

## Variant: *NM\_000546.5(TP53):c.892G>T (p.Glu298Ter)*

Version: 1.1

CA000484 [↗](#)

93323 (ClinVar) [↗](#)

**Gene:** TP53 ([HGNC:7157](#))

**Condition:** Li-Fraumeni syndrome ([MONDO:0018875](#))

**Inheritance Mode:** Autosomal dominant inheritance

**UID:** 42caf4f2-7ef5-45d4-a593-271e92b9ca88

**Approved on:** 2024-08-05

**Published on:** 2024-08-05

### *HGVS expressions*

**NM\_000546.5:c.892G>T**

NM\_000546.5(TP53):c.892G>T (p.Glu298Ter)

NC\_000017.11:g.7673728C>A

CM000679.2:g.7673728C>A

NC\_000017.10:g.7577046C>A

CM000679.1:g.7577046C>A

NC\_000017.9:g.7517771C>A

NG\_017013.2:g.18823G>T

ENST00000503591.2:c.892G>T

ENST00000508793.6:c.892G>T

ENST00000509690.6:c.496G>T

ENST00000514944.6:c.613G>T

ENST00000604348.6:c.871G>T

ENST00000269305.9:c.892G>T

ENST00000269305.8:c.892G>T

ENST00000359597.8:c.892G>T

ENST00000413465.6:c.782+453G>T

ENST00000420246.6:c.892G>T

ENST00000445888.6:c.892G>T

ENST00000455263.6:c.892G>T

ENST00000504290.5:c.496G>T

ENST00000504937.5:c.496G>T

ENST00000509690.5:c.496G>T

ENST00000510385.5:c.496G>T

ENST00000610292.4:c.775G>T

ENST00000610538.4:c.775G>T

ENST00000610623.4:c.415G>T

ENST00000615910.4:c.859G>T

ENST00000617185.4:c.892G>T

ENST00000618944.4:c.415G>T

ENST00000619186.4:c.415G>T

ENST00000619485.4:c.775G>T

ENST00000620739.4:c.775G>T

ENST00000622645.4:c.775G>T

ENST00000635293.1:c.775G>T

NM\_001126112.2:c.892G>T

NM\_001126113.2:c.892G>T

NM\_001126114.2:c.892G>T  
NM\_001126115.1:c.496G>T  
NM\_001126116.1:c.496G>T  
NM\_001126117.1:c.496G>T  
NM\_001126118.1:c.775G>T  
NM\_001276695.1:c.775G>T  
NM\_001276696.1:c.775G>T  
NM\_001276697.1:c.415G>T  
NM\_001276698.1:c.415G>T  
NM\_001276699.1:c.415G>T  
NM\_001276760.1:c.775G>T  
NM\_001276761.1:c.775G>T  
NM\_001276695.2:c.775G>T  
NM\_001276696.2:c.775G>T  
NM\_001276697.2:c.415G>T  
NM\_001276698.2:c.415G>T  
NM\_001276699.2:c.415G>T  
NM\_001276760.2:c.775G>T  
NM\_001276761.2:c.775G>T  
NM\_000546.6:c.892G>T  
NM\_001126112.3:c.892G>T  
NM\_001126113.3:c.892G>T  
NM\_001126114.3:c.892G>T  
NM\_001126115.2:c.496G>T  
NM\_001126116.2:c.496G>T  
NM\_001126117.2:c.496G>T  
NM\_001126118.2:c.775G>T  
NM\_001276695.3:c.775G>T  
NM\_001276696.3:c.775G>T  
NM\_001276697.3:c.415G>T  
NM\_001276698.3:c.415G>T  
NM\_001276699.3:c.415G>T  
NM\_001276760.3:c.775G>T  
NM\_001276761.3:c.775G>T

**Pathogenic**

Met criteria codes **4**

PS4\_Supporting PVS1 PP4  
PM2\_Supporting

Not Met criteria codes **14**

PS1 PS2 PS3 PP1 PP3 PM1  
PM5 BA1 BS1 BS4 BS3  
BS2 BP7 BP4

Evidence Links **0**

Expert Panel

[TP53 VCEP](#)

Criteria Specification Information

[Criteria Specification:](#) *ClinGen TP53 Expert Panel Specifications to the ACMG/AMP Variant Interpretation Guidelines for TP53 Version 2.0.0*

[Criteria Specification Approval History](#)









[Criteria Specifications for this VCEP](#)

Evidence submitted by expert panel









**TP53 VCEP**







The NM\_000546.6 c.892G>T(p.Glu298Ter) is a TP53 nonsense variant inducing a premature termination codon upstream of p.Lys351. The variant is predicted to undergo nonsense-mediated decay (PVS1). This variant has been observed in 1 family meeting Revised Chompret criteria. This proband was under the age of 40 diagnosed with a HER2+ breast cancer. Based on this evidence, this variant scores 1 total point meeting the TP53 VCEP phenotype scoring criteria of 1-1.5 points. (PS4\_Supporting; Internal lab contributors: SCV000278127.7). At least one individual with this variant was found to have a variant allele fraction 25-35%, which is a significant predictor of variant pathogenicity (PP4, PMID: 34906512, SCV000278127.7). This variant is absent from gnomAD v4.1.0 (PM2\_Supporting). In summary, this variant meets the criteria to be classified as pathogenic for Li Fraumeni Syndrome based on the ACMG/AMP criteria applied, as specified by the ClinGen TP53 VCEP: PVS1, PS4\_Supporting, PP4, PM2\_Supporting. (Bayesian Points: 11; VCEP specifications version 2.0; 7/24/2024)

#### Met criteria codes

<b>PS4_Supporting</b>			This variant has been observed in 1 family meeting Revised Chompret criteria. This proband was under the age of 40 diagnosed with a HER2+ breast cancer. Based on this evidence, this variant scores 1 total point meeting the TP53 VCEP phenotype scoring criteria of 1-1.5 points. (PS4_Supporting; Internal lab contributors: SCV000278127.7).
<b>PVS1</b>			The NM_000546.6 c.892C>T (p.Glu298Ter) is a TP53 nonsense variant upstream of p.Lys351. The variant is predicted to undergo nonsense-mediated decay (PVS1).
<b>PP4</b>			At least one individual with this variant was found to have a variant allele fraction 25-35%, which is a significant predictor of variant pathogenicity (PP4, PMID: 34906512, SCV000278127.7).
<b>PM2_Supporting</b>			This variant is absent from gnomAD v4.1.0 (PM2_Supporting).

#### Not Met criteria codes

<b>PS1</b>			No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
<b>PS2</b>			No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
<b>PS3</b>			No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
<b>PP1</b>			No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
<b>PP3</b>			No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
<b>PM1</b>			No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
<b>PM5</b>			No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline

<b>BA1</b>			No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
<b>BS1</b>			No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
<b>BS4</b>			No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
<b>BS3</b>			No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
<b>BS2</b>			No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
<b>BP7</b>			No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline
<b>BP4</b>			No code specific comments provided, please refer to the summary above or general recommendations provided in the guideline

Curation History [↗](#)




Showing 1 to 2 of 2 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.