

# Package ‘ginmapper’

May 7, 2025

**Type** Package

**Title** Gene Identifier Mapper

**Version** 1.4.0

**Description** Provides functionalities to translate gene or protein identifiers between state-of-art biological databases: CARD (<https://card.mcmaster.ca/>), NCBI Protein, Nucleotide and Gene (<https://www.ncbi.nlm.nih.gov/>), UniProt (<https://www.uniprot.org/>) and KEGG (<https://www.kegg.jp/>). Also offers complementary functionality like NCBI identical proteins or UniProt similar genes clusters retrieval.

**License** GPL-3 + file LICENSE

**Encoding** UTF-8

**VignetteBuilder** knitr

**biocViews** Annotation, KEGG, Genetics, ThirdPartyClient, Software

**Imports** KEGGREST, UniProt.ws, XML, rentrez, httr, utils, memoise, cachem, jsonlite, rvest

**Suggests** RUnit, BiocGenerics, markdown, knitr

**git\_url** <https://git.bioconductor.org/packages/ginmapper>

**git\_branch** RELEASE\_3\_21

**git\_last\_commit** a4cc299

**git\_last\_commit\_date** 2025-04-15

**Repository** Bioconductor 3.21

**Date/Publication** 2025-05-07

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|                |                                              |
|----------------|----------------------------------------------|
| changeCARDPath | <i>Change path to download CARD database</i> |
|----------------|----------------------------------------------|

---

**Description**

Defines a new path to the folder where to download CARD database. By default, this path is the temporary folder tempdir().

**Usage**

```
changeCARDPath(path=tempdir())
```

**Arguments**

|      |                                                         |
|------|---------------------------------------------------------|
| path | valid route to desired folder. Default value: tempdir() |
|------|---------------------------------------------------------|

**Value**

Updated path for CARD database to be downloaded.

---

|              |                                                       |
|--------------|-------------------------------------------------------|
| getCARD2KEGG | <i>Translates a CARD database ID to KEGG database</i> |
|--------------|-------------------------------------------------------|

---

**Description**

Retrieves the ID translation of an specific instance of CARD database ID to the KEGG Genes database.

**Usage**

```
getCARD2KEGG(cardId, exhaustiveMapping = FALSE, detailedMapping = FALSE,  
byIdenticalProteins = TRUE, bySimilarGenes = TRUE)
```

**Arguments**

|                   |                                                                                                                                                                                                                                                                                                                       |
|-------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| cardId            | valid CARD database ARO ID or a vector of them                                                                                                                                                                                                                                                                        |
| exhaustiveMapping | logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one.                                                                                                                                                                                                     |
| detailedMapping   | logical value (TRUE or FALSE) indicating to return a vector of KEGG IDs if FALSE, or a named vector if TRUE. This named vector details the origin of the translated IDs, dividing them in DT, if it is a Direct Translation, or 1.0, 0.9 or 0.5, if it has been retrieved through any of the identical gene clusters. |

**byIdenticalProteins**

logical value (TRUE or FALSE) indicating to translate also through NCBI Identical Proteins. TRUE value could mean higher translation time and FALSE, fewer chances to translate the ID

**bySimilarGenes**

logical value (TRUE or FALSE) indicating to translate also through similar UniProt genes (100%, 90% or 50% of identity). Using TRUE value may take a long time, only advised if there is not a direct translation possible.

**Value**

A character vector of the same length as `cardId`. If parameter `exhaustiveMapping` is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

**Examples**

```
# Translate CARD ID '3000938' to KEGG
getCARD2KEGG('3000938', byIdenticalProteins = FALSE, bySimilarGenes = FALSE)

# Translate CARD IDs '3000938' and '3002557' to KEGG
getCARD2KEGG(c('3000938', '3002557'))

# Get all possible translations of IDs '3000938', '3002511' with detailed origin
getCARD2KEGG(c('3000938', '3002511'), exhaustiveMapping = TRUE,
detailedMapping = TRUE)
```

---

getCARD2NCBIGene

*Translates a CARD database ID to NCBI Gene database*


---

**Description**

Retrieves the ID translation of an specific instance of CARD database ID to the NCBI Gene database.

**Usage**

```
getCARD2NCBIGene(cardId, exhaustiveMapping = FALSE)
```

**Arguments**

`cardId` valid CARD database ARO ID or a vector of them

`exhaustiveMapping`

logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one.

**Value**

A character vector of the same length as `cardId`. If parameter `exhaustiveMapping` is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

### Examples

```
# Translate CARD ID '3002525' to NCBI Gene
getCARD2NCBIGene('3002525')

# Translate CARD IDs '3002524' and '3002525' to NCBI Gene
getCARD2NCBIGene(c('3002524', '3002525'))

# Get all possible translations of IDs '3005061' and '3002524'
getCARD2NCBIGene(c('3005061', '3002524'), exhaustiveMapping = TRUE)
```

---

`getCARD2NCBINucleotide`*Translates a CARD database ID to NCBI Nucleotide database*

---

### Description

Retrieves the ID translation of an specific instance of CARD database ID to the NCBI Nucleotide database. Note that, due to the nature of NCBI Nucleotide database, a complete genome is often retrieved.

### Usage

```
getCARD2NCBINucleotide(cardId)
```

### Arguments

|        |                                                |
|--------|------------------------------------------------|
| cardId | valid CARD database ARO ID or a vector of them |
|--------|------------------------------------------------|

### Value

A character vector of the same length as cardId.

### Examples

```
# Translate CARD ID '3002535' to NCBI Nucleotide
getCARD2NCBINucleotide('3002535')

# Translate CARD ID 'ARO:3002535' to NCBI Nucleotide
getCARD2NCBINucleotide('ARO:3002535')

# Translate CARD IDs '3002535' and '3003988' to NCBI Nucleotide
getCARD2NCBINucleotide(c('3002535', '3003988'))
```

---

|                     |                                                               |
|---------------------|---------------------------------------------------------------|
| getCARD2NCBIProtein | <i>Translates a CARD database ID to NCBI Protein database</i> |
|---------------------|---------------------------------------------------------------|

---

**Description**

Retrieves the ID translation of an specific instance of CARD database ID to the NCBI Protein database.

**Usage**

```
getCARD2NCBIProtein(cardId)
```

**Arguments**

|        |                                                |
|--------|------------------------------------------------|
| cardId | valid CARD database ARO ID or a vector of them |
|--------|------------------------------------------------|

**Value**

A character vector of the same length as cardId.

**Examples**

```
# Translate CARD ID '3002535' to NCBI Protein
getCARD2NCBIProtein('3002535')

# Translate CARD ID 'ARO:3002535' to NCBI Protein
getCARD2NCBIProtein('ARO:3002535')

# Translate CARD IDs '3002535' and '3003988' to NCBI Protein
getCARD2NCBIProtein(c('3002535', '3003988'))
```

---

|                 |                                                          |
|-----------------|----------------------------------------------------------|
| getCARD2UniProt | <i>Translates a CARD database ID to UniProt database</i> |
|-----------------|----------------------------------------------------------|

---

**Description**

Retrieves the ID translation of an specific instance of CARD database ID to the UniProt database.

**Usage**

```
getCARD2UniProt(cardId, exhaustiveMapping = FALSE, detailedMapping = FALSE)
```

**Arguments**

`cardId`                valid CARD database ARO ID or a vector of them

`exhaustiveMapping`    logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one.

`detailedMapping`      logical value (TRUE or FALSE) indicating to return a vector of UniProt IDs if FALSE, or a named vector if TRUE. This named vector details the origin of the translated IDs, dividing them in DT, if it is a Direct Translation, or 1.0, 0.9 or 0.5, if it has been retrieved through any of the identical gene clusters.

**Value**

A character vector of the same length as `cardId`. If parameter `exhaustiveMapping` is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

**Examples**

```
# Translate CARD ID '3002867' to UniProt
getCARD2UniProt('3002867')

# Translate CARD IDs '3002867' and '3002535' to UniProt
getCARD2UniProt(c('3002867', '3002535'))

# Get all possible translations of IDs '3002867', '3002535' with detailed origin
getCARD2UniProt(c('3002867', '3002535'), detailedMapping = TRUE,
exhaustiveMapping = TRUE)
```

---

|                |                                                |
|----------------|------------------------------------------------|
| getCARDVersion | <i>Prints CARD database version being used</i> |
|----------------|------------------------------------------------|

---

**Description**

Prints the downloaded CARD version and release date.

**Usage**

```
getCARDVersion()
```

**Value**

Downloaded CARD database version.

---

getKEGG2CARD

*Translates a KEGG database ID to CARD database*


---

### Description

Retrieves the ID translation of an specific instance of KEGG Genes database ID to the CARD database.

### Usage

```
getKEGG2CARD(keggId, exhaustiveMapping = FALSE, detailedMapping = FALSE,
bySimilarGenes = TRUE)
```

### Arguments

|                   |                                                                                                                                                                                                                                                                                                                       |
|-------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| keggId            | valid KEGG Genes database ID or a vector of them                                                                                                                                                                                                                                                                      |
| exhaustiveMapping | logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one.                                                                                                                                                                                                     |
| detailedMapping   | logical value (TRUE or FALSE) indicating to return a vector of CARD IDs if FALSE, or a named vector if TRUE. This named vector details the origin of the translated IDs, dividing them in DT, if it is a Direct Translation, or 1.0, 0.9 or 0.5, if it has been retrieved through any of the identical gene clusters. |
| bySimilarGenes    | logical value (TRUE or FALSE) indicating to translate also through similar UniProt genes (100%, 90% or 50% of identity). Using TRUE value may take a long time, only advised if there is not a direct translation possible.                                                                                           |

### Value

A character vector of the same length as keggId. If parameter exhaustiveMapping is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

### Examples

```
# Translate KEGG ID 'ag:ACC85616' to CARD
getKEGG2CARD('ag:ACC85616')

# Translate KEGG IDs 'ag:ACC85616' and 'ag:CAJ47134' to CARD
getKEGG2CARD(c('ag:ACC85616', 'ag:CAJ47134'))

# Get all possible translations of IDs 'ag:ACC85616', 'ag:CAJ47134'
# with detailed origin
getKEGG2CARD(c('ag:ACC85616', 'ag:CAJ47134'), detailedMapping = TRUE,
exhaustiveMapping = TRUE)
```

---

|                  |                                                            |
|------------------|------------------------------------------------------------|
| getKEGG2NCBIGene | <i>Translates a KEGG database ID to NCBI Gene database</i> |
|------------------|------------------------------------------------------------|

---

## Description

Retrieves the ID translation of an specific instance of KEGG Genes database ID to the NCBI Gene database.

## Usage

```
getKEGG2NCBIGene(keggId, exhaustiveMapping = FALSE, detailedMapping = FALSE,
  bySimilarGenes = TRUE)
```

## Arguments

|                   |                                                                                                                                                                                                                                                                                                                            |
|-------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| keggId            | valid KEGG Genes database ID or a vector of them                                                                                                                                                                                                                                                                           |
| exhaustiveMapping | logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one.                                                                                                                                                                                                          |
| detailedMapping   | logical value (TRUE or FALSE) indicating to return a vector of NCBI Gene IDs if FALSE, or a named vector if TRUE. This named vector details the origin of the translated IDs, dividing them in DT, if it is a Direct Translation, or 1.0, 0.9 or 0.5, if it has been retrieved through any of the identical gene clusters. |
| bySimilarGenes    | logical value (TRUE or FALSE) indicating to translate also through similar UniProt genes (100%, 90% or 50% of identity). Using TRUE value may take a long time, only advised if there is not a direct translation possible.                                                                                                |

## Value

A character vector of the same length as keggId. If parameter exhaustiveMapping is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

## Examples

```
# Translate KEGG ID 'aag:5579347' to NCBI Gene
getKEGG2NCBIGene('aag:5579347', detailedMapping = TRUE)

# Translate KEGG IDs 'aag:5579347' and 'abh:M3Q_2831' to NCBI Gene
getKEGG2NCBIGene(c('aag:5579347', 'abh:M3Q_2831'))

# Get all possible translations of IDs 'aag:5579347', 'abh:M3Q_2831' with
# detailed origin
getKEGG2NCBIGene(c('aag:5579347', 'abh:M3Q_2831'), exhaustiveMapping = TRUE,
  detailedMapping = TRUE)
```

---

```
getKEGG2NCBINucleotide
```

*Translates a KEGG database ID to NCBI Nucleotide database*

---

## Description

Retrieves the ID translation of an specific instance of KEGG Genes database ID to the NCBI Nucleotide database. Note that, due to the nature of NCBI Nucleotide database, a complete genome is often retrieved.

## Usage

```
getKEGG2NCBINucleotide(keggId, exhaustiveMapping = FALSE,
  detailedMapping = FALSE, bySimilarGenes = TRUE)
```

## Arguments

|                   |                                                                                                                                                                                                                                                                                                                                  |
|-------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| keggId            | valid KEGG Genes database ID or a vector of them                                                                                                                                                                                                                                                                                 |
| exhaustiveMapping | logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one.                                                                                                                                                                                                                |
| detailedMapping   | logical value (TRUE or FALSE) indicating to return a vector of NCBI Nucleotide IDs if FALSE, or a named vector if TRUE. This named vector details the origin of the translated IDs, dividing them in DT, if it is a Direct Translation, or 1.0, 0.9 or 0.5, if it has been retrieved through any of the identical gene clusters. |
| bySimilarGenes    | logical value (TRUE or FALSE) indicating to translate also through similar UniProt genes (100%, 90% or 50% of identity). Using TRUE value may take a long time, only advised if there is not a direct translation possible.                                                                                                      |

## Value

A character vector of the same length as keggId. If parameter exhaustiveMapping is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

## Examples

```
# Translate KEGG ID 'llo:LLO_2673' to NCBI Nucleotide
getKEGG2NCBINucleotide('llo:LLO_2673', bySimilarGenes = FALSE)

# Translate KEGG IDs 'llo:LLO_2673' and 'abh:M3Q_2831' to NCBI Nucleotide
getKEGG2NCBINucleotide(c('llo:LLO_2673', 'abh:M3Q_2831'))

# Get all possible translations of IDs 'aag:5579347', 'abh:M3Q_2831' with
# detailed origin
getKEGG2NCBINucleotide(c('aag:5579347', 'abh:M3Q_2831'),
  exhaustiveMapping = TRUE, detailedMapping = TRUE)
```

---

|                     |                                                               |
|---------------------|---------------------------------------------------------------|
| getKEGG2NCBIProtein | <i>Translates a KEGG database ID to NCBI Protein database</i> |
|---------------------|---------------------------------------------------------------|

---

## Description

Retrieves the ID translation of an specific instance of KEGG Genes database ID to the NCBI Protein database.

## Usage

```
getKEGG2NCBIProtein(keggId, exhaustiveMapping = FALSE, detailedMapping = FALSE,  
bySimilarGenes = TRUE)
```

## Arguments

|                   |                                                                                                                                                                                                                                                                                                                               |
|-------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| keggId            | valid KEGG Genes database ID or a vector of them                                                                                                                                                                                                                                                                              |
| exhaustiveMapping | logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one.                                                                                                                                                                                                             |
| detailedMapping   | logical value (TRUE or FALSE) indicating to return a vector of NCBI Protein IDs if FALSE, or a named vector if TRUE. This named vector details the origin of the translated IDs, dividing them in DT, if it is a Direct Translation, or 1.0, 0.9 or 0.5, if it has been retrieved through any of the identical gene clusters. |
| bySimilarGenes    | logical value (TRUE or FALSE) indicating to translate also through similar UniProt genes (100%, 90% or 50% of identity). Using TRUE value may take a long time, only advised if there is not a direct translation possible.                                                                                                   |

## Value

A character vector of the same length as keggId. If parameter exhaustiveMapping is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

## Examples

```
# Translate KEGG ID 'llo:LLO_2673' to NCBI Protein  
getKEGG2NCBIProtein('llo:LLO_2673')  
  
# Translate KEGG IDs 'llo:LLO_2673' and 'abh:M3Q_2831' to NCBI Protein  
getKEGG2NCBIProtein(c('llo:LLO_2673', 'abh:M3Q_2831'))  
  
# Get all possible translations of IDs 'llo:LLO_2673', 'abh:M3Q_2831' with  
# detailed origin  
getKEGG2NCBIProtein(c('llo:LLO_2673', 'abh:M3Q_2831'),  
exhaustiveMapping = TRUE, detailedMapping = TRUE)
```

---

|                 |                                                                |
|-----------------|----------------------------------------------------------------|
| getKEGG2UniProt | <i>Translates a KEGG Genes database ID to UniProt database</i> |
|-----------------|----------------------------------------------------------------|

---

**Description**

Retrieves the ID translation of an specific instance of KEGG Genes database ID to the UniProt database.

**Usage**

```
getKEGG2UniProt(keggId, exhaustiveMapping = FALSE)
```

**Arguments**

|                   |                                                                                                                   |
|-------------------|-------------------------------------------------------------------------------------------------------------------|
| keggId            | valid KEGG Genes database ID or a vector of them                                                                  |
| exhaustiveMapping | logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one. |

**Value**

A character vector of the same length as keggId. If parameter exhaustiveMapping is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

**Examples**

```
# Translate KEGG Genes ID 'llo:LL0_2673' to UniProt
getKEGG2UniProt('llo:LL0_2673')

# Translate KEGG IDs 'llo:LL0_2673' and 'abh:M3Q_2831' to NCBI UniProt
getKEGG2UniProt(c('llo:LL0_2673', 'abh:M3Q_2831'))

# Get all possible translations of IDs 'llo:LL0_2673', 'abh:M3Q_2831'
getKEGG2UniProt(c('llo:LL0_2673', 'abh:M3Q_2831'), exhaustiveMapping = TRUE)
```

---

|                |                                                |
|----------------|------------------------------------------------|
| getKEGGVersion | <i>Prints KEGG database version being used</i> |
|----------------|------------------------------------------------|

---

**Description**

Prints the accessed KEGG version and its corresponding release date. It also prints the API version release date.

**Usage**

```
getKEGGVersion()
```

**Value**

KEGG database and API version.

---

|                  |                                                            |
|------------------|------------------------------------------------------------|
| getNCBIGene2CARD | <i>Translates a NCBI Gene database ID to CARD database</i> |
|------------------|------------------------------------------------------------|

---

**Description**

Retrieves the ID translation of an specific instance of NCBI Gene database ID to the CARD database.

**Usage**

```
getNCBIGene2CARD(ncbiId, exhaustiveMapping = FALSE)
```

**Arguments**

|                   |                                                                                                                   |
|-------------------|-------------------------------------------------------------------------------------------------------------------|
| ncbiId            | valid NCBI Gene database ID or a vector of them                                                                   |
| exhaustiveMapping | logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one. |

**Value**

A character vector of the same length as ncbiId. If parameter exhaustiveMapping is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

**Examples**

```
# Translate NCBI Gene ID '3510143' to CARD
getNCBIGene2CARD('3510143')

# Translate NCBI Gene IDs '3510143' and '29426913' to CARD
getNCBIGene2CARD(c('3510143', '29426913'))

# Get all possible translations of IDs '3510143' and '29426913'
getNCBIGene2CARD(c('3510143', '29426913'), exhaustiveMapping = TRUE)
```

---

getNCBIGene2KEGG

*Translates a NCBI Gene database ID to KEGG database*


---

## Description

Retrieves the ID translation of an specific instance of NCBI Gene database ID to the KEGG Genes database.

## Usage

```
getNCBIGene2KEGG(ncbiId, exhaustiveMapping = FALSE, detailedMapping = FALSE,
  byIdenticalProteins = TRUE, bySimilarGenes = TRUE)
```

## Arguments

|                     |                                                                                                                                                                                                                                                                                                                       |
|---------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| ncbiId              | valid NCBI Gene database ID or a vector of them                                                                                                                                                                                                                                                                       |
| exhaustiveMapping   | logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one.                                                                                                                                                                                                     |
| detailedMapping     | logical value (TRUE or FALSE) indicating to return a vector of KEGG IDs if FALSE, or a named vector if TRUE. This named vector details the origin of the translated IDs, dividing them in DT, if it is a Direct Translation, or 1.0, 0.9 or 0.5, if it has been retrieved through any of the identical gene clusters. |
| byIdenticalProteins | logical value (TRUE or FALSE) indicating to translate also through NCBI Identical Proteins. TRUE value could mean higher translation time and FALSE, fewer chances to translate the ID                                                                                                                                |
| bySimilarGenes      | logical value (TRUE or FALSE) indicating to translate also through similar UniProt genes (100%, 90% or 50% of identity). Using TRUE value may take a long time, only advised if there is not a direct translation possible.                                                                                           |

## Value

A character vector of the same length as ncbiId. If parameter exhaustiveMapping is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

## Examples

```
# Translate NCBI Gene ID '76524190' to KEGG
getNCBIGene2KEGG('AFU38919', bySimilarGenes = FALSE, byIdenticalProteins = FALSE)

# Translate NCBI Gene IDs '76524190' and 'AFU38919' to KEGG
getNCBIGene2KEGG(c('76524190', 'AFU38919'))

# Get all possible translations of IDs '76524190' and 'AFU38919' with
# detailed origin
```

```
getNCBIGene2KEGG(c('76524190', 'AFU38919'), detailedMapping = TRUE,  
exhaustiveMapping = TRUE)
```

---

```
getNCBIGene2NCBINucleotide
```

*Translates a NCBI Gene database ID to NCBI Nucleotide database*

---

## Description

Retrieves the ID translation of an specific instance of NCBI Gene database ID to the NCBI Nucleotide database. Note that, due to the nature of NCBI Nucleotide database, a complete genome is often retrieved.

## Usage

```
getNCBIGene2NCBINucleotide(id, exhaustiveMapping=FALSE)
```

## Arguments

|                   |                                                                                                                   |
|-------------------|-------------------------------------------------------------------------------------------------------------------|
| id                | valid NCBI Gene database ID or a vector of them                                                                   |
| exhaustiveMapping | logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one. |

## Value

A character vector of the same length as id. If parameter exhaustiveMapping is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

## Examples

```
# Translate NCBI Gene ID '76524190' to NCBI Nucleotide  
getNCBIGene2NCBINucleotide('76524190')  
  
# Translate NCBI Gene IDs '76524190' and '1272' to NCBI Nucleotide  
getNCBIGene2NCBINucleotide(c('76524190', '1272'))  
  
# Get all possible translations of IDs '76524190' and '1272'  
getNCBIGene2NCBINucleotide(c('76524190', '1272'), exhaustiveMapping = TRUE)
```

---

`getNCBIGene2NCBIProtein`*Translates a NCBI Gene database ID to NCBI Protein database*

---

**Description**

Retrieves the ID translation of an specific instance of NCBI Gene database ID to the NCBI Protein database.

**Usage**

```
getNCBIGene2NCBIProtein(id, exhaustiveMapping=FALSE)
```

**Arguments**

|                                |                                                                                                                   |
|--------------------------------|-------------------------------------------------------------------------------------------------------------------|
| <code>id</code>                | valid NCBI Gene database ID or a vector of them                                                                   |
| <code>exhaustiveMapping</code> | logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one. |

**Value**

A character vector of the same length as `id`. If parameter `exhaustiveMapping` is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

**Examples**

```
# Translate NCBI Gene ID '76524190' to NCBI Protein
getNCBIGene2NCBIProtein('76524190')

# Translate NCBI Gene IDs '76524190' and '1272' to NCBI Protein
getNCBIGene2NCBIProtein(c('76524190', '1272'))

# Get all possible translations of IDs '76524190', '1272'
getNCBIGene2NCBIProtein(c('76524190', '1272'), exhaustiveMapping = TRUE)
```

---

`getNCBIGene2UniProt`*Translates a NCBI Gene database ID to UniProt database*

---

**Description**

Retrieves the ID translation of an specific instance of NCBI Gene database ID to the UniProt database.

**Usage**

```
getNCBIGene2UniProt(ncbiId, exhaustiveMapping = FALSE,
  detailedMapping = FALSE, byIdenticalProteins = TRUE)
```

**Arguments**

**ncbiId** valid NCBI Gene database ID or a vector of them

**exhaustiveMapping** logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one.

**detailedMapping** logical value (TRUE or FALSE) indicating to return a vector of UniProt IDs if FALSE, or a named vector if TRUE. This named vector details the origin of the translated IDs, dividing them in DT, if it is a Direct Translation, or 1.0, 0.9 or 0.5, if it has been retrieved through any of the identical gene clusters.

**byIdenticalProteins** logical value (TRUE or FALSE) indicating to translate also through NCBI Identical Proteins. TRUE value could mean higher translation time and FALSE, fewer chances to translate the ID

**Value**

A character vector of the same length as `ncbiId`. If parameter `exhaustiveMapping` is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

**Examples**

```
# Translate NCBI Gene ID '76524190' to UniProt
getNCBIGene2UniProt('76524190')

# Translate NCBI Gene IDs '76524190' and '1272' to UniProt
getNCBIGene2UniProt(c('76524190', '1272'))

# Get all possible translations of IDs '76524190', '1272' with detailed origin
getNCBIGene2UniProt(c('76524190', '1272'), detailedMapping = TRUE,
  exhaustiveMapping = TRUE)
```

---

```
getNCBIIdenticalProteins
```

*Search NCBI Identical Protein database by a given NCBI ID*

---

**Description**

Retrieves identical NCBI proteins to that of a given ID.

**Usage**

```
getNCBIIdenticalProteins(ncbiId, format='ids')
```

**Arguments**

|        |                                                                                                                                                                                                    |
|--------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| ncbiId | valid NCBI ID or a vector of them                                                                                                                                                                  |
| format | return format of the response. Default value ids returns a vector of the identical proteins IDs; dataframe value returns a dataframe with different information about the found identical proteins |

**Value**

Depending on parameter format, it will return a list of same length as ncbiId with identical NCBI Protein IDs or with dataframes containing different information about the identical proteins found: Id, Source, Nucleotide.Accession, Start, Stop, Strand, Protein, Protein.Name, Organism, Strain and Assembly.

**Examples**

```
# Get IDs on identical proteins to the one with ID 'AHA80958'
getNCBIIIdenticalProteins('AHA80958', format = 'ids')

# Get IDs on identical proteins to the ones with IDs 'AHA80958' and 'CAA76794'
getNCBIIIdenticalProteins(c('AHA80958', 'CAA76794'), format = 'ids')
```

---

```
getNCBINucleotide2CARD
```

*Translates a NCBI Nucleotide database ID to CARD database*

---

**Description**

Retrieves the ID translation of an specific instance of NCBI Nucleotide database ID to the CARD database.

**Usage**

```
getNCBINucleotide2CARD(ncbiId, exhaustiveMapping = FALSE)
```

**Arguments**

|                   |                                                                                                                   |
|-------------------|-------------------------------------------------------------------------------------------------------------------|
| ncbiId            | valid NCBI Nucleotide database ID or a vector of them                                                             |
| exhaustiveMapping | logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one. |

**Value**

A character vector of the same length as ncbiId. If parameter exhaustiveMapping is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

**Examples**

```
# Translate NCBI Nucleotide ID 'AY034138' to CARD
getNCBINucleotide2CARD('AY034138')

# Translate NCBI Nucleotide IDs 'AY034138' and 'X54723.1' to CARD
getNCBINucleotide2CARD(c('AY034138', 'X54723.1'))

# Get all possible translations of IDs 'AY034138', 'X54723.1'
getNCBINucleotide2CARD(c('AY034138', 'X54723.1'), exhaustiveMapping = TRUE)
```

---

```
getNCBINucleotide2KEGG
```

*Translates a NCBI Nucleotide database ID to KEGG database*

---

**Description**

Retrieves the ID translation of an specific instance of NCBI Nucleotide database ID to the KEGG Genes database.

**Usage**

```
getNCBINucleotide2KEGG(ncbiId, exhaustiveMapping = FALSE,
  detailedMapping = FALSE, byIdenticalProteins = TRUE, bySimilarGenes = TRUE)
```

**Arguments**

|                     |                                                                                                                                                                                                                                                                                                                       |
|---------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| ncbiId              | valid NCBI Nucleotide database ID or a vector of them                                                                                                                                                                                                                                                                 |
| exhaustiveMapping   | logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one.                                                                                                                                                                                                     |
| detailedMapping     | logical value (TRUE or FALSE) indicating to return a vector of KEGG IDs if FALSE, or a named vector if TRUE. This named vector details the origin of the translated IDs, dividing them in DT, if it is a Direct Translation, or 1.0, 0.9 or 0.5, if it has been retrieved through any of the identical gene clusters. |
| byIdenticalProteins | logical value (TRUE or FALSE) indicating to translate also through NCBI Identical Proteins. TRUE value could mean higher translation time and FALSE, fewer chances to translate the ID                                                                                                                                |
| bySimilarGenes      | logical value (TRUE or FALSE) indicating to translate also through similar UniProt genes (100%, 90% or 50% of identity). Using TRUE value may take a long time, only advised if there is not a direct translation possible.                                                                                           |

**Value**

A character vector of the same length as ncbiId. If parameter exhaustiveMapping is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

**Examples**

```
# Translate NCBI Nucleotide ID 'NC_013861.1' to KEGG
getNCBINucleotide2KEGG('NC_013861.1', bySimilarGenes = FALSE, byIdenticalProteins = FALSE)

# Translate NCBI Nucleotide IDs 'AY536519' and 'NC_013861.1' to KEGG
getNCBINucleotide2KEGG(c('AY536519', 'NC_013861.1'))

# Get all possible translations of IDs 'AY536519', 'NC_013861.1'
# with detailed origin
getNCBINucleotide2KEGG(c('AY536519', 'NC_013861.1'), detailedMapping = TRUE,
exhaustiveMapping = TRUE)
```

---

```
getNCBINucleotide2NCBIGene
```

*Translates a NCBI Nucleotide database ID to NCBI Gene database*

---

**Description**

Retrieves the ID translation of an specific instance of NCBI Nucleotide database ID to the NCBI Gene database.

**Usage**

```
getNCBINucleotide2NCBIGene(id, exhaustiveMapping=FALSE)
```

**Arguments**

`id` valid NCBI Nucleotide database ID or a vector of them  
`exhaustiveMapping` logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one.

**Value**

A character vector of the same length as `id`. If parameter `exhaustiveMapping` is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

**Examples**

```
# Translate NCBI Nucleotide ID 'Z21488' to NCBI Gene
getNCBINucleotide2NCBIGene('Z21488')

# Get all possible translations of ID 'Z21488'
getNCBINucleotide2NCBIGene(c('Z21488'), exhaustiveMapping = TRUE)
```

---

`getNCBINucleotide2NCBIProtein`*Translates a NCBI Nucleotide database ID to NCBI Protein database*

---

**Description**

Retrieves the ID translation of an specific instance of NCBI Nucleotide database ID to the NCBI Protein database.

**Usage**

```
getNCBINucleotide2NCBIProtein(id, exhaustiveMapping=FALSE)
```

**Arguments**

|                                |                                                                                                                   |
|--------------------------------|-------------------------------------------------------------------------------------------------------------------|
| <code>id</code>                | valid NCBI Nucleotide database ID or a vector of them                                                             |
| <code>exhaustiveMapping</code> | logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one. |

**Value**

A character vector of the same length as `id`. If parameter `exhaustiveMapping` is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

**Examples**

```
# Translate NCBI Nucleotide ID 'AY536519' to NCBI Protein
getNCBINucleotide2NCBIProtein('AY536519')

# Translate NCBI Nucleotide IDs 'AY536519' and 'X54723.1' to NCBI Protein
getNCBINucleotide2NCBIProtein(c('AY536519', 'X54723.1'))

# Get all possible translations of IDs 'AY536519', 'X54723.1'
getNCBINucleotide2NCBIProtein(c('AY536519', 'X54723.1'),
exhaustiveMapping = TRUE)
```

---

`getNCBINucleotide2UniProt`*Translates a NCBI Nucleotide database ID to UniProt database*

---

**Description**

Retrieves the ID translation of an specific instance of NCBI Nucleotide database ID to the UniProt database.

**Usage**

```
getNCBINucleotide2UniProt(ncbiId, exhaustiveMapping = FALSE,
  detailedMapping = FALSE, byIdenticalProteins = TRUE)
```

**Arguments**

**ncbiId**                valid NCBI Nucleotide database ID or a vector of them

**exhaustiveMapping**       logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one.

**detailedMapping**       logical value (TRUE or FALSE) indicating to return a vector of UniProt IDs if FALSE, or a named vector if TRUE. This named vector details the origin of the translated IDs, dividing them in DT, if it is a Direct Translation, or 1.0, 0.9 or 0.5, if it has been retrieved through any of the identical gene clusters.

**byIdenticalProteins**     logical value (TRUE or FALSE) indicating to translate also through NCBI Identical Proteins. TRUE value could mean higher translation time and FALSE, fewer chances to translate the ID

**Value**

A character vector of the same length as `ncbiId`. If parameter `exhaustiveMapping` is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

**Examples**

```
# Translate NCBI Nucleotide ID 'AY536519' to UniProt
getNCBINucleotide2UniProt('AY536519')

# Translate NCBI Nucleotide IDs 'AY536519' and 'X54723.1' to UniProt
getNCBINucleotide2UniProt(c('AY536519', 'X54723.1'))

# Get all possible translations of IDs 'AY536519', 'X54723.1'
# with detailed origin
getNCBINucleotide2UniProt(c('AY536519', 'X54723.1'), detailedMapping = TRUE,
  exhaustiveMapping = TRUE)
```

---

|                     |                                                               |
|---------------------|---------------------------------------------------------------|
| getNCBIProtein2CARD | <i>Translates a NCBI Protein database ID to CARD database</i> |
|---------------------|---------------------------------------------------------------|

---

**Description**

Retrieves the ID translation of an specific instance of NCBI Protein database ID to the CARD database.

**Usage**

```
getNCBIProtein2CARD(ncbiId, exhaustiveMapping = FALSE)
```

**Arguments**

**ncbiId**                valid NCBI Protein database ID or a vector of them  
**exhaustiveMapping**       logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one.

**Value**

A character vector of the same length as **ncbiId**. If parameter **exhaustiveMapping** is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

**Examples**

```
# Translate NCBI Protein ID 'AAK64581' to CARD
getNCBIProtein2CARD('AAK64581')

# Translate NCBI Protein IDs 'AAK64581' and 'WP_010896559.1' to CARD
getNCBIProtein2CARD(c('AAK64581', 'WP_010896559.1'))

# Get all possible translations of IDs 'AAK64581', 'WP_010896559.1'
getNCBIProtein2CARD(c('AAK64581', 'WP_010896559.1'), exhaustiveMapping = TRUE)
```

---

|                     |                                                               |
|---------------------|---------------------------------------------------------------|
| getNCBIProtein2KEGG | <i>Translates a NCBI Protein database ID to KEGG database</i> |
|---------------------|---------------------------------------------------------------|

---

**Description**

Retrieves the ID translation of an specific instance of NCBI Protein database ID to the KEGG Genes database.

**Usage**

```
getNCBIProtein2KEGG(ncbiId, exhaustiveMapping = FALSE, detailedMapping = FALSE,
byIdenticalProteins = TRUE, bySimilarGenes = TRUE)
```

**Arguments**

**ncbiId**                valid NCBI Protein database ID or a vector of them  
**exhaustiveMapping**       logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one.

**detailedMapping**

logical value (TRUE or FALSE) indicating to return a vector of KEGG IDs if FALSE, or a named vector if TRUE. This named vector details the origin of the translated IDs, dividing them in DT, if it is a Direct Translation, or 1.0, 0.9 or 0.5, if it has been retrieved through any of the identical gene clusters.

**byIdenticalProteins**

logical value (TRUE or FALSE) indicating to translate also through NCBI Identical Proteins. TRUE value could mean higher translation time and FALSE, fewer chances to translate the ID

**bySimilarGenes**

logical value (TRUE or FALSE) indicating to translate also through similar UniProt genes (100%, 90% or 50% of identity). Using TRUE value may take a long time, only advised if there is not a direct translation possible.

**Value**

A character vector of the same length as ncbiId. If parameter exhaustiveMapping is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

**Examples**

```
# Translate NCBI Protein ID 'AFU38919' to KEGG
getNCBIProtein2KEGG('AFU38919', exhaustiveMapping = FALSE,
detailedMapping = FALSE, byIdenticalProteins = FALSE, bySimilarGenes = FALSE)

# Translate NCBI Protein IDs 'AFU38919' and 'WP_010896559.1' to KEGG
getNCBIProtein2KEGG(c('AFU38919', 'WP_010896559.1'),
byIdenticalProteins = FALSE)

# Get all possible translations of ID 'AFU38919', 'WP_010896559.1'
# with detailed origin
getNCBIProtein2KEGG(c('AFU38919', 'WP_010896559.1'), exhaustiveMapping = TRUE,
detailedMapping = TRUE)
```

---

```
getNCBIProtein2NCBIGene
```

*Translates a NCBI Protein database ID to NCBI Gene database*

---

**Description**

Retrieves the ID translation of an specific instance of NCBI Protein database ID to the NCBI Gene database.

**Usage**

```
getNCBIProtein2NCBIGene(id, exhaustiveMapping=FALSE)
```

**Arguments**

`id` valid NCBI Protein database ID or a vector of them  
`exhaustiveMapping` logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one.

**Value**

A character vector of the same length as `id`. If parameter `exhaustiveMapping` is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

**Examples**

```
# Translate NCBI Protein ID 'CAA79696' to NCBI Gene
getNCBIProtein2NCBIGene('CAA79696')

# Translate NCBI Protein IDs 'CAA79696' and 'WP_001082319' to NCBI Gene
getNCBIProtein2NCBIGene(c('CAA79696', 'WP_001082319'))

# Get all possible translations of IDs 'CAA79696', 'WP_001082319'
getNCBIProtein2NCBIGene(c('CAA79696', 'WP_001082319'), exhaustiveMapping=TRUE)
```

---

`getNCBIProtein2NCBINucleotide`

*Translates a NCBI Protein database ID to NCBI Nucleotide database*

---

**Description**

Retrieves the ID translation of an specific instance of NCBI Protein database ID to the NCBI Nucleotide database. Note that, due to the nature of NCBI Nucleotide database, a complete genome is often retrieved.

**Usage**

```
getNCBIProtein2NCBINucleotide(id, exhaustiveMapping=FALSE)
```

**Arguments**

`id` valid NCBI Protein database ID or a vector of them  
`exhaustiveMapping` logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one.

**Value**

A character vector of the same length as `id`. If parameter `exhaustiveMapping` is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

**Examples**

```
# Translate NCBI Protein ID 'AAS48620' to NCBI Nucleotide
getNCBIProtein2NCBINucleotide('AAS48620')

# Translate NCBI Protein IDs 'AAS48620' and 'CAA38525' to NCBI Nucleotide
getNCBIProtein2NCBINucleotide(c('AAS48620', 'CAA38525'))

# Get all possible translations of IDs 'AAS48620', 'CAA38525'
getNCBIProtein2NCBINucleotide(c('AAS48620', 'CAA38525'), exhaustiveMapping=TRUE)
```

---

```
getNCBIProtein2UniProt
```

*Translates a NCBI Protein database ID to UniProt database*

---

**Description**

Retrieves the ID translation of an specific instance of NCBI Protein database ID to the UniProt database.

**Usage**

```
getNCBIProtein2UniProt(ncbiId, exhaustiveMapping = FALSE,
  detailedMapping = FALSE, byIdenticalProteins = TRUE)
```

**Arguments**

|                     |                                                                                                                                                                                                                                                                                                                          |
|---------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| ncbiId              | valid NCBI Protein database ID or a vector of them                                                                                                                                                                                                                                                                       |
| exhaustiveMapping   | logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one.                                                                                                                                                                                                        |
| detailedMapping     | logical value (TRUE or FALSE) indicating to return a vector of UniProt IDs if FALSE, or a named vector if TRUE. This named vector details the origin of the translated IDs, dividing them in DT, if it is a Direct Translation, or 1.0, 0.9 or 0.5, if it has been retrieved through any of the identical gene clusters. |
| byIdenticalProteins | logical value (TRUE or FALSE) indicating to translate also through NCBI Identical Proteins. TRUE value could mean higher translation time and FALSE, fewer chances to translate the ID                                                                                                                                   |

**Value**

A character vector of the same length as ncbiId. If parameter exhaustiveMapping is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

### Examples

```
# Translate NCBI Protein ID 'WP_010896559.1' to UniProt
getNCBIProtein2UniProt('WP_010896559.1')

# Translate NCBI Protein IDs 'AAS48620' and 'WP_010896559' to UniProt
getNCBIProtein2UniProt(c('AAS48620', 'WP_010896559'))

# Get all possible translations of IDs 'AAS48620', 'WP_010896559'
# with detailed origin
getNCBIProtein2UniProt(c('AAS48620', 'WP_010896559'), detailedMapping = TRUE,
exhaustiveMapping = TRUE)
```

---

|                |                                                  |
|----------------|--------------------------------------------------|
| getNCBIVersion | <i>Prints NCBI databases versions being used</i> |
|----------------|--------------------------------------------------|

---

### Description

Prints the build version and release date of the Protein, Gene, Nucleotide and Identical Proteins NCBI databases.

### Usage

```
getNCBIVersion()
```

### Value

NCBI databases versions.

---

|                 |                                                           |
|-----------------|-----------------------------------------------------------|
| getUniProt2CARD | <i>Translates an UniProt database ID to CARD database</i> |
|-----------------|-----------------------------------------------------------|

---

### Description

Retrieves the ID translation of an specific instance of UniProt database ID to the CARD database.

### Usage

```
getUniProt2CARD(upId, exhaustiveMapping = FALSE, detailedMapping = FALSE,
bySimilarGenes = TRUE)
```

**Arguments**

- upId** valid UniProt database ID or a vector of them
- exhaustiveMapping** logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one.
- detailedMapping** logical value (TRUE or FALSE) indicating to return a vector of CARD IDs if FALSE, or a named vector if TRUE. This named vector details the origin of the translated IDs, dividing them in DT, if it is a Direct Translation, or 1.0, 0.9 or 0.5, if it has been retrieved through any of the identical gene clusters.
- bySimilarGenes** logical value (TRUE or FALSE) indicating to translate also through similar UniProt genes (100%, 90% or 50% of identity). Using TRUE value may take a long time, only advised if there is not a direct translation possible.

**Value**

A character vector of the same length as upId. If parameter exhaustiveMapping is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

**Examples**

```
# Translate UniProt ID 'A0A1S7BGS4' to CARD
getUniProt2CARD('A0A1S7BGS4')

# Translate UniProt IDs 'A0A1S7BGS4' and 'A0A6H2TXZ6' to CARD
getUniProt2CARD(c('A0A1S7BGS4', 'A0A6H2TXZ6'))

# Get all possible translations of IDs 'A0A1S7BGS4', 'A0A6H2TXZ6'
# with detailed origin
getUniProt2CARD(c('A0A1S7BGS4', 'A0A6H2TXZ6'), detailedMapping = TRUE,
exhaustiveMapping = TRUE)
```

---

getUniProt2KEGG

*Translates an UniProt database ID to KEGG database*


---

**Description**

Retrieves the ID translation of an specific instance of UniProt database ID to the KEGG Genes database.

**Usage**

```
getUniProt2KEGG(upId, exhaustiveMapping = FALSE, bySimilarGenes = TRUE,
detailedMapping = FALSE)
```

**Arguments**

|                   |                                                                                                                                                                                                                                                                                                                       |
|-------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| upId              | valid UniProt database ID or a vector of them.                                                                                                                                                                                                                                                                        |
| exhaustiveMapping | logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one.                                                                                                                                                                                                     |
| bySimilarGenes    | logical value (TRUE or FALSE) indicating to translate also through similar UniProt genes (100%, 90% or 50% of identity). Using TRUE value may take a long time, only advised if there is not a direct translation possible.                                                                                           |
| detailedMapping   | logical value (TRUE or FALSE) indicating to return a vector of KEGG IDs if FALSE, or a named vector if TRUE. This named vector details the origin of the translated IDs, dividing them in DT, if it is a Direct Translation, or 1.0, 0.9 or 0.5, if it has been retrieved through any of the identical gene clusters. |

**Value**

A character vector of the same length as upId. If parameter exhaustiveMapping is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

**Examples**

```
# Translate UniProt ID 'G9JVE6' to KEGG
getUniProt2KEGG('G9JVE6')

# Translate UniProt IDs 'A0A1S7BGS4' and 'G9JVE6' to KEGG
getUniProt2KEGG(c('A0A1S7BGS4', 'G9JVE6'))

# Get all possible translations of ID 'A0A2R4PHC7' with detailed origin
getUniProt2KEGG(c('test_id', 'A0A2R4PHC7'), detailedMapping = TRUE,
exhaustiveMapping = TRUE)
```

---

|                     |                                                                |
|---------------------|----------------------------------------------------------------|
| getUniProt2NCBIGene | <i>Translates an UniProt database ID to NCBI Gene database</i> |
|---------------------|----------------------------------------------------------------|

---

**Description**

Retrieves the ID translation of an specific instance of UniProt database ID to the NCBI Gene database.

**Usage**

```
getUniProt2NCBIGene(upId, exhaustiveMapping = FALSE, detailedMapping = FALSE,
bySimilarGenes = TRUE)
```

**Arguments**

|                   |                                                                                                                                                                                                                                                                                                                            |
|-------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| upId              | valid UniProt database ID or a vector of them                                                                                                                                                                                                                                                                              |
| exhaustiveMapping | logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one.                                                                                                                                                                                                          |
| detailedMapping   | logical value (TRUE or FALSE) indicating to return a vector of NCBI Gene IDs if FALSE, or a named vector if TRUE. This named vector details the origin of the translated IDs, dividing them in DT, if it is a Direct Translation, or 1.0, 0.9 or 0.5, if it has been retrieved through any of the identical gene clusters. |
| bySimilarGenes    | logical value (TRUE or FALSE) indicating to translate also through similar UniProt genes (100%, 90% or 50% of identity). Using TRUE value may take a long time, only advised if there is not a direct translation possible.                                                                                                |

**Value**

A character vector of the same length as upId. If parameter exhaustiveMapping is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

**Examples**

```
# Translate UniProt ID 'G9JVE6' to NCBI Gene
getUniProt2NCBIGene('G9JVE6')

# Translate UniProt IDs 'A0SNL9' and 'G9JVE6' to NCBI Gene
getUniProt2NCBIGene(c('A0SNL9', 'G9JVE6'))

# Get all possible translations of IDs 'A0SNL9', 'G9JVE6' with detailed origin
getUniProt2NCBIGene(c('A0SNL9', 'G9JVE6'), detailedMapping = TRUE,
exhaustiveMapping = TRUE)
```

---

```
getUniProt2NCBINucleotide
```

*Translates an UniProt database ID to NCBI Nucleotide database*

---

**Description**

Retrieves the ID translation of an specific instance of UniProt database ID to the NCBI Nucleotide database. Note that, due to the nature of NCBI Nucleotide database, a complete genome is often retrieved.

**Usage**

```
getUniProt2NCBINucleotide(upId, exhaustiveMapping = FALSE,
detailedMapping = FALSE, bySimilarGenes = TRUE)
```

**Arguments**

|                   |                                                                                                                                                                                                                                                                                                                                  |
|-------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| upId              | valid UniProt database ID or a vector of them                                                                                                                                                                                                                                                                                    |
| exhaustiveMapping | logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one.                                                                                                                                                                                                                |
| detailedMapping   | logical value (TRUE or FALSE) indicating to return a vector of NCBI Nucleotide IDs if FALSE, or a named vector if TRUE. This named vector details the origin of the translated IDs, dividing them in DT, if it is a Direct Translation, or 1.0, 0.9 or 0.5, if it has been retrieved through any of the identical gene clusters. |
| bySimilarGenes    | logical value (TRUE or FALSE) indicating to translate also through similar UniProt genes (100%, 90% or 50% of identity). Using TRUE value may take a long time, only advised if there is not a direct translation possible.                                                                                                      |

**Value**

A character vector of the same length as upId. If parameter exhaustiveMapping is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

**Examples**

```
# Translate UniProt ID 'G9JVE6' to NCBI Nucleotide
getUniProt2NCBINucleotide('G9JVE6')

# Translate UniProt IDs 'A0SNL9' and 'G9JVE6' to NCBI Nucleotide
getUniProt2NCBINucleotide(c('A0SNL9', 'G9JVE6'))

# Get all possible translations of IDs 'A0SNL9', 'G9JVE6' with detailed origin
getUniProt2NCBINucleotide(c('A0SNL9', 'G9JVE6'), detailedMapping = TRUE,
exhaustiveMapping = TRUE)
```

---

```
getUniProt2NCBIProtein
```

*Translates an UniProt database ID to NCBI Protein database*

---

**Description**

Retrieves the ID translation of an specific instance of UniProt database ID to the NCBI Protein database.

**Usage**

```
getUniProt2NCBIProtein(upId, exhaustiveMapping = FALSE, detailedMapping = FALSE,
bySimilarGenes = TRUE)
```

**Arguments**

- upId                    valid UniProt database ID or a vector of them
- exhaustiveMapping    logical value (TRUE or FALSE) indicating to retrieve all possible translations or only the first and fastest one.
- detailedMapping      logical value (TRUE or FALSE) indicating to return a vector of NCBI Protein IDs if FALSE, or a named vector if TRUE. This named vector details the origin of the translated IDs, dividing them in DT, if it is a Direct Translation, or 1.0, 0.9 or 0.5, if it has been retrieved through any of the identical gene clusters.
- bySimilarGenes       logical value (TRUE or FALSE) indicating to translate also through similar UniProt genes (100%, 90% or 50% of identity). Using TRUE value may take a long time, only advised if there is not a direct translation possible.

**Value**

A character vector of the same length as upId. If parameter exhaustiveMapping is TRUE, it will return a list instead of a character vector to avoid mixing the identifiers.

**Examples**

```
# Translate UniProt ID 'G9JVE6' to NCBI Protein
getUniProt2NCBIProtein('G9JVE6')

# Translate UniProt IDs 'A0SNL9' and 'G9JVE6' to NCBI Protein
getUniProt2NCBIProtein(c('A0SNL9', 'G9JVE6'))

# Get all possible translations of IDs 'A0SNL9', 'G9JVE6' with detailed origin
getUniProt2NCBIProtein(c('A0SNL9', 'G9JVE6'), detailedMapping = TRUE,
exhaustiveMapping = TRUE)
```

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getUniProtSimilarGenes

*Get similar UniProt genes by a given identity percentage*

---

**Description**

Retrieves the cluster of genes with a certain percentage of identity where a given UniProt gene belongs.

**Usage**

```
getUniProtSimilarGenes(upId, clusterIdentity = '1.0', clusterNames = FALSE)
```

**Arguments**

|                 |                                                                                                                                                                                 |
|-----------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| upId            | valid UniProt ID or a vector of them                                                                                                                                            |
| clusterIdentity | identity of the cluster to be retrieved. Default value 1.0 returns the 100% identical genes cluster, values 0.9 and 0.5, the 90% and 50% identical genes clusters, respectively |
| clusterNames    | logical value (TRUE or FALSE) indicating if cluster names are retrieved                                                                                                         |

**Value**

A list of the same length as upId. If parameter clusterNames is TRUE it will return a named list.

**Examples**

```
# Get UniProt IDs of 50% identical genes of UniProt gene 'G0L217'
getUniProtSimilarGenes('G0L217', clusterIdentity = '0.5')

# Get UniProt IDs of 50% identical genes of UniProt genes 'G0L217' and 'A0SNL9'
getUniProtSimilarGenes(c('G0L217', 'A0SNL9'), clusterIdentity = '0.5')
```

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|                   |                                                   |
|-------------------|---------------------------------------------------|
| getUniProtVersion | <i>Prints UniProt database version being used</i> |
|-------------------|---------------------------------------------------|

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**Description**

Prints the accessed UniProt version and its corresponding release date.

**Usage**

```
getUniProtVersion()
```

**Value**

UniProt database version.

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|                    |                                                               |
|--------------------|---------------------------------------------------------------|
| updateCARDDataBase | <i>Downloads the most up-to-date version of CARD database</i> |
|--------------------|---------------------------------------------------------------|

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**Description**

Cleans previous CARD database versions if there were, and downloads its latest version.

**Usage**

```
updateCARDDataBase()
```

**Value**

Downloaded CARD database.

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