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





SANTA BARBARA • SANTA CRUZ

PK DEFICIENCY AND IDENTITY MARKER REPORT

MARIE HARRIMAN 169 CLINTON ROAD ANTRIM, NH 03440

Case: CAT118055

Date Received: 06-Sep-2019

Print Date: 10-Sep-2019

Report ID: 6896-9964-9784-3024
Verify report at www.vgl.ucdavis.edu/myvgl/verify.htm

Cat: GGLEGACY LOLA OF ATLASTCATS Reg: SBV 062317 013

DOB: 06/23/2017 Sex: Female Breed: Maine Coon Microchip: 933000120133102 Color: Brown Classic Torbie with White

Sire: SGC GGLEGACY ANGELUNEAU Reg: SBT 090814 027

Dam: GGLEGACY ANGELJUNO Reg: SBV 052716 009

PYRUVATE KINASE DEFICIENCY TEST RESULT

N/N

Result Codes:

N/N	no copies of I	PK deficiency,	cat is normal
1 N/ 1 N	no copies of i	i i i dellelelle y,	cat is morniar

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	S	FCA220	L
FCA223	UX	FCA678	NP
FCA698	AT		

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

MAINE COON HCM (HYPERTROPHIC CARDIOMYOPATHY) TEST REPORT

MARIE HARRIMAN 169 CLINTON ROAD ANTRIM, NH 03440

Case: CAT118055

Date Received: 06-Sep-2019

Print Date: 10-Sep-2019

Report ID: 6403-9638-5524-5140

Verify report at www.vgl.ucdavis.edu/myvgl/verify.htm

Cat: GGLEGACY LOLA OF ATLASTCATS

Reg: SBV 062317 013

DOB: 06/23/2017 Sex: Female Breed: Maine Coon Microchip: 933000120133102 Color: Brown Classic Torbie with White

Sire: SGC GGLEGACY ANGELUNEAU Reg: SBT 090814 027

Dam: GGLEGACY ANGELJUNO Reg: SBV 052716 009

Maine Coon HCM Test Result

N/N

Result Codes:

N/N	Normal.
N/HCMmc	One copy of the A31P mutation is present. Cat is 1.8 times more likely to develop HCM than cats without the mutation.
HCMmc/HCMmc	Two copies of the A31P mutation are present. Cat is 18 times more likely to develop HCM than cats without the mutation.

This test only detects the A31P mutation associated with HCM in Maine Coon cats and outcrosses as described by Meurs et al. 2005. The A31P mutation is not the sole cause of HCM in Maine Coons. The other causes are not known at this time.

For more information on Maine Coon HCM test results, please go to: www.vgl.ucdavis.edu/services/cat/MaineCoonHCM.php

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MAINE COON SPINAL MUSCULAR ATROPHY TEST REPORT

MARIE HARRIMAN 169 CLINTON ROAD ANTRIM, NH 03440 *Case:* CAT118055

Date Received: 06-Sep-2019

Print Date: 10-Sep-2019

Report ID: 4416-0577-6159-9004

Verify report at www.vgl.ucdavis.edu/myvgl/verify.htm

Cat: GGLEGACY LOLA OF ATLASTCATS

ATLASTCATS Reg: SBV 062317 013

DOB: 06/23/2017 Sex: Female Breed: Maine Coon Microchip: 933000120133102 Color: Brown Classic Torbie with White

Sire: SGC GGLEGACY ANGELUNEAU Reg: SBT 090814 027

Dam: GGLEGACY ANGELJUNO Reg: SBV 052716 009

SMA Result

N/N

Result Codes:

N/N No copies of SMA are present.

N/S 1 copy of SMA is present. Cat is normal but is a carrier. Breedings between carriers will be expected to produce

25% affected, 50% carriers and 25% normal kittens.

S/S 2 copies of SMA are present, cat is affected.

This test is specific for the mutation associated with SMA in Maine Coon cats and outcrosses.

For more information on SMA test results, please go to: www.vgl.ucdavis.edu/services/cat/SMA.php