

GENETIC TEST:**Becker muscular dystrophy / Duchenne muscular dystrophy (Full sequencing DMD gene through Myopathy gene panel)**

FULL NAME:	Becker muscular dystrophy / Duchenne muscular dystrophy (Full sequencing DMD gene through Myopathy gene panel)
TEST TYPE:	Clinical
TEST SPECIALTY:	Molecular Genetics
TEST PURPOSE:	Carrier diagnosis, Post-natal Diagnosis
SPECIMEN:	Peripheral (whole) blood on EDTA, Muscle biopsy
METHOD CATEGORY:	Sequence analysis: entire coding region
METHOD TECHNIQUE:	Next Generation Sequencing (NGS) Bi-directional Sanger Sequence analysis
RIZIV CODE:	565471-565482
ACCREDITATION (ISO 15189):	2023-11-09 / 2024-05-08

EQA:	• Duchenne / Becker Muscular Dystrophy, • Duchenne / Becker Muscular Dystrophy, • Duchenne / Becker Muscular Dystrophy
TURNAROUND TIME (MAXIMUM):	4 months
CREATED:	27 Jul 2018 - 11:04
CHANGED:	22 Jan 2024 - 11:35
URL:	https://labogidsmedgen.uza.be/analyses/duchenne-becker-spierdystrofie-stap-2-ui...

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