

Package ‘AlphaMissense.v2023.hg19’

December 10, 2025

Type Package

Title AlphaMissense v2023 Pathogenicity Scores AnnotationHub Resource
Metadata for hg19

Version 3.18.2

Encoding UTF-8

Description Store Google DeepMind AlphaMissense v2023 hg19 pathogenicity scores AnnotationHub Resource Metadata. Provide provenance and citation information for Google DeepMind AlphaMissense v2023 hg19 pathogenicity score AnnotationHub resources. Illustrate in a vignette how to access those resources.

VignetteBuilder knitr

License Artistic-2.0

Depends GenomicScores, AnnotationHub

Suggests knitr, rmarkdown, BiocStyle

biocViews AnnotationData, PackageType, AnnotationHub, Organism,
Homo_sapiens

organism Homo sapiens

provider Google DeepMind

provider_version v2023

git_url <https://git.bioconductor.org/packages/AlphaMissense.v2023.hg19>

git_branch devel

git_last_commit f8b2919

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Author Robert Castelo [aut, cre]

Maintainer Robert Castelo <robert.castelo@upf.edu>

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AlphaMissense.v2023.hg19

AlphaMissense Pathogenicity Scores AnnotationHub Resource Metadata for Homo sapiens (hg19)

Description

This annotation package stores AlphaMissense v2023 pathogenicity scores AnnotationHub resource metadata for Homo sapiens, genome version hg19.

Note

This data package was made from resources at Google DeepMind based on version v2023.

Author(s)

Robert Castelo <robert.castelo@upf.edu> [cre]

See Also

[GenomicScores](#)

Examples

```
library(GenomicScores)

gsco <- getGScores("AlphaMissense.v2023.hg19")
name(gsco)
type(gsco)
provider(gsco)
providerVersion(gsco)
organism(gsco)
referenceGenome(gsco)
seqlevelsStyle(gsco)
seqinfo(gsco)
head(seqnames(gsco))
head(seqlengths(gsco))
citation(gsco)
```

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