

*Variant: NM_001033855.3(DCLRE1C):c.103C>G
(p.His35Asp)*

Version: 1.0

CA117007 [↗](#)

4674 (ClinVar) [↗](#)

Gene: DCLRE1C ([HGNC:64421](#))

Condition: severe combined immunodeficiency due to DCLRE1C deficiency ([MONDO:0011225](#))

Inheritance Mode: Autosomal recessive inheritance

UUID: 50baf4df-88c8-469b-a6eb-3b8a7fecc340

Approved on: 2024-01-23

Published on: 2024-01-23

HGVS expressions

NM_001033855.3:c.103C>G

NM_001033855.3(DCLRE1C):c.103C>G (p.His35Asp)

NC_000010.11:g.14953908G>C

CM000672.2:g.14953908G>C

NC_000010.10:g.14995907G>C

CM000672.1:g.14995907G>C

NC_000010.9:g.15035913G>C

NG_007276.1:g.5188C>G

ENST00000378241.6:c.103C>G

ENST00000456122.2:c.103C>G

ENST00000489161.2:c.103C>G

ENST00000492201.6:c.103C>G

ENST00000697047.1:c.103C>G

ENST00000697070.1:c.103C>G

ENST00000697071.1:c.103C>G

ENST00000697072.1:c.103C>G

ENST00000697073.1:c.103C>G

ENST00000697074.1:c.103C>G

ENST00000697075.1:c.103C>G

ENST00000697076.1:c.103C>G

ENST00000697077.1:c.103C>G

ENST00000697078.1:c.103C>G

ENST00000697080.1:c.103C>G

ENST00000697081.1:c.103C>G

ENST00000697082.1:c.103C>G

ENST00000697083.1:c.103C>G

ENST00000697084.1:c.103C>G

ENST00000697085.1:c.103C>G

ENST00000697087.1:c.103C>G

ENST00000697088.1:c.103C>G

ENST00000697089.1:c.103C>G

ENST00000697090.1:n.26C>G

ENST00000697091.1:n.164C>G

ENST00000378278.7:c.103C>G

ENST00000357717.6:c.-102C>G

ENST00000378241.5:c.-470C>G
ENST00000378246.6:c.-187C>G
ENST00000378249.5:c.-135C>G
ENST00000378254.5:c.-389C>G
ENST00000378255.5:c.-711C>G
ENST00000378258.5:c.-343C>G
ENST00000378278.6:c.103C>G
ENST00000378289.8:c.103C>G
ENST00000396817.6:c.-665C>G
ENST00000418843.5:c.-426C>G
ENST00000456122.1:c.-594C>G
NM_001033855.2:c.103C>G
NM_001033857.2:c.-343C>G
NM_001033858.2:c.-665C>G
NM_001289076.1:c.-102C>G
NM_001289077.1:c.-389C>G
NM_001289078.1:c.-135C>G
NM_001289079.1:c.-711C>G
NM_022487.3:c.-187C>G
NR_110297.1:n.525C>G
NM_001350965.1:c.103C>G
NM_001350966.1:c.-135C>G
NM_001350967.1:c.-343C>G
NR_146960.1:n.525C>G
NR_146961.1:n.525C>G
NR_146962.1:n.525C>G
NM_001033857.3:c.-343C>G
NM_001033858.3:c.-665C>G
NM_001289076.2:c.-102C>G
NM_001289077.2:c.-389C>G
NM_001289078.2:c.-135C>G
NM_001289079.2:c.-711C>G
NM_001350965.2:c.103C>G
NM_001350966.2:c.-135C>G
NM_001350967.2:c.-343C>G
NM_022487.4:c.-187C>G
NR_110297.2:n.189C>G
NR_146961.2:n.189C>G

Pathogenic

Met criteria codes **5**

PM2_Supporting PP1 PP4_Moderate
PS3_Moderate PM3_Strong

Not Met criteria codes **2**

PP3 BS2

Evidence Links **0**

Expert Panel

[Severe Combined Immunodeficiency Disease VCEP](#)

Criteria Specification Information

[Criteria Specification:](#) *ClinGen Severe Combined Immunodeficiency Disease Expert Panel Specifications to the ACMG/AMP Variant Interpretation Guidelines for DCLRE1C Version 1.0.0*











[Criteria Specification Approval History](#)

[Criteria Specifications for this VCEP](#)




Severe Combined Immunodeficiency Disease VCEP

The c.103C>G (NM_001033855.3) variant in DCLRE1C is a missense variant predicted to cause the substitution of Histidine by Aspartic Acid at amino acid 35 (p.His35Asp). The highest population minor allele frequency in gnomAD v4 is 0.000003100 (8/1111884 alleles) in the European (non-Finnish) population, which is lower than the ClinGen SCID VCEP threshold (<0.00003266) for PM2_Supporting, meeting this criterion (PM2_Supporting). No homozygotes have been observed in gnomAD. Activity levels in % of WT activity = Recombination: Mean (SD): 0 (0.3) and DNA repair (36h after IR): Mean (SD): 27.29 (16.57). PS3 is Met at a moderate level (PMID: 25917813). This variant has been detected in at least 4 individuals with SCID. Of those individuals, 02 were compound heterozygous for the variants: c.2T>C (p.Met1Thr) VUS according to SCID VCEP, 0.25pts AND p.L187*; Pathogenic according to the SCID VCEP specifications, 1 point. 02 individuals were homozygous for the variants (1 point). The total is 2.25 points, PM3_Strong. (PMIDs: 24481607, 15731174, and 32441320). At least one patient with this variant displayed T-B-NK+ (0.5 pts) + Diagnostic criteria for SCID/Leaky SCID/Omenn syndrome met (0.5 pts) + Family history of SCID (0.5 pts) + SCID gene panel or exome/genome sequencing conducted (0.5 pts), totaling 2 points, which is highly specific for SCID (PP4_Moderate, PMID: 24481607). In summary, this variant is classified as a Pathogenic for autosomal recessive SCID based on ACMG/AMP criteria applied, as specified by the ClinGen SCID VCEP (specification version 1.0): PM3_Strong, PS3_Moderate, PP1_Supporting, PP4_Moderate, and PM2_Supporting.

Met criteria codes

PM2_Supporting			The highest population minor allele frequency in gnomAD v4 is 0.000003100 (8/1111884 alleles) in the European (non-Finnish) population, which is lower than the ClinGen SCID VCEP threshold (<0.00003266) for PM2_Supporting, meeting this criterion (PM2_Supporting). No homozygotes have been observed in gnomAD.
PP1			The variant has been reported to segregate with OS in - at least - 02 affected family members (proband + one brother) from one family (PP1_Supporting). The patient also had another affected brother without genomic information; PMID: 15731174.
PP4_Moderate			At least one patient with this variant displayed T-B-NK+ (0.5 pts) + Diagnostic criteria for SCID/Leaky SCID/Omenn syndrome met (0.5 pts) + Family history of SCID (0.5 pts) + SCID gene panel or exome/genome sequencing conducted (0.5 pts), totalizing 2 points, which is highly specific for SCID (PP4_Moderate, PMID: 24481607).
PS3_Moderate			Activity levels in % of WT activity = Recombination: Mean (SD): 0 (0.3) and DNA repair (36h after IR): Mean (SD): 27.29 (16.57). PS3 is Met at a moderate level (PMID: 25917813).
PM3_Strong			This variant has been detected in at least 4 individuals with SCID. Of those individuals, 02 were compound heterozygous for the variants: c.2T>C (p.Met1Thr) VUS according to SCID VCEP. Phase is confirmed in another paper that describes the same patient: 0.25pts AND p.L187*; Pathogenic according to the SCID VCEP specifications, 1 point. 02 individuals were homozygous for the variants (1 point). The total is 2.25 points, PM3_Strong. (PMIDs: 24481607, 15731174, and 32441320).

Not Met criteria codes

PP3			Do not apply to missense variants.
BS2			No homozygotes have been observed in gnomAD.

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.