

Variant: NM_000022.4(ADA):c.445C>T (p.Arg149Trp)

Version: 1.0

CA266014 [↗](#)

68264 (ClinVar) [↗](#)

Gene: ADA (HGNC:100)

Condition: adenosine deaminase deficiency (MONDO:0007064)

Inheritance Mode: Autosomal recessive inheritance

UUID: 66e50984-088e-48e7-b034-9faab22a49dc

Approved on: 2024-01-16

Published on: 2024-01-16

HGVS expressions

NM_000022.4:c.445C>T

NM_000022.4(ADA):c.445C>T (p.Arg149Trp)

NC_000020.11:g.44625602G>A

CM000682.2:g.44625602G>A

NC_000020.10:g.43254243G>A

CM000682.1:g.43254243G>A

NC_000020.9:g.42687657G>A

NG_007385.1:g.31134C>T

ENST00000492931.6:n.536C>T

ENST00000536076.2:c.292C>T

ENST00000536532.6:c.445C>T

ENST00000537820.2:c.445C>T

ENST00000539235.6:c.219-2524C>T

ENST00000695889.1:c.219-2672C>T

ENST00000695890.1:n.2248C>T

ENST00000695891.1:c.219-2672C>T

ENST00000695927.1:c.523C>T

ENST00000695949.1:c.442C>T

ENST00000695957.1:c.362+854C>T

ENST00000695991.1:c.217-2672C>T

ENST00000695992.1:c.445C>T

ENST00000695993.1:c.445C>T

ENST00000695994.1:c.445C>T

ENST00000695995.1:c.217-2524C>T

ENST00000695996.1:n.516C>T

ENST00000695997.1:n.433+854C>T

ENST00000696003.1:n.537C>T

ENST00000696004.1:n.537C>T

ENST00000696006.1:c.445C>T

ENST00000696007.1:c.329+854C>T

ENST00000696008.1:n.361C>T

ENST00000696009.1:n.556C>T

ENST00000696017.1:c.442C>T

ENST00000696034.1:c.445C>T

ENST00000696035.1:n.555C>T

ENST00000696036.1:n.1135C>T

ENST00000696037.1:n.2122C>T

ENST00000696038.1:c.*191C>T
ENST00000696039.1:n.733C>T
ENST00000696058.1:c.445C>T
ENST00000696059.1:c.*390C>T
ENST00000696060.1:c.445C>T
ENST00000696061.1:c.442C>T
ENST00000696062.1:c.508C>T
ENST00000696063.1:c.520C>T
ENST00000696064.1:c.292C>T
ENST00000696065.1:c.66-2672C>T
ENST00000696075.1:c.*415C>T
ENST00000696076.1:c.445C>T
ENST00000696077.1:c.442C>T
ENST00000696078.1:c.445C>T
ENST00000696079.1:c.445C>T
ENST00000696080.1:c.445C>T
ENST00000696082.1:c.523C>T
ENST00000696083.1:n.87C>T
ENST00000696084.1:n.546C>T
ENST00000696104.1:c.362+854C>T
ENST00000696105.1:c.445C>T
ENST00000372874.9:c.445C>T
ENST00000372874.8:c.445C>T
ENST00000464097.5:n.119C>T
ENST00000492931.5:n.529C>T
ENST00000536532.5:c.445C>T
ENST00000537820.1:c.445C>T
ENST00000539235.5:c.219-2524C>T
NM_000022.2:c.445C>T
NM_000022.3:c.445C>T
NM_001322050.1:c.73+854C>T
NM_001322051.1:c.445C>T
NR_136160.1:n.596C>T
NM_001322050.2:c.73+854C>T
NM_001322051.2:c.445C>T
NR_136160.2:n.537C>T

Likely Pathogenic

Met criteria codes **3**

PM2 PP4_Moderate PS3_Moderate

Not Met criteria codes **2**

PM5 BS1

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 ADA Version 1.0.0*






[Criteria Specification Approval History](#)

[Criteria Specifications for this VCEP](#)





Severe Combined Immunodeficiency Disease VCEP

The NM_000022.4:c.445C>T (p.Arg149Trp) variant in ADA is a missense variant predicted to cause substitution of arginine by tryptophan at amino acid 149 (p.Arg149Trp). This variant has been reported in an individual with severe combined immunodeficiency and elevated pre-treatment erythrocyte dAXP (2pts, PMID: 10200056) (PP4_Moderate). In experimental studies, the activity of this variant was reported as being 0.012% of wild-type activity (PMID: 9758612) (PS3_Moderate). The highest subpopulation allele frequency 0.00009294 (2/21520) in African/African American population, which is lower than the ADA cutoff gnomAD popmax filtering allele frequency <0.0001742). So PM2 is met. In summary, this variant meets the criteria to be classified as likely pathogenic for SCID. ACMG/AMP criteria applied, as specified by the ClinGen SCID-VCEP: PM2_Supporting, PP4_Moderate, PS3_Moderate (SCID VCEP specifications version 1.0).

Met criteria codes

PM2		Overall allele frequency on gnomAD is 0.00001648 (4/242700 alleles) the highest subpopulation allele frequency 0.00009294 (2/21520) in African/African American population (https://gnomad.broadinstitute.org/variant/20-43254243-G-A - No homozygous), which is lower than the ADA cutoff gnomAD popmax filtering allele frequency <0.0001742). So PM2 is met.
PP4_Moderate	 	PMID: 10200056: Patient CS with diagnosis of SCID and pretreatment RBC dAXP of 41.5% (normal is <0.2% as per PMID). Under ClinGen SCID VCEP ADA PP4 specifications, this is worth 2 points. 2 or more points is sufficient to meet PP4_Moderate criteria, therefore PP4 is applied at moderate strength.
PS3_Moderate	 	PMID: 9758612: activity of Arg149Trp is in group 1 which has 0.012% activity vs WT. This is below the threshold set for ADA PS3_Moderate by the ClinGen SCID VCEP (PS3_Moderate < 0.05% of wild type activity (group I)). Therefore PS3 is met at moderate strength.

Not Met criteria codes

PM5	 	Two additional missense variants in the same codon: * NM_000022.4(ADA):c.446G>A (p.Arg149Gln): VUS according SCID VCEP * NM_000022.4(ADA):c.446G>T (p.Arg149Leu), VUS according SCID VCEP PM5 is not met.
BS1	 	Allele frequency in gnomAD does not exceed threshold set by SCID VCEP ADA specifications (gnomAD popmax filtering allele frequency >0.00161), therefore BS1 is not met

	▼	▼
--	---	---

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.