

# Resource Summary Report

Generated by [ASWG](#) on Aug 4, 2026

## Ensembl Variation

RRID:SCR\_001630

Type: Tool

### Proper Citation

Ensembl Variation (RRID:SCR\_001630)

### Resource Information

**URL:** <https://github.com/Ensembl>

**Proper Citation:** Ensembl Variation (RRID:SCR\_001630)

**Description:** Public database that stores areas of genome that differ between individual genomes (variants) and, where available, associated disease and phenotype information. Different types of variants for several species: single nucleotide polymorphisms (SNPs), short nucleotide insertions and/or deletions, and longer variants classified as structural variants (including CNVs). Effects of variants on the Ensembl transcripts and regulatory features for each species are predicted. You can run same analysis on your own data using Variant Effect Predictor. These data are integrated with other data sources in Ensembl, and can be accessed using the API or website. For several different species in Ensembl, they import variation data (SNPs, CNVs, allele frequencies, genotypes, etc) from a variety of sources (e.g. dbSNP). Imported variants and alleles are subjected to quality control process to flag suspect data. In human, they calculate linkage disequilibrium for each variant, by population.

**Abbreviations:** Ensembl Variation

**Synonyms:** ensembl variation

**Resource Type:** production service resource, data analysis service, service resource, database, analysis service resource, data or information resource

**Defining Citation:** [PMID:23203987](#), [PMID:20562413](#), [PMID:20459810](#), [PMID:20459805](#)

**Keywords:** genome, disease, phenotype, genomic variant, single nucleotide polymorphism nucleotide, insertion, deletion, structural variant, copy number variation, inversion, translocation, somatic variant, allele frequency, genotype, disease phenotype, inherited disease

**Availability:** Free, Available for download, Freely available

**Resource Name:** Ensembl Variation

**Resource ID:** SCR\_001630

**Alternate IDs:** nlx\_153897

**License:** Apache 2.0 License

**Record Creation Time:** 20220129T080208+0000

**Record Last Update:** 20260804T043129+0000

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## Ratings and Alerts

No rating or validation information has been found for Ensembl Variation.

No alerts have been found for Ensembl Variation.

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## Data and Source Information

**Source:** [SciCrunch Registry](#)

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## Usage and Citation Metrics

We found 4 mentions in open access literature.

**Listed below are recent publications.** The full list is available at [ASWG](#) .

Alatwi E, et al. (2024) The role of genetic polymorphisms in the sulfation of pregnenolone by human cytosolic sulfotransferase SULT2B1a. Scientific reports, 14(1), 8050.

Alatwi E, et al. (2023) The role of genetic polymorphisms in the sulfation of pregnenolone by human cytosolic sulfotransferase SULT2B1a. Research square.

Srivastava M, et al. (2021) Transcriptome-wide high-throughput mapping of protein-RNA occupancy profiles using POP-seq. Scientific reports, 11(1), 1175.

Pua CJ, et al. (2016) Development of a Comprehensive Sequencing Assay for Inherited Cardiac Condition Genes. Journal of cardiovascular translational research, 9(1), 3.