

Variant: NM_000277.2(PAH):c.355C>T (p.Pro119Ser)

CA220582 [↗](#)

92741 (ClinVar) [↗](#)

Gene: PAH ([HGNC:5053](#))
Condition: phenylketonuria ([MONDO:0009861](#))
Inheritance Mode: Autosomal recessive inheritance
UUID: 38a183ee-7e53-4c24-a250-50958f390c78
Approved on: 2018-07-29
Published on: 2018-12-10

HGVS expressions

NM_000277.2(PAH):c.355C>T (p.Pro119Ser)
NM_000277.2:c.355C>T
NC_000012.12:g.102877548G>A
CM000674.2:g.102877548G>A
NC_000012.11:g.103271326G>A
CM000674.1:g.103271326G>A
NC_000012.10:g.101795456G>A
NG_008690.1:g.45055C>T
NG_008690.2:g.85863C>T
NM_000277.1:c.355C>T
NM_001354304.1:c.355C>T
NM_000277.3:c.355C>T
ENST00000307000.7:c.340C>T
ENST00000549111.5:n.451C>T
ENST00000550978.6:n.339C>T
ENST00000551337.5:c.355C>T
ENST00000551988.5:n.444C>T
ENST00000553106.5:c.355C>T

Likely Pathogenic

Met criteria codes **3**

PM3_Strong PP3 PP4_Moderate

Not Met criteria codes **1**

PM2

Evidence Links **2**

Expert Panel

Phenylketonuria VCEP [↗](#)

Criteria Specification Information **!**

[↗](#) Criteria Specifications for this VCEP

Evidence submitted by expert panel

Phenylketonuria VCEP

PAH-specific ACMG/AMP criteria applied: PP3: in silico analysis supportive of damaging effect; PM3_Strong: In trans with R261Q (PMID 21147011), and in trans with IVS2+1G>A (PMID 12655554) (PMID:21147011; PMID:12655554); PP4_Moderate: BH4 deficiency excluded (PMID:21147011). In summary this variant meets criteria to be

classified as likely pathogenic for phenylketonuria in an autosomal recessive manner based on the ACMG/AMP criteria applied as specified by the PAH Expert Panel: (PP3, PM3_Strong, PP4_Moderate).

Met criteria codes

PM3_Strong			In trans with R261Q (PMID 21147011), and in trans with IVS2+1G>A (PMID 12655554) in trans with IVS2+1G>A PubMed:12655554  in trans with R261Q PubMed:21147011 
PP3			in silico analysis supportive of damaging effect
PP4_Moderate			BH4 deficiency excluded BH4 defect excluded at NBS and by sequencing of BH4 associated genes PubMed:21147011 

Not Met criteria codes

PM2			PAH specific specifications state PM2 criteria as 0.02% (AF=0.0002)
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Curation History



 

 

See Report	Preferred Variant Title	Classification 	Condition	Published Date	Version 	Criteria Specification	Gene
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No matching records found

The information on this website is not intended for direct diagnostic use or medical decision-making without review by a genetics professional. Individuals should not change their health behavior solely on the basis of information contained on this website. If you have questions about the information contained on this website, please see a health care professional.