## **DNA** Test Report

Owner Info		
First Name	Last Name	
Roxann	Vass	
Pet Info		
Registered Name	Date of Birth	
Creekcats Hollywood	4/2/2023	
Nickname (Call Name)	Sample ID	
Creekcats Hollywood	FWBCDLX	
Sex	Registration	
emale	N/A	
Country of Origin	Microchip ID	
JS	N/A	
Dwner Reported Breed	Tattoo ID	
Ragdoll	N/A	

### **DNA Test Report**

Sample ID: FWBCDLX Test Date: 5/16/2023 Optimal Selection - Feline

#### Genetic Diversity (Heterozygosity)

#### Creekcats Hollywood's Percentage of Heterozygosity

35%

Creekcats Hollywood's genome analysis shows an average level of genetic heterozygosity when compared with other Ragdolls. **Typical Range for Ragdolls** 32 - 37%

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#### Health Conditions Known in This Breed

Genetic Condition	Gene	Risk Variant	Copies	Result
Factor XII Deficiency (Variant 1)	F12	Deletion	0	Clear
Factor XII Deficiency (Variant 2)	F12	Deletion	0	Clear
Hypertrophic Cardiomyopathy (Discovered in the Ragdoll)	MYBPC	C>T	0	Clear
Polycystic Kidney Disease (PKD)	PKD1	C>A	0	Clear
Other Conditions Tested				
Genetic Condition	Gene	Risk Variant	Copies	Result
Acute Intermittent Porphyria (Variant 1)	HMBS	Deletion	0	Clear
Acute Intermittent Porphyria (Variant 2)	HMBS	G>A	0	Clear
Acute Intermittent Porphyria (Variant 3)	HMBS	Insertion	0	Clear
Acute Intermittent Porphyria (Variant 4)	HMBS	Deletion	0	Clear
Acute Intermittent Porphyria (Variant 5)	HMBS	G>A	0	Clear
Autoimmune Lymphoproliferative Syndrome	FASL	Insertion	0	Clear
Burmese Head Defect (Discovered in the Burmese)	ALX1	Deletion	0	Clear
Chediak-Higashi Syndrome (Discovered in the Persian)	LYST	Insertion	0	Clear
Congenital Adrenal Hyperplasia	CYP11B1	G>A	0	Clear
Congenital Erythropoietic Porphyria	UROS	G>A	0	Clear
Congenital Myasthenic Syndrome (Discovered in the Devon Rex and Sphynx)	COLQ	G>A	0	Clear
Cystinuria Type 1A	SCL3A1	C>T	0	Clear
Cystinuria Type B (Variant 1)	SCL7A9	C>T	0	Clear
Cystinuria Type B (Variant 2)	SCL7A9	G>A	0	Clear
Cystinuria Type B (Variant 3)	SCL7A9	T>A	0	Clear
Dihydropyrimidinase Deficiency	DPYS	G>A	0	Clear

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#### Other Conditions Tested (continued)

Genetic Condition	Gene	Risk Variant	Copies	Result
Earfold and Osteochondrodysplasia (Discovered in the Scottish Fold)	TRPV4	G>T	0	Clear
Familial Episodic Hypokalemic Polymyopathy (Discovered in the Burmese)	WNK4	C>T	0	Clear
Glutaric Aciduria Type II	ETFDH	T>G	0	Clear
Glycogen Storage Disease (Discovered in the Norwegian Forest Cat)	GBE1	Insertion	0	Clear
GM1 Gangliosidosis	GLB1	G>C	0	Clear
GM2 Gangliosidosis	GM2A	Deletion	0	Clear
GM2 Gangliosidosis Type II (Discovered in Domestic Shorthair cats)	HEXB	Insertion	0	Clear
GM2 Gangliosidosis Type II (Discovered in Japanese domestic cats)	HEXB	C>T	0	Clear
GM2 Gangliosidosis Type II (Discovered in the Burmese)	HEXB	0>0	0	Clear
Hemophilia B (Variant 1)	F9	C>T	0	Clear
Hemophilia B (Variant 2)	F9	G>A	0	Clear
Hyperoxaluria Type II	GRHPR	G>A	0	Clear
Hypertrophic Cardiomyopathy (Discovered in the Maine Coon)	MYBPC	G>C	0	Clear
Hypotrichosis (Discovered in the Birman)	FOXN1	Deletion	0	Clear
Lipoprotein Lipase Deficiency	LPL	G>A	0	Clear
MDR1 Medication Sensitivity	ABCB1	Deletion	0	Clear
Mucopolysaccharidosis Type I	IDUA	Deletion	0	Clear
Mucopolysaccharidosis Type VI	ARSB	T>C	0	Clear
Mucopolysaccharidosis Type VI Modifier	ARSB	G>A	0	Clear
Mucopolysaccharidosis Type VII (Variant 1)	GUSB	G>A	0	Clear
Mucopolysaccharidosis Type VII (Variant 2)	USB	C>T	0	Clear
Myotonia Congenita	CLCN1	G>T	0	Clear

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#### Other Conditions Tested (continued)

Genetic Condition	Gene	Risk Variant	Copies	Result
Progressive Retinal Atrophy (Discovered in the Abyssinian)	CEP290	T>G	0	Clear
Progressive Retinal Atrophy (Discovered in the Bengal)	KIF3B	G>A	0	Clear
Progressive Retinal Atrophy (Discovered in the Persian)	AIPL1	C>T	0	Clear
Pyruvate Kinase Deficiency	PKLR	G>A	0	Clear
Sphingomyelinosis (Variant 1)	NPC1	G>C	0	Clear
Sphingomyelinosis (Variant 2)	NPC2	G>A	0	Clear
Spinal Muscular Atrophy (Discovered in the Maine Coon)	LIX1	Deletion	0	Clear
Vitamin D-Dependent Rickets	CYP27B1	G>T	0	Clear

c variant - Causes AB Blood Type

### **DNA Test Report**

#### Blood Type Blood Type Genotype А A/A (Most common) **Breeding Risk Transfusion Risk** Moderate Low Creekcats Hollywood has the most common If breeding, Creekcats Hollywood has a low blood type. She can be transfused with Type risk of blood type incompatibility with nursing A blood. kittens. Variant Tested Description Copies b variant 1 (Common b variant) 0 (Discovered in Turkish breeds) b variant 2 0 b variant 3 (Discovered in Ragdolls) 0

(Discovered in Ragdolls)

0

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#### Coat Color

Gene	Variant	Copies	Result
ASIP	A <sup>Pb</sup>	0	No effect
ASIP	а	2	Solid color hairs likely
KIT	w <sup>g</sup>	1	No effect
KIT	W or w <sup>s</sup>	1	Partly or fully white coat likely
MC1R	e	0	No effect
MC1R	e <sup>r</sup>	0	No effect
MLPH	d	2	Lightened coat color likely
TYR	C <sup>a</sup>	0	No effect
TYR	c <sup>b</sup>	0	No effect
TYR	c <sup>°</sup>	2	Siamese colorpoint pattern likely
TYR	c <sup>m</sup>	0	No effect
TYRP	b	0	No effect
TYRP	b	0	No effect
	ASIP ASIP KIT KIT MC1R MC1R MLPH TYR TYR TYR	ASIPAASIPaKITw <sup>9</sup> KITWorw <sup>9</sup> MCIReMCIRe <sup>1</sup> MCIRc <sup>8</sup> TYRc <sup>8</sup> TYRc <sup>9</sup> TYRc <sup>9</sup> TYRb	ASIP A <sup>Pb</sup> O   ASIP a 2   KIT w <sup>g</sup> 1   KIT Worw <sup>s</sup> 1   MCIR e O   MCIR e <sup>r</sup> O   MCIR of O   TYR c <sup>a</sup> O   TYR c <sup>b</sup> O   TYR c <sup>b</sup> O   TYR c <sup>b</sup> O   TYR b O

#### Coat Type

Genetic Trait	Gene	Variant	Copies	Result
Glitter	Confidential	_	0	No effect
Long Hair (Discovered in many breeds)	FGF5	M4	2	Long coat likely
Long Hair (Discovered in the Norwegian Forest Cat)	FGF5	M2	0	No effect
Long Hair (Discovered in the Ragdoll and Maine Coon)	FGF5	M3	0	No effect
Long Hair (Discovered in the Ragdoll)	FGF5	M1	0	No effect
Lykoi Coat (Variant 1)	HR	hr <sup>Ca</sup>	0	No effect

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Sample ID: FWBCDLX Test Date: 5/16/2023 Optimal Selection - Feline

### Coat Type (continued)

Genetic Trait	Gene	Variant	Copies	Result
Lykoi Coat (Variant 2)	HR	hr <sup>VA</sup>	0	No effect
Hairlessness (Discovered in the Sphynx)	KRT71	re <sup>hr</sup>	0	No effect
Rexing (Discovered in the Devon Rex)	KRT71	re <sup>dr</sup>	0	No effect
Rexing (Discovered in the Cornish Rex and German Rex)	LPAR6	r	0	No effect
Tail Length				
Genetic Trait	Gene	Variant	Copies	Result
Short Tail (Variant 3)	HES7	jb	0	No effect
Short Tail (Variant 1)	т	C1199del	0	No effect
Short Tail (Variant 2)	т	T988del	0	No effect
Extra Toes				
Genetic Trait	Gene	Variant	Copies	Result
Polydactyly (Variant 1)	LIMBR1	HW	0	No effect
Polydactyly (Variant 2)	LIMBR1	UK1	0	No effect
Polydactyly (Variant 3)	LIMBR1	UK2	0	No effect