UNIVERSITY OF CALIFORNIA, DAVIS

BERKELEY + DAVIS + IRVINE + LOS ANGELES + MERCED + RIVERSIDE + SAN DIEGO + SAN FRANCISCO

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



SANTA BARBARA + SANTA CRUZ

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

AQHA GENETIC DISEASE PANEL TEST RESULTS

AMERICAN QUARTER HORSE ASSOCIATION P.O. BOX 200 AMARILLO, TX 79168-0001 Case: QHA47404

Date Received: 28-Feb-2012

Report Date:

06-Mar-2012

Report ID:

Reg: 4807356

9763-7779-4969-5191

Verify report at https://www.vgi.ucdavis.edu/myvgl/verify.htmi

Horse: ROCKIN W

YOB: 06

Breed: QH

Sex: S

Alt. ID: 5632119

Sire: DUAL REY

Reg: 3258332

Dam: BOON SAN KITTY

Reg: 3955171

GBED	N/N
HERDA	N/HRD
НҮРР	N/N
МН	N/N
PSSM1	N/N

N/N - Normal - Does not possess the disease-causing GBED gene

N/HRD - Carrier - horse carries one copy of the HERDA gene

N/N - Normal - Does not possess the disease-causing HYPP gene

N/N - Normal - horse does not have the MH gene

N/N - Normal - horse does not have the PSSM1 gene

GBED - Glycogen Branching Enzyme Deficiency. Fatal disease of newborn foals caused by defect in glycogen storage. Affects heart and skeletal muscles and brain. Inherited as recessive disease.

HERDA - Hereditary Equine Regional Dermal Asthenia. Skin disease characterized by hyperextensible skin, scarring, and severe lesions along the back of affected horses. Typical onset is around 2 years of age. Inherited as a recessive disease.

HYPP - Hyperkalemic Periodic Paralysis. Muscle disease caused by defect in sodium channel gene that causes involuntary muscle contraction and increased level of potassium in blood. Inherited as dominant disease. Two copies of defective gene produce more severe signs than one copy.

MH - Malignant Hyperthermia. Rare but life-threatening skeletal muscle disease triggered by exposure to volatile anesthetics (halothane), depolarizing muscle relaxants (succinylcholine), and stress. Presumed inheritance as dominant disease.

PSSM1 - Polysaccharide Storage Myopathy Type 1. Muscle disease characterized by accumulation of abnormal complex sugars in skeletal muscles. Signs include muscle pain, stiffness, skin twitching, sweating, weakness and reluctance to move. Inherited as a dominant disease.

GBED testing performed under a license agreement with the University of Minnesota.

HERDA testing performed under a license agreement with the University of California, Davis.

PSSM1 testing performed under a license agreement with the American Quarter Horse Association.





Saturday, January 4, 2025

ROCKIN W SYNDICATE PO BOX 130 GUTHRIE TX 79236 Customer ID: 3673036

Transaction #: 42785915-11

Registration Key: 5632119

Dear ROCKIN W SYNDICATE:

The results for the genetic test(s) that you ordered for ROCKIN W, 4807356 are below.

MYHM N/N

For more information regarding these genetic diseases, please refer to AQHA.com\genetics and AQHA rules REG109.3-REG109.7, which are available in the AQHA Official Handbook of Rules and Regulations at www.agha.com.

If you have any questions, please contact AQHA at 806-376-4811 8 a.m. to 5 p.m. Central, Monday through Friday, or submit a Contact Us form on AQHA.com.

Sincerely,

AQHA Member Experience Team





EQUINE JUVENILE SPINOCEREBELLAR ATAXIA TEST REPORT

Provided Information:

Name: ROCKIN W

Registration: 4807356

Case: NQ51867

Date Received: 25-Jul-2019
Report Issue Date: 18-Sep-2024

Report ID: 3550-0530-5567-0017

Verify report at vgl.ucdavis.edu/verify

DOB: 01/01/2006 Sex: Stallion Breed: Quarter Horse

RESULT

INTERPRETATION

Equine Juvenile
Spinocerebellar Ataxia
No copies of the allele associated with equine juvenile spinocerebellar ataxia (EJSCA) detected.