# MafDb.gnomADex.r2.1.GRCh38

March 26, 2025

MafDb.gnomADex.r2.1.GRCh38-package

Annotation package for minor allele frequency data from exomes of the Genome Aggregation Database

# **Description**

This annotation package stores minor allele frequency (MAF) data derived from the exome variant set release 2.1 of the Genome Aggregation Database (gnomAD). The data are exposed to the user in the form of a GScores 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 GScores objects are found in the GenomicScores 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.

Please consult the gnomAD FAQ page at http://gnomad.broadinstitute.org/faq before you use these data for your own research.

### Format

MafDb.gnomADex.r2.1.GRCh38 GScores object containing MAF values from gnomAD exomes downloaded on April 2019

# Author(s)

R. Castelo

#### **Source**

Karczewski et al. Variation across 141,456 human exomes and genomes reveals the spectrum of loss-of-function intolerance across human protein-coding genes. bioRxiv, 531210, 2019.

The Genome Aggregation Database (gnomAD), Cambridge, MA (URL: http://gnomad.broadinstitute.org) [April, 2019, accessed]

## See Also

GScores-class gscores GenomicScores

### **Examples**

```
library(SNPlocs.Hsapiens.dbSNP149.GRCh38)
library(MafDb.gnomADex.r2.1.GRCh38)
ls("package:MafDb.gnomADex.r2.1.GRCh38")
mafdb <- MafDb.gnomADex.r2.1.GRCh38</pre>
mafdb
citation(mafdb)
populations(mafdb)
## lookup allele frequencies for rs1129038, a SNP associated to blue and brown eye colors
## as reported by 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.
## Human Genetics, 123(2):177-87, 2008 [http://www.ncbi.nlm.nih.gov/pubmed/18172690]
snpdb <- SNPlocs.Hsapiens.dbSNP149.GRCh38</pre>
rng <- snpsById(snpdb, ids="rs1129038")</pre>
rng
gscores(mafdb, rng)
gscores(mafdb, GRanges("15:28111713"))
```

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