

Variant: *NM_000527.5(LDLR):c.858C>A (p.Ser286Arg)*

Version: 1.0

[CA029904](#)

[251488 \(ClinVar\)](#)

Gene: LDLR ([HGNC:3949](#))

Condition: hypercholesterolemia, familial ([MONDO:0007750](#))

Inheritance Mode: Semidominant inheritance

UUID: 7d35f7d3-b40a-4137-ace3-7d0c87f989f6

Approved on: 2022-01-19

Published on: 2022-06-30

HGVS expressions

NM_000527.5:c.858C>A

NM_000527.5(LDLR):c.858C>A (p.Ser286Arg)

NC_000019.10:g.11107432C>A

CM000681.2:g.11107432C>A

NC_000019.9:g.11218108C>A

CM000681.1:g.11218108C>A

NC_000019.8:g.11079108C>A

NG_009060.1:g.23052C>A

ENST00000252444.10:c.1116C>A

ENST00000559340.2:c.858C>A

ENST00000560467.2:c.858C>A

ENST00000558518.6:c.858C>A

ENST00000252444.9:c.1112C>A

ENST00000455727.6:c.354C>A

ENST00000535915.5:c.735C>A

ENST00000545707.5:c.477C>A

ENST00000557933.5:c.858C>A

ENST00000558013.5:c.858C>A

ENST00000558518.5:c.858C>A

ENST00000558528.1:n.373C>A

ENST00000560467.1:c.458C>A

NM_000527.4:c.858C>A

NM_001195798.1:c.858C>A

NM_001195799.1:c.735C>A

NM_001195800.1:c.354C>A

NM_001195803.1:c.477C>A

NM_001195798.2:c.858C>A

NM_001195799.2:c.735C>A

NM_001195800.2:c.354C>A

NM_001195803.2:c.477C>A

Likely Pathogenic

Met criteria codes **5**

PP1 PP3 PP4 PM2

PS4_Supporting

Expert Panel

[Familial Hypercholesterolemia VCEP](#)

Criteria Specification Information

Not Met criteria codes **21**

[Criteria Specifications for this VCEP](#)

PS1 PS2 PS3 BA1 PP2
PM1 PM3 PM5 PM4 PM6
BS2 BS1 BS4 BS3 BP4 BP3
BP1 BP2 BP5 BP7 PVS1

Evidence Links **0**

Evidence submitted by expert panel

Familial Hypercholesterolemia VCEP

NM_000527.5(LDLR):c.858C>A (p.Ser286Arg) variant is classified as Likely pathogenic for Familial Hypercholesterolemia by applying evidence codes (PM2, PS4_Supporting, PP1, PP3 and PP4) as defined by the ClinGen Familial Hypercholesterolemia Expert Panel LDLR-specific variant curation guidelines (<https://doi.org/10.1016/j.gim.2021.09.012>). The supporting evidence is as follows: PM2 - PopMax MAF = 0.00006155 (0.006155%) in European (non-Finnish) exomes+genomes (gnomAD v2.1.1). PS4_Supporting - Variant meets PM2 and is identified in 4 unrelated index cases who fulfil DLCN ≥ 6 (1 case in PMID: 32770674; 2 cases from Robarts Research Institute) or SB definitive (1 case from Molecular Genetics Laboratory of Centre for Cardiovascular Surgery and Transplantation). PP1 - Variant segregates with FH phenotype in 2 informative meioses from 2 unrelated families (1 case from Molecular Genetics Laboratory of Centre for Cardiovascular Surgery and Transplantation and 1 case from Robarts Research Institute). PP3 - REVEL = 0.766. PP4 - Variant meets PM2 and is identified in 4 index cases who fulfil DLCN ≥ 6 (1 case in PMID: 32770674; 2 cases from Robarts Research Institute) or SB definitive (1 case from Molecular Genetics Laboratory of Centre for Cardiovascular Surgery and Transplantation).

Met criteria codes

PP1	✓	Variant segregates with FH phenotype in 2 informative meioses from 2 unrelated families (1 case from Molecular Genetics Laboratory of Centre for Cardiovascular Surgery and Transplantation and 1 case from Robarts Research Institute)
PP3	✓	REVEL = 0.766.
PP4	✓	Variant meets PM2 and is identified in 4 index cases who fulfil DLCN ≥ 6 (1 case in PMID: 32770674; 2 cases from Robarts Research Institute) or SB definitive (1 case from Molecular Genetics Laboratory of Centre for Cardiovascular Surgery and Transplantation).
PM2	✓	PopMax MAF = 0.00006155 (0.006155%) in European (non-Finnish) exomes+genomes (gnomAD v2.1.1).
PS4_Supporting	✓	Variant meets PM2 and is identified in 4 unrelated index cases who fulfil DLCN ≥ 6 (1 case in PMID: 32770674; 2 cases from Robarts Research Institute) or SB definitive (1 case from Molecular Genetics Laboratory of Centre for Cardiovascular Surgery and Transplantation).

Not Met criteria codes

PS1	✗	No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
------------	---	--

PS2	✘	No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
PS3	✘	No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
BA1	✘	No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
PP2	✘	No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
PM1	✘	No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
PM3	✘	No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
PM5	✘	No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
PM4	✘	No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
PM6	✘	No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
BS2	✘	No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
BS1	✘	No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
BS4	✘	No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
BS3	✘	No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
BP4	✘	No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
BP3	✘	No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline

BP1	✘	No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
BP2	✘	No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
BP5	✘	No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
BP7	✘	No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
PVS1	✘	No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline

Curation History [↗](#)

Showing 1 to 1 of 1 rows

--

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.