

Result certificate #012345

Detection of mutations in SLC7A9 gene related to cystinurie type B development in cats

Customer: Jan Novák, Dlouhá 1, 30000 Plzeň, Czech Republic

Sample:

Sample: 08-12346 Date received: 01.01.2008 Sample type: buccal swab

Information provided by the customer

Name: Madame Théophile DEMO

Breed: Persian catDate of birth: 31.12.1909

Reg. number: (CZ)ABCD EF 123/45/XYZ

Microchip: 123456789012345

Sex: female

Date of sampling: 01.01.2008

The identity of the animal has been checked.

Result: PREVIEW RESULT LINE

Explanation

Presence or absence of c.706G>A, c.1175C>T, c.881T>A mutations in SLC7A9 gene related to cystinurie type B development in cats were tested. Mutation c.881T>A has been identified in Maine Coon, Siamese and Siberian cats and in the Sphynx breed. Cystinuria is a disorder of amino acid transport in the kidneys. In affected cats the accumulation of cystine in the urine occurs, the formation of cystine crystals and the formation of urinary stones. These irritate the urinary tract, causing stagnation of urine and increasing the susceptibility of the urinary tract to infection. Clinically, cystinuria is manifested by dark and distinctly foul-smelling urine with an admixture of blood. Urination is painful and often unsuccessful.

The N/N result means that the cat is not at risk of cystinuria B development. The presence of mutation in homozygous (result P/P) or heterozygous (result N/P) state means that the cat is at risk of cystinuria B development. Not enough data are yet available to determine the mode of inheritance. Autosomal recessive or autosomal dominant inheritance is assumed.

Method: SOPAgriseq_feline, ngs

Date of issue: 06.01.2008

Date of testing: 12.06.2008 - 06.01.2008

Approved by: Mgr. Martina Šafrová, Laboratory Manager



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