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Archive!

## Ensembl Views

**AlignSliceView**: pairwise or multiple genomic alignment for a genomic region  
**AlignView**: alignment for a set of nucleotide or protein sequences  
**BLASTView**: do a BLAST or SSAHA sequence similarity search  
**ContigView**: genome annotation for a 1 bp – 1 Mb region  
**CytoView**: genome annotation for a 200 kb – 50 Mb region  
**DASConfView**: upload your data to the Ensembl DAS server or add your own DAS server  
**DotterView**: dot plot for a conserved block  
**DomainView**: Ensembl genes containing a given InterPro domain  
**ExonView**: exon, intron and flanking sequence and supporting evidence for a transcript  
**ExportView**: export sequence and/or annotation for a feature or region  
**FamilyView**: information for a protein family  
**FastaView**: sequence and associated details for feature external to the Ensembl core databases  
**FeatureView**: locations to which a given feature (e.g. microarray probeset, mRNA, protein) has been mapped  
**GeneRegulationView**: regulatory factors for a gene  
**GeneSeqAlignView**: pairwise or multiple alignment of genomic sequence for a gene  
**GeneSeqView**: genomic sequence for a gene  
**GeneSNPView**: SNPs for a gene  
**GeneSpliceView**: alternative splicing for a gene  
**GeneTreeView**: gene tree with orthologues and paralogues for a gene  
**GeneView**: information for a gene  
**GOView**: information for a GO term, including hierarchy  
**HelpView**: help information for an Ensembl view

**HistoryView**: retrieve the history for a batch of Ensembl stable IDs  
**IDHistoryView**: history for an Ensembl stable ID  
**KaryoView**: annotate a chromosome or karyotype with your own data  
**LDTableView**: LD values between SNPs in table format  
**LDView**: LD values between SNPs, including the possibility of dumping for upload into Haploview software  
**MapView**: explore a chromosome  
**MarkerView**: information for a marker  
**MartView**: data mine with BioMart  
**MultiContigView**: simultaneous display of genome annotation for multiple species  
**ProtView**: information for a protein, including protein sequence  
**SearchView**: search in Ensembl  
**SequenceAlignView**: alignment of sequences of different individuals / strains  
**SNPView**: information for a SNP  
**SyntenyView**: syntenic regions between species pairs  
**TranscriptSNPView**: comparison of SNPs in individuals / strains / populations for a transcript  
**TransView**: information for a transcript, including spliced transcript sequence

## Questions or Problems?

Contact [helpdesk@ensembl.org](mailto:helpdesk@ensembl.org)

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## Glossary of Terms

**Assembly:** the whole of sequenced pieces of DNA of an organism put back together to create a representation of the original chromosomes from which the DNA originated

**BLAST:** Basic Local Alignment Search Tool, a program that finds sequence similarity

**CDS:** coding sequence

**Contig:** a contiguous stretch of DNA sequence without gaps that has been assembled based on direct sequencing information

**DAS:** Distributed Annotation System, a communication protocol used to exchange biological annotations

**EST:** expressed sequence tag

**Genebuild:** the process according to which Ensembl gene predictions are created and annotated

**GENSCAN:** *ab initio* gene prediction program

**GO term:** Gene Ontology term; the Gene Ontology Consortium produces an organised hierarchy of terms, used to describe biological processes, cellular components, and molecular functions (see also 'Databases & Websites')

**Havana:** Human And Vertebrate Analysis and Annotation; the Havana team is part of the Vega consortium (see 'Databases and websites')

**Manual annotation:** annotation of nucleotide or protein sequences by a human being

**NCBI:** National Center for Biotechnology Information

**SNP:** single nucleotide polymorphism

**SSAHA:** Sequence Search and Alignment by Hashing Algorithm, a program that finds sequence similarity

**UCSC:** University of California Santa Cruz

**UTR:** untranslated region

## Databases & Websites

**CCDS:** Consensus CDS database  
(<http://www.ncbi.nlm.nih.gov/CCDS/>)

**dbSNP:** SNP repository at the NCBI  
(<http://www.ncbi.nlm.nih.gov/projects/SNP/>)

**DAS registration server:** registry of available DAS sources  
(<http://www.dasregistry.org>)

**EMBL-Bank/GenBank/DDBJ:** nucleotide sequence repositories  
(<http://www.ebi.ac.uk/embl/>)

**GO:** Gene Ontology database  
(<http://www.ebi.ac.uk/GO/>)

**HapMap:** multi-country effort to identify and catalog genetic similarities and differences in human beings  
(<http://www.hapmap.org/>)

**InterPro:** database of protein families, domains and functional sites  
(<http://www.ebi.ac.uk/interpro/>)

**NCBI Map Viewer:** genome browser of the NCBI  
(<http://www.ncbi.nlm.nih.gov/mapview/>)

**RefSeq:** partially manually curated protein and nucleotide sequence database at the NCBI  
(<http://www.ncbi.nlm.nih.gov/RefSeq/>)

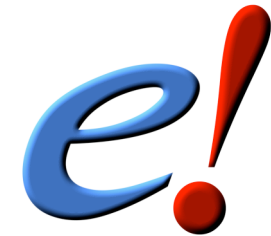
**UCSC Genome Browser:** genome browser of the UCSC  
(<http://genome.ucsc.edu/>)

**UniProt/Swiss-Prot:** manually curated protein sequence database  
(<http://www.ebi.ac.uk/swissprot/>)

**UniProt/TrEMBL:** computer-annotated protein sequence database  
(<http://www.ebi.ac.uk/trembl/>)

**Vega:** Vertebrate Genome Annotation database, containing manually annotated vertebrate genome sequence  
(<http://vega.sanger.ac.uk/>)

## Quick Reference Guide to the



## Ensembl Genome Browser

