# Package 'miRNATarget'

June 6, 2024

Title gene target tabale of miRNA for human/mouse used for MiRaGE package
Version 1.42.0
Author Y-h. Taguchi <tag@granular.com>
Description gene target tabale of miRNA for human/mouse used for MiRaGE package
Maintainer Y-h. Taguchi <tag@granular.com>
Depends R (>= 2.10), Biobase
License GPL
biocViews ExperimentData, Homo\_sapiens\_Data
git\_url https://git.bioconductor.org/packages/miRNATarget
git\_branch RELEASE\_3\_19
git\_last\_commit\_b454d34
git\_last\_commit\_date 2024-04-30
Repository Bioconductor 3.19

Date/Publication 2024-06-06

## Contents

conv_id 4
HS_conv_id
HS_refseq_to_affy_hc_g110 6
HS_refseq_to_affy_hg_focus
HS_refseq_to_affy_hg_u133a 8
HS_refseq_to_affy_hg_u133a_2 9
HS_refseq_to_affy_hg_u133b 10
HS_refseq_to_affy_hg_u133_plus_2
HS_refseq_to_affy_hg_u95a
HS_refseq_to_affy_hg_u95av2
HS_refseq_to_affy_hg_u95b
HS_refseq_to_affy_hg_u95c
HS_refseq_to_affy_hg_u95d

HS_refseq_to_affy_hg_u95e	. 17
$HS_{refseq_to_affy_huex_1_0_{st_v2}}$	
HS_refseq_to_affy_hugenefi	
$HS_refseq_to_affy_hugene_1_0_st_v1$	
HS_refseq_to_affy_u133_x3p	
HS_refseq_to_agilent_cgh_44b	
HS_refseq_to_agilent_wholegenome	
HS_refseq_to_canonical_transcript_stable_id	
HS_refseq_to_ccds	
HS_refseq_to_codelink	
HS_refseq_to_embl	
HS_refseq_to_ensembl_gene_id	
HS_refseq_to_ensembl_peptide_id	
HS_refseq_to_ensembl_transcript_id	. 29
HS_refseq_to_entrezgene	. 30
HS_refseq_to_hgnc_id	31
HS_refseq_to_hgnc_symbol	31
HS_refseq_to_hgnc_transcript_name	
HS_refseq_to_HS_refseq_to_efg_agilent_sureprint_g3_ge_8x60k	
HS_refseq_to_HS_refseq_to_efg_agilent_wholegenome_4x44k_v1	
HS_refseq_to_HS_refseq_to_efg_agilent_wholegenome_4x44k_v2	
HS_refseq_to_HS_refseq_to_ensembl_exon_id	
HS_refseq_to_HS_refseq_to_uniprot_genename_transcript_name	
HS_refseq_to_HS_refseq_to_uniprot_sptrembl	
HS_refseq_to_HS_refseq_to_uniprot_swissprot	
HS_refseq_to_HS_refseq_to_uniprot_swissprot_accession	
HS_refseq_to_HS_refseq_to_wikigene_id	
HS_refseq_to_illumina_humanht_12	
HS_refseq_to_illumina_humanwg_6_v1	
HS_refseq_to_illumina_humanwg_6_v2	
HS_refseq_to_illumina_humanwg_6_v3	. 45
HS_refseq_to_interpro	46
HS_refseq_to_ipi	46
HS_refseq_to_merops	
HS_refseq_to_pdb	
$HS_refseq_to_pfam$	
HS_refseq_to_phalanx_onearray	
HS_refseq_to_protein_id	50
HS_refseq_to_refseq_dna	
HS_refseq_to_refseq_genomic	
HS_refseq_to_refseq_peptide	
HS_refseq_to_rfam	
HS_refseq_to_rfam_gene_name	
HS_refseq_to_rfam_transcript_name	
HS_refseq_to_smart	56
HS_refseq_to_tigrfam	57
HS_refseq_to_ucsc	58
HS_refseq_to_unigene	58

HS_refseq_to_uniprot_genename	59
HS_refseq_to_wikigene_name	
id_conv	61
MM_conv_id	62
MM_refseq_to_affy_mg_u74a	
MM_refseq_to_affy_mg_u74av2	
MM_refseq_to_affy_mg_u74b	
MM_refseq_to_affy_mg_u74bv2	
$MM_refseq_to_affy_mg_u74c$	
MM_refseq_to_affy_mg_u74cv2	
MM_refseq_to_affy_moe430a	
MM_refseq_to_affy_moe430b	
MM_refseq_to_affy_moex_1_0_st_v1	
· ·	
MM_refseq_to_affy_mogene_1_0_st_v1	
MM_refseq_to_affy_mouse430a_2	
MM_refseq_to_affy_mouse430_2	
MM_refseq_to_affy_mu11ksuba	
MM_refseq_to_affy_mu11ksubb	
MM_refseq_to_agilent_wholegenome	77
MM_refseq_to_canonical_transcript_stable_id	78
MM_refseq_to_ccds	79
MM_refseq_to_codelink	79
MM_refseq_to_embl	
MM_refseq_to_ensembl_gene_id	
MM_refseq_to_ensembl_peptide_id	
MM_refseq_to_ensembl_transcript_id	
MM_refseq_to_entrezgene	
MM_refseq_to_fantom	
MM_refseq_to_illumina_mousewg_6_v1	
MM_refseq_to_illumina_mousewg_6_v2	
MM_refseq_to_interpro	
MM_refseq_to_ipi	
MM_refseq_to_merops	
MM_refseq_to_mgi_id	
MM_refseq_to_mgi_symbol	
MM_refseq_to_mgi_transcript_name	
MM_refseq_to_MM_refseq_to_efg_agilent_sureprint_g3_ge_8x60k	92
MM_refseq_to_MM_refseq_to_efg_agilent_wholegenome_4x44k_v1	93
MM_refseq_to_MM_refseq_to_efg_agilent_wholegenome_4x44k_v2	94
MM_refseq_to_MM_refseq_to_ensembl_exon_id	
MM_refseq_to_MM_refseq_to_uniprot_genename_transcript_name	
MM_refseq_to_MM_refseq_to_uniprot_sptrembl	
MM_refseq_to_MM_refseq_to_uniprot_swissprot	
MM_refseq_to_MM_refseq_to_uniprot_swissprot_accession	
MM_refseq_to_MM_refseq_to_wikigene_id	
MM_refseq_to_pdb	
MM_refseq_to_pfam	
MM_refseq_to_phalanx_onearray	102

116

MM_refseq_to_protein_id
MM_refseq_to_refseq_dna
MM_refseq_to_refseq_peptide
MM_refseq_to_rfam
MM_refseq_to_rfam_gene_name
MM_refseq_to_rfam_transcript_name
MM_refseq_to_smart
MM_refseq_to_tigrfam
MM_refseq_to_ucsc
MM_refseq_to_unigene
MM_refseq_to_uniprot_genename
MM_refseq_to_wikigene_name112
TBL2
TBL2_HS
TBL2_MM

#### Index

conv	

miRNA conservation table of mouse/human, based upon TargetScan-Mouse/Human 6.1

## Description

This miRNA conservation table is for MiRaGE package. Actual name of data files is HS\_conv\_id for human and MM\_conv\_id for mouse, but name of loaded data frame is "conv\_id".

#### Usage

data(MM\_conv\_id)

## Format

The data set is a data frame. The number of rows is the number of miRNAs. The number of column is two.

#### Details

The first column includes miRNA names based upon miRBase Release 18 and the second column includes integers which describe miRNA conservation. 2 stands for broadly conserved (in MiRaGE, it corresponds to "conserved"), 1 stands for conserved (in MiRaGE, broadly conserved and conserved correspond to "weakly conserved"), 0 stands for others (in MiRaGE, "all" corresponds to all miRNAs).

## Note

How to generate this table, please see functions in MiRaGE package and vignette.

#### HS\_conv\_id

## References

For more details about TargetScan, access to http://www.targetscan.org/

#### Examples

data(MM\_conv\_id)

HS\_conv\_id

miRNA conservation table of human

## Description

This miRNA conservation table of human is for MiRaGE package, based upon TargetScanHuman 6.1

#### Usage

data(HS\_conv\_id)

#### Format

The data set is a data frame. The number of rows is the number of miRNAs. The number of column is two.

#### Details

The first column includes miRNA names based upon miRBase Release 18 and the second column includes integers which describe miRNA conservation. 2 stands for broadly conserved (in MiRaGE, it corresponds to "conserved"), 1 stands for conserved (in MiRaGE, broadly conserved and conserved correspond to "weakly conserved"), 0 stands for others (in MiRaGE, "all" corresponds to all miRNAs). Please note that the name of object loaded is not "HS\_conv\_id" but "conv\_id", because of the requirements by MiRaGE package.

#### Note

How to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about TargetScanHuman, access to http://www.targetscan.org/vert\_61/

## Examples

data(HS\_conv\_id)

#### HS\_refseq\_to\_affy\_hc\_g110

Conversion table between RefSeq and affy\_hc\_g110 for human

## Description

This gene id conversion table between RefSeq and affy\_hc\_g110 is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

```
data(HS_refseq_to_affy_hc_g110)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_hc\_g110. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_affy\_hc\_g110" but "id\_conv", because of the requirements by Mi-RaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

```
data(HS_refseq_to_affy_hc_g110)
```

HS\_refseq\_to\_affy\_hg\_focus

Conversion table between RefSeq and affy\_hg\_focus for human

## Description

This gene id conversion table between RefSeq and affy\_hg\_focus is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

data(HS\_refseq\_to\_affy\_hg\_focus)

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_hg\_focus. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_affy\_hg\_focus" but "id\_conv", because of the requirements by Mi-RaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(HS\_refseq\_to\_affy\_hg\_focus)

#### HS\_refseq\_to\_affy\_hg\_u133a

Conversion table between RefSeq and affy\_hg\_u133a for human

## Description

This gene id conversion table between RefSeq and affy\_hg\_u133a is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

```
data(HS_refseq_to_affy_hg_u133a)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_hg\_u133a. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_affy\_hg\_u133a" but "id\_conv", because of the requirements by Mi-RaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

```
data(HS_refseq_to_affy_hg_u133a)
```

HS\_refseq\_to\_affy\_hg\_u133a\_2

Conversion table between RefSeq and affy\_hg\_u133a\_2 for human

## Description

This gene id conversion table between RefSeq and affy\_hg\_u133a\_2 is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

```
data(HS_refseq_to_affy_hg_u133a_2)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_hg\_u133a\_2. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_affy\_hg\_u133a\_2" but "id\_conv", because of the requirements by Mi-RaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

```
data(HS_refseq_to_affy_hg_u133a_2)
```

#### HS\_refseq\_to\_affy\_hg\_u133b

Conversion table between RefSeq and affy\_hg\_u133b for human

## Description

This gene id conversion table between RefSeq and affy\_hg\_u133b is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

```
data(HS_refseq_to_affy_hg_u133b)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_hg\_u133b. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_affy\_hg\_u133b" but "id\_conv", because of the requirements by Mi-RaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

```
data(HS_refseq_to_affy_hg_u133b)
```

HS\_refseq\_to\_affy\_hg\_u133\_plus\_2

Conversion table between RefSeq and affy\_hg\_u133\_plus\_2 for human

## Description

This gene id conversion table between RefSeq and affy\_hg\_u133\_plus\_2 is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

data(HS\_refseq\_to\_affy\_hg\_u133\_plus\_2)

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_hg\_u133\_plus\_2. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_affy\_hg\_u133\_plus\_2" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(HS\_refseq\_to\_affy\_hg\_u133\_plus\_2)

#### HS\_refseq\_to\_affy\_hg\_u95a

Conversion table between RefSeq and affy\_hg\_u95a for human

## Description

This gene id conversion table between RefSeq and affy\_hg\_u95a is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

```
data(HS_refseq_to_affy_hg_u95a)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_hg\_u95a. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_affy\_hg\_u95a" but "id\_conv", because of the requirements by Mi-RaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(HS\_refseq\_to\_affy\_hg\_u95a)

HS\_refseq\_to\_affy\_hg\_u95av2

Conversion table between RefSeq and affy\_hg\_u95av2 for human

## Description

This gene id conversion table between RefSeq and affy\_hg\_u95av2 is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

data(HS\_refseq\_to\_affy\_hg\_u95av2)

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_hg\_u95av2. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_affy\_hg\_u95av2" but "id\_conv", because of the requirements by Mi-RaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

```
data(HS_refseq_to_affy_hg_u95av2)
```

#### HS\_refseq\_to\_affy\_hg\_u95b

Conversion table between RefSeq and affy\_hg\_u95b for human

## Description

This gene id conversion table between RefSeq and affy\_hg\_u95b is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

```
data(HS_refseq_to_affy_hg_u95b)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_hg\_u95b. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_affy\_hg\_u95b" but "id\_conv", because of the requirements by Mi-RaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(HS\_refseq\_to\_affy\_hg\_u95b)

HS\_refseq\_to\_affy\_hg\_u95c

Conversion table between RefSeq and affy\_hg\_u95c for human

## Description

This gene id conversion table between RefSeq and affy\_hg\_u95c is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

```
data(HS_refseq_to_affy_hg_u95c)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_hg\_u95c. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_affy\_hg\_u95c" but "id\_conv", because of the requirements by Mi-RaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(HS\_refseq\_to\_affy\_hg\_u95c)

#### HS\_refseq\_to\_affy\_hg\_u95d

Conversion table between RefSeq and affy\_hg\_u95d for human

## Description

This gene id conversion table between RefSeq and affy\_hg\_u95d is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

data(HS\_refseq\_to\_affy\_hg\_u95d)

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_hg\_u95d. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_affy\_hg\_u95d" but "id\_conv", because of the requirements by Mi-RaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(HS\_refseq\_to\_affy\_hg\_u95d)

HS\_refseq\_to\_affy\_hg\_u95e

Conversion table between RefSeq and affy\_hg\_u95e for human

## Description

This gene id conversion table between RefSeq and affy\_hg\_u95e is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

data(HS\_refseq\_to\_affy\_hg\_u95e)

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_hg\_u95e. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_affy\_hg\_u95e" but "id\_conv", because of the requirements by Mi-RaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(HS\_refseq\_to\_affy\_hg\_u95e)

HS\_refseq\_to\_affy\_huex\_1\_0\_st\_v2

Conversion table between RefSeq and affy\_huex\_1\_0\_st\_v2 for human

#### Description

This gene id conversion table between RefSeq and affy\_huex\_1\_0\_st\_v2 is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

data(HS\_refseq\_to\_affy\_huex\_1\_0\_st\_v2)

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_huex\_1\_0\_st\_v2. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_affy\_huex\_1\_0\_st\_v2" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

```
data(HS_refseq_to_affy_huex_1_0_st_v2)
```

HS\_refseq\_to\_affy\_hugenef1

Conversion table between RefSeq and affy\_hugenefl for human

## Description

This gene id conversion table between RefSeq and affy\_hugenefl is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

```
data(HS_refseq_to_affy_hugenefl)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_hugenefl. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_affy\_hugenefl" but "id\_conv", because of the requirements by Mi-RaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

```
data(HS_refseq_to_affy_hugenefl)
```

HS\_refseq\_to\_affy\_hugene\_1\_0\_st\_v1

Conversion table between RefSeq and affy\_hugene\_1\_0\_st\_v1 for human

## Description

This gene id conversion table between RefSeq and affy\_hugene\_1\_0\_st\_v1 is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

data(HS\_refseq\_to\_affy\_hugene\_1\_0\_st\_v1)

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_hugene\_1\_0\_st\_v1. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_affy\_hugene\_1\_0\_st\_v1" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(HS\_refseq\_to\_affy\_hugene\_1\_0\_st\_v1)

HS\_refseq\_to\_affy\_u133\_x3p

Conversion table between RefSeq and affy\_u133\_x3p for human

## Description

This gene id conversion table between RefSeq and affy\_u133\_x3p is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

```
data(HS_refseq_to_affy_u133_x3p)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_u133\_x3p. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_affy\_u133\_x3p" but "id\_conv", because of the requirements by Mi-RaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

```
data(HS_refseq_to_affy_u133_x3p)
```

HS\_refseq\_to\_agilent\_cgh\_44b

Conversion table between RefSeq and agilent\_cgh\_44b for human

## Description

This gene id conversion table between RefSeq and agilent\_cgh\_44b is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

```
data(HS_refseq_to_agilent_cgh_44b)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding agilent\_cgh\_44b. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_agilent\_cgh\_44b" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(HS\_refseq\_to\_agilent\_cgh\_44b)

HS\_refseq\_to\_agilent\_wholegenome

Conversion table between RefSeq and agilent\_wholegenome for human

## Description

This gene id conversion table between RefSeq and agilent\_wholegenome is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

data(HS\_refseq\_to\_agilent\_wholegenome)

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding agilent\_wholegenome. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_agilent\_wholegenome" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(HS\_refseq\_to\_agilent\_wholegenome)

HS\_refseq\_to\_canonical\_transcript\_stable\_id *Conversion table between RefSeq and canonical\_transcript\_stable\_id* 

for human

## Description

This gene id conversion table between RefSeq and canonical\_transcript\_stable\_id is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

data(HS\_refseq\_to\_canonical\_transcript\_stable\_id)

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

#### Details

The first column includes RefSeq gene id and the second column includes corresponding canonical\_transcript\_stable\_id. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_canonical\_transcript\_stable\_id" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(HS\_refseq\_to\_canonical\_transcript\_stable\_id)

HS\_refseq\_to\_ccds Conversion table between RefSeq and ccds for human

#### Description

This gene id conversion table between RefSeq and ccds is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

```
data(HS_refseq_to_ccds)
```

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding ccds. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_ccds" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(HS\_refseq\_to\_ccds)

HS\_refseq\_to\_codelink Conversion table between RefSeq and codelink for human

## Description

This gene id conversion table between RefSeq and codelink is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

```
data(HS_refseq_to_codelink)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding codelink. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_codelink" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

#### Examples

```
data(HS_refseq_to_codelink)
```

HS\_refseq\_to\_embl *Conversion table between RefSeq and embl for human* 

## Description

This gene id conversion table between RefSeq and embl is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

```
data(HS_refseq_to_embl)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

#### Details

The first column includes RefSeq gene id and the second column includes corresponding embl. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_embl" but "id\_conv", because of the requirements by MiRaGE package.

#### 26

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

#### Examples

data(HS\_refseq\_to\_embl)

HS\_refseq\_to\_ensembl\_gene\_id

Conversion table between RefSeq and ensembl\_gene\_id for human

## Description

This gene id conversion table between RefSeq and ensembl\_gene\_id is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

data(HS\_refseq\_to\_ensembl\_gene\_id)

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

#### Details

The first column includes RefSeq gene id and the second column includes corresponding ensembl\_gene\_id. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_ensembl\_gene\_id" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

#### Examples

data(HS\_refseq\_to\_ensembl\_gene\_id)

HS\_refseq\_to\_ensembl\_peptide\_id

Conversion table between RefSeq and ensembl\_peptide\_id for human

#### Description

This gene id conversion table between RefSeq and ensembl\_peptide\_id is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

data(HS\_refseq\_to\_ensembl\_peptide\_id)

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding ensembl\_peptide\_id. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_ensembl\_peptide\_id" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(HS\_refseq\_to\_ensembl\_peptide\_id)

HS\_refseq\_to\_ensembl\_transcript\_id

Conversion table between RefSeq and ensembl\_transcript\_id for human

## Description

This gene id conversion table between RefSeq and ensembl\_transcript\_id is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

data(HS\_refseq\_to\_ensembl\_transcript\_id)

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding ensembl\_transcript\_id. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_ensembl\_transcript\_id" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(HS\_refseq\_to\_ensembl\_transcript\_id)

#### HS\_refseq\_to\_entrezgene

Conversion table between RefSeq and entrezgene for human

## Description

This gene id conversion table between RefSeq and entrezgene is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

```
data(HS_refseq_to_entrezgene)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding entrezgene. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_entrezgene" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

```
data(HS_refseq_to_entrezgene)
```

HS\_refseq\_to\_hgnc\_id Conversion table between RefSeq and hgnc\_id for human

## Description

This gene id conversion table between RefSeq and hgnc\_id is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

```
data(HS_refseq_to_hgnc_id)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding hgnc\_id. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_hgnc\_id" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

#### Examples

```
data(HS_refseq_to_hgnc_id)
```

HS\_refseq\_to\_hgnc\_symbol

Conversion table between RefSeq and hgnc\_symbol for human

## Description

This gene id conversion table between RefSeq and hgnc\_symbol is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

```
data(HS_refseq_to_hgnc_symbol)
```

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding hgnc\_symbol. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_hgnc\_symbol" but "id\_conv", because of the requirements by Mi-RaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(HS\_refseq\_to\_hgnc\_symbol)

HS\_refseq\_to\_hgnc\_transcript\_name

Conversion table between RefSeq and hgnc\_transcript\_name for human

## Description

This gene id conversion table between RefSeq and hgnc\_transcript\_name is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

data(HS\_refseq\_to\_hgnc\_transcript\_name)

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

32

## Details

The first column includes RefSeq gene id and the second column includes corresponding hgnc\_transcript\_name. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_hgnc\_transcript\_name" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(HS\_refseq\_to\_hgnc\_transcript\_name)

HS_refseq_to_HS_refseq_to_efg_agilent_sureprint_g3_ge_8x60k					
	Conversion	table	between	RefSeq	and
	HS_refseq_to_efg_agilent_sureprint_g3_ge_8x60k for human				

## Description

This gene id conversion table between RefSeq and HS\_refseq\_to\_efg\_agilent\_sureprint\_g3\_ge\_8x60k is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

data(HS\_refseq\_to\_HS\_refseq\_to\_efg\_agilent\_sureprint\_g3\_ge\_8x60k)

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding HS\_refseq\_to\_efg\_agilent\_sureprint It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_HS\_refseq\_to\_efg\_agilent\_sureprint\_g3\_ge\_8x60k" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(HS\_refseq\_to\_HS\_refseq\_to\_efg\_agilent\_sureprint\_g3\_ge\_8x60k)

HS_refseq_to_HS_refseq_to_efg_agilent_wholegenome_4x44k_v1					
	Conversion	table	between	RefSeq	and
	HS_refseq_to_efg_agilent_wholegenome_4x44k_v1 for human				

#### Description

This gene id conversion table between RefSeq and HS\_refseq\_to\_efg\_agilent\_wholegenome\_4x44k\_v1 is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

data(HS\_refseq\_to\_HS\_refseq\_to\_efg\_agilent\_wholegenome\_4x44k\_v1)

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding HS\_refseq\_to\_efg\_agilent\_wholegen It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_HS\_refseq\_to\_efg\_agilent\_wholegenome\_4x44k\_v1" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

#### Examples

data(HS\_refseq\_to\_HS\_refseq\_to\_efg\_agilent\_wholegenome\_4x44k\_v1)

HS\_refseq\_to\_HS\_refseq\_to\_efg\_agilent\_wholegenome\_4x44k\_v2 *Conversion* table between RefSeq and HS\_refseq\_to\_efg\_agilent\_wholegenome\_4x44k\_v2 for human

## Description

This gene id conversion table between RefSeq and HS\_refseq\_to\_efg\_agilent\_wholegenome\_4x44k\_v2 is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

data(HS\_refseq\_to\_HS\_refseq\_to\_efg\_agilent\_wholegenome\_4x44k\_v2)

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

#### Details

The first column includes RefSeq gene id and the second column includes corresponding HS\_refseq\_to\_efg\_agilent\_wholegen It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_HS\_refseq\_to\_efg\_agilent\_wholegenome\_4x44k\_v2" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(HS\_refseq\_to\_HS\_refseq\_to\_efg\_agilent\_wholegenome\_4x44k\_v2)

HS\_refseq\_to\_HS\_refseq\_to\_ensembl\_exon\_id

Conversion table between RefSeq and HS\_refseq\_to\_ensembl\_exon\_id for human

## Description

This gene id conversion table between RefSeq and HS\_refseq\_to\_ensembl\_exon\_id is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

data(HS\_refseq\_to\_HS\_refseq\_to\_ensembl\_exon\_id)

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

#### Details

The first column includes RefSeq gene id and the second column includes corresponding HS\_refseq\_to\_ensembl\_exon\_id. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_HS\_refseq\_to\_ensembl\_exon\_id" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(HS\_refseq\_to\_HS\_refseq\_to\_ensembl\_exon\_id)

HS\_refseq\_to\_HS\_refseq\_to\_uniprot\_genename\_transcript\_name Conversion table between RefSeq and HS\_refseq\_to\_uniprot\_genename\_transcript\_name for human

## Description

This gene id conversion table between RefSeq and HS\_refseq\_to\_uniprot\_genename\_transcript\_name is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

data(HS\_refseq\_to\_HS\_refseq\_to\_uniprot\_genename\_transcript\_name)

### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

### Details

The first column includes RefSeq gene id and the second column includes corresponding HS\_refseq\_to\_uniprot\_genename\_tr It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_HS\_refseq\_to\_uniprot\_genename\_transcript\_name" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(HS\_refseq\_to\_HS\_refseq\_to\_uniprot\_genename\_transcript\_name)

HS\_refseq\_to\_HS\_refseq\_to\_uniprot\_sptrembl

Conversion table between RefSeq and HS\_refseq\_to\_uniprot\_sptrembl for human

### Description

This gene id conversion table between RefSeq and HS\_refseq\_to\_uniprot\_sptrembl is for MiRaGE package, based upon BioMart Gene ID Converter

### Usage

data(HS\_refseq\_to\_HS\_refseq\_to\_uniprot\_sptrembl)

### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

### Details

The first column includes RefSeq gene id and the second column includes corresponding HS\_refseq\_to\_uniprot\_sptrembl. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_HS\_refseq\_to\_uniprot\_sptrembl" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(HS\_refseq\_to\_HS\_refseq\_to\_uniprot\_sptrembl)

HS_refseq_to_HS_refseq_to_uniprot_swissprot					
	Conversion	table	between	RefSeq	and
	HS_refseq_to_uniprot_swissprot for human				

## Description

This gene id conversion table between RefSeq and HS\_refseq\_to\_uniprot\_swissprot is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

data(HS\_refseq\_to\_HS\_refseq\_to\_uniprot\_swissprot)

### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

### Details

The first column includes RefSeq gene id and the second column includes corresponding HS\_refseq\_to\_uniprot\_swissprot. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_HS\_refseq\_to\_uniprot\_swissprot" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(HS\_refseq\_to\_HS\_refseq\_to\_uniprot\_swissprot)

 HS\_refseq\_to\_HS\_refseq\_to\_uniprot\_swissprot\_accession

 Conversion
 table
 between
 RefSeq
 and

 HS\_refseq\_to\_uniprot\_swissprot\_accession for human

### Description

This gene id conversion table between RefSeq and HS\_refseq\_to\_uniprot\_swissprot\_accession is for MiRaGE package, based upon BioMart Gene ID Converter

### Usage

data(HS\_refseq\_to\_HS\_refseq\_to\_uniprot\_swissprot\_accession)

### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

### Details

The first column includes RefSeq gene id and the second column includes corresponding HS\_refseq\_to\_uniprot\_swissprot\_action in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_HS\_refseq\_to\_uniprot\_swissprot\_accession" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(HS\_refseq\_to\_HS\_refseq\_to\_uniprot\_swissprot\_accession)

HS\_refseq\_to\_HS\_refseq\_to\_wikigene\_id

Conversion table between RefSeq and HS\_refseq\_to\_wikigene\_id for human

# Description

This gene id conversion table between RefSeq and HS\_refseq\_to\_wikigene\_id is for MiRaGE package, based upon BioMart Gene ID Converter

# Usage

data(HS\_refseq\_to\_HS\_refseq\_to\_wikigene\_id)

# Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding HS\_refseq\_to\_wikigene\_id. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_HS\_refseq\_to\_wikigene\_id" but "id\_conv", because of the requirements by MiRaGE package.

### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(HS\_refseq\_to\_HS\_refseq\_to\_wikigene\_id)

HS\_refseq\_to\_illumina\_humanht\_12

Conversion table between RefSeq and illumina\_humanht\_12 for human

### Description

This gene id conversion table between RefSeq and illumina\_humanht\_12 is for MiRaGE package, based upon BioMart Gene ID Converter

# Usage

data(HS\_refseq\_to\_illumina\_humanht\_12)

# Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding illumina\_humanht\_12. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_illumina\_humanht\_12" but "id\_conv", because of the requirements by MiRaGE package.

### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(HS\_refseq\_to\_illumina\_humanht\_12)

HS\_refseq\_to\_illumina\_humanwg\_6\_v1

Conversion table between RefSeq and illumina\_humanwg\_6\_v1 for human

# Description

This gene id conversion table between RefSeq and illumina\_humanwg\_6\_v1 is for MiRaGE package, based upon BioMart Gene ID Converter

# Usage

data(HS\_refseq\_to\_illumina\_humanwg\_6\_v1)

# Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding illumina\_humanwg\_6\_v1. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_illumina\_humanwg\_6\_v1" but "id\_conv", because of the requirements by MiRaGE package.

### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(HS\_refseq\_to\_illumina\_humanwg\_6\_v1)

HS\_refseq\_to\_illumina\_humanwg\_6\_v2

Conversion table between RefSeq and illumina\_humanwg\_6\_v2 for human

# Description

This gene id conversion table between RefSeq and illumina\_humanwg\_6\_v2 is for MiRaGE package, based upon BioMart Gene ID Converter

# Usage

data(HS\_refseq\_to\_illumina\_humanwg\_6\_v2)

# Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding illumina\_humanwg\_6\_v2. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_illumina\_humanwg\_6\_v2" but "id\_conv", because of the requirements by MiRaGE package.

### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(HS\_refseq\_to\_illumina\_humanwg\_6\_v2)

HS\_refseq\_to\_illumina\_humanwg\_6\_v3

Conversion table between RefSeq and illumina\_humanwg\_6\_v3 for human

# Description

This gene id conversion table between RefSeq and illumina\_humanwg\_6\_v3 is for MiRaGE package, based upon BioMart Gene ID Converter

# Usage

data(HS\_refseq\_to\_illumina\_humanwg\_6\_v3)

# Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding illumina\_humanwg\_6\_v3. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_illumina\_humanwg\_6\_v3" but "id\_conv", because of the requirements by MiRaGE package.

### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(HS\_refseq\_to\_illumina\_humanwg\_6\_v3)

HS\_refseq\_to\_interpro Conversion table between RefSeq and interpro for human

### Description

This gene id conversion table between RefSeq and interpro is for MiRaGE package, based upon BioMart Gene ID Converter

### Usage

```
data(HS_refseq_to_interpro)
```

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

### Details

The first column includes RefSeq gene id and the second column includes corresponding interpro. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_interpro" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

# References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

### Examples

data(HS\_refseq\_to\_interpro)

HS\_refseq\_to\_ipi Conversion table between RefSeq and ipi for human

## Description

This gene id conversion table between RefSeq and ipi is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

data(HS\_refseq\_to\_ipi)

### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

### Details

The first column includes RefSeq gene id and the second column includes corresponding ipi. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_ipi" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

### Examples

data(HS\_refseq\_to\_ipi)

HS\_refseq\_to\_merops Conversion table between RefSeq and merops for human

### Description

This gene id conversion table between RefSeq and merops is for MiRaGE package, based upon BioMart Gene ID Converter

### Usage

```
data(HS_refseq_to_merops)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

#### Details

The first column includes RefSeq gene id and the second column includes corresponding merops. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_merops" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

### Examples

```
data(HS_refseq_to_merops)
```

HS\_refseq\_to\_pdb Conversion table between RefSeq and pdb for human

# Description

This gene id conversion table between RefSeq and pdb is for MiRaGE package, based upon BioMart Gene ID Converter

### Usage

```
data(HS_refseq_to_pdb)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

# Details

The first column includes RefSeq gene id and the second column includes corresponding pdb. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_pdb" but "id\_conv", because of the requirements by MiRaGE package.

### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

#### Examples

data(HS\_refseq\_to\_pdb)

48

HS\_refseq\_to\_pfam Conversion table between RefSeq and pfam for human

# Description

This gene id conversion table between RefSeq and pfam is for MiRaGE package, based upon BioMart Gene ID Converter

### Usage

```
data(HS_refseq_to_pfam)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

### Details

The first column includes RefSeq gene id and the second column includes corresponding pfam. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_pfam" but "id\_conv", because of the requirements by MiRaGE package.

### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

#### Examples

data(HS\_refseq\_to\_pfam)

HS\_refseq\_to\_phalanx\_onearray

Conversion table between RefSeq and phalanx\_onearray for human

### Description

This gene id conversion table between RefSeq and phalanx\_onearray is for MiRaGE package, based upon BioMart Gene ID Converter

### Usage

data(HS\_refseq\_to\_phalanx\_onearray)

# Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding phalanx\_onearray. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_phalanx\_onearray" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

# References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

### Examples

data(HS\_refseq\_to\_phalanx\_onearray)

HS\_refseq\_to\_protein\_id

Conversion table between RefSeq and protein\_id for human

### Description

This gene id conversion table between RefSeq and protein\_id is for MiRaGE package, based upon BioMart Gene ID Converter

### Usage

```
data(HS_refseq_to_protein_id)
```

### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

### Details

The first column includes RefSeq gene id and the second column includes corresponding protein\_id. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_protein\_id" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

### Examples

```
data(HS_refseq_to_protein_id)
```

HS\_refseq\_to\_refseq\_dna

Conversion table between RefSeq and refseq\_dna for human

### Description

This gene id conversion table between RefSeq and refseq\_dna is for MiRaGE package, based upon BioMart Gene ID Converter

# Usage

```
data(HS_refseq_to_refseq_dna)
```

### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding refseq\_dna. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_refseq\_dna" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

#### Examples

data(HS\_refseq\_to\_refseq\_dna)

HS\_refseq\_to\_refseq\_genomic

Conversion table between RefSeq and refseq\_genomic for human

# Description

This gene id conversion table between RefSeq and refseq\_genomic is for MiRaGE package, based upon BioMart Gene ID Converter

### Usage

data(HS\_refseq\_to\_refseq\_genomic)

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

# Details

The first column includes RefSeq gene id and the second column includes corresponding refseq\_genomic. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_refseq\_genomic" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(HS\_refseq\_to\_refseq\_genomic)

HS\_refseq\_to\_refseq\_peptide

Conversion table between RefSeq and refseq\_peptide for human

## Description

This gene id conversion table between RefSeq and refseq\_peptide is for MiRaGE package, based upon BioMart Gene ID Converter

### Usage

data(HS\_refseq\_to\_refseq\_peptide)

### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

# Details

The first column includes RefSeq gene id and the second column includes corresponding refseq\_peptide. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_refseq\_peptide" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(HS\_refseq\_to\_refseq\_peptide)

HS\_refseq\_to\_rfam Conversion table between RefSeq and rfam for human

# Description

This gene id conversion table between RefSeq and rfam is for MiRaGE package, based upon BioMart Gene ID Converter

### Usage

```
data(HS_refseq_to_rfam)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

### Details

The first column includes RefSeq gene id and the second column includes corresponding rfam. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_rfam" but "id\_conv", because of the requirements by MiRaGE package.

### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

#### Examples

data(HS\_refseq\_to\_rfam)

HS\_refseq\_to\_rfam\_gene\_name

Conversion table between RefSeq and rfam\_gene\_name for human

### Description

This gene id conversion table between RefSeq and rfam\_gene\_name is for MiRaGE package, based upon BioMart Gene ID Converter

# Usage

data(HS\_refseq\_to\_rfam\_gene\_name)

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

### Details

The first column includes RefSeq gene id and the second column includes corresponding rfam\_gene\_name. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_rfam\_gene\_name" but "id\_conv", because of the requirements by Mi-RaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(HS\_refseq\_to\_rfam\_gene\_name)

HS\_refseq\_to\_rfam\_transcript\_name

Conversion table between RefSeq and rfam\_transcript\_name for human

# Description

This gene id conversion table between RefSeq and rfam\_transcript\_name is for MiRaGE package, based upon BioMart Gene ID Converter

# Usage

data(HS\_refseq\_to\_rfam\_transcript\_name)

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

### Details

The first column includes RefSeq gene id and the second column includes corresponding rfam\_transcript\_name. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_rfam\_transcript\_name" but "id\_conv", because of the requirements by MiRaGE package.

### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

#### Examples

data(HS\_refseq\_to\_rfam\_transcript\_name)

HS\_refseq\_to\_smart Conversion table between RefSeq and smart for human

### Description

This gene id conversion table between RefSeq and smart is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

```
data(HS_refseq_to_smart)
```

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

### Details

The first column includes RefSeq gene id and the second column includes corresponding smart. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_smart" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

### Examples

```
data(HS_refseq_to_smart)
```

HS\_refseq\_to\_tigrfam Conversion table between RefSeq and tigrfam for human

### Description

This gene id conversion table between RefSeq and tigrfam is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

```
data(HS_refseq_to_tigrfam)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

# Details

The first column includes RefSeq gene id and the second column includes corresponding tigrfam. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_tigrfam" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

### Examples

data(HS\_refseq\_to\_tigrfam)

HS\_refseq\_to\_ucsc Conversion table between RefSeq and ucsc for human

### Description

This gene id conversion table between RefSeq and ucsc is for MiRaGE package, based upon BioMart Gene ID Converter

### Usage

```
data(HS_refseq_to_ucsc)
```

### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

### Details

The first column includes RefSeq gene id and the second column includes corresponding ucsc. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_ucsc" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

# References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

### Examples

data(HS\_refseq\_to\_ucsc)

HS\_refseq\_to\_unigene Conversion table between RefSeq and unigene for human

# Description

This gene id conversion table between RefSeq and unigene is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

```
data(HS_refseq_to_unigene)
```

### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

#### Details

The first column includes RefSeq gene id and the second column includes corresponding unigene. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_unigene" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

#### Examples

```
data(HS_refseq_to_unigene)
```

```
HS_refseq_to_uniprot_genename
```

Conversion table between RefSeq and uniprot\_genename for human

#### Description

This gene id conversion table between RefSeq and uniprot\_genename is for MiRaGE package, based upon BioMart Gene ID Converter

### Usage

```
data(HS_refseq_to_uniprot_genename)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

# Details

The first column includes RefSeq gene id and the second column includes corresponding uniprot\_genename. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_uniprot\_genename" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

#### Examples

data(HS\_refseq\_to\_uniprot\_genename)

HS\_refseq\_to\_wikigene\_name

Conversion table between RefSeq and wikigene\_name for human

# Description

This gene id conversion table between RefSeq and wikigene\_name is for MiRaGE package, based upon BioMart Gene ID Converter

# Usage

data(HS\_refseq\_to\_wikigene\_name)

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

#### Details

The first column includes RefSeq gene id and the second column includes corresponding wikigene\_name. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "HS\_refseq\_to\_wikigene\_name" but "id\_conv", because of the requirements by MiRaGE package.

# Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

#### Examples

data(HS\_refseq\_to\_wikigene\_name)

60

id\_conv

### Description

This gene id conversion table between RefSeq and gene id/probe id is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

data(MM\_refseq\_to\_wikigene\_name)

### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

#### Details

The first column includes RefSeq gene id and the second column includes corresponding gene id. It can be made out of getBM function in biomaRt package. Actual names of data file is HS\_refseq\_to\_[gene id/probe id] for human and MM\_refseq\_to\_[gene id/probe id] for mouse. [gene id/probe id] stands for various gene id / probe id, but loaded data frame has the name "id\_conv".

#### Note

How to generate this table, please see functions in MiRaGE package and vignette.

### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(MM\_refseq\_to\_wikigene\_name)

MM\_conv\_id

### Description

This miRNA conservation table of mouse is for MiRaGE package, based upon TargetScanMouse 6.1

### Usage

data(MM\_conv\_id)

# Format

The data set is a data frame. The number of rows is the number of miRNAs. The number of column is two.

## Details

The first column includes miRNA names based upon miRBase Release 18 and the second column includes integers which describe miRNA conservation. 2 stands for broadly conserved (in MiRaGE, it corresponds to "conserved"), 1 stands for conserved (in MiRaGE, broadly conserved and conserved correspond to "weakly conserved"), 0 stands for others (in MiRaGE, "all" corresponds to all miRNAs). Please note that the name of object loaded is not "MM\_conv\_id" but "conv\_id", because of the requirements by MiRaGE package.

### Note

How to generate this table, please see functions in MiRaGE package and vignette.

### References

For more details about TargetScanMouse, access to http://www.targetscan.org/mmu\_61/

### Examples

data(MM\_conv\_id)

MM\_refseq\_to\_affy\_mg\_u74a

Conversion table between RefSeq and affy\_mg\_u74a for mouse

## Description

This gene id conversion table between RefSeq and affy\_mg\_u74a is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

```
data(MM_refseq_to_affy_mg_u74a)
```

### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

# Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_mg\_u74a. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_affy\_mg\_u74a" but "id\_conv", because of the requirements by Mi-RaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

```
data(MM_refseq_to_affy_mg_u74a)
```

### MM\_refseq\_to\_affy\_mg\_u74av2

Conversion table between RefSeq and affy\_mg\_u74av2 for mouse

### Description

This gene id conversion table between RefSeq and affy\_mg\_u74av2 is for MiRaGE package, based upon BioMart Gene ID Converter

### Usage

data(MM\_refseq\_to\_affy\_mg\_u74av2)

### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

# Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_mg\_u74av2. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_affy\_mg\_u74av2" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

```
data(MM_refseq_to_affy_mg_u74av2)
```

MM\_refseq\_to\_affy\_mg\_u74b

Conversion table between RefSeq and affy\_mg\_u74b for mouse

### Description

This gene id conversion table between RefSeq and affy\_mg\_u74b is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

```
data(MM_refseq_to_affy_mg_u74b)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

# Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_mg\_u74b. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_affy\_mg\_u74b" but "id\_conv", because of the requirements by Mi-RaGE package.

# Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

```
data(MM_refseq_to_affy_mg_u74b)
```

### MM\_refseq\_to\_affy\_mg\_u74bv2

Conversion table between RefSeq and affy\_mg\_u74bv2 for mouse

# Description

This gene id conversion table between RefSeq and affy\_mg\_u74bv2 is for MiRaGE package, based upon BioMart Gene ID Converter

### Usage

data(MM\_refseq\_to\_affy\_mg\_u74bv2)

### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

# Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_mg\_u74bv2. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_affy\_mg\_u74bv2" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

```
data(MM_refseq_to_affy_mg_u74bv2)
```

MM\_refseq\_to\_affy\_mg\_u74c

Conversion table between RefSeq and affy\_mg\_u74c for mouse

## Description

This gene id conversion table between RefSeq and affy\_mg\_u74c is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

```
data(MM_refseq_to_affy_mg_u74c)
```

### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

# Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_mg\_u74c. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_affy\_mg\_u74c" but "id\_conv", because of the requirements by Mi-RaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

```
data(MM_refseq_to_affy_mg_u74c)
```

### MM\_refseq\_to\_affy\_mg\_u74cv2

Conversion table between RefSeq and affy\_mg\_u74cv2 for mouse

# Description

This gene id conversion table between RefSeq and affy\_mg\_u74cv2 is for MiRaGE package, based upon BioMart Gene ID Converter

### Usage

```
data(MM_refseq_to_affy_mg_u74cv2)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

# Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_mg\_u74cv2. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_affy\_mg\_u74cv2" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

```
data(MM_refseq_to_affy_mg_u74cv2)
```

MM\_refseq\_to\_affy\_moe430a

Conversion table between RefSeq and affy\_moe430a for mouse

## Description

This gene id conversion table between RefSeq and affy\_moe430a is for MiRaGE package, based upon BioMart Gene ID Converter

# Usage

data(MM\_refseq\_to\_affy\_moe430a)

### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

# Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_moe430a. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_affy\_moe430a" but "id\_conv", because of the requirements by Mi-RaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(MM\_refseq\_to\_affy\_moe430a)

### MM\_refseq\_to\_affy\_moe430b

Conversion table between RefSeq and affy\_moe430b for mouse

### Description

This gene id conversion table between RefSeq and affy\_moe430b is for MiRaGE package, based upon BioMart Gene ID Converter

### Usage

data(MM\_refseq\_to\_affy\_moe430b)

### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

# Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_moe430b. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_affy\_moe430b" but "id\_conv", because of the requirements by Mi-RaGE package.

# Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(MM\_refseq\_to\_affy\_moe430b)

MM\_refseq\_to\_affy\_moex\_1\_0\_st\_v1

Conversion table between RefSeq and affy\_moex\_1\_0\_st\_v1 for mouse

### Description

This gene id conversion table between RefSeq and affy\_moex\_1\_0\_st\_v1 is for MiRaGE package, based upon BioMart Gene ID Converter

# Usage

data(MM\_refseq\_to\_affy\_moex\_1\_0\_st\_v1)

### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

# Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_moex\_1\_0\_st\_v1. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_affy\_moex\_1\_0\_st\_v1" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

```
data(MM_refseq_to_affy_moex_1_0_st_v1)
```

MM\_refseq\_to\_affy\_mogene\_1\_0\_st\_v1

Conversion table between RefSeq and affy\_mogene\_1\_0\_st\_v1 for mouse

# Description

This gene id conversion table between RefSeq and affy\_mogene\_1\_0\_st\_v1 is for MiRaGE package, based upon BioMart Gene ID Converter

# Usage

data(MM\_refseq\_to\_affy\_mogene\_1\_0\_st\_v1)

# Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_mogene\_1\_0\_st\_v1. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_affy\_mogene\_1\_0\_st\_v1" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(MM\_refseq\_to\_affy\_mogene\_1\_0\_st\_v1)

MM\_refseq\_to\_affy\_mouse430a\_2

Conversion table between RefSeq and affy\_mouse430a\_2 for mouse

# Description

This gene id conversion table between RefSeq and affy\_mouse430a\_2 is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

data(MM\_refseq\_to\_affy\_mouse430a\_2)

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

# Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_mouse430a\_2. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_affy\_mouse430a\_2" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

```
data(MM_refseq_to_affy_mouse430a_2)
```

## MM\_refseq\_to\_affy\_mouse430\_2

Conversion table between RefSeq and affy\_mouse430\_2 for mouse

## Description

This gene id conversion table between RefSeq and affy\_mouse430\_2 is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

```
data(MM_refseq_to_affy_mouse430_2)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

# Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_mouse430\_2. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_affy\_mouse430\_2" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

```
data(MM_refseq_to_affy_mouse430_2)
```

## MM\_refseq\_to\_affy\_mu11ksuba

Conversion table between RefSeq and affy\_mul1ksuba for mouse

# Description

This gene id conversion table between RefSeq and affy\_mu11ksuba is for MiRaGE package, based upon BioMart Gene ID Converter

# Usage

data(MM\_refseq\_to\_affy\_mu11ksuba)

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

# Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_mu11ksuba. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_affy\_mu11ksuba" but "id\_conv", because of the requirements by Mi-RaGE package.

# Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(MM\_refseq\_to\_affy\_mu11ksuba)

## MM\_refseq\_to\_affy\_mu11ksubb

Conversion table between RefSeq and affy\_mul1ksubb for mouse

# Description

This gene id conversion table between RefSeq and affy\_mu11ksubb is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

data(MM\_refseq\_to\_affy\_mu11ksubb)

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

# Details

The first column includes RefSeq gene id and the second column includes corresponding affy\_mu11ksubb. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_affy\_mu11ksubb" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(MM\_refseq\_to\_affy\_mu11ksubb)

MM\_refseq\_to\_agilent\_wholegenome

Conversion table between RefSeq and agilent\_wholegenome for mouse

## Description

This gene id conversion table between RefSeq and agilent\_wholegenome is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

data(MM\_refseq\_to\_agilent\_wholegenome)

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

# Details

The first column includes RefSeq gene id and the second column includes corresponding agilent\_wholegenome. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_agilent\_wholegenome" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(MM\_refseq\_to\_agilent\_wholegenome)

for mouse

# Description

This gene id conversion table between RefSeq and canonical\_transcript\_stable\_id is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

data(MM\_refseq\_to\_canonical\_transcript\_stable\_id)

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding canonical\_transcript\_stable\_id. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_canonical\_transcript\_stable\_id" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(MM\_refseq\_to\_canonical\_transcript\_stable\_id)

MM\_refseq\_to\_ccds Conversion table between RefSeq and ccds for mouse

## Description

This gene id conversion table between RefSeq and ccds is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

```
data(MM_refseq_to_ccds)
```

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding ccds. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_ccds" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

# References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(MM\_refseq\_to\_ccds)

MM\_refseq\_to\_codelink Conversion table between RefSeq and codelink for mouse

## Description

This gene id conversion table between RefSeq and codelink is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

```
data(MM_refseq_to_codelink)
```

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

# Details

The first column includes RefSeq gene id and the second column includes corresponding codelink. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_codelink" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

```
data(MM_refseq_to_codelink)
```

MM\_refseq\_to\_embl Conversion table between RefSeq and embl for mouse

# Description

This gene id conversion table between RefSeq and embl is for MiRaGE package, based upon BioMart Gene ID Converter

# Usage

```
data(MM_refseq_to_embl)
```

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

#### Details

The first column includes RefSeq gene id and the second column includes corresponding embl. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_embl" but "id\_conv", because of the requirements by MiRaGE package.

## 80

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

#### Examples

data(MM\_refseq\_to\_embl)

MM\_refseq\_to\_ensembl\_gene\_id

Conversion table between RefSeq and ensembl\_gene\_id for mouse

# Description

This gene id conversion table between RefSeq and ensembl\_gene\_id is for MiRaGE package, based upon BioMart Gene ID Converter

# Usage

data(MM\_refseq\_to\_ensembl\_gene\_id)

### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding ensembl\_gene\_id. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_ensembl\_gene\_id" but "id\_conv", because of the requirements by MiRaGE package.

# Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

#### Examples

data(MM\_refseq\_to\_ensembl\_gene\_id)

MM\_refseq\_to\_ensembl\_peptide\_id

Conversion table between RefSeq and ensembl\_peptide\_id for mouse

# Description

This gene id conversion table between RefSeq and ensembl\_peptide\_id is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

data(MM\_refseq\_to\_ensembl\_peptide\_id)

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

# Details

The first column includes RefSeq gene id and the second column includes corresponding ensembl\_peptide\_id. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_ensembl\_peptide\_id" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(MM\_refseq\_to\_ensembl\_peptide\_id)

Conversion table between RefSeq and ensembl\_transcript\_id for mouse

# Description

This gene id conversion table between RefSeq and ensembl\_transcript\_id is for MiRaGE package, based upon BioMart Gene ID Converter

# Usage

data(MM\_refseq\_to\_ensembl\_transcript\_id)

# Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding ensembl\_transcript\_id. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_ensembl\_transcript\_id" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(MM\_refseq\_to\_ensembl\_transcript\_id)

## MM\_refseq\_to\_entrezgene

Conversion table between RefSeq and entrezgene for mouse

# Description

This gene id conversion table between RefSeq and entrezgene is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

```
data(MM_refseq_to_entrezgene)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

# Details

The first column includes RefSeq gene id and the second column includes corresponding entrezgene. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_entrezgene" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

```
data(MM_refseq_to_entrezgene)
```

MM\_refseq\_to\_fantom Conversion table between RefSeq and fantom for mouse

#### Description

This gene id conversion table between RefSeq and fantom is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

```
data(MM_refseq_to_fantom)
```

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

# Details

The first column includes RefSeq gene id and the second column includes corresponding fantom. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_fantom" but "id\_conv", because of the requirements by MiRaGE package.

# Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

### Examples

data(MM\_refseq\_to\_fantom)

MM\_refseq\_to\_illumina\_mousewg\_6\_v1

Conversion table between RefSeq and illumina\_mousewg\_6\_v1 for mouse

#### Description

This gene id conversion table between RefSeq and illumina\_mousewg\_6\_v1 is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

```
data(MM_refseq_to_illumina_mousewg_6_v1)
```

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding illumina\_mousewg\_6\_v1. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_illumina\_mousewg\_6\_v1" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(MM\_refseq\_to\_illumina\_mousewg\_6\_v1)

MM\_refseq\_to\_illumina\_mousewg\_6\_v2

Conversion table between RefSeq and illumina\_mousewg\_6\_v2 for mouse

# Description

This gene id conversion table between RefSeq and illumina\_mousewg\_6\_v2 is for MiRaGE package, based upon BioMart Gene ID Converter

# Usage

data(MM\_refseq\_to\_illumina\_mousewg\_6\_v2)

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding illumina\_mousewg\_6\_v2. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_illumina\_mousewg\_6\_v2" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

#### Examples

```
data(MM_refseq_to_illumina_mousewg_6_v2)
```

MM\_refseq\_to\_interpro Conversion table between RefSeq and interpro for mouse

# Description

This gene id conversion table between RefSeq and interpro is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

```
data(MM_refseq_to_interpro)
```

# Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

#### Details

The first column includes RefSeq gene id and the second column includes corresponding interpro. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_interpro" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

```
data(MM_refseq_to_interpro)
```

MM\_refseq\_to\_ipi Conversion table between RefSeq and ipi for mouse

## Description

This gene id conversion table between RefSeq and ipi is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

```
data(MM_refseq_to_ipi)
```

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

#### Details

The first column includes RefSeq gene id and the second column includes corresponding ipi. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_ipi" but "id\_conv", because of the requirements by MiRaGE package.

# Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(MM\_refseq\_to\_ipi)

MM\_refseq\_to\_merops Conversion table between RefSeq and merops for mouse

## Description

This gene id conversion table between RefSeq and merops is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

```
data(MM_refseq_to_merops)
```

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding merops. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_merops" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

# References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(MM\_refseq\_to\_merops)

MM\_refseq\_to\_mgi\_id Conversion table between RefSeq and mgi\_id for mouse

## Description

This gene id conversion table between RefSeq and mgi\_id is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

```
data(MM_refseq_to_mgi_id)
```

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

#### Details

The first column includes RefSeq gene id and the second column includes corresponding mgi\_id. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_mgi\_id" but "id\_conv", because of the requirements by MiRaGE package.

# Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

# References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

#### Examples

data(MM\_refseq\_to\_mgi\_id)

```
MM_refseq_to_mgi_symbol
```

Conversion table between RefSeq and mgi\_symbol for mouse

## Description

This gene id conversion table between RefSeq and mgi\_symbol is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

```
data(MM_refseq_to_mgi_symbol)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

# Details

The first column includes RefSeq gene id and the second column includes corresponding mgi\_symbol. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_mgi\_symbol" but "id\_conv", because of the requirements by Mi-RaGE package.

#### 90

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

```
data(MM_refseq_to_mgi_symbol)
```

MM\_refseq\_to\_mgi\_transcript\_name

Conversion table between RefSeq and mgi\_transcript\_name for mouse

## Description

This gene id conversion table between RefSeq and mgi\_transcript\_name is for MiRaGE package, based upon BioMart Gene ID Converter

# Usage

data(MM\_refseq\_to\_mgi\_transcript\_name)

### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

#### Details

The first column includes RefSeq gene id and the second column includes corresponding mgi\_transcript\_name. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_mgi\_transcript\_name" but "id\_conv", because of the requirements by MiRaGE package.

# Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

#### Examples

data(MM\_refseq\_to\_mgi\_transcript\_name)

MM\_refseq\_to\_MM\_refseq\_to\_efg\_agilent\_sureprint\_g3\_ge\_8x60k
Conversion table between RefSeq and
MM\_refseq\_to\_efg\_agilent\_sureprint\_g3\_ge\_8x60k for mouse

## Description

This gene id conversion table between RefSeq and MM\_refseq\_to\_efg\_agilent\_sureprint\_g3\_ge\_8x60k is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

data(MM\_refseq\_to\_MM\_refseq\_to\_efg\_agilent\_sureprint\_g3\_ge\_8x60k)

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding MM\_refseq\_to\_efg\_agilent\_sureprint It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_MM\_refseq\_to\_efg\_agilent\_sureprint\_g3\_ge\_8x60k" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(MM\_refseq\_to\_efg\_agilent\_sureprint\_g3\_ge\_8x60k)

MM_refseq_to_MM_refseq_to_efg_agilent_wholegenome_4x44k_v1						
	Conversion	table	between	RefSeq	and	
	MM_refseq_to_efg_agilent_wholegenome_4x44k_v1 for mouse					

## Description

This gene id conversion table between RefSeq and MM\_refseq\_to\_efg\_agilent\_wholegenome\_4x44k\_v1 is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

data(MM\_refseq\_to\_MM\_refseq\_to\_efg\_agilent\_wholegenome\_4x44k\_v1)

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding MM\_refseq\_to\_efg\_agilent\_wholeg It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_MM\_refseq\_to\_efg\_agilent\_wholegenome\_4x44k\_v1" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(MM\_refseq\_to\_MM\_refseq\_to\_efg\_agilent\_wholegenome\_4x44k\_v1)

 MM\_refseq\_to\_efg\_agilent\_wholegenome\_4x44k\_v2

 Conversion
 table
 between
 RefSeq
 and

 MM\_refseq\_to\_efg\_agilent\_wholegenome\_4x44k\_v2 for mouse

## Description

This gene id conversion table between RefSeq and MM\_refseq\_to\_efg\_agilent\_wholegenome\_4x44k\_v2 is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

data(MM\_refseq\_to\_MM\_refseq\_to\_efg\_agilent\_wholegenome\_4x44k\_v2)

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding MM\_refseq\_to\_efg\_agilent\_wholeg It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_MM\_refseq\_to\_efg\_agilent\_wholegenome\_4x44k\_v2" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(MM\_refseq\_to\_MM\_refseq\_to\_efg\_agilent\_wholegenome\_4x44k\_v2)

MM_refseq_to_MM_refse	eq_to_ensembl_exc	on_id			
	Conversion	table	between	RefSeq	and
MM_refseq_to_ensembl_exon_id for mouse					

## Description

This gene id conversion table between RefSeq and MM\_refseq\_to\_ensembl\_exon\_id is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

data(MM\_refseq\_to\_MM\_refseq\_to\_ensembl\_exon\_id)

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding MM\_refseq\_to\_ensembl\_exon\_id. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_MM\_refseq\_to\_ensembl\_exon\_id" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(MM\_refseq\_to\_MM\_refseq\_to\_ensembl\_exon\_id)

 MM\_refseq\_to\_MM\_refseq\_to\_uniprot\_genename\_transcript\_name

 Conversion
 table
 between
 RefSeq
 and

 MM\_refseq\_to\_uniprot\_genename\_transcript\_name for mouse

## Description

This gene id conversion table between RefSeq and MM\_refseq\_to\_uniprot\_genename\_transcript\_name is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

data(MM\_refseq\_to\_MM\_refseq\_to\_uniprot\_genename\_transcript\_name)

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding MM\_refseq\_to\_uniprot\_genename\_ It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_MM\_refseq\_to\_uniprot\_genename\_transcript\_name" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(MM\_refseq\_to\_MM\_refseq\_to\_uniprot\_genename\_transcript\_name)

MM_refseq_to_MM_refseq_to_uniprot_sptrembl						
	Conversion	table	between	RefSeq	and	
	MM_refseq_to_uniprot_sptrembl for mouse					

## Description

This gene id conversion table between RefSeq and MM\_refseq\_to\_uniprot\_sptrembl is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

data(MM\_refseq\_to\_MM\_refseq\_to\_uniprot\_sptrembl)

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding MM\_refseq\_to\_uniprot\_sptrembl. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_MM\_refseq\_to\_uniprot\_sptrembl" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(MM\_refseq\_to\_MM\_refseq\_to\_uniprot\_sptrembl)

and

## Description

This gene id conversion table between RefSeq and MM\_refseq\_to\_uniprot\_swissprot is for Mi-RaGE package, based upon BioMart Gene ID Converter

#### Usage

data(MM\_refseq\_to\_MM\_refseq\_to\_uniprot\_swissprot)

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding MM\_refseq\_to\_uniprot\_swissprot. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_MM\_refseq\_to\_uniprot\_swissprot" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(MM\_refseq\_to\_MM\_refseq\_to\_uniprot\_swissprot)

MM_refseq_to_MM_refseq_to_uniprot_swissprot_accession						
	Conversion	table	between	RefSeq	and	
	MM_refseq_to_uniprot_swissprot_accession for mouse					

## Description

This gene id conversion table between RefSeq and MM\_refseq\_to\_uniprot\_swissprot\_accession is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

data(MM\_refseq\_to\_MM\_refseq\_to\_uniprot\_swissprot\_accession)

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding MM\_refseq\_to\_uniprot\_swissprot\_a It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_MM\_refseq\_to\_uniprot\_swissprot\_accession" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(MM\_refseq\_to\_MM\_refseq\_to\_uniprot\_swissprot\_accession)

MM\_refseq\_to\_MM\_refseq\_to\_wikigene\_id

Conversion table between RefSeq and MM\_refseq\_to\_wikigene\_id for mouse

## Description

This gene id conversion table between RefSeq and MM\_refseq\_to\_wikigene\_id is for MiRaGE package, based upon BioMart Gene ID Converter

# Usage

data(MM\_refseq\_to\_MM\_refseq\_to\_wikigene\_id)

# Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding MM\_refseq\_to\_wikigene\_id. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_MM\_refseq\_to\_wikigene\_id" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(MM\_refseq\_to\_MM\_refseq\_to\_wikigene\_id)

MM\_refseq\_to\_pdb Conversion table between RefSeq and pdb for mouse

## Description

This gene id conversion table between RefSeq and pdb is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

```
data(MM_refseq_to_pdb)
```

# Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding pdb. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_pdb" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(MM\_refseq\_to\_pdb)

MM\_refseq\_to\_pfam Conversion table between RefSeq and pfam for mouse

## Description

This gene id conversion table between RefSeq and pfam is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

data(MM\_refseq\_to\_pfam)

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

#### Details

The first column includes RefSeq gene id and the second column includes corresponding pfam. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_pfam" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

# References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

#### Examples

data(MM\_refseq\_to\_pfam)

```
MM_refseq_to_phalanx_onearray
```

Conversion table between RefSeq and phalanx\_onearray for mouse

#### Description

This gene id conversion table between RefSeq and phalanx\_onearray is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

```
data(MM_refseq_to_phalanx_onearray)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

# Details

The first column includes RefSeq gene id and the second column includes corresponding phalanx\_onearray. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_phalanx\_onearray" but "id\_conv", because of the requirements by MiRaGE package.

# 102

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(MM\_refseq\_to\_phalanx\_onearray)

```
MM_refseq_to_protein_id
```

Conversion table between RefSeq and protein\_id for mouse

# Description

This gene id conversion table between RefSeq and protein\_id is for MiRaGE package, based upon BioMart Gene ID Converter

# Usage

```
data(MM_refseq_to_protein_id)
```

### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

#### Details

The first column includes RefSeq gene id and the second column includes corresponding protein\_id. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_protein\_id" but "id\_conv", because of the requirements by MiRaGE package.

# Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(MM\_refseq\_to\_protein\_id)

## MM\_refseq\_to\_refseq\_dna

Conversion table between RefSeq and refseq\_dna for mouse

## Description

This gene id conversion table between RefSeq and refseq\_dna is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

```
data(MM_refseq_to_refseq_dna)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

# Details

The first column includes RefSeq gene id and the second column includes corresponding refseq\_dna. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_refseq\_dna" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

```
data(MM_refseq_to_refseq_dna)
```

MM\_refseq\_to\_refseq\_peptide

Conversion table between RefSeq and refseq\_peptide for mouse

# Description

This gene id conversion table between RefSeq and refseq\_peptide is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

data(MM\_refseq\_to\_refseq\_peptide)

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding refseq\_peptide. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_refseq\_peptide" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(MM\_refseq\_to\_refseq\_peptide)

MM\_refseq\_to\_rfam Conversion table between RefSeq and rfam for mouse

# Description

This gene id conversion table between RefSeq and rfam is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

```
data(MM_refseq_to_rfam)
```

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding rfam. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_rfam" but "id\_conv", because of the requirements by MiRaGE package.

## Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

#### Examples

data(MM\_refseq\_to\_rfam)

MM\_refseq\_to\_rfam\_gene\_name

Conversion table between RefSeq and rfam\_gene\_name for mouse

## Description

This gene id conversion table between RefSeq and rfam\_gene\_name is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

data(MM\_refseq\_to\_rfam\_gene\_name)

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding rfam\_gene\_name. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_rfam\_gene\_name" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(MM\_refseq\_to\_rfam\_gene\_name)

MM\_refseq\_to\_rfam\_transcript\_name

Conversion table between RefSeq and rfam\_transcript\_name for mouse

# Description

This gene id conversion table between RefSeq and rfam\_transcript\_name is for MiRaGE package, based upon BioMart Gene ID Converter

# Usage

data(MM\_refseq\_to\_rfam\_transcript\_name)

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding rfam\_transcript\_name. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_rfam\_transcript\_name" but "id\_conv", because of the requirements by MiRaGE package.

# Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

#### Examples

data(MM\_refseq\_to\_rfam\_transcript\_name)

MM\_refseq\_to\_smart Conversion table between RefSeq and smart for mouse

## Description

This gene id conversion table between RefSeq and smart is for MiRaGE package, based upon BioMart Gene ID Converter

# Usage

```
data(MM_refseq_to_smart)
```

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding smart. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_smart" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## 108

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

```
data(MM_refseq_to_smart)
```

MM\_refseq\_to\_tigrfam Conversion table between RefSeq and tigrfam for mouse

## Description

This gene id conversion table between RefSeq and tigrfam is for MiRaGE package, based upon BioMart Gene ID Converter

### Usage

```
data(MM_refseq_to_tigrfam)
```

### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding tigrfam. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_tigrfam" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

# Examples

data(MM\_refseq\_to\_tigrfam)

MM\_refseq\_to\_ucsc Conversion table between RefSeq and ucsc for mouse

## Description

This gene id conversion table between RefSeq and ucsc is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

```
data(MM_refseq_to_ucsc)
```

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding ucsc. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_ucsc" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

## References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(MM\_refseq\_to\_ucsc)

MM\_refseq\_to\_unigene Conversion table between RefSeq and unigene for mouse

## Description

This gene id conversion table between RefSeq and unigene is for MiRaGE package, based upon BioMart Gene ID Converter

#### Usage

```
data(MM_refseq_to_unigene)
```

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

## Details

The first column includes RefSeq gene id and the second column includes corresponding unigene. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_unigene" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

#### Examples

data(MM\_refseq\_to\_unigene)

MM\_refseq\_to\_uniprot\_genename

Conversion table between RefSeq and uniprot\_genename for mouse

## Description

This gene id conversion table between RefSeq and uniprot\_genename is for MiRaGE package, based upon BioMart Gene ID Converter

## Usage

```
data(MM_refseq_to_uniprot_genename)
```

## Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

#### Details

The first column includes RefSeq gene id and the second column includes corresponding uniprot\_genename. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_uniprot\_genename" but "id\_conv", because of the requirements by MiRaGE package.

#### Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(MM\_refseq\_to\_uniprot\_genename)

MM\_refseq\_to\_wikigene\_name

Conversion table between RefSeq and wikigene\_name for mouse

# Description

This gene id conversion table between RefSeq and wikigene\_name is for MiRaGE package, based upon BioMart Gene ID Converter

# Usage

data(MM\_refseq\_to\_wikigene\_name)

#### Format

The data set is a data frame. The number of rows is the number of gene id. The number of column is two.

#### Details

The first column includes RefSeq gene id and the second column includes corresponding wikigene\_name. It can be made out of getBM function in biomaRt package. Please note that the name of object loaded is not "MM\_refseq\_to\_wikigene\_name" but "id\_conv", because of the requirements by MiRaGE package.

# Note

For more details on how to generate this table, please see functions in MiRaGE package and vignette.

#### References

For more details about BioMart Gene ID Converter, access to http://www.ensembl.org/biomart/martview/

## Examples

data(MM\_refseq\_to\_wikigene\_name)

112

TBL2

## Description

This target gene table of miRNA is for MiRaGE package. Actual name of data file is either TBL2\_MM (for mouse) or TBL2\_HS (for human), but name of data frame loaded is "TBL2"

## Usage

data(TBL2\_MM)

# Format

The data set is a data frame. The number of rows is the number of miRNAs. The number of column is the number of RefSeq gene id.

#### Details

This target gene table is generated by simple seed match between 3' UTR RefSeq gene and 5' UTR of miRNAs. 3' UTR RefSeq gene sequence is taken from genome table. miRNA mature sequences are taken from miRBase Release 18. Row names are miRNA names and column names are RefSeq gene id.

#### Note

How to generate this table, please see functions in MiRaGE package and vignette.

## References

M, Yoshizawa, Y-h. Taguchi and J. Yasuda, Inference of Gene Regulation via miRNAs During ES Cell Differentiation Using MiRaGE Method, Int. J. Mol. Sci. 2011, 12(12), 9265-9276

# Examples

data(TBL2\_MM)

TBL2\_HS

# Description

This target gene table miRNA of human is for MiRaGE package

## Usage

data(TBL2\_HS)

# Format

The data set is a data frame. The number of rows is the number of miRNAs. The number of column is the number of RefSeq gene id.

## Details

This target gene table is generated by simple seed match between 3' UTR RefSeq gene and 5' UTR of miRNAs. 3' UTR RefSeq gene sequence is taken from genome table, from hg19 genome sequence. miRNA mature sequences are taken from miRBase Release 18. Row names are miRNA names and column names are RefSeq gene id.

## Note

How to generate this table, please see functions in MiRaGE package and vignette.

# References

M, Yoshizawa, Y-h. Taguchi and J. Yasuda, Inference of Gene Regulation via miRNAs During ES Cell Differentiation Using MiRaGE Method, Int. J. Mol. Sci. 2011, 12(12), 9265-9276

## Examples

data(TBL2\_HS)

TBL2\_MM

Target gene table miRNA of mouse

# Description

This target gene table miRNA of mouse is for MiRaGE package

#### Usage

data(TBL2\_MM)

# TBL2\_MM

# Format

The data set is a data frame. The number of rows is the number of miRNAs. The number of column is the number of RefSeq gene id.

# Details

This target gene table is generated by simple seed match between 3' UTR RefSeq gene and 5' UTR of miRNAs. 3' UTR RefSeq gene sequence is taken from genome table, from mm9 genome sequence. miRNA mature sequences are taken from miRBase Release 18. Row names are miRNA names and column names are RefSeq gene id.

## Note

How to generate this table, please see functions in MiRaGE package and vignette.

# References

M, Yoshizawa, Y-h. Taguchi and J. Yasuda, Inference of Gene Regulation via miRNAs During ES Cell Differentiation Using MiRaGE Method, Int. J. Mol. Sci. 2011, 12(12), 9265-9276

# Examples

data(TBL2\_MM)

# Index

\* data conv\_id, 4 HS\_conv\_id, 5 HS\_refseq\_to\_affy\_hc\_g110, 6 HS\_refseq\_to\_affy\_hg\_focus, 7 HS\_refseq\_to\_affy\_hg\_u133\_plus\_2, 11 HS\_refseq\_to\_affy\_hg\_u133a, 8 HS\_refseq\_to\_affy\_hg\_u133a\_2,9 HS\_refseq\_to\_affy\_hg\_u133b, 10 HS\_refseq\_to\_affy\_hg\_u95a, 12 HS\_refseq\_to\_affy\_hg\_u95av2, 13 HS\_refseq\_to\_affy\_hg\_u95b, 14 HS\_refseq\_to\_affy\_hg\_u95c, 15 HS\_refseq\_to\_affy\_hg\_u95d, 16 HS\_refseq\_to\_affy\_hg\_u95e, 17 HS\_refseq\_to\_affy\_huex\_1\_0\_st\_v2, 18 HS\_refseq\_to\_affy\_hugene\_1\_0\_st\_v1, 20HS\_refseq\_to\_affy\_hugenef1, 19 HS\_refseq\_to\_affy\_u133\_x3p, 21 HS\_refseq\_to\_agilent\_cgh\_44b, 22 HS\_refseq\_to\_agilent\_wholegenome, 23 HS\_refseq\_to\_canonical\_transcript\_stable\_id, HS\_refseq\_to\_illumina\_humanwg\_6\_v3, 24 HS\_refseq\_to\_ccds, 25 HS\_refseq\_to\_codelink, 25 HS\_refseq\_to\_embl, 26 HS\_refseq\_to\_ensembl\_gene\_id, 27 HS\_refseq\_to\_ensembl\_peptide\_id, 28 HS\_refseq\_to\_ensembl\_transcript\_id, 29 HS\_refseq\_to\_entrezgene, 30 HS\_refseq\_to\_hgnc\_id, 31 HS\_refseq\_to\_hgnc\_symbol, 31 HS\_refseq\_to\_hgnc\_transcript\_name,

#### 32

HS\_refseq\_to\_HS\_refseq\_to\_efg\_agilent\_sureprint\_g3\_ge\_ HS\_refseq\_to\_HS\_refseq\_to\_efg\_agilent\_wholegenome\_4x44 34 HS\_refseq\_to\_HS\_refseq\_to\_efg\_agilent\_wholegenome\_4x44 35 HS\_refseq\_to\_HS\_refseq\_to\_ensembl\_exon\_id, 36 HS\_refseq\_to\_HS\_refseq\_to\_uniprot\_genename\_transcript\_ 37 HS\_refseq\_to\_HS\_refseq\_to\_uniprot\_sptrembl, 38 HS\_refseq\_to\_HS\_refseq\_to\_uniprot\_swissprot, 39 HS\_refseq\_to\_HS\_refseq\_to\_uniprot\_swissprot\_accession, 40 HS\_refseq\_to\_HS\_refseq\_to\_wikigene\_id, 41 HS\_refseq\_to\_illumina\_humanht\_12, 42 HS\_refseq\_to\_illumina\_humanwg\_6\_v1, HS\_refseq\_to\_illumina\_humanwg\_6\_v2, 44 45 HS\_refseq\_to\_interpro, 46 HS\_refseq\_to\_ipi, 46 HS\_refseq\_to\_merops, 47 HS\_refseq\_to\_pdb, 48 HS\_refseq\_to\_pfam, 49 HS\_refseq\_to\_phalanx\_onearray, 49 HS\_refseq\_to\_protein\_id, 50 HS\_refseq\_to\_refseq\_dna, 51 HS\_refseq\_to\_refseq\_genomic, 52 HS\_refseq\_to\_refseq\_peptide, 53 HS\_refseq\_to\_rfam, 54 HS\_refseq\_to\_rfam\_gene\_name, 54

HS\_refseq\_to\_rfam\_transcript\_name, 55 HS\_refseq\_to\_smart, 56 HS\_refseq\_to\_tigrfam, 57 HS\_refseq\_to\_ucsc, 58 HS\_refseq\_to\_unigene, 58 HS\_refseq\_to\_uniprot\_genename, 59 HS\_refseq\_to\_wikigene\_name, 60 id\_conv, 61 MM\_conv\_id, 62 MM\_refseq\_to\_affy\_mg\_u74a, 63 MM\_refseq\_to\_affy\_mg\_u74av2, 64 MM\_refseq\_to\_affy\_mg\_u74b, 65 MM\_refseq\_to\_affy\_mg\_u74bv2,66 MM\_refseq\_to\_affy\_mg\_u74c, 67 MM\_refseq\_to\_affy\_mg\_u74cv2, 68 MM\_refseq\_to\_affy\_moe430a, 69 MM\_refseq\_to\_affy\_moe430b, 70 MM\_refseq\_to\_affy\_moex\_1\_0\_st\_v1, 71 MM\_refseq\_to\_affy\_mogene\_1\_0\_st\_v1, 72 MM\_refseq\_to\_affy\_mouse430\_2,74 MM\_refseq\_to\_affy\_mouse430a\_2,73 MM\_refseq\_to\_affy\_mu11ksuba, 75 MM\_refseq\_to\_affy\_mu11ksubb, 76 MM\_refseq\_to\_agilent\_wholegenome, 77 MM\_refseq\_to\_canonical\_transcript\_stable\_id, 78 MM\_refseq\_to\_ccds, 79 MM\_refseq\_to\_codelink, 79 MM\_refseq\_to\_embl, 80 MM\_refseq\_to\_ensembl\_gene\_id, 81 MM\_refseq\_to\_ensembl\_peptide\_id, 82 MM\_refseq\_to\_ensembl\_transcript\_id, 83  $MM_refseq_to_entrezgene, 84$ MM\_refseq\_to\_fantom, 85 MM\_refseq\_to\_illumina\_mousewg\_6\_v1, 85 MM\_refseq\_to\_illumina\_mousewg\_6\_v2, 86 MM\_refseq\_to\_interpro, 87 MM\_refseq\_to\_ipi, 88 MM\_refseq\_to\_merops, 89 MM\_refseq\_to\_mgi\_id, 89

MM\_refseq\_to\_mgi\_symbol, 90 MM\_refseq\_to\_mgi\_transcript\_name, 91 MM\_refseq\_to\_MM\_refseq\_to\_efg\_agilent\_sureprint\_g3\_ge\_ 92 MM\_refseq\_to\_MM\_refseq\_to\_efg\_agilent\_wholegenome\_4x44 93 MM\_refseq\_to\_MM\_refseq\_to\_efg\_agilent\_wholegenome\_4x44 94 MM\_refseq\_to\_MM\_refseq\_to\_ensembl\_exon\_id, 95 MM\_refseq\_to\_MM\_refseq\_to\_uniprot\_genename\_transcript\_ 96 MM\_refseq\_to\_MM\_refseq\_to\_uniprot\_sptrembl, 97 MM\_refseq\_to\_MM\_refseq\_to\_uniprot\_swissprot, 98 MM\_refseq\_to\_MM\_refseq\_to\_uniprot\_swissprot\_accession, 99 MM\_refseq\_to\_MM\_refseq\_to\_wikigene\_id, 100 MM\_refseq\_to\_pdb, 101 MM\_refseq\_to\_pfam, 101 MM\_refseq\_to\_phalanx\_onearray, 102 MM\_refseq\_to\_protein\_id, 103 MM\_refseq\_to\_refseq\_dna, 104 MM\_refseq\_to\_refseq\_peptide, 105 MM\_refseq\_to\_rfam, 106 MM\_refseq\_to\_rfam\_gene\_name, 106 MM\_refseq\_to\_rfam\_transcript\_name, 107 MM\_refseq\_to\_smart, 108 MM\_refseq\_to\_tigrfam, 109 MM\_refseq\_to\_ucsc, 110 MM\_refseq\_to\_unigene, 110 MM\_refseq\_to\_uniprot\_genename, 111 MM\_refseq\_to\_wikigene\_name, 112 TBL2, 113 TBL2\_HS, 114 TBL2\_MM, 114 conv\_id, 4 HS\_conv\_id, 5 HS\_refseq\_to\_affy\_hc\_g110, 6 HS\_refseq\_to\_affy\_hg\_focus, 7 HS\_refseq\_to\_affy\_hg\_u133\_plus\_2, 11 HS\_refseq\_to\_affy\_hg\_u133a, 8

HS\_refseq\_to\_affy\_hg\_u133a\_2,9

```
HS_refseq_to_affy_hg_u133b, 10
                                               HS_refseq_to_ipi, 46
                                               HS_refseq_to_merops, 47
HS_refseq_to_affy_hg_u95a, 12
                                               HS_refseq_to_pdb, 48
HS_refseq_to_affy_hg_u95av2, 13
HS_refseq_to_affy_hg_u95b, 14
                                               HS_refseq_to_pfam, 49
                                               HS_refseq_to_phalanx_onearray, 49
HS_refseq_to_affy_hg_u95c, 15
                                               HS_refseq_to_protein_id, 50
HS_refseq_to_affy_hg_u95d, 16
                                               HS_refseq_to_refseq_dna, 51
HS_refseq_to_affy_hg_u95e, 17
                                               HS_refseq_to_refseq_genomic, 52
HS_refseq_to_affy_huex_1_0_st_v2, 18
                                               HS_refseq_to_refseq_peptide, 53
HS_refseq_to_affy_hugene_1_0_st_v1, 20
                                               HS_refseq_to_rfam, 54
HS_refseq_to_affy_hugenef1, 19
                                               HS_refseq_to_rfam_gene_name, 54
HS_refseq_to_affy_u133_x3p, 21
                                               HS_refseq_to_rfam_transcript_name, 55
HS_refseq_to_agilent_cgh_44b, 22
                                               HS_refseq_to_smart, 56
HS_refseq_to_agilent_wholegenome, 23
                                               HS_refseq_to_tigrfam, 57
HS_refseq_to_canonical_transcript_stable_id,
                                               HS_refseq_to_ucsc, 58
        24
                                               HS_refseq_to_unigene, 58
HS_refseq_to_ccds, 25
                                               HS_refseq_to_uniprot_genename, 59
HS_refseq_to_codelink, 25
                                               HS_refseq_to_wikigene_name, 60
HS_refseq_to_embl, 26
HS_refseq_to_ensembl_gene_id, 27
                                               id_conv, 61
HS_refseq_to_ensembl_peptide_id, 28
HS_refseq_to_ensembl_transcript_id, 29
                                               MM conv id. 62
HS_refseq_to_entrezgene, 30
                                               MM_refseq_to_affy_mg_u74a, 63
HS_refseq_to_hgnc_id, 31
                                               MM_refseq_to_affy_mg_u74av2, 64
HS_refseq_to_hgnc_symbol, 31
                                               MM_refseq_to_affy_mg_u74b, 65
HS_refseq_to_hgnc_transcript_name, 32
                                               MM_refseq_to_affy_mg_u74bv2, 66
HS_refseq_to_HS_refseq_to_efg_agilent_surepript_gafseq_t60&ffy_mg_u74c,67
        33
                                               MM_refseq_to_affy_mg_u74cv2,68
HS_refseq_to_HS_refseq_to_efg_agilent_wholegengere_fsed_to_affy_moe430a, 69
        34
                                               MM_refseq_to_affy_moe430b, 70
\label{eq:HS_refseq_to_HS_refseq_to_efg_agilent\_wholegeggggfgefeetsed_tsddk_tsd_affy\_moex\_1\_0\_st\_v1, 71
        35
                                               MM_refseq_to_affy_mogene_1_0_st_v1,72
HS_refseq_to_HS_refseq_to_ensembl_exon_id,
                                               MM_refseq_to_affy_mouse430_2,74
        36
                                               MM_refseq_to_affy_mouse430a_2,73
HS_refseq_to_HS_refseq_to_uniprot_genename_tnams_ferenew_affy_mu11ksuba,75
        37
                                               MM_refseq_to_affy_mu11ksubb, 76
HS_refseq_to_HS_refseq_to_uniprot_sptrembl,
                                               MM_refseq_to_agilent_wholegenome, 77
        38
                                               MM_refseq_to_canonical_transcript_stable_id,
HS_refseq_to_HS_refseq_to_uniprot_swissprot,
                                                       78
        39
                                               MM_refseq_to_ccds, 79
HS_refseq_to_HS_refseq_to_uniprot_swissprot_aqqesetseq_to_codelink,79
        40
                                               MM_refseq_to_embl, 80
HS_refseq_to_HS_refseq_to_wikigene_id,
                                               MM_refseq_to_ensembl_gene_id, 81
        41
                                               MM_refseq_to_ensembl_peptide_id, 82
HS_refseq_to_illumina_humanht_12, 42
                                               MM_refseq_to_ensembl_transcript_id, 83
HS_refseq_to_illumina_humanwg_6_v1, 43
                                               MM_refseq_to_entrezgene, 84
HS_refseq_to_illumina_humanwg_6_v2, 44
                                               MM_refseq_to_fantom, 85
HS_refseq_to_illumina_humanwg_6_v3, 45
                                               MM_refseq_to_illumina_mousewg_6_v1, 85
                                               MM_refseq_to_illumina_mousewg_6_v2, 86
HS_refseq_to_interpro, 46
```

# INDEX

MM\_refseq\_to\_interpro, 87 MM\_refseq\_to\_ipi, 88 MM\_refseq\_to\_merops, 89 MM\_refseq\_to\_mgi\_id, 89 MM\_refseq\_to\_mgi\_symbol, 90 MM\_refseq\_to\_mgi\_transcript\_name, 91 MM\_refseq\_to\_MM\_refseq\_to\_efg\_agilent\_sureprint\_g3\_ge\_8x60k, 92 MM\_refseq\_to\_efg\_agilent\_wholegenome\_4x44k\_v1, 93 MM\_refseq\_to\_MM\_refseq\_to\_efg\_agilent\_wholegenome\_4x44k\_v2, 94 MM\_refseq\_to\_MM\_refseq\_to\_ensembl\_exon\_id, 95 MM\_refseq\_to\_MM\_refseq\_to\_uniprot\_genename\_transcript\_name, 96 MM\_refseq\_to\_MM\_refseq\_to\_uniprot\_sptrembl, 97 MM\_refseq\_to\_MM\_refseq\_to\_uniprot\_swissprot, 98 MM\_refseq\_to\_MM\_refseq\_to\_uniprot\_swissprot\_accession, 99 MM\_refseq\_to\_MM\_refseq\_to\_wikigene\_id, 100 MM\_refseq\_to\_pdb, 101 MM\_refseq\_to\_pfam, 101 MM\_refseq\_to\_phalanx\_onearray, 102 MM\_refseq\_to\_protein\_id, 103 MM\_refseq\_to\_refseq\_dna, 104 MM\_refseq\_to\_refseq\_peptide, 105 MM\_refseq\_to\_rfam, 106 MM\_refseq\_to\_rfam\_gene\_name, 106 MM\_refseq\_to\_rfam\_transcript\_name, 107 MM\_refseq\_to\_smart, 108 MM\_refseq\_to\_tigrfam, 109 MM\_refseq\_to\_ucsc, 110 MM\_refseq\_to\_unigene, 110 MM\_refseq\_to\_uniprot\_genename, 111 MM\_refseq\_to\_wikigene\_name, 112 TBL2, 113 TBL2\_HS, 114

TBL2\_MM, 114