SIFT.Hsapiens.dbSNP137
March 20, 2024

Description
Database of PROVEAN/SIFT predictions for Homo sapiens dbSNP build 137

Details
The SIFT tool is no longer actively maintained. A few of the original authors have started the PROVEAN (Protein Variation Effect Analyzer) project. PROVEAN is a software tool which predicts whether an amino acid substitution or indel has an impact on the biological function of a protein. PROVEAN is useful for filtering sequence variants to identify nonsynonymous or indel variants that are predicted to be functionally important.

See the web pages for a complete description of the methods.

- SIFT Home: http://sift.jcvi.org/

Though SIFT is not under active development, the PROVEAN team still provides the SIFT scores in the pre-computed downloads. This package, SIFT.Hsapiens.dbSNP137, contains both SIFT and PROVEAN scores. One notable difference between this and the previous SIFT database package is that keys in SIFT.Hsapiens.dbSNP132 are rs IDs whereas in SIFT.Hsapiens.dbSNP137 they are NCBI dbSNP IDs.

Methods
- Methods: See '?PROVEANDb-class' in the VariantAnnotation package for a complete listing of available methods.
- Creation of Database Tables: This package includes PROVEAN/SIFT predictions for dbSNP build 137 human coding non-synonymous SNPs.
- Source Files:
Source: http://provean.jcvi.org/downloads.php
Software: PROVEAN 1.1, SIFT 4.0.3
Databases: PSI-BLAST
Source Files: dbsnp137.coding.variants.prediction.tsv.gz PROVEAN/SIFT predictions for coding snps in dbSNP build 137
Description: This package contains PROVEAN/SIFT annotations human SNPs included in dbSNP build 137.

Column descriptions

These names are displayed when columns is called on the PROVEANDb object (i.e., columns(SIFT.Hsapiens.dbSNP137))

- **DBSNPID**: NCBI dbSNP ID
- **VARIANT**: comma separated values of <chromosome>,<position>,<reference allele>,<variant allele>, <comment(optional)>
- **PROTEINID**: Ensembl protein ID
- **LENGTH**: length of the protein
- **STRAND**: ‘+’, ‘-’ or NA
- **CODONCHANGE**: codon change including flanking codons
- **POS**: position of amino acid residue affected
- **RESIDUEREF**: reference amino acid residue
- **RESIDUEALT**: variant amino acid residue
- **TYPE**: synonymous | nonsynonymous | frameshift | ...
- **PROVEANSCORE**: PROVEAN score (see http://provean.jcvi.org/about.php#about_1)
- **PROVEANPRED**: deleterious or neutral (cutoff=-2.5)
- **PROVEANNUMSEQ**: number of sequences used for prediction
- **PROVEANNUMCLUST**: number of clusters used for prediction
- **SIFTSCORE**: SIFT score (range 0 to 1)
- **SIFTPRED**: tolerated or damaging (cutoff=0.05)
- **SIFTMEDIAN**: median sequence information used to measure the diversity of the sequences used for prediction
- **SIFTNUMSEQ**: number of sequences used for prediction

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References

The PROVEAN tool has replaced SIFT: [http://provean.jcvi.org/about.php](http://provean.jcvi.org/about.php)


Ng PC, Henikoff S. Predicting the Effects of Amino Acid Substitutions on Protein Function Annu Rev Genomics Hum Genet. 2006;7:61-80.


See Also

PROVEANDb-class

Examples

```r
library(SIFT.Hsapiens.dbSNP137)

## metadata
metadata(SIFT.Hsapiens.dbSNP137)

## keys are the DBSNPID (NCBI dbSNP ID)
dbsnp <- keys(SIFT.Hsapiens.dbSNP137)
head(dbsnp)
columns(SIFT.Hsapiens.dbSNP137)

## Return all columns. Note that the key, DBSNPID, is always returned.
select(SIFT.Hsapiens.dbSNP137, dbsnp[10])

## subset on keys and cols
cols <- c("VARIANT", "PROVEANPRED", "SIFTPRED")
select(SIFT.Hsapiens.dbSNP137, dbsnp[20:23], cols)
```
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