lyons LA. Characterization of an Early-Onset, Autosomal Recessive, Progressive Retinal Degeneration in Bengal Cats. *Invest Ophthalmol Vis Sci.* 2015 Aug;56(9):5299-308. doi: 10.1167/iovs.15-16585. PubMed PMID: 26258614; PubMed Central PMCID: PMC4539567.

2: Lyons LA, Foe IT, Rah HC, Grahn RA. Chocolate coated cats: TYRP1 mutations for brown color in domestic cats. *Mamm Genome.* 2005 May;16(5):356-66. PubMed PMID: 16104383.

3: Bighignoli B, Niini T, Grahn RA, Pedersen NC, Millon LV, Polli M, Longeri M, Lyons LA. Cytidine monophospho-N-acetylneuraminic acid hydroxylase (CMAH) mutations associated with the domestic cat AB blood group. *BMC Genet.* 2007 Jun 6;8:27. PubMed PMID: 17553163; PubMed Central PMCID: PMC1913925.

ФЕДЕРАЛЬНОЕ ГОСУДАРСТВЕННОЕ БЮДЖЕТНОЕ УЧРЕЖДЕНИЕ
"ВСЕРОССИЙСКИЙ ЦЕНТР КАЧЕСТВА И СТАНДАРТИЗАЦИИ
ЛЕКАРСТВЕННЫХ СРЕДСТВ ДЛЯ ЖИВОТНЫХ И КОРМОВ"
ФГБУ "ВГНКИ"

ЦЕНТР
МОЛЕКУЛЯРНОЙ
ДИАГНОСТИКИ
ФГБУ "ВГНКИ" № 7003050007
ИНН. ЦЛВД 006014

Центр Молекулярной Диагностики

123 022, г. Москва, Звенигородское шоссе, д. 5
тел./факс (499) 259 27 18 Адрес в Интернет: cmd-вгнки.рф

E-mail: cmd@vgnki.ru

ВЫЯВЛЕНИЕ МУТАЦИИ, ОТВЕТСТВЕННОЙ ЗА РАЗВИТИЕ ДЕФИЦИТ
ПИРУВАТКИНАЗЫ У КОШЕК ПОРОД АБИССИНСКАЯ, СОМАЛИ.
ERYTHROCYTE PYRUVATE KINASE DEFICIENCY DNA TEST FOR ABYSSINIAN
AND SOMALI CATS



Номер карточки / Test	212-06	Дата поступления / Date:	11 июня 2014 г.
ФИО владельца / Owner Na	Лопатина Татьяна		
Порода /Breed	Дата рождения / Date of birth:		
BEN	01.05.2013		
Кличка / Name	Victoria Grand Felicia	Пол / Sex	♀
Результат	Генетическая мутация отсутствует / Genetic mutation, responsible for Pyruvate Kinase Deficiency		





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PK DEFICIENCY AND IDENTITY MARKER REPORT

TATIANA LOPATINA STR. DEKABRISTOV, 28-2-42 MOSCOW, 127273 RUSSIAN FEDERATION		Case: CAT104188 Date Received: 26-Mar-2018 Print Date: 27-Mar-2018 Report ID: 6686-6438-0698-6100 Verify report at www.vgl.ucdavis.edu/myvgl/verify.html
Cat: LEGION RAFFAELLO SANTI DOB: 07/13/2016 Sex: Male Breed: Bengal Color: n 24		Reg: RU 0220.130716.20872.LO.BEN
Sire: PRASLIN PERSEUS OF LEGION Dam: AZARTIS KAMELLA OF LEGION		Reg: SBT 091514 001 Reg: CBC-RU 0277/149/BEN/2015

PYRUVATE KINASE DEFICIENCY TEST RESULT

N/N

Result Codes:

N/N no copies of PK deficiency, cat is normal

N/K 1 copy of PK deficiency, cat is normal but is a carrier

K/K 2 copies of PK deficiency, cat is or will be affected. Severity of symptoms cannot be predicted*

Erythrocyte Pyruvate Kinase Deficiency (PK deficiency) is an inherited, autosomal recessive, hemolytic anemia. Breedings between carriers will be expected to produce 25% affected kittens. Go to our website for a list of breeds at risk of PK deficiency due to a significant frequency of the mutation.

*If your cat is diagnosed as homozygous for PK deficiency, we recommend that you contact your veterinarian for information on disease progression and management.

For more information on PK Deficiency test results, please go to:
www.vgl.ucdavis.edu/services/pkdeficiency.php

IDENTITY MARKERS

LOCUS	TYPE	LOCUS	TYPE
FCA075	PS	FCA220	J1/L
FCA223	GV	FCA678	MN
FCA698	Zc		

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SANTA BARBARA • SANTA CRUZ

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BENGAL PRA TEST REPORT

TATIANA LOPATINA STR. DEKABRISTOV, 28-2-42 MOSCOW, 127273 RUSSIAN FEDERATION	Case: CAT87719 Date Received: 15-Aug-2016 Print Date: 17-Aug-2016 Report ID: 7625-5284-3070-1011 <small>Verify report at www.vgl.ucdavis.edu/myvgl/verify.html</small>
Cat: QUICK HUNTERS X LADY Reg: RU 0220.220513.10554.LO.BEN DOB: 05/22/2013 Sex: Female Breed: Bengal Color: Brown (black) spotted tabby	
Sire: DESTINYBENGALS BLACK JAGUAR Reg: Dam: QUICK HUNTERS MARCELLA Reg:	

BENGAL PRA RESULT

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Cat: LEGION RAFFAELLO SANTI DOB: 07/13/2016 Sex: Male Breed: Bengal Color: n 24	Reg: RU 0220.130716.20872.LO.BEN
Sire: PRASLIN PERSEUS OF LEGION Dam: AZARTIS KAMELLA OF LEGION	Reg: SBT 091514 001 Reg: CBC-RU 0277/149/BEN/2015

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