# Description of the biomaRt package

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# Contents

1	Intr	oduction	2					
2	obje 2.1 2.2	Mart-class	2 2 2					
3	Functions							
	3.1	martConnect	2					
	3.2	martDisconnect	3					
	3.3	getGene	3					
	3.4	getGO	5					
	3.5	getOMIM	9					
	3.6	getFeature	10					
	3.7	getSequence	11					
	3.8	getSNP	11					
	3.9	0 1	12					
	3.10	getAffyArrays	13					
	3.11	getHomolog	13					

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# 1 Introduction

The BioConductor *biomaRt* package enables to directly query databases based on biomaRt such as Ensembl, a software system which produces and maintains automatic annotation on metazoan genomes. This way you can annotate the features on your array with the latest annotations starting from identifiers such as affy id's, locuslink, RefSeq and more. Annotation includes gene names, GO and OMIM annotation (depending on species).

# 2 objects

#### 2.1 Mart-class

An object of the Mart class represents a connection to a BioMart. And has the following slots:

- ensembl: stores the RMySQLConnection to Ensembl
- vega: stores the RMySQLConnection to VEGA
- sequence: stores the RMySQLConnection to sequence mart
- snp: stores the RMySQLConnection to snp mart
- arrayToSpecies: Stores mapping from affy arrays to species
- arrayToEnsembl: Stores mapping from affy arrays to Ensembl tables

#### 2.2 martTable-class

An object of the martTable class is the output of most biomaRt funtions and has the following slots:

- id: stores the id used for querying
- table: is a list of vectors storing the retrieved data

# 3 Functions

#### 3.1 martConnect

A first step in using the biomaRt package is to connect to a BioMart database. The function martConnect establishes a connection with one of the following BioMart databases: snp, ensembl and vega.

Examples:

#### > library(biomaRt)

Loading required package: Biobase
Welcome to Bioconductor
Vignettes contain introduct

Vignettes contain introductory material. To view,

simply type: openVignette()

For details on reading vignettes, see

the openVignette help page.

Loading required package: RMySQL Loading required package: DBI

> mart <- martConnect()</pre>

- Connected to ensembl\_mart\_30, ensembl\_mart\_30, ensembl\_mart\_30 and ensembl\_mart\_30 -

#### 3.2 martDisconnect

You can only hold a limited number of connections with different BioMarts. The function martDisconnect can be used to close a mart connection.

Examples:

> martDisconnect(mart)

[1] TRUE

## 3.3 getGene

The function getGene uses a query id to look up identification and chromosomal information of the corresponding gene. Depending on the selected output, this function returns a martTable or an object of class Gene. When no information about the identifier is found in Ensembl, an empty Gene object will be created. If however multiple genes match a certain identifier, then an object of class MultiGene will be return containing information of all matches. Currently the getGene function takes identifiers from Entrez-Gene, Affymetrix, RefSeq and embl. Besides the id argument, this function also has a species, array and type argument.

The id argument is either a vector of identifiers or a single identifier to be annotated.

The species argument should have the species from which the identifier comes as value. For the value of species, we use the full name of the species where separate words are separated by an underscore, e.g. 'gallusgallus'. A list of possible species to choose from can be obtained by executing the function getSpecies.

The array argument takes affy array identifiers as values. A list of possible identifiers supported by the package can be obtained by executing the function getAffyArrays.

The mart argument is a mart connection, which was obtained using the method martConnect

The type takes the values of 'entrezgene', 'refseq' and 'embl' to clarify which type of identifier is specified in the id argument.

The output can be changed using the output argument. One can choose between a mart-Table (default) and an output of Gene/Multi-Gene objects. Depending on the identifier, different additional arguments will have to be given, summarized below:

```
• Affy id's: id, array, mart
   • Entrez-Gene: id, type, species, mart
   • RefSeq: id, type, species, mart
   • embl: id, type, species, mart
   Examples:
> mart <- martConnect()
- Connected to ensembl_mart_30, ensembl_mart_30, ensembl_mart_30 and ensembl_mart_30 -
> getGene(id = "1939_at", array = "hgu95av2", mart = mart)
An object of class "martTable"
Slot "id":
[1] "1939_at"
Slot "table":
$symbol
[1] "TP53"
$description
[1] "Cellular tumor antigen p53 (Tumor suppressor p53) (Phosphoprotein p53) (Antigen NY-
$band
[1] "p13.1"
$chromosome
[1] "17"
$start
[1] 7512464
$end
[1] 7531642
$martID
[1] "ENSG00000141510"
```

```
> getGene(id = 672, type = "entrezgene", species = "homo_sapiens",
      mart = mart)
An object of class "martTable"
Slot "id":
[1] "672"
Slot "table":
$symbol
[1] "BRCA1"
$description
[1] "Breast cancer type 1 susceptibility protein (RING finger protein 53). [Source:Unipr
$band
[1] "q21.31"
$chromosome
[1] "17"
$start
[1] 38449844
$end
[1] 38530934
$martID
[1] "ENSG00000012048"
```

# 3.4 getGO

The function getGO uses a query id to look up GO annotation of the corresponding gene. It return an object of class GO. When no information about the identifier is found in Ensembl, an empty GO object will be created. Currently the getGO function takes identifiers from Entrez-Gene, Affymetrix, RefSeq and embl. Besides the id argument, this function also has a species, array and type argument.

The id argument is either a vector of identifiers or a single identifier to be annotated.

The species argument should have the species from which the identifier comes as value. A list of possible species to choose from can be obtained by executing the function getSpecies. The array argument takes affy array identifiers as values. A list of possible identifiers supported by the package can be obtained by executing the function getAffyArrays.

The mart argument is a mart connection, which was obtained using the method martConnect

A last argument of this function is the type argument which, takes the values of 'entrez-gene', 'refseq' and 'embl' to clarify which type of identifier is specified in the id argument.

Depending on the identifier, different additional arguments will have to be given, summarized below:

• Affy id's: id, array, mart

```
• Entrez-Gene: id, type, species, mart
   • RefSeq: id, type, species, mart
   • embl: id, type, species, mart
   Examples:
> getGO(id = "1939_at", array = "hgu95av2", mart = mart)
An object of class "martTable"
Slot "id":
 [1] "1939_at" "1939_at" "1939_at" "1939_at" "1939_at" "1939_at" "1939_at"
 [8] "1939_at" "1939_at" "1939_at" "1939_at" "1939_at" "1939_at" "1939_at"
[15] "1939_at" "1939_at" "1939_at" "1939_at" "1939_at" "1939_at" "1939_at"
[22] "1939_at" "1939_at" "1939_at" "1939_at" "1939_at" "1939_at"
Slot "table":
$GOID
 [1] "GD:0000739" "GD:0003700" "GD:0004518" "GD:0005507" "GD:0005515"
 [6] "GD:0005524" "GD:0008270" "GD:0000075" "GD:0006284" "GD:0006289"
[11] "GD:0006310" "GD:0006355" "GD:0006915" "GD:0007050" "GD:0007569"
[16] "GD:0008283" "GD:0008628" "GD:0008630" "GD:0008635" "GD:0030154"
[21] "GD:0030308" "GD:0045786" "GD:0046902" "GD:0051097" "GD:0005730"
[26] "GD:0005739" NA
$description
 [1] "DNA strand annealing activity"
 [2] "transcription factor activity"
 [3] "nuclease activity"
 [4] "copper ion binding"
 [5] "protein binding"
 [6] "ATP binding"
 [7] "zinc ion binding"
 [8] "cell cycle checkpoint"
 [9] "base-excision repair"
[10] "nucleotide-excision repair"
[11] "DNA recombination"
[12] "regulation of transcription, DNA-dependent"
[13] "apoptosis"
```

```
[14] "cell cycle arrest"
```

- [15] "cell aging"
- [16] "cell proliferation"
- [17] "induction of apoptosis by hormones"
- [18] "DNA damage response, signal transduction resulting in induction of apoptosis"
- [19] "caspase activation via cytochrome c"
- [20] "cell differentiation"
- [21] "negative regulation of cell growth"
- [22] "negative regulation of cell cycle"
- [23] "regulation of mitochondrial membrane permeability"
- [24] "negative regulation of helicase activity"
- [25] "nucleolus"
- [26] "mitochondrion"
- [27] NA

#### \$evidence

- [1] "IDA" "IDA" "TAS" "IDA" "IPI" "IDA" "TAS" "TAS" "TAS" "IMP" "TAS" "IDA"
- [13] "IDA" "TAS" "IMP" "TAS" "TAS" "TAS" "IDA" "TAS" "IMP" "IEA" "TAS" "TAS"
- [25] "IDA" "IDA" NA

#### \$martID

- [1] "ENSG00000141510" "ENSG00000141510" "ENSG000000141510" "ENSG00000141510"
- [5] "ENSG00000141510" "ENSG00000141510" "ENSG00000141510" "ENSG00000141510"
- [9] "ENSG00000141510" "ENSG00000141510" "ENSG00000141510" "ENSG00000141510"
- [13] "ENSG00000141510" "ENSG00000141510" "ENSG000000141510" "ENSG00000141510"
- [17] "ENSG00000141510" "ENSG00000141510" "ENSG00000141510" "ENSG00000141510"
- [21] "ENSG00000141510" "ENSG00000141510" "ENSG00000141510" "ENSG00000141510"
- [25] "ENSG00000141510" "ENSG00000141510" "ENSG00000141510"
- > getGO(id = 672, type = "entrezgene", species = "homo\_sapiens",
  + mart = mart)

An object of class "martTable"

Slot "id":

- [1] "672" "672" "672" "672" "672" "672" "672" "672" "672" "672" "672" "672" "672"
- [13] "672" "672" "672" "672" "672" "672" "672" "672" "672" "672" "672" "672" "672"
- [25] "672" "672"

#### Slot "table":

#### \$GOID

- [1] "G0:0003684" "G0:0003713" "G0:0004842" "G0:0005515" "G0:0008270"
- [6] "GD:0015631" "GD:0016563" "GD:0006357" "GD:0006359" "GD:0006978"
- [11] "G0:0016567" "G0:0042127" "G0:0042981" "G0:0045739" "G0:0045786"
- [16] "GD:0046600" "GD:0000151" "GD:0005615" "GD:0005634" "GD:0005667"
- [21] "G0:0008274" "G0:0005554" "G0:0000075" "G0:0008372" "G0:0005622"

#### [26] "GD:0008270"

#### \$description

- [1] "damaged DNA binding"
- [2] "transcription coactivator activity"
- [3] "ubiquitin-protein ligase activity"
- [4] "protein binding"
- [5] "zinc ion binding"
- [6] "tubulin binding"
- [7] "transcriptional activator activity"
- [8] "regulation of transcription from Pol II promoter"
- [9] "regulation of transcription from Pol III promoter"
- [10] "DNA damage response, signal transduction by p53 class mediator resulting in transc
- [11] "protein ubiquitination"
- [12] "regulation of cell proliferation"
- [13] "regulation of apoptosis"
- [14] "positive regulation of DNA repair"
- [15] "negative regulation of cell cycle"
- [16] "negative regulation of centriole replication"
- [17] "ubiquitin ligase complex"
- [18] "extracellular space"
- [19] "nucleus"
- [20] "transcription factor complex"
- [21] "gamma-tubulin ring complex"
- [22] "molecular\_function unknown"
- [23] "cell cycle checkpoint"
- [24] "cellular\_component unknown"
- [25] "intracellular"
- [26] "zinc ion binding"

#### \$evidence

- [1] "NR" "TAS" "IEA" "IPI" "TAS" "NAS" "TAS" "TAS" "TAS" "TAS" "IEA" "TAS"
- [13] "TAS" "NAS" "IEA" "NAS" "IEA" "TAS" "TAS" "TAS" "NAS" "ND" "NAS" "ND"
- [25] "IEA" "IEA"

#### \$martID

- [1] "ENSG00000012048" "ENSG00000012048" "ENSG00000012048" "ENSG00000012048"
- [5] "ENSG00000012048" "ENSG00000012048" "ENSG00000012048" "ENSG00000012048"
- [9] "ENSG00000012048" "ENSG00000012048" "ENSG00000012048" "ENSG00000012048"
- [13] "ENSG00000012048" "ENSG00000012048" "ENSG00000012048" "ENSG00000012048"
- [17] "ENSG00000012048" "ENSG00000012048" "ENSG00000012048" "ENSG00000012048"
- [21] "ENSG0000012048" "ENSG00000012048" "ENSG00000012048" "ENSG00000012048"
- [25] "ENSG00000012048" "ENSG00000012048"

# 3.5 getOMIM

The function getOMIM uses a query id to look up OMIM annotation of the corresponding gene. It return an object of class OMIM. When no information about the identifier is found in Ensembl, an empty OMIM object will be created. Currently the getOMIM function takes identifiers from entrezgene, affy, RefSeq and embl. Besides the id argument, this function also has an array ,type and mart argument.

The id argument is either a vector of identifiers or a single identifier to be annotated.

The array argument takes affy array identifiers as values. A list of possible identifiers supported by the package can be obtained by executing the function getAffyArrays.

The type argument takes the values of 'entrezgene', 'refseq' and 'embl' to clarify which type of identifier is specified in the id argument. If the argumant array is used then biomaRt knows the identifiers given corresponf to affy id's. The mart argument is a mart connection, which was obtained using the method martConnect

Depending on the identifier, different additional arguments will have to be given, summarized below:

```
• Affy id's: id, array, mart
```

• Entrez-Gene: id, type, mart

• RefSeq: id, type, mart

• embl: id, type, mart

Examples:

```
> getOMIM(id = "1939_at", array = "hgu95av2", mart = mart)
An object of class "martTable"
Slot "id":
[1] "1939_at" "1939_at"
Slot "table":
$OMIMID
[1] 191170 191170
$disease
[1] "Colorectal cancer, 114500 (3)" "Li-Fraumeni syndrome (3)"
$martID
[1] "ENSG000000141510" "ENSG000000141510"
> getOMIM(id = 672, type = "entrezgene", mart = mart)
```

```
An object of class "martTable"
Slot "id":
[1] "672" "672"

Slot "table":
$0MIMID
[1] 113705 113705

$disease
[1] "Breast cancer-1 (3)" "Ovarian cancer (3)"
$martID
[1] "ENSG00000012048" "ENSG00000012048"
```

# 3.6 getFeature

The function getFeature looks up affy identifiers on a given affy array which correspond to a given symbol. As output this function returns a martTable. Currently the getFeature function takes identifiers from affy only. Besides the symbol argument, this function also has array and mart argument.

The mart argument is a mart connection, which was obtained using the method martConnect

A last argument of this function is the type argument which, takes the values of 'affy', 'locuslink', 'refseq' and 'embl' to clarify which type of identifier is specified in the id argument.

Examples:

```
> getFeature(symbol = "P53", array = "hgu95av2", mart = mart)
An object of class "martTable"
Slot "id":
[1] "36079_at"
                "1939_at"
                            "1974_s_at" "31618_at" "1711_at"
                                                                 "33749_at"
[7] "1860_at"
                "34822_at"
Slot "table":
$symbol
[1] "TP53I3" "TP53"
                        "TP53"
                                  "TP53"
                                            "TP53BP1" "TP53AP1" "TP53BP2"
[8] "TP53BP2"
$description
[1] "tumor protein p53 inducible protein 3 [Source:RefSeq_peptide;Acc:NP_671713]"
[2] "Cellular tumor antigen p53 (Tumor suppressor p53) (Phosphoprotein p53) (Antigen NY-
```

[3] "Cellular tumor antigen p53 (Tumor suppressor p53) (Phosphoprotein p53) (Antigen NY-[4] "Cellular tumor antigen p53 (Tumor suppressor p53) (Phosphoprotein p53) (Antigen NY-

- [5] "Tumor suppressor p53-binding protein 1 (p53-binding protein 1) (53BP1). [Source:Uni
- [6] "TP53 activated protein 1 [Source:RefSeq\_peptide;Acc:NP\_009164]"
- [7] "Apoptosis stimulating of p53 protein 2 (Tumor suppressor p53-binding protein 2) (p5
- [8] "Apoptosis stimulating of p53 protein 2 (Tumor suppressor p53-binding protein 2) (p5

#### \$martID

- [1] "ENSG00000115129" "ENSG00000141510" "ENSG00000141510" "ENSG00000141510"
- [5] "ENSG00000067369" "ENSG00000182165" "ENSG00000143514" "ENSG00000143514"

# 3.7 getSequence

The function getSequence retrieves the sequence given it's chromosome, start and end position. As output this function returns a martTable. The mart argument is a mart connection, which was obtained using the method martConnect and should in this case be the sequence mart.

Examples:

```
> getSequence(species = "gallus_gallus", chromosome = 1, start = 400,
+ end = 500, mart = mart)

An object of class "martTable"
Slot "id":
[1] "1_400_500"

Slot "table":
$chromosome
[1] 1

$start
[1] 400

$end
[1] 500
```

# \$sequence

[1] "GTGACATTTCCAGCATTCAGTGTCAAAGCCTAGCTTCATTTTTGAATGTATTGAGGGGCAGATGTCCATCTCATGAATCAT

# 3.8 getSNP

The function getSNP retrieves all SNP's between a given a start and end position on a gives chromosome. As output this function returns a martTable. The mart argument is a mart connection, which was obtained using the method martConnect and should in this case be

the snp mart.

Examples:

```
> getSNP(chromosome = 8, start = 148350, end = 148612, species = "homo_sapiens",
+ mart = mart)
```

An object of class "martTable"

Slot "id":

- [1] "TSC1421398" "TSC1421399" "TSC1421400" NA "TSC1421401"
- [6] NA "TSC1421402" "TSC1737607" NA NA

Slot "table":

\$snpStart

[1] 148394 148411 148462 148471 148499 148525 148533 148535 148539 148601

\$allele

[1] "C/A" "A/G" "C/T" "T/G" "G/A" "G/A" "G/A" "C/T" "C/T" "G/A"

\$coding

[1] NA NA NA NA NA NA NA NA NA

\$intronic

[1] NA NA NA NA NA NA NA NA NA

\$syn

[1] NA NA NA NA NA NA NA NA NA

\$utr5

[1] NA NA NA NA NA NA NA NA NA

\$utr3

[1] 1 1 1 1 1 1 1 1 1 1

# 3.9 getSpecies

The function getSpecies looks up which species are present in the BioMart. This function currently works only for ensembl.

Examples:

> getSpecies(mart = mart)

```
[1] "anopheles_gambiae" "fugu_rubripes"
[3] "gallus_gallus" "drosophila_melanogaster"
[5] "xenopus_tropicalis" "caenorhabditis_elegans"
[7] "danio_rerio" "homo_sapiens"
[9] "pan_troglodytes" "mus_musculus"
[11] "apis_mellifera" "canis_familiaris"
[13] "tetraodon_nigroviridis" "rattus_norvegicus"
[15] "saccharomyces_cerevisiae"
```

# 3.10 getAffyArrays

The function getAffyArrays retrieves the Affymetrix array identifiers which are present in ensembl and which can be queried using the biomaRt package.

Examples:

# > getAffyArrays(mart = mart)

	V1	V2
1	hgu133plus2	homo_sapiens
2	hgu133a2	homo_sapiens
3	hgu133a	homo_sapiens
4	hgu133b	homo_sapiens
5	hgu95av2	homo_sapiens
6	hgu95b	homo_sapiens
7	hgu95c	homo_sapiens
8	hgu95d	homo_sapiens
9	hgu95e	homo_sapiens
10	mgu74av2	mus_musculus
11	mgu74bv2	mus_musculus
12	mgu74cv2	mus_musculus
13	$mouse430_2$	mus_musculus
14	${\tt mouse430a\_2}$	mus_musculus
15	mu11ksuba	mus_musculus
16	mu11ksubb	mus_musculus
17	rat230_2	rattus_norvegicus
18	rgu34a	rattus_norvegicus
19	rgu34b	rattus_norvegicus
20	rgu34c	rattus_norvegicus

# 3.11 getHomolog

This function retrieves homologs of genes of one species in another species Example:

```
> getHomolog(id = 1:20, fromspecies = "homo_sapiens", tospecies = "mus_musculus",
+ fromtype = "entrezgene", totype = "refseq", mart = mart)
```

An object of class "martTable"

NA

Slot "id":

- [1] "2" "2" "2" "2" "3" "4" "5" "6" "7" "8" "9" "10" "11" "12" "12"
- [16] "12" "12" "12" "13" "14" "15" "16" "17" "18" "19" "20"

## Slot "table":

## \$MappedID

[25] NA

[1]	"NM_175628"	"NM_008646"	"NM_008645"	"NM_007376"	NA	NA
[7]	NA	NA	NA	NA	NA	NA
[13]	NA	"NM_009252"	"NM_173024"	"NM_008458"	"NM_011458"	"NM_009253"
[19]	NA	NA	"NM_009591"	"NM_146217"	NA	"NM_172961"