

GENETIC TEST:
Homocystinuria (hot spot mutation - c.677C>T)

FULL NAME:	Homocystinuria (hot spot mutation - c.677C>T)
TEST TYPE:	Clinical
TEST SPECIALTY:	Molecular Genetics
TEST PURPOSE:	Post-natal Diagnosis
SPECIMEN:	Peripheral (whole) blood on EDTA
METHOD CATEGORY:	Targeted variant analysis
METHOD TECHNIQUE:	Next Generation Sequencing (NGS)
RIZIV CODE:	565331-565342
EQA:	• MTHFR (Set 01-D)
TURNAROUND TIME (MAXIMUM):	2 months
CREATED:	30 Aug 2019 - 11:14
CHANGED:	13 Dec 2022 - 11:54

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