

For in vitro diagnostic use



APO-E (CYS112ARG) T3932C POLYMORPHISM

ORDERING INFORMATIONS

REF: GEN-008-25 RDM Code: 2255489/R Tests: 25 Reactions: 31 REF: GEN-008-50 RDM Code: 1735881/R Tests: 50 Reactions: 62 CND Code: W0106010499 Manufacturer: BioMol Laboratories s.r.l.

CONTENTS OF THE KIT

The kit consists of: reagents for Real-Time PCR amplification *reagents for the extraction of genomic DNA are not supplied in the kit

PRODUCT CHARACTERISTICS

Detection of T3932C polymorphism (also called C112R, Cys-Arg) of the APO-E gene by Real-Time PCR technique. Kit optimized for Real-Time PCR instrumentation Biorad CFX96 Dx, Biorad Opus Dx and Agilent AriaDx, Hyris bCUBE e Hyris bCUBE3 with Hyris bAPP.

SCIENTIFIC BACKGROUND

The genetic origin of the three variants of the human apolipoprotein E (apoE) protein, known as E2, E3, and E4, was understood in 1981. The underlying genetic variants of these protein isoforms, known as ϵ_2 , ϵ_3 , and ϵ_4 , are allelic forms of the APOE gene, resulting from different haplotypes at the APOE locus (19q13.31). In particular, APOE is polymorphic with three main alleles (e2, e3 and e4): APOE- ϵ_2 (cys112, cys158), APOE- ϵ_3 (cys112, arg158) and APOE- ϵ_4 (arg112, arg158). Although these allelic forms differ from each other by only one or two amino acids at positions 112 and 158, these differences alter the structure and function of APOE.

CLINICAL SIGNIFICANCE

The combination of the various polymorphisms is responsible for some risk conditions:

- ϵ_2 (rs7412-T, rs429358-T) has an allele frequency of about 7%. This apolipoprotein variant binds poorly to cell surface receptors while E3 and E4 bind well. Individuals with an e2/e2 combination may have an increased risk of early vascular disease. The e2 allele has also been implicated in Parkinson's disease.

- ϵ 3 (rs7412-C, rs429358-T) has an allele frequency of approximately 79%. It is considered the "neutral" Apo E genotype.

- ϵ 4 (rs7412-C, rs429358-C) has an allele frequency of approximately 14%. ϵ 4 has been implicated in atherosclerosis, Alzheimer's disease, decreased cognition, decreased hippocampal volume, time to disease progression in multiple sclerosis, poor

§ The APOE E4 Allele Confers Increased Risk of Ischemic Stroke Among Greek Carriers. Adv Clin Exp Med. 2016 May-Jun; 25 (3):471-8.

§ Plasma levels of apolipoprotein E, APOE genotype and risk of dementia and ischemic heart disease: A review Atherosclerosis. 2016 Dec; 255: 145-155.

§ APOE epsilon 4 allele predicts faster cognitive decline in mild Alzheimer disease. Neurology 70:1842–1849. Cosentino S, Scarmeas N, Helzner E, Clymour MM, Brandt J, et al. (2008).

§ Genetics of healthy aging and longevity. Hum Genet. 2013 Dec; 132 (12):1323-38. doi

BIOMOL LABORATORIES S.R.L.

Via Arcora 110 (Palazzo Gecos) 80013 Casalnuovo di Napoli, NA info@biomollaboratories.com

biomollaboratories.it







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DESCRIPTION	LABEL	VOLUME		STORAGE
		GEN-008-25	GEN-008-50	
Mix oligonucleotides and probes	Mix T3932C APO-E 10X	1 x 85 µl	1 x 170 µl	-20°C
Mix buffer and Taq polymerase enzyme	Mix Real-Time PCR 2X	1 x 425 µl	1 x 850 µl	-20°C
Deionized H ₂ O	Deionized H ₂ 0	2 x 1 ml	2 x 1 ml	-20°C
Genomic DNA or recombinant DNA	Control +1	1 x 22 µl	1 x 22 µl	-20°C
Genomic DNA or recombinant DNA	Control +2	1 x 22 µl	1 x 22 µl	-20°C
Genomic DNA or recombinant DNA	Control +3	1 x 22 µl	1 x 22 µl	-20°C

TECHNICAL CHARACTERISTICS

COD. GEN-008-25 / COD. GEN-008-50

STABILITY	18 months
REAGENTS STATUS	Ready to use
BIOLOGICAL MATRIX	Genomic DNA extracted from whole blood, tissue, cells
POSITIVE CONTROL	Recombinant DNA for at least 3 analytical sessions
VALIDATED INSTRUMENTS	Biorad CFX96 Dx, Biorad Opus Dx e Agilent AriaDx, Hyris bCUBE, Hyris bCUBE3 with Hyris bAPP.
TECHNOLOGY	Real-time PCR; oligonucleotides and specific probes; 2 FAM/HEX fluorescence channels
RUNNING TIME	85 min
THERMAL CYCLING PROFILE	1 cycle at 95 °C (10 min); 45 cycles at 95 °C (15 sec) + 60 °C (60 sec)
ANALYTICAL SPECIFICITY	Absence of non-specific pairings of oligonucleotides and probes; absence of cross-reactivity
ANALYTICAL SENSITIVITY : LIMIT OF DETECTION (LOD)	≥ 0,016 ng of DNA
ANALYTICAL SENSITIVITY : LIMIT OF BLANK (LOB)	0% NCN
REPRODUCIBILITY	99,9%
DIAGNOSTIC SPECIFICITY / DIAGNOSTIC SENSITIVITY	100%/98%

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