



Results for Lulu

Lulu's demographic profile:

Call Name:	Lulu	Owner:	Ashley Edens
Registered Name:	MMA CHROMED BUCKLE BUNNY	Registration #:	ASDM-TX-1701254
Breed:	Miniature Australian Shepherd	Microchip/Tattoo:	OAD2191516
Sex:	Female	Kit #:	11949
Approx. DOB:	September, 2016	Report Date:	January 18, 2019

Lulu's genetic health profile:

- ✓ Lulu is not at-risk for any of the diseases tested
- ✓ Lulu is not a carrier for any of the diseases tested

Lulu's appearance profile:

- ✓ Lulu is a **Female**
- ✓ Lulu's coat is likely **Straight, Long and Black with tan points** in color
- ✓ Lulu's face likely **Has a Mask on the Muzzle** and a **Black nose**
- ✓ Lulu's tail is likely **Normal** in length

These tests were developed and performed by Paw Print Genetics®, Spokane WA.

Explanation of Results

Normal	A "normal" result means that your dog does not have the mutation that causes the associated genetic disease.
Carrier	A "carrier" result indicates that your dog has inherited one copy of the mutation that has been reported to cause this genetic disease. Your dog may not be clinically affected by this mutation because two copies of the mutation are usually required to cause disease.
At-Risk	An "at-risk" result indicates that your dog may have inherited one or two copies of the mutation that has been reported to cause this genetic disease. Depending on the mode of genetic inheritance for this particular disease, inheriting one or two mutant copies of the gene may result in the disease. <i>You may want to consider ordering follow-up testing to confirm the results of this initial screen for any dog that is "at-risk" for a disease.</i>
Failed	A "failed" result indicates that we were unable to obtain a genotype for your dog for this specific disease or trait and does not mean that your dog is a carrier or at-risk for this

disease. There are a variety of reasons why a specific test may fail. However, we promise to provide at least 150 results to you for your dog.

Please review our [testing terms and disclaimers](#) regarding your results.

Blood and Clotting

Coagulation factor VII deficiency	Normal
Elliptocytosis	Normal
Glanzmann's thrombasthenia (Great Pyrenees type)	Normal
Glanzmann's thrombasthenia (Otterhound type)	Normal
Glycogen storage disease VII (Wachtelhund type)	Normal
Hemophilia A (Boxer type)	Normal
Hemophilia A (German Shepherd Dog, type 1)	Normal
Hemophilia A (German Shepherd Dog, type 2)	Normal
Hemophilia B (Cairn Terrier type)	Normal
Hemophilia B (Lhasa Apso type)	Normal
Hemophilia B (Rhodesian Ridgeback type)	Normal
Leukocyte adhesion deficiency, type III	Normal
May-Hegglin anomaly	Normal
P2RY12 receptor platelet disorder	Normal
Prekallikrein deficiency	Normal
Pyruvate kinase deficiency (Basenji type)	Normal
Pyruvate kinase deficiency (Beagle type)	Normal
Pyruvate kinase deficiency (Labrador Retriever type)	Normal
Pyruvate kinase deficiency (Pug type)	Normal
Pyruvate kinase deficiency (Terrier type)	Normal
Thrombopathia (American Eskimo Dog type)	Normal
Thrombopathia (Basset Hound type)	Normal
Thrombopathia (Newfoundland type)	Normal
Von Willebrand disease I	Normal
Von Willebrand disease II	Failed
Von Willebrand disease III (Kooikerhondje type)	Normal
Von Willebrand disease III (Scottish Terrier type)	Normal

Cancer

Renal cystadenocarcinoma and nodular dermatofibrosis	Normal
--	--------

Dental

Amelogenesis imperfecta	Normal
-------------------------	--------

Drug Metabolism

Multidrug resistance 1

Normal

Eyes

Collie eye anomaly	Normal
Cone degeneration	Normal
Cone degeneration (German Shorthaired Pointer type)	Normal
Congenital stationary night blindness	Normal
Dry eye curly coat syndrome	Normal
Early retinal degeneration	Normal
GM1 Gangliosidosis (Alaskan Husky type)	Normal
GM1 Gangliosidosis (Portuguese Water Dog type)	Normal
GM1 Gangliosidosis (Shiba Inu type)	Normal
GM2 Gangliosidosis (Poodle type)	Normal
Hereditary cataracts	Normal
Hereditary cataracts (Australian Shepherd type)	Normal
Juvenile Laryngeal Paralysis and Polyneuropathy	Normal
Multifocal retinopathy 1	Normal
Multifocal retinopathy 2	Normal
Multifocal retinopathy 3	Normal
Primary lens luxation	Normal
Primary open angle glaucoma	Normal
Progressive retinal atrophy (Basenji type)	Normal
Progressive retinal atrophy (Bullmastiff/Mastiff type)	Normal
Progressive retinal atrophy (Irish Setter type)	Normal
Progressive retinal atrophy (Sloughi type)	Normal
Progressive retinal atrophy, Cone-rod dystrophy 1	Normal
Progressive retinal atrophy, Cone-rod dystrophy 3	Normal
Progressive retinal atrophy, Golden Retriever 1	Normal
Progressive retinal atrophy, Golden Retriever 2	Normal
Progressive retinal atrophy, PRA1 (Papillon type)	Normal
Progressive retinal atrophy, Progressive rod-cone degeneration	Normal
Progressive retinal atrophy, Rod-cone dysplasia 3	Normal
Progressive retinal atrophy, generalized	Normal

Heart

Dilated cardiomyopathy

Failed

Immune System

Complement 3 deficiency	Normal
Leukocyte adhesion deficiency, type I	Normal
Leukocyte adhesion deficiency, type III	Normal
Primary ciliary dyskinesia	Normal

Severe combined immunodeficiency disease (Terrier type)	Normal
Severe combined immunodeficiency disease (Wetterhoun type)	Normal
Severe combined immunodeficiency disease, X-linked (Basset Hound type)	Normal
Severe combined immunodeficiency disease, X-linked (Corgi type)	Normal
Trapped neutrophil syndrome	Normal

Liver/Gastrointestinal

Gallbladder mucoceles	Normal
Glycogen storage disease IIIa	Normal
Intestinal cobalamin malabsorption (Beagle type)	Normal
Intestinal cobalamin malabsorption (Border Collie type)	Normal

Metabolic

Adult-onset neuronal ceroid lipofuscinosis	Normal
GM1 Gangliosidosis (Alaskan Husky type)	Normal
GM1 Gangliosidosis (Portuguese Water Dog type)	Normal
GM1 Gangliosidosis (Shiba Inu type)	Normal
GM2 Gangliosidosis (Japanese Chin type)	Normal
GM2 Gangliosidosis (Poodle type)	Normal
Globoid cell leukodystrophy (Irish Setter type)	Normal
Globoid cell leukodystrophy (Terrier type)	Normal
Glycogen storage disease IIIa	Normal
Glycogen storage disease Ia	Normal
Glycogen storage disease VII (Wachtelhund type)	Normal
Intestinal cobalamin malabsorption (Beagle type)	Normal
Intestinal cobalamin malabsorption (Border Collie type)	Normal
L-2-hydroxyglutaric aciduria (Staffordshire Bull Terrier type)	Normal
Mucopolysaccharidosis I	Normal
Mucopolysaccharidosis IIIA (Dachshund type)	Normal
Mucopolysaccharidosis IIIA (New Zealand Huntaway type)	Normal
Mucopolysaccharidosis VII (Shepherd type)	Normal
Neuronal ceroid lipofuscinosis 1	Normal
Neuronal ceroid lipofuscinosis 10	Normal
Neuronal ceroid lipofuscinosis 2	Normal
Neuronal ceroid lipofuscinosis 4A	Normal
Neuronal ceroid lipofuscinosis 5	Normal
Neuronal ceroid lipofuscinosis 6	Normal
Neuronal ceroid lipofuscinosis 8 (Australian Shepherd type)	Normal
Neuronal ceroid lipofuscinosis 8 (Setter type)	Normal
Pompe disease	Normal
Pyruvate dehydrogenase deficiency	Normal
Pyruvate kinase deficiency (Basenji type)	Normal
Pyruvate kinase deficiency (Beagle type)	Normal
Pyruvate kinase deficiency (Labrador Retriever type)	Normal
Pyruvate kinase deficiency (Pug type)	Normal

Pyruvate kinase deficiency (Terrier type)

Normal

Midline Defect

Juvenile Laryngeal Paralysis and Polyneuropathy

Normal

Spinal dysraphism (Weimaraner type)

Normal

Musculoskeletal

Adult-onset neuronal ceroid lipofuscinosis

Normal

Alaskan Malamute polyneuropathy

Normal

Chondrodysplasia (Karelian Bear Dog and Norwegian Elkhound type)

Normal

Congenital myasthenic syndrome (Labrador Retriever type)

Normal

Congenital myasthenic syndrome (Old Danish Pointer type)

Normal

Degenerative myelopathy

Normal

Exercise-induced collapse

Normal

GM1 Gangliosidosis (Alaskan Husky type)

Normal

GM1 Gangliosidosis (Portuguese Water Dog type)

Normal

GM1 Gangliosidosis (Shiba Inu type)

Normal

Glycogen storage disease IIIa

Normal

Glycogen storage disease VII (Wachtelhund type)

Normal

Greyhound polyneuropathy

Normal

Inherited myopathy of Great Danes

Normal

Juvenile Laryngeal Paralysis and Polyneuropathy

Normal

Mucopolysaccharidosis I

Normal

Mucopolysaccharidosis VII (Shepherd type)

Normal

Muscular Dystrophy (Golden Retriever Type)

Normal

Myostatin deficiency (Whippet and Longhaired Whippet type)

Normal

Myotonia congenita (Australian Cattle Dog type)

Normal

Myotonia congenita (Schnauzer type)

Normal

Myotubular myopathy 1

Normal

Osteogenesis imperfecta (Beagle type)

Normal

Osteogenesis imperfecta (Golden Retriever type)

Normal

Polyneuropathy (Leonberger and Saint Bernard type)

Normal

Pompe disease

Normal

Skeletal dysplasia 2

Normal

Vitamin D dependent rickets, type II (Pomeranian type)

Normal

Neurologic

Adult-onset neuronal ceroid lipofuscinosis

Normal

Alaskan Husky encephalopathy

Normal

Alaskan Malamute polyneuropathy

Normal

Benign familial juvenile epilepsy

Normal

Canine multiple system degeneration (Chinese Crested type)

Normal

Canine multiple system degeneration (Kerry Blue Terrier type)

Normal

Cerebellar ataxia (Finnish Hound type)	Normal
Congenital myasthenic syndrome (Labrador Retriever type)	Normal
Congenital myasthenic syndrome (Old Danish Pointer type)	Normal
Degenerative myelopathy	Normal
Exercise-induced collapse	Normal
GM1 Gangliosidosis (Alaskan Husky type)	Normal
GM1 Gangliosidosis (Portuguese Water Dog type)	Normal
GM1 Gangliosidosis (Shiba Inu type)	Normal
GM2 Gangliosidosis (Japanese Chin type)	Normal
GM2 Gangliosidosis (Poodle type)	Normal
Globoid cell leukodystrophy (Irish Setter type)	Normal
Globoid cell leukodystrophy (Terrier type)	Normal
Greyhound polyneuropathy	Normal
Juvenile Laryngeal Paralysis and Polyneuropathy	Normal
L-2-hydroxyglutaric aciduria (Staffordshire Bull Terrier type)	Normal
Late onset ataxia	Normal
Mucopolysaccharidosis I	Normal
Mucopolysaccharidosis IIIA (Dachshund type)	Normal
Mucopolysaccharidosis IIIA (New Zealand Huntaway type)	Normal
Myotonia congenita (Australian Cattle Dog type)	Normal
Myotonia congenita (Schnauzer type)	Normal
Narcolepsy (Dachshund type)	Normal
Narcolepsy (Doberman Pinscher type)	Normal
Narcolepsy (Labrador Retriever type)	Normal
Neonatal cerebellar cortical degeneration	Normal
Neonatal encephalopathy with seizures	Normal
Neuronal ceroid lipofuscinosis 1	Normal
Neuronal ceroid lipofuscinosis 10	Normal
Neuronal ceroid lipofuscinosis 2	Normal
Neuronal ceroid lipofuscinosis 4A	Normal
Neuronal ceroid lipofuscinosis 5	Normal
Neuronal ceroid lipofuscinosis 6	Normal
Neuronal ceroid lipofuscinosis 8 (Australian Shepherd type)	Normal
Neuronal ceroid lipofuscinosis 8 (Setter type)	Normal
Polyneuropathy (Leonberger and Saint Bernard type)	Normal
Sensory ataxic neuropathy	Normal
Spinocerebellar ataxia	Normal
Startle disease	Normal

Neuromuscular

Globoid cell leukodystrophy (Irish Setter type)	Normal
Globoid cell leukodystrophy (Terrier type)	Normal

Reproduction

Primary ciliary dyskinesia

Normal

Respiratory

Primary ciliary dyskinesia

Normal

Skin and Hair

Anhidrotic ectodermal dysplasia

Normal

Dry eye curly coat syndrome

Normal

Dystrophic epidermolysis bullosa

Normal

Ectodermal dysplasia

Normal

Epidermolytic hyperkeratosis

Normal

Hereditary footpad hyperkeratosis (Irish Terrier and Kromfohrländer type)

Normal

Hereditary nasal parakeratosis

Normal

Ichthyosis (Golden Retriever type)

Normal

Renal cystadenocarcinoma and nodular dermatofibrosis

Normal

Urinary Tract

Cystinuria (Australian Cattle Dog type)

Normal

Cystinuria (Miniature Pinscher type)

Normal

Cystinuria (Newfoundland type)

Normal

Familial nephropathy (Cocker Spaniel type)

Normal

Familial nephropathy (English Springer Spaniel type)

Normal

Hereditary nephritis (Samoyed type)

Normal

Hyperuricosuria

Normal

Persistent Müllerian duct syndrome

Normal

Primary ciliary dyskinesia

Normal

Primary hyperoxaluria

Normal

Renal cystadenocarcinoma and nodular dermatofibrosis

Normal