

KTBR 0412 9

Bastu (or Mia?), Mixed Breed

Registered Name: Bastu (or Mia?)

Call Name: Bastu

Microchip: 982000363813269

Breed: Mixed Breed

Gender: Female

Owner: Charles Warden

Country: United States

Testing date: 2019/11/25

Test results - Known disorders in the breed - page 1

| Disorder | Туре | Mode of Inheritance | Result |
|---|------------------------|---------------------|--------|
| Acute Intermittent Porphyria; HMBS mutation: c.107_110delACAG | Metabolic Disorders | Autosomal Dominant | Clear |
| Acute Intermittent Porphyria; HMBS mutation: c.826-1G>A | Metabolic Disorders | Autosomal Dominant | Clear |
| Chylomicronemia, Lipoprotein Lipase Deficiency | Metabolic Disorders | Autosomal Recessive | Clear |
| Congenital Adrenal Hyperplasia | Endocrine Disorders | Autosomal Recessive | Clear |
| Cystinuria; SCL3A1 mutation | Metabolic Disorders | Autosomal Recessive | Clear |
| Cystinuria; SCL7A9 mutation: c.1175C>T | Metabolic Disorders | Autosomal Recessive | Clear |
| Cystinuria; SCL7A9 mutation: c.706G>A | Metabolic Disorders | Autosomal Recessive | Clear |
| Cystinuria; SCL7A9 mutation: c.881A>T | Metabolic Disorders | Autosomal Recessive | Clear |
| Dihydropyrimidinuria | Metabolic Disorders | Autosomal Recessive | Clear |
| | | | |

On behalf of Genoscoper Laboratories,

SIGNATURE

Jonas Donner, PhD, Head of Research and Development at Genoscoper Laboratories

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Call Name: Bastu Country: United States

Microchip: 982000363813269 **Testing date:** 2019/11/25

Breed: Mixed Breed Gender: Female

Test results - Known disorders in the breed - page 2

| Disorder | Туре | Mode of Inheritance | Result |
|--|-------------------------|---------------------|--------|
| Factor XII Deficiency | Blood Disorders | Autosomal Recessive | Clear |
| Feline GM1 Gangliosidosis | Neurologic Disorders | Autosomal Recessive | Clear |
| GM2 Gangliosidosis, Domestic Shorthair mutation HEXB: c.1467_1491inv | Neurologic Disorders | Autosomal Recessive | Clear |
| GM2 Gangliosidosis; Domestic Shorthair GM2A Mutation | Neurologic Disorders | Autosomal Recessive | Clear |
| Hemophilia B, mutation F9: c.1014C>T | Blood Disorders | X-linked Recessive | Clear |
| Hemophilia B, mutation F9: c.247G>A | Blood Disorders | X-linked Recessive | Clear |
| Hyperoxaluria | Renal Disorders | Autosomal Recessive | Clear |
| Mucopolysaccharidosis Type I | Metabolic Disorders | Autosomal Recessive | Clear |
| Mucopolysaccharidosis Type VI (MPS VI), Typical Form | Metabolic Disorders | Autosomal Recessive | Clear |

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Test results - Known disorders in the breed - page 3

| Disorder | Туре | Mode of Inheritance | Result |
|---|-------------------------|---------------------|--------|
| Mucopolysaccharidosis Type VII, mutation GUSB: c.1074G>A | Metabolic Disorders | Autosomal Recessive | Clear |
| Mucopolysaccharidosis VII; GUSB mutation C1424T | Metabolic Disorders | Autosomal Recessive | Clear |
| Myotonia Congenita | Muscular Disorders | Autosomal Recessive | Clear |
| Niemann-Pick C2, NPC Disease, Sphingomyelinosis NPC2 Mutation | Neurologic Disorders | Autosomal Recessive | Clear |
| Niemann–Pick C1, NPC Disease, Sphingomyelinosis NPC1 Mutation | Neurologic Disorders | Autosomal Recessive | Clear |
| Polycystic Kidney Disease | Renal Disorders | Autosomal Dominant | Clear |
| Vitamin D-Dependent Rickets (VDDR-1A); CYP27B mutation: c.G637T | Metabolic Disorders | Autosomal Recessive | Clear |
| | | | |

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Optimal Selection POWERED BY GEN STOPER

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Test results - Traits - page 1

Blood Type

| Trait | Genotype | Description |
|-------------------------|----------|-----------------------|
| Blood Type (3 variants) | N/N | Cat has blood type A. |

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Test results - Traits - page 2

Coat Color

| Trait | Genotype | Description |
|---|----------|---|
| Color Locus B: Chocolate and Cinnamon (2 variants) | B/B | The cat produces black pigment. |
| Color Locus C: Pointed Coloration and Albinism (3 variants) | C/C | This cat has full color with no color restrictions. |
| Color Locus A: Agouti and Charcoal (2 variants) | a/a | The cat is likely to have non-agouti (solid) coat color. |
| any FERV1 insertion in the KIT gene | N/N | The cat has no copy of the FERV1 insertion in the KIT gene. The cat's coat color is not affected by White spotting or Dominant White mutations. |
| MLPH T83del (d allele) | D/d | The coat color is not diluted; the cat carries one copy of the dilute gene. |
| MC1R c.250G>A (e allele) | E/E | The cat does not have e allele for Amber color found in Norwegian Forest Cat. |
| MC1R (er allele) | -/- | The cat does not have er allele for Russet color found in Burmese. |

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Test results - Traits - page 3

Coat Type

| Trait | Genotype | Description |
|----------------------------|----------|---|
| Long Hair (4 variants) | N/M3 | The cat carries one copy of the long hair mutations and is likely to have short hair. |
| LPAR6 c.250_253_delTTTG | N/N | The cat does not have Cornish Rex curly coat. |
| KRT71 c.445-1C | N/N | The cat does not have Selkirk Rex curly coat. |

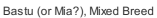
Morphology

| Trait | Genotype | Description |
|--|----------|---|
| Short tail, T-box mutations (3 variants) | N/N | The cat does not have any of the tested bobtail mutations originally found in Manx. |
| Polydactyly (3 variants) | N/N | The cat does not have any of the tested mutations causing extra digits. |
| HES7 c.T5C | T/T | The cat has no bobtail mutation originally found in Japanese Bobtail. |

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Test results - Additional disorders found in other breeds - page 1

Blood Disorders

| Disorder | Mode of Inheritance | Result |
|---|---------------------|--------|
| Erythrocyte Pyruvate Kinase (PK) Deficiency | Autosomal Recessive | Clear |

Cardiac Disorders

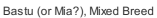
| Disorder | Mode of Inheritance | Result |
|--|---|--------|
| Hypertrophic Cardiomyopathy, MYBPC3 mutation: A31P found in Maine Coon | Autosomal Dominant (Incomplete Penetrance) | Clear |
| Hypertrophic Cardiomyopathy, MYBPC3 mutation: c.2460C>T found in Ragdoll | Autosomal Dominant (Incomplete Penetrance) | Clear |

Immunologic Disorders

| Disorder | Mode of Inheritance | Result |
|---|---------------------|--------|
| Autoimmune Lymphoproliferative Syndrome | Autosomal Recessive | Clear |
| Congenital Hypotrichosis with Short Life Expectancy | Autosomal Recessive | Clear |

Metabolic Disorders

| Disorder | Mode of Inheritance | Result |
|--|---------------------|--------|
| Acute Intermittent Porphyria | Autosomal Dominant | Clear |
| Acute Intermittent Porphyria; HMBS mutation: c.844delGAG | Autosomal Dominant | Clear |
| Congenital Erythropoietic Porphyria, mutation UROS: c.331G>A | Autosomal Recessive | Clear |





Test results - Additional disorders found in other breeds - page 2

Muscular Disorders

| Disorder | Mode of Inheritance | Result |
|--|---------------------|--------|
| Congenital Myasthenic Syndrome (CMS) | Autosomal Recessive | Clear |
| Periodic Hypokalemic Polymyopathy, Burmese Hypokalemia, or Familial Episodic Hypokalaemic Polymyopathy | Autosomal Recessive | Clear |
| Spinal Muscular Atrophy (SMA)/Spinal Muscular Dystrophy | Autosomal Recessive | Clear |

Neurologic Disorders

| Disorder | Mode of Inheritance | Result |
|---|---------------------|--------|
| GM2 Gangliosidosis, Japanese Domestic mutation HEXB: c.667C>T | Autosomal Recessive | Clear |

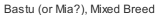
Neuromuscular Disorders

| Disorder | Mode of Inheritance | Result |
|----------------------------------|---------------------|--------|
| Glycogen Storage Disease Type IV | Autosomal Recessive | Clear |

Ocular Disorders

| Disorder | Mode of Inheritance | Result |
|----------------------------|---------------------|--------|
| Bengal Progressive Atrophy | Autosomal Recessive | Clear |
| Retinal Dystrophy (rdAc) | Autosomal Recessive | Clear |

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APPENDIX

Explanation of the results of the tested disorders

Autosomal recessive inheritance (ARI)

Clear - A cat carries no copies of the tested mutation and has no or reduced likelihood of developing and passing on the disease/condition.

Carrier - A cat carries one copy of the tested mutation. Carriers typically have a normal, healthy appearance but pass on the mutation to approximately 50% of their offspring.

At risk - A cat carries two copies of the tested mutation and is at high or increased risk of developing the disease/condition.

Autosomal dominant inheritance (ADI)

Clear - A cat carries no copies of the tested mutation and has no or reduced likelihood of developing and passing on the disease/condition.

At risk - A cat carries one or two copies of the tested mutation and is at high or increased risk of developing the disease/condition.

X-linked recessive inheritance (X-linked)

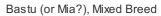
Clear - A cat carries no copies of the tested mutation and has no or reduced likelihood of developing and passing on the disease/condition.

Carrier - Female carriers typically have a normal, healthy appearance but carry one copy of the tested mutation on one of their X chromosomes. As males only have one X chromosome, there are no male carriers.

At risk - Female cats at risk carry two mutated copies of the tested mutation. Males carry one copy of the tested mutation on their single X chromosome. Cats at risk are at high or increased risk of developing the disease/condition.

Please note that the descriptions above are generalized based on typically observed inheritance patterns. When obtaining a 'carrier' or 'at risk' test result, always refer to the corresponding online test documentation for more detailed information on the condition and any exceptions.







OPTIMAL SELECTION™ FELINE DNA TEST TERMS AND CONDITIONS

Optimal Selection™ Genetic Breeding Analysis is a proprietary process designed and intended to be used on purebred cats solely to 1) Help quantify the genetic compatibility of potential breeding pairs and 2) To identify specific alleles or DNA mutations that are associated with certain inherited diseases or traits. No other purpose is authorized or permitted. It is not intended to diagnose diseases or predict behavior in any particular cat.

Upon receipt of your cat's DNA sample, Wisdom Health will analyze your cat's DNA to determine chromosomal similarities and differences in the genetic profile of a potential sire and dam and provide a match analysis. Your cat's DNA will also be analyzed for the presence of specific alleles that are associated with inherited conditions identified as occurring in your cat's breed. Wisdom Health's testing procedures are designed to provide reliable and accurate results, but are not guaranteed. By submitting your cat's sample(s) for Optimal Selection™ analysis it is understood that you agree that the sample(s), analysis, results and related information may be used confidentially by Mars in conjunction with other samples to increase the understanding of the breed's genetic structure, as well as for internal, research and development, or statistical purposes and may be shared with third parties for these purposes.

Samples may be disposed of or stored at Wisdom Health's option and will not be returned. Please view the full Mars Privacy Policy here: http://www.mars.com/global/policies/privacy/pp-english.aspx. It is also understood that future releases of the Optimal Selection™ test may refine results as more information is obtained regarding the breed structure and/or if new genetic markers are included.

Optimal Selection™ genetic assessments for individual cats and potential mates will be available online to the person(s) who registered the sample. A cat's results, photo and other information may be shared by the owner with other individuals whom they choose or transferred to a new owner if the cat changes ownership. The content of such online services 1) may be altered due to changes, additions, or removals of a cat's information in the Optimal Selection™ database or due to changes in technical or other design of such services and 2) includes information about third parties and other Wisdom Health clients' cats, which Wisdom Health is not responsible or liable for. Wisdom Health has right to terminate access to online services one year from the purchase date, unless a longer period has been agreed upon.

You agree to Wisdom Health instructions related to ordering process, payment, sampling and sample delivery. You also certify that the animal described in your order is the same animal whose sample is submitted for analysis, and that all information is accurate. You warrant that you are entitled to obtain and supply samples to Wisdom Health.

In the unlikely event that it is not possible to provide an analysis (for example due to an insufficient DNA sample) or that an error in the analysis occurs, liability by Wisdom Health or related companies and individuals is disclaimed and damages in any event are limited to the payment actually received by Wisdom Health for the specified analysis at issue. Wisdom Health's study of the complexities of the feline genome is ongoing with the goal of continuing to provide the most advanced and complete analysis possible.

Wisdom Health reserves the right to use any third party of its choice to undertake the testing, analysis or laboratory services for the analysis.