

Standards and guidelines for the interpretation of sequence variants: a joint consensus recommendation of the American College of Medical Genetics and Genomics and the Association for Molecular Pathology

Sue Richards, PhD¹, Nazneen Aziz, PhD^{2,16}, Sherri Bale, PhD³, David Bick, MD⁴, Soma Das, PhD⁵, Julie Gastier-Foster, PhD^{6,7,8}, Wayne W. Grody, MD, PhD^{9,10,11}, Madhuri Hegde, PhD¹², Elaine Lyon, PhD¹³, Elaine Spector, PhD¹⁴, Karl Voelkerding, MD¹³ and Heidi L. Rehm, PhD¹⁵; on behalf of the ACMG Laboratory Quality Assurance Committee

Das ACMG Klassifizierungssystem dient der Einteilung von Sequenzvarianten für mendelische Erkrankungen, unabhängig ob Varianten per Sangersequenzierung oder per NGS (Next Generation Sequencing) nachgewiesen wurden.

Varianten werden entsprechend der IARC Empfehlungen (Plon et al.; *Hum Mutat* 2008) in fünf Klassen eingeteilt:

Klasse 1	benign	Normvariante ohne klinische Relevanz
Klasse 2	likely benign	Wahrscheinliche Normvariante
Klasse 3	variant of uncertain significance (VUS)	Variante unklarer klinischer Relevanz
Klasse 4	likely pathogenic	Wahrscheinlich pathogene Variante
Klasse 5	pathogenic	Pathogene Variante

Die folgenden 3 Tabellen basieren auf der o.g. Primärliteratur und geben einen schematischen Überblick über das ACMG-Klassifizierungssystem. Für detaillierte Informationen empfehlen wir die Lektüre der kompletten Veröffentlichung.

Tabelle 1 listet Kriterien für eine pathogene Wirkung auf. In Tabelle 2 sind Kriterien aufgeführt die für eine benigne Wirkung sprechen. Um eine Sequenzvariante zu klassifizieren werden die entsprechenden Kriterien nach dem Regeln von Tabelle 3 kombiniert, um zu einer eindeutigen Klassifizierung zu gelangen.

Criteria for classifying pathogenic variants (Tabelle 1)

Evidence of pathogenicity		Category
Very strong	PVS1	<p>Null variant (nonsense, frameshift, canonical ± 1 or 2 splice sites, initiation codon, single or multiexon deletion) in a gene where LOF is a known mechanism of disease.</p> <p><i>Caveats:</i></p> <ul style="list-style-type: none"> Beware of genes where LOF is not a known disease mechanism (e.g., GFAP, MYH7) Use caution interpreting LOF variants at the extreme 3' end of a gene Use caution with splice variants that are predicted to lead to exon skipping but leave the remainder of the protein intact Use caution in the presence of multiple transcripts
Strong	PS1	<p>Same amino acid change as a previously established pathogenic variant regardless of nucleotide change</p> <ul style="list-style-type: none"> Example: Val→Leu caused by either G>C or G>T in the same codon Caveat: Beware of changes that impact splicing rather than at the amino acid/protein level
	PS2	<p>De novo (both maternity and paternity confirmed) in a patient with the disease and no family history</p> <p><i>Note: Confirmation of paternity only is insufficient. Egg donation, surrogate motherhood, errors in embryo transfer, and so on, can contribute to non maternity.</i></p>
	PS3	<p>Well-established in vitro or in vivo functional studies supportive of a damaging effect on the gene or gene product</p> <p><i>Note: Functional studies that have been validated and shown to be reproducible and robust in a clinical diagnostic laboratory setting are considered the most well established.</i></p>
	PS4	<p>The prevalence of the variant in affected individuals is significantly increased compared with the prevalence in controls</p> <p><i>Note 1: Relative risk or OR, as obtained from case–control studies, is >5.0, and the confidence interval around the estimate of relative risk or OR does not include 1.0. See the article for detailed guidance.</i></p> <p><i>Note 2: In instances of very rare variants where case–control studies may not reach statistical significance, the prior observation of the variant in multiple unrelated patients with the same phenotype, and its absence in controls, may be used as moderate level of evidence.</i></p>
Moderate	PM1	<p>Located in a mutational hot spot and/or critical and well-established functional domain (e.g., active site of an enzyme) without benign variation.</p>
	PM2	<p>Absent from controls (or at extremely low frequency if recessive) (Table 6) in Exome Sequencing Project, 1000 Genomes Project, or Exome Aggregation Consortium</p> <ul style="list-style-type: none"> Caveat: Population data for insertions/deletions may be poorly called by next-generation sequencing.
	PM3	<p>For recessive disorders, detected in trans with a pathogenic variant</p> <p><i>Note: This requires testing of parents (or offspring) to determine phase.</i></p>
	PM4	<p>Protein length changes as a result of in-frame deletions/insertions in a nonrepeat region or stop-loss variants</p>
	PM5	<p>Novel missense change at an amino acid residue where a different missense change determined to be pathogenic has been seen before</p> <ul style="list-style-type: none"> Example: Arg156His is pathogenic; now you observe Arg156Cys Caveat: Beware of changes that impact splicing rather than at the amino acid/protein level.
	PM6	<p>Assumed de novo, but without confirmation of paternity and maternity</p>

Supporting	PP1	Co segregation with disease in multiple affected family members in a gene definitively known to cause the disease <i>Note: May be used as stronger evidence with increasing segregation data</i>
	PP2	Missense variant in a gene that has a low rate of benign missense variation and in which missense variants are a common mechanism of disease
	PP3	Multiple lines of computational evidence support a deleterious effect on the gene or gene product (conservation, evolutionary, splicing impact, etc.) <ul style="list-style-type: none"> <i>Caveat: Because many in-silico algorithms use the same or very similar input for their predictions, each algorithm should not be counted as an independent criterion. PP3 can be used only once in any evaluation of a variant.</i>
	PP4	Patient's phenotype or family history is highly specific for a disease with a single genetic etiology
	PP5	Reputable source recently reports variant as pathogenic, but the evidence is not available to the laboratory to perform an independent evaluation

Criteria for classifying benign variants (Tabelle 2)

Evidence of benign impact		Category
Stand alone	BA1	Allele frequency is >5% in Exome Sequencing Project, 1000 Genomes Project, or Exome Aggregation Consortium

Strong	BS1	Allele frequency is greater than expected for disorder (see Table 6)
	BS2	Observed in a healthy adult individual for a recessive (homozygous), dominant (heterozygous), or X-linked (hemizygous) disorder, with full penetrance expected at an early age
	BS3	Well-established in vitro or in vivo functional studies show no damaging effect on protein function or splicing
	BS4	Lack of segregation in affected members of a family <ul style="list-style-type: none"> <i>Caveat: The presence of phenocopies for common phenotypes (i.e., cancer, epilepsy) can mimic lack of segregation among affected individuals. Also, families may have more than one pathogenic variant contributing to an autosomal dominant disorder, further confounding an apparent lack of segregation.</i>

Supporting	BP1	Missense variant in a gene for which primarily truncating variants are known to cause disease
	BP2	Observed in trans with a pathogenic variant for a fully penetrant dominant gene/disorder or observed in cis with a pathogenic variant in any inheritance pattern
	BP3	In-frame deletions/insertions in a repetitive region without a known function
	BP4	Multiple lines of computational evidence suggest no impact on gene or gene product (conservation, evolutionary, splicing impact, etc.) <ul style="list-style-type: none"> <i>Caveat: Because many in silico algorithms use the same or very similar input for their predictions, each algorithm cannot be counted as an independent criterion. BP4 can be used only once in any evaluation of a variant.</i>
	BP5	Variant found in a case with an alternate molecular basis for disease
	BP6	Reputable source recently reports variant as benign, but the evidence is not available to the laboratory to perform an independent evaluation
	BP7	A synonymous (silent) variant for which splicing prediction algorithms predict no impact to the splice consensus sequence nor the creation of a new splice site AND the nucleotide is not highly conserved

Rules for combining criteria to classify sequence variants

(Tabelle 3)

Pathogenic	(i) 1 Very strong (PVS1) AND <i>a) ≥1 Strong (PS1–PS4) OR</i> <i>b) ≥2 Moderate (PM1–PM6) OR</i> <i>c) 1 Moderate (PM1–PM6) and 1 supporting (PP1–PP5) OR</i> <i>d) ≥2 Supporting (PP1–PP5)</i>
	(ii) ≥2 Strong (PS1–PS4) OR
	(iii) 1 Strong (PS1–PS4) AND <i>a) ≥3 Moderate (PM1–PM6) OR</i> <i>b) 2 Moderate (PM1–PM6) AND ≥2 Supporting (PP1–PP5) OR</i> <i>c) 1 Moderate (PM1–PM6) AND ≥4 supporting (PP1–PP5)</i>
Likely pathogenic	(i) 1 Very strong (PVS1) AND 1 moderate (PM1–PM6) OR
	(ii) 1 Strong (PS1–PS4) AND 1–2 moderate (PM1–PM6) OR
	(iii) 1 Strong (PS1–PS4) AND ≥2 supporting (PP1–PP5) OR
	(iv) ≥3 Moderate (PM1–PM6) OR
	(v) 2 Moderate (PM1–PM6) AND ≥2 supporting (PP1–PP5) OR
	(vi) 1 Moderate (PM1–PM6) AND ≥4 supporting (PP1–PP5)
Benign	(i) 1 Stand-alone (BA1) OR
	(ii) ≥2 Strong (BS1–BS4)
Likely benign	(i) 1 Strong (BS1–BS4) and 1 supporting (BP1–BP7) OR
	(ii) ≥2 Supporting (BP1–BP7)
Uncertain significance	(i) Other criteria shown above are not met OR
	(ii) the criteria for benign and pathogenic are contradictory