

- No Criteria Specification was provided in the publication event for the variant classifaction.
- The **Criteria Specification** shown below is a best-estimate based on the VCEP, gene, disease, and publication date.

Variant: NM_000277.2(PAH):c.503delA (p.Tyr168Serfs)

CA229588

102705 (ClinVar)

Gene: PAH ([HGNC:5053](#))
Condition: phenylketonuria ([MONDO:0009861](#))
Inheritance Mode: Autosomal recessive inheritance
UUID: 7a494430-377e-4473-a5f7-e0698ec51cab
Approved on: 2018-08-10
Published on: 2018-12-10

HGVS expressions

NM_000277.2(PAH):c.503delA (p.Tyr168Serfs)
NM_000277.2:c.503delA
NC_000012.12:g.102866602del
CM000674.2:g.102866602del
NC_000012.11:g.103260380del
CM000674.1:g.103260380del
NC_000012.10:g.101784510del
NG_008690.1:g.56001del
NG_008690.2:g.96809del
ENST00000553106.6:c.503del
ENST00000307000.7:c.488del
ENST00000549111.5:n.599del
ENST00000551988.5:n.530+10860del
ENST00000553106.5:c.503del
NM_000277.1:c.503del
NM_000277.2:c.503del
NM_001354304.1:c.503del
NM_000277.3:c.503del
NM_001354304.2:c.503del
NM_000277.3(PAH):c.503del (p.Tyr168fs)
More

Pathogenic

Met criteria codes 3
PM2 PP4 PVS1

Evidence Links 1

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: PVS1: Frameshift variant; PM2: Extremely low frequency. ExAC MAF: 0.00001.; PP4: Detected in PKU patient in international phase II clinical trial for sapropterin. (PMID:23430918). In summary this variant meets criteria to be classified as pathogenic for phenylketonuria in an autosomal recessive manner based on the ACMG/AMP criteria applied as specified by the PAH Expert Panel: (PVS1, PM2, PP4).
Met criteria codes
PM2
Extremely low frequency. ExAC MAF: 0.00001.
PP4
Detected in PKU patient in international phase II clinical trial for sapropterin.

c.503delA (p.Y168>Sfs) detected in study. [PubMed:23430918](#)

PVS1
Frameshift variant

Curation History

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.



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