SNPlocs.Hsapiens.dbSNP155.GRCh38

September 21, 2022

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The SNPlocs. Hsapiens. dbSNP155. GRCh38 package

Description

Locations and alleles for single-nucleotide variants extracted from dbSNP Human Build 155.

Details

The 948,979,291 SNPs in this package were extracted from the RefSNP JSON files for chromosomes 1-22, X, Y, and MT, located at https://ftp.ncbi.nih.gov/snp/latest_release/JSON/ (these files were created by NCBI on May 25, 2021). These SNPs are compatible with packages **BSgenome.Hsapiens.NCBI.GRCh38** and **BSgenome.Hsapiens.UCSC.hg38** i.e. they can be "injected" in the BSgenome objects defined in these packages.

SNP positions and alleles are reported with respect to the *plus* strand.

Only SNPs of type *snv* (*single-nucleotide* variant a.k.a. *single-base substitution*) were kept. Other variant types supported by dbSNP are: *delins* (indel), *ins* (insertion), *del* (deletion), and *mnv* (multiple nucleotide variation). These other variants are NOT included in SNPlocs.Hsapiens.dbSNP155.GRCh38 but are available in the XtraSNPlocs.Hsapiens.dbSNP155.GRCh38 package.

Note

The SNPs in this package can be "injected" in BSgenome. Hsapiens. NCBI. GRCh38 or BSgenome. Hsapiens. UCSC. hg38, and will land at the correct position.

See ?injectSNPs in the **BSgenome** software package for more information about the SNP injection mechanism.

Author(s)

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References

```
SNP Home at NCBI: https://www.ncbi.nlm.nih.gov/snp dbSNP Human Build 155 Release announcement (June 22, 2021): https://www.ncbi.nlm.nih.gov/mailman/pipermail/dbsnp-announce/2021q2/000229.html
The GRCh38.pl3 assembly: https://www.ncbi.nlm.nih.gov/assembly/GCF_000001405.39/
```

The hg38 genome at UCSC (based on GRCh38.p13, as of April 2022, but the UCSC folks could change this in the future and base hg38 on a more recent patch release of GRCh38): http://genome.ucsc.edu/cgi-bin/hgGateway?db=hg38

See Also

- The XtraSNPlocs.Hsapiens.dbSNP155.GRCh38 package for SNPs of type other than snv.
- snpcount in the **BSgenome** software package for how to access the data stored in this package.
- IUPAC_CODE_MAP in the **Biostrings** package.
- The GPos class in the GenomicRanges package.
- injectSNPs in the **BSgenome** software package for SNP injection.
- The **VariantAnnotation** software package to annotate variants with respect to location and amino acid coding.

Examples

```
## -----
## A. BASIC USAGE
## -----
snps <- SNPlocs.Hsapiens.dbSNP155.GRCh38</pre>
snpcount(snps)
## Get the positions and alleles of all SNPs on chromosome 22:
chr22_snps <- snpsBySeqname(snps, "22")</pre>
chr22_snps
## Get the positions and alleles of all SNPs on chromosomes 22 and MT:
snpsBySeqname(snps, c("22", "MT"))
## -----
## B. EXTRACT SNP INFORMATION FOR A SET OF RS IDS
my_rsids <- c("rs2639606", "rs75264089", "rs73396229", "rs55871206",
           "rs10932221", "rs56219727", "rs73709730", "rs55838886",
           "rs3734153", "rs79381275", "rs1516535")
## Note that the 1st call to snpsById() takes a long time but subsequent
## calls are expected to be slightly faster.
my_snps <- snpsById(snps, my_rsids)</pre>
my_snps
```

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