

Result certificate #240186

Detection of mutation c.[1297T>C;1299C>T] of L2HGDH gene causing L2HGA in Staffordshire bull terriers

Customer: Jaroslav Zavadil, Vlasatice 355, 69130 Vlasatice , Czech Republic Sample: Sample: 23-42420 Date received: 06.09.2023 Sample type: buccal swab Information provided by the customer Name: Viktory z Hambalek Bušín Breed: Staffordshire Bull Terrier Microchip: 203164000137927 Reg. number: CMKU/SBT/16390/21 Date of birth: 27.07.2021 Sex: female The identity of the animal has been checked by Bc. Eva Šandarová, Genomia s.r.o.

Result: Mutation was not detected (N/N)

Legend: N/N = wild-type genotype. N/P = carrier of the mutation. P/P = mutated genotype (individual will be most probably affected with the disease). (N = negative, P = positive)

Explanation

Presence or absence of mutation c.[1297T>C;1299C>T] in exon 10 of L2HGDH gene causing L-2hydroxyglutaric aciduria in Staffordshire bull terriers were tested. Double nucleotide substitution leads to double amino acid replacement in L2-hydroglutaric dehydrogenase enzyme: leucine to proline in position 433 and histidine to tyrosine in position 434. These mutations cause L-2-hydroxyglutaric aciduria disease (L-2-HGA) in Staffordshire bull terriers. L-2-HGA is a neurological disorder manifested by psychomotor retardation, seizures and ataxia. Accumulation of L-2-hydroxyglutaric acid in cerebrospinal fluid, plasma and urine is very typical.

Mutation that causes L-2-HGA is inherited as an autosomal recessive trait. That means the disease affects dogs with P/P genotype only. The dogs with N/P genotype are considered carriers of the disease (heterozygotes). In offspring of two heterozygous animals following genotype distribution can be expected: 25 % N/N, 25 % P/P and 50 % N/P.

Method: SOP175-L2HGA, real-time PCR-ASA

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