MafDb.ESP6500SI.V2.SSA137.GRCh38

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MafDb.ESP6500SI.V2.SSA137.GRCh38-package

Annotation package for minor allele frequency data from the NHLBI ESP project

Description

This annotation package stores minor allele frequency (MAF) data values from the release ESP6500SI-V2 of the NHLBI Exome Sequencing project (ESP). The data are exposed to the user in the form of a MafDb object, named after the package and loaded into main memory only as different chromosomes and populations are being queried. The class definition and methods to access MafDb objects are found in the VariantFiltering software package. To minimize disk space and memory requirements, MAF values larger or equal than 0.1 are stored using two significant digits, while MAF values smaller than 0.1 are stored using one significant digit.

WARNING: The positions associated to these MAF data are based on the GRCh38 release of the human genome and they were lifted by the NHLBI ESP from GRCh37. This means, the variants were not called directly on the GRCh38 human genome reference sequence. Moreover, nonSNVs positions were lifted to single nucleotide positions and, for this reason, have been discarded in this package.

Format

MafDb.ESP6500SI.V2.SSA137.GRCh38 MafDb object containing MAF values from 6503 exomes downloaded in March 2017 from http://evs.gs.washington.edu/evs_bulk_data/ESP6500SI-V2-SSA137.GRCh38-liftover.snps_indels.vcf.tar.gz. See the inst/extdata/README file from the source code for more information on how to update these data.

Author(s)

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Source


Exome Variant Server, NHLBI GO Exome Sequencing Project (ESP), Seattle, WA (URL: http://evs.gs.washington.edu/EVS) [March, 2017, accessed]
MafDb-class mafByOverlaps mafById VariantFiltering

Examples

```r
library(MafDb.ESP6500SI.V2.SSA137.GRCh38)

ls("package:MafDb.ESP6500SI.V2.SSA137.GRCh38")

mafdb <- MafDb.ESP6500SI.V2.SSA137.GRCh38
mafdb

populations(mafdb)

## lookup allele frequencies for rs1129038, an SNP associated to blue and brown eye colors
## as reported in Eiberg et al. Blue eye color in humans may be caused by a perfectly associated
## founder mutation in a regulatory element located within the HERC2 gene inhibiting OCA2 expression.
rng <- GRanges("15", IRanges(28356859, 28356859))
mafByOverlaps(mafdb, rng)
mafByOverlaps(mafdb, "15:28356859-28356859")
mafByOverlaps(mafdb, "15:28356859")
mafById(mafdb, "rs1129038")
```
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