

# Package ‘InterMineR’

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**Title** R Interface with InterMine-Powered Databases

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**Description** Databases based on the InterMine platform such as FlyMine, modMine (modENCODE), RatMine, YeastMine, HumanMine and TargetMine are integrated databases of genomic, expression and protein data for various organisms. Integrating data makes it possible to run sophisticated data mining queries that span domains of biological knowledge. This R package provides interfaces with these databases through web services. It makes most from the correspondence of the data frame object in R and the table object in databases, while hiding the details of data exchange through XML or JSON.

**Depends** R (>= 3.4.1)

**License** LGPL

**LazyData** true

**VignetteBuilder** knitr

**Imports** Biostrings, RCurl, XML, xml2, RJSONIO, sqldf, igraph, httr, S4Vectors, IRanges, GenomicRanges, SummarizedExperiment, methods

**Suggests** BiocStyle, Gviz, knitr, rmarkdown, GeneAnswers, GO.db, org.Hs.eg.db

**BugReports** <https://github.com/intermine/intermineR/issues>

**biocViews** GeneExpression, SNP, GeneSetEnrichment, DifferentialExpression, GeneRegulation, GenomeAnnotation, GenomeWideAssociation, FunctionalPrediction, AlternativeSplicing, ComparativeGenomics, FunctionalGenomics, Proteomics, SystemsBiology, Microarray, MultipleComparison, Pathways, GO, KEGG, Reactome, Visualization

**NeedsCompilation** no

## R topics documented:

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## Description

InterMine-powered databases such as FlyMine, modENCODE, RatMine, YeastMine, HumanMine and TargetMine are integrated databases of genomic, expression and protein data for various organisms. Integrating data makes it possible to run sophisticated data mining queries that span domains of biological knowledge. This R package provides interface with these databases through webservices. It makes most from the correspondence of the data frame object in R and the table object in databases while hiding the details of data exchange through XML or JSON.

## Details

Package:	InterMineR
Type:	Package
Version:	0.99.4
Date:	2016-01-05
License:	LGPL
Depends:	Biostrings, RCurl, XML, RJSONIO, sqldf, igraph
Suggests:	Gviz

## Author(s)

InterMine Team

Maintainer: InterMine Team <info@intermine.org>

## References

<http://intermine.readthedocs.org/en/latest/web-services/>

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convertToGeneAnswers    *Convert InterMineR enrichment analysis results to GeneAnswers object*

---

## Description

convertToGeneAnswers constitutes a wrapper function for converting the results of `doEnrichment` function to a `GeneAnswers-class` object. This way the user can utilize the functions of `GeneAnswers` package to visualize the results of InterMineR enrichment analysis.

## Usage

```
convertToGeneAnswers(  
  enrichmentResult,  
  geneInput,  
  geneInputType,  
  geneExprProfile,  
  annLib,  
  categoryType,  
  enrichCategoryChildName  
)
```

## Arguments

enrichmentResult                a list containing the results of `doEnrichment` function.  
geneInput                      a data.frame containing the gene identifiers used for the InterMineR enrichment analysis, and possible values associated with them.  
geneInputType                  a character string specifying the InterMineR gene identifier type of the values assigned to geneInput argument. InterMineR identifiers for each Mine can be retrieved with `getModel` function.  
geneExprProfile                a data.frame containing gene expression information (optional)  
annLib                         name of given annotation library file or user provided annotation list.  
categoryType                  name of given annotation category or NULL for user provided annotation list.  
enrichCategoryChildName        a character string specifying the InterMineR annotation category identifier. This argument must be assigned manually if:  
                                  1) 'enrichIdentifier' column is missing from the widgets of the Mine used by `doEnrichment`,  
                                  2) the value of 'enrichIdentifier' for the enrichment widget used by `doEnrichment` is NA.  
                                  InterMineR identifiers for each Mine can be retrieved with `getModel` function.

**Value**

a `GeneAnswers-class` object.

**Author(s)**

InterMine Team

**References**

<https://bioconductor.org/packages/release/bioc/html/GeneAnswers.html>

**See Also**

`doEnrichment`, `GeneAnswers-class`, `getWidgets`, `getModel`

**Examples**

```
# load human genes which are associated with Diabetes (retrieved from HumanMine)
data("PL_DiabetesGenes")

# get Gene.primaryIdentifiers (ENTREZ Gene identifier)
hsa_gene_entrez = as.character(PL_DiabetesGenes$Gene.primaryIdentifier)

# perform enrichment analysis with InterMineR
hsa_enrichResult = doEnrichment(
  im = initInterMine(listMines()["HumanMine"]),
  genelist = "PL_DiabetesGenes",
  widget = "go_enrichment_for_gene",
  correction = "Benjamini Hochberg"
)

# convert InterMineR enrichment analysis results to a GeneAnswers object
# Do not run unless you have installed and loaded GeneAnswers package!

# load GeneAnswers
library(GeneAnswers)

hsa_geneanswers = convertToGeneAnswers(
  enrichmentResult = hsa_enrichResult,
  geneInput = data.frame(GeneID = as.character(hsa_gene_entrez),
    stringsAsFactors = FALSE),
  geneInputType = "Gene.primaryIdentifier",
  annLib = 'org.Hs.eg.db',
  categoryType = "GO"
  # enrichCategoryChildName = "Gene.goAnnotation.ontologyTerm.parents.identifier"
)
```

## Description

convertToGRanges constitutes a wrapper function for converting genomic locations and their respective annotations to objects of the [GRanges](#) class.

## Usage

```
convertToGRanges(  
  dataset,  
  seqnames,  
  start,  
  end,  
  names,  
  strand,  
  columnsAsMetadata = NULL,  
  listAsMetadata = NULL,  
  seqnamesInterMineChromosome = TRUE  
)
```

## Arguments

dataset	a data.frame containing all the genomic information retrieved by InterMineR. Its columns are used to create the <a href="#">GRanges</a> object.
seqnames	the name of a column from dataset or a character vector of length equal to the number of rows of the dataset. Defines the values assigned as seqnames to the <a href="#">GRanges</a> object.
start	the name of a column from dataset or a vector of length equal to the number of rows of the dataset. Defines the genomic coordinates assigned as start to the ranges argument of the <a href="#">GRanges</a> object.
end	the name of a column from dataset or a vector of length equal to the number of rows of the dataset. Defines the genomic coordinates assigned as end to the ranges argument of the <a href="#">GRanges</a> object.
names	the name of a column from dataset or a character vector of length equal to the number of rows of the dataset. Defines the values assigned as names to the ranges argument of the <a href="#">GRanges</a> object.
strand	the name of a column from dataset or a character vector of length equal to the number of rows of the dataset. Defines the values assigned as strand information to the <a href="#">GRanges</a> object.
columnsAsMetadata	a character vector containing the names of the dataset columns that are passed as metadata in the <a href="#">GRanges</a> object.
listAsMetadata	a list of vectors, each of which has length equal to the number of rows of the dataset. The values of the list are passed as metadata to the <a href="#">GRanges</a> object.
seqnamesInterMineChromosome	a logical value indicating whether the values passed as seqnames are InterMineR chromosome.primaryIdentifiers (e.g. "2R", "3R", "X") or not.

## Details

The InterMineR package provides a flexible interface to the InterMine web services, which allow for rapid retrieval of various genomic information.

`convertToGRanges` function is designed to facilitate the conversion of genomic locations and their respective annotations, in the format by which they are retrieved using InterMineR, to an object of the [GRanges](#) class.

### Value

An object of the [GRanges](#) class containing genomic locations and annotations retrieved by InterMineR queries.

### Author(s)

InterMine Team

### See Also

[GRanges](#), [convertToRangedSummarizedExperiment](#)

### Examples

```
# get FlyMine
im.fly = initInterMine(listMines()["FlyMine"])

# modify template query for Transcription Factor (TF) Binding sites
qTF_Binding = getTemplateQuery(im.fly, "ChromLocation_TFBindingSiteLocationGeneFactor")

qTF_Binding$where[[4]]$value = "1000000"
qTF_Binding$where[[5]]$value = "20000000"

rTF_Binding = runQuery(im.fly, qTF_Binding)

# assign random values for strand of the genomic location retrieved, in InterMine format
rTF_Binding$gene.strand = sample(c("1", "-1", ""), nrow(rTF_Binding), replace = TRUE)

# convert to GRanges object
test = convertToGRanges(
  dataset = rTF_Binding,
  seqnames = rTF_Binding$TFBindingSite.chromosome.primaryIdentifier,
  names = rTF_Binding$TFBindingSite.factor.name,
  start = rTF_Binding$TFBindingSite.chromosomeLocation.start,
  end = rTF_Binding$TFBindingSite.chromosomeLocation.end,
  strand = "gene.strand",
  columnsAsMetadata = c(
    "TFBindingSite.gene.regulatoryRegions.dataSets.dataSource.name",
    "TFBindingSite.factor.primaryIdentifier"),
  listAsMetadata = list(
    c(factor.primaryIdentifier = rTF_Binding$TFBindingSite.factor.primaryIdentifier)
  )
)

# check results
test
```

---

**convertToRangedSummarizedExperiment**

*Convert experimental results retrieved by InterMineR queries to an object of the [RangedSummarizedExperiment](#) class*

---

**Description**

convertToRangedSummarizedExperiment constitutes a wrapper function for converting genomic information and experimental data from InterMine to an object of the [RangedSummarizedExperiment](#) class.

**Usage**

```
convertToRangedSummarizedExperiment(  
  im,  
  dataset,  
  SampleColumn,  
  GeneColumn,  
  ValueColumn,  
  OrganismValue,  
  colsForSampleMetadata,  
  exonsForRowRanges = FALSE  
)
```

**Arguments**

im	a list containing the base URL and API token.
dataset	a data.frame retrieved with InterMineR queries that contains experimental data from high-throughput assays.
SampleColumn	a character string or an integer indicating which column of the dataset contains the samples.
GeneColumn	a character string or an integer indicating which column of the dataset contains the genes.
ValueColumn	a character string or an integer indicating which column of the dataset contains the experimental data.
OrganismValue	a character string with the name of the organism from which the genomic information are retrieved.
colsForSampleMetadata	an integer vector indicating the columns of the dataset which will be assigned as sample metadata in the colData argument of the <a href="#">RangedSummarizedExperiment</a> .
exonsForRowRanges	a logical value indicating whether the rowRanges argument of the <a href="#">RangedSummarizedExperiment</a> should be assigned with a GRangeslist containing only annotations about the genes (default) or annotations about all exons of each gene.

**Details**

The InterMineR package provides a flexible interface to InterMine web services, which allow for rapid retrieval of various genomic information.

`convertToRangedSummarizedExperiment` function facilitates the conversion of genomic information and experimental data from high-throughput assays, which are retrieved by using InterMineR queries, to an object of the [RangedSummarizedExperiment](#) class.

It is noteworthy that the `reshape` function is used to convert the experimental data from the InterMineR format (long format) to the matrix (wide format) assigned to the `assays` argument of the [SummarizedExperiment](#).

### Value

an object of the [RangedSummarizedExperiment](#) class containing genomic information and experimental data of high-throughput assays, which are retrieved with the InterMineR queries system.

### Author(s)

InterMine Team

### References

[SummarizedExperiment](#) for Coordinating Experimental Assays, Samples, and Regions of Interest

### See Also

[RangedSummarizedExperiment](#), `link{convertToGRanges}`

### Examples

```
# 10 Drosophila melanogaster genes of interest
Drosophila.genes = c("BEAF-32", "Antp", "bcd", "caup", "tup", "E2f2", "dsx", "so", "toy", "Lim1")

# retrieve microarray time course experimental data for Drosophila.genes
# get FlyMine instance
im.fly = initInterMine(listMines()["FlyMine"])

# get FlyMine microarray time course template query
queryForData = getTemplateQuery(im.fly, "Gene_TimeCourseExpression")

test.data = list(NULL)
ind.null = c()

for(i in seq(length(Drosophila.genes))){
  # set value in gene constraint
  queryForData$where[[3]]$value = as.character(Drosophila.genes[i])

  # run query and save the results of genes that exist in the Microarray time course dataset
  r = runQuery(im.fly, queryForData)

  ind.null = c(ind.null, is.null(r))

  test.data[[i]] = r
}

# remove genes for which no experimental data were retrieved
test.data = test.data[which(!ind.null)]

# rbind data together
```

```
test.data = do.call(rbind, test.data)

# using integer index for columns in arguments
test1 = convertToRangedSummarizedExperiment(
  im = im.fly,
  dataset = test.data,
  SampleColumn = 2,
  GeneColumn = 1,
  ValueColumn = 3,
  OrganismValue = "Drosophila melanogaster",
  colsForSampleMetadata = 4:7,
  exonsForRowRanges = TRUE
)

test1

# using directly column names in arguments
test2 = convertToRangedSummarizedExperiment(
  im = im.fly,
  dataset = test.data,
  SampleColumn = "Gene.microArrayResults.assays.sample2",
  GeneColumn = "Gene.symbol",
  ValueColumn = "Gene.microArrayResults.value",
  OrganismValue = "Drosophila melanogaster",
  colsForSampleMetadata = 4:7,
  exonsForRowRanges = TRUE
)

test2
```

---

**doEnrichment***Perform enrichment analysis*

---

**Description**

Retrieve enrichment analysis results from InterMine platform. Enrichment widgets provide a statistical summary of what makes a list distinct from the background population over a certain domain. They return a list of members of the domain ranked by p-value (low to high).

**Usage**

```
doEnrichment(
  im,
  genelist = NULL,
  ids = NULL,
  widget = NULL,
  population = NULL,
  maxp = 0.05,
  correction = "Benjamini Hochberg",
  filter = NULL,
  organism = NULL
)
```

## Arguments

<code>im</code>	a list containing the base URL and API token.
<code>genelist</code>	The name of the list to investigate, optional unless <code>ids</code> (identifiers) is NULL.
<code>ids</code>	a character vector containing the identifiers (list of genes, proteins, SNPs, etc.) for the enrichment analysis, optional unless <code>genelist</code> is NULL. Alternatively, a comma-separated character string of InterMine object IDs can be assigned directly to this argument. Use <code>getGeneIds</code> function to retrieve unique Gene.id values for a list of gene identifiers.
<code>widget</code>	The name of the enrichment widget to display. Use <code>getWidgets</code> function to retrieve available enrichment type widgets for the respective Mine.
<code>population</code>	The name of the list to use as the background population.
<code>maxp</code>	The maximum p-value of results to display. The range is 0.0 - 1.0
<code>correction</code>	The error correction algorithm to use. Possible options are "Benjamini Hochberg" (default), "Holm-Bonferroni", "Bonferroni" or "None".
<code>filter</code>	An optional filter that some widgets accept. Use <code>getWidgets</code> function to retrieve available filters of the respective enrichment widget.
<code>organism</code>	a character string defining the name of the organism (e.g. "Homo sapiens"). This argument is optional and can be used to limit the analysis to the list of identifiers for a specific organism. Can be useful for InterMine instances that contain data from multiple organisms.

## Details

The public gene lists for each Mine are available at the websites obtained by the following command: `paste0(listMines()["Mine_Name"], "/bag.do?subtab=view")`

Each type of enrichment widget can be applied to a gene list with specific annotation. To apply the enrichment analysis to the appropriate list of genes, use the targets column of the respective enrichment widgets as they are retrieved by `getWidgets` function.

## Value

`doEnrichment` function returns a list containing the following values:

<code>data</code>	A data.frame containing the results of the enrichment analysis performed in InterMine platform. The statistically significant results are ordered by increasing p-values
<code>populationCount</code>	a numeric value indicating the size of the reference population
<code>notAnalyzed</code>	a numeric value indicating the number of input features that were not included in the enrichment analysis
<code>im</code>	the list containing the base URL and API token
<code>parameters</code>	a character vector containing all the parameters used for the enrichment analysis, except <code>im</code>

## Author(s)

InterMine Team

**See Also**

[getWidgets](#), [getGeneIds](#)

**Examples**

```
# FlyMine
enrichResults.FlyMine <- doEnrichment(
  im = initInterMine(mine = listMines()["FlyMine"]),
  genelist = 'PL_FlyTF_site_specific_TFs',
  widget = "go_enrichment_for_gene"
)

# use ids instead of genelist
data("PL_FlyTF_site_specific_TFs")

enrichResults.FlyMine <- doEnrichment(
  im = initInterMine(mine = listMines()["FlyMine"]),
  #genelist = 'PL_FlyTF_site_specific_TFs',
  ids = PL_FlyTF_site_specific_TFs$Gene.primaryIdentifier,
  widget = "go_enrichment_for_gene"
)

# HumanMine
enrichResults.HumanMine <- doEnrichment(
  im = initInterMine(mine = listMines()["HumanMine"]),
  genelist = "PL_DiabetesGenes",
  widget = "go_enrichment_for_gene"
)

# use ids instead of genelist
data("PL_DiabetesGenes")

enrichResults.HumanMine <- doEnrichment(
  im = initInterMine(mine = listMines()["HumanMine"]),
  #genelist = "PL_DiabetesGenes",
  ids = PL_DiabetesGenes$Gene.primaryIdentifier,
  widget = "go_enrichment_for_gene"
)
```

**getDatasets**

*Retrieve information about the available datasets of a Mine instance.*

**Description**

This function retrieves information about the available datasets of a mine instance. It is necessary to specify the type of feature (e.g. Gene, TFBindingSite) for this mine, and a value of this feature OR of one of its child\_name values.

**Usage**

```
getDatasets(
  im,
  type,
```

```

    child_name,
    value,
    op
)

```

## Arguments

<code>im</code>	a list containing the base URL and API token.
<code>type</code>	a character string defining the type of feature for which the information about the available datasets will be retrieved. Use the <a href="#">listDatasets</a> function to retrieve types with available datasets on each mine.
<code>child_name</code>	a character string defining the <code>child_name</code> values of <code>type</code> , which will be used to retrieve the information about the available datasets. Use <a href="#">getModel</a> function to retrieve the available <code>child_name</code> values for each type.
<code>value</code>	a character string defining the value of the <code>child_name</code> .
<code>op</code>	a character string defining the <code>op</code> argument.

## Details

The `getDatasets` function uses an InterMineR query to retrieve the information of the available datasets for:

1. a specific mine (e.g. FlyMine)
2. a specific type of feature (e.g. Gene, TFBindingSite) for this mine, and
3. a specific value of this feature OR of one of its `child_name` values.

For all genes of *Drosophila melanogaster* (`Gene.organism = Drosophila melanogaster`) OR for a specific gene (`Gene LOOKUP PPARG`)

## Value

a data.frame containing information about about the available datasets for a specific type of feature (e.g. Gene) on a specific mine instance (HumanMine, FlyMine, etc.).

## Author(s)

InterMine Team

## See Also

[listDatasets](#), [getModel](#)

## Examples

```

# Define mines
im.fly = initInterMine(listMines()["FlyMine"])
im.human = initInterMine(listMines()["HumanMine"])

# get information about all available datasets of Drosophila melanogaster genes
dme_gene_datasets = getDatasets(
    im = im.fly,
    type = "Gene",
    child_name = "organism.name",
    value = "Drosophila melanogaster",

```

```

op = "="
)

# get available datasets for PPARG gene from FlyMine
pparg_related = getDatasets(
  im = im.fly,
  type = "Gene",
  value = "PPARG",
  op = "LOOKUP"
)

# get available datasets for PPARG gene from HumanMine
pparg_related.2 = getDatasets(
  im = im.human,
  type = "Gene",
  value = "PPARG",
  op = "LOOKUP"
)

```

**getGeneIds***Get Gene.id values for a list of gene identifiers***Description**

This function takes as input a list of gene identifiers and retrieves their unique Gene.id values for a specific organism.

**Usage**

```
getGeneIds(im, genes, organism)
```

**Arguments**

<code>im</code>	a list containing the base URL and API token.
<code>genes</code>	a character vector containing the gene identifiers.
<code>organism</code>	a character string for the name of the Organism.

**Details**

This function is designed to work in conjunction with the [doEnrichment](#) function.

**Value**

`getGeneIds` returns a list containing a data.frame of unique gene identifiers as well as the Gene.id values as a comma-separated string, ready to be used as input for the `ids` argument of the `doEnrichment` function.

Furthermore, in case of genes that return multiple identifiers or no identifiers, these are also reported separately in a data.frame and a character vector respectively.

**unique.results**

data.frame with unique gene identifiers

```

doEnrichment.string
    comma-separated character string of Gene.id values
multiple.results
    data.frame with genes returning multiple identifiers
genes.with.no.results
    character vector with genes returning no identifiers

```

**Note**

doEnrichment.string can be passed as input in the [doEnrichment](#) function.

**Author(s)**

InterMine Team

**See Also**

[getWidgets](#), [doEnrichment](#)

**Examples**

```

# HumanMine and Homo sapiens genes
hsa.genes <- c("ABCC8", "ACE", "AKT2", "APPL1", "AQP2", "AVP", "AVPR2", "BLK", "CAPN10", "CCR5")

list.hsa <- getGeneIds(
    im = initInterMine(listMines()["HumanMine"]),
    organism = "Homo sapiens",
    genes = hsa.genes # must be a character vector!!!
)

# FlyMine and Drosophila melanogaster genes
fly.genes <- c("fkh", "vvl", "BEAF-32", "Antp", "Scr", "abd-A", "bcd", "Ubx", "zen", "ara")

list.fly <- getGeneIds(
    im = initInterMine(listMines()["FlyMine"]),
    organism = "Drosophila melanogaster",
    genes = fly.genes # must be a character vector!!!
)

```

**Description**

Returns a representation of the data model for the mine. This describes the kind of data held, and the properties that data can have. This information can be used to build queries against that data, and to interpret the information received.

**Usage**

```
getModel(im, timeout = 3)
```

**Arguments**

- |         |   |
|---------|---|
| im      | a list containing the base URL and API token.   |
| timeout | an integer, representing the number of seconds to wait for the webservice to respond. |

**Details**

The details of the data model for the various mines are available at the websites obtained by running the following command: `paste(listMines()$URL, "/tree.do", sep="")`

**Value**

a multi-level list, representing the data model for the mine. The first-level is a list of the InterMine objects (e.g., Gene, Exon). Each second-level list, corresponding to an InterMine object, contains three data.frame objects: attributes, references and collections. Each attribute is a property of the InterMine object. Each reference or collection is itself an InterMine object, acting as a member object of the InterMine object.

**Author(s)**

InterMine Team

**Examples**

```
# Retrieve data model for FlyMine
im.fly = initInterMine(listMines()["FlyMine"])

model = getModel(im.fly)
```

---

**getRelease**

*Get the current release information of InterMine*

---

**Description**

Returns a string describing the release of the mine.

**Usage**

```
getRelease(im, timeout = 3)
```

**Arguments**

- |         |   |
|---------|---|
| im      | a list containing the base URL and API token.   |
| timeout | an integer, representing the number of seconds to wait for the webservice to respond. |

**Value**

a string, describing the release of the mine.

**Author(s)**

InterMine Team

**Examples**

```
im <- initInterMine(listMines()["HumanMine"])

getRelease(im)
```

*getTemplateQuery*

*Get the query contained in a template*

**Description**

Get a template query for a mine. A template contain a saved query with a view and constraint. The user can modify this query to obtain the desired result. The view is a vector containing the output columns of the query. The constraint is a matrix containing the following columns: path (the path of the constraint), op (the constraint operator, one of '=', '!=', 'LOOKUP', 'ONE OF', 'NONE OF', '>', '<', '>=', '<='), value (the constraint value), code (the name of the constraint), extraValue (optional, required for LOOKUP constraints).

**Usage**

```
getTemplateQuery(im, name, timeout=3)
```

**Arguments**

im	a list, containing the base URL and API token.
name	a string, representing the name of the pre-defined template.
timeout	an integer, representing the number of seconds to wait for the webservice to respond.

**Value**

a list, representing the query contained in the pre-defined template. The list should contain at least two elements, view and constrain.

**Author(s)**

InterMine Team

**Examples**

```
# Get template queries from HumanMine
im <- initInterMine(listMines()["HumanMine"])

queryGeneIden <- getTemplateQuery(im, "Gene_Identifiers")
```

---

getTemplates	<i>Get the information (name and title) of the templates pre-defined in InterMine</i>
--------------	---

---

## Description

Get the information (name and title) of the templates pre-defined in InterMine. A template contain a query with fixed set of output columns, and at least one editable constraint, and possibly more.

## Usage

```
getTemplates(im, format = "data.frame", timeout = 3)
```

## Arguments

im	a list, containing the base URL and API token.
format	a string with values being either "data.frame" or "list", representing the output format of the template information.
timeout	an integer, representing the number of seconds to wait for the webservice to respond.

## Value

a data.frame or list object, representing the information (name and title) for the pre-define templates in the mine.

## Author(s)

InterMine Team

## Examples

```
# Get HumanMine template queries
im <- initInterMine(listMines()["HumanMine"])

templates <- getTemplates(im)
```

---

getVersion	<i>Get the version information of InterMine</i>
------------	---

---

## Description

Returns an integer representing the capabilities of the webservice.

## Usage

```
getVersion(im, timeout = 3)
```

**Arguments**

- `im` a list containing the base URL and API token.  
`timeout` an integer, representing the number of seconds to wait for the webservice to respond.

**Value**

an integer, representing the capabilities of the webservice.

**Author(s)**

InterMine Team

**Examples**

```
im <- initInterMine(listMines()["HumanMine"])

getVersion(im)
```

`getWidgets`

*Get the widgets of InterMine*

**Description**

Returns a representation of the available widgets for the mine. Each entry for the widget includes details of its type, the kind of data it processes, and any filters it accepts.

**Usage**

```
getWidgets(im)
```

**Arguments**

- `im` a list containing the base URL and API token.

**Value**

A data.frame containing information about the widgets that are available for each mine.

**Note**

The names of the widgets of the enrichment type can be passed as arguments to `doEnrichment` function.

**Author(s)**

InterMine Team

**See Also**

`getGeneIds`, `doEnrichment`

## Examples

```
# Get available FlyMine widgets
FlyMine.widgets <- getWidgets(im = initInterMine(mine = listMines()["FlyMine"]))
# Get available HumanMine widgets
HumanMine.widgets <- getWidgets(im = initInterMine(mine = listMines()["HumanMine"]))
```

---

initInterMine

*Initialize the list containing the base URL and API token.*

---

## Description

Initialize the InterMine list with the base URL of the webservice of the database and the API token.

Some resources such as lists are normally privately associated with the individual user that created them and require authentication for access. To access these private resources, each request needs to be authenticated, using an API key token. You can get an API token from the web-app of the service you intend to access: visit the MyMine tab after logging-in and click on API Key.

## Usage

```
initInterMine(mine = listMines()["HumanMine"], token="")
```

## Arguments

mine	a string, representing the base URL of the webservice of the database.
token	a string, representing the API token in order to use private functions such as list and enrichment.

## Value

A list containing the base URL and API token.

## Author(s)

InterMine Team

## References

[/urlhttp://intermine.readthedocs.io/en/latest/web-services/](http://intermine.readthedocs.io/en/latest/web-services/)

## Examples

```
im <- initInterMine(mine = listMines()["HumanMine"], "TOKEN") #replace TOKEN with your token
```

---

**InterMineR-class**

*InterMineR class contains the input values for performing queries in an InterMine instance.*

---

**Description**

InterMineR constitutes a class used to store the information which are required for performing a query for biological data in an InterMine instance. Specifically, it contains information about:

- 1) the type of data which are to be returned from the InterMine instance,
- 2) the type of sorting performed on these data, and
- 3) the constraints used to perform the query for the data of interest.

**Creating Objects**

Objects can be created using the function [setQuery](#).

**Slots**

**name** Assign with a character string giving a name to the query. Pre-fixed with "".

**description** Assign with a character string describing the purpose and the type of data retrieved by the query. Pre-fixed with "".

**select** a character vector defining the type of data to be returned.

**orderBy** a list the name of the column and the type of sorting which will be used to order the retrieved data.frame.

**where** a list containing the constraints used to restrict the query. Each constraint constitutes a list as well.

**Details**

InterMineR class specifies an object in which the input values of a query can be stored. A single constraint within the object can be assigned with multiple values.

**Author(s)**

InterMine Team

**See Also**

[setConstraints](#), [setQuery](#), [InterMineR-methods](#)

---

InterMineR-methods      *Methods for accessing InterMineR-class objects.*

---

## Description

InterMineR constitutes a class used to store the information which are required for performing a query for biological data in an InterMine instance. The methods for accessing the slots of an InterMineR-class object are presented here.

## Methods

### Class-specific methods:

`getName(InterMineR-class)`: Access the name slot of InterMineR-class object.  
`getDescription(InterMineR-class)`: Access the description slot of InterMineR-class object.  
`getSelect(InterMineR-class)`: Access the select slot of InterMineR-class object.  
`getOrderBy(InterMineR-class)`: Access the orderBy slot of InterMineR-class object.  
`getWhere(InterMineR-class)`: Access the where slot of InterMineR-class object.

## Author(s)

InterMine Team

## See Also

[setConstraints](#), [setQuery](#), [InterMineR-class](#)

## Examples

```
# get mine instance
im.fly = initInterMine(listMines()["FlyMine"])

# get GO_Gene template query
qGO_Gene = getTemplateQuery(im.fly, "GO_Gene")

# create 'InterMineR' object
go.query = setQuery(
    inheritQuery = qGO_Gene
)

class(go.query)

# access name:
getName(go.query)

# access description:
getDescription(go.query)

# access select:
getSelect(go.query)

# access orderBy:
```

```
getOrderBy(go.query)

# access where:
getWhere(go.query)
```

**listDatasets**

*Retrieve all types of InterMine features that possess dataSet as a child\_name.*

**Description**

This function retrieves all types of features from the data model of each Mine, that possess dataSet as a child\_name. For these types of features, there are available datasets in the respective mine instance.

**Usage**

```
listDatasets(im)
```

**Arguments**

im	a list containing the base URL and API token.
----	---

**Value**

a character vector containing the all types of features from the data model of the mine, that possess dataSet as a child\_name.

**Author(s)**

InterMine Team

**See Also**

[getDatasets](#)

**Examples**

```
# HumanMine
listDatasets(im = initInterMine(listMines()["HumanMine"]))

# FlyMine
listDatasets(im = initInterMine(listMines()["FlyMine"]))
```

---

**listMines***List the available InterMine-powered databases*

---

**Description**

InterMine-powered databases such as FlyMine, modENCODE, RatMine, YeastMine, HumanMine and TargetMine are integrated databases of genomic, expression and protein data for various organisms. The function listMines() lists the current available databases.

**Usage**

```
listMines()
```

**Value**

A character vector containing the name and the base URL of the web service of the database.

**Author(s)**

InterMine Team

**References**

<http://registry.intermine.org/>

**Examples**

```
listMines()
```

---

**newQuery***Initialize a new list query*

---

**Description**

A query needs to have at least view, constraints and constraintLogic. The view is a vector containing the columns of the query output. The constraint is a matrix containing the following columns: path (the path of the constraint), op (the constraint operator, one of '=', '!=', 'LOOKUP', 'ONE OF', 'NONE OF', '>', '<', '>=', '<='), value (the constraint value), code (the name of the constraint), extraValue (optional, required for LOOKUP constraints). The constraintLogic by default is "AND" operation, e.g., "A and B", where A and B are the codes in the constraints.

**Usage**

```
newQuery(name="", view=character(), sortOrder="", longDescription="",
constraintLogic=NULL)
```

### Arguments

<code>name</code>	a string, representing the name of the query.
<code>view</code>	a character vector, representing the fields to be selected from InterMine.
<code>sortOrder</code>	a string, representing the field according to which the query result is sorted and the sort order ("asc" or "desc"), following the format "FIELD ORDER".
<code>longDescription</code>	a string, representing the description of the query.
<code>constraintLogic</code>	a string, representing the logical relationship between the constraints, e.g., "A or B" where "A" and "B" are the codes in the constraints.

### Value

a list representing the query.

### Author(s)

InterMine Team

### See Also

[runQuery](#)

### Examples

```
nq <- newQuery()
```

*PL\_DiabetesGenes*

*PL\_DiabetesGenes data*

### Description

A dataset containing identifiers of genes associated with all forms of Diabetes according to OMIM  
<http://www.omim.org/>.

### Usage

```
data("PL_DiabetesGenes")
```

### Format

A data frame with 68 observations on the following 6 variables:

- Gene.symbol** Gene symbol
- Gene.name** Gene whole name
- Gene.primaryIdentifier** InterMine Gene.primaryIdentifier (ENTREZ identifier)
- Gene.secondaryIdentifier** InterMine Gene.secondaryIdentifier (ENSEMBLE identifier)
- Gene.length** Gene length in base pairs
- Gene.organism.name** Gene organism name

**Source**

<http://www.humanmine.org/humanmine/bag.do?subtab=view>

**Examples**

```
data(PL_DiabetesGenes)
```

---

PL\_FlyTF\_site\_specific\_TFs

*PL\_FlyTF\_site\_specific\_TFs data*

---

**Description**

A dataset containing identifiers of transcription factors with experimental evidence for both DNA-binding and transcriptional regulatory function. This data comes from [www.flyTF.org](http://www.flyTF.org), the drosophila transcription factor database version 2.

**Usage**

```
data("PL_FlyTF_site_specific_TFs")
```

**Format**

A data frame with 171 observations on the following 5 variables:

**Index** Index of observations

**Gene.secondaryIdentifier** InterMine Gene.secondaryIdentifier

**Gene.symbol** Gene symbol

**Gene.primaryIdentifier** InterMine Gene.primaryIdentifier

**Gene.organism.name** Gene organism name

**Source**

<http://www.flymine.org/flymine/bag.do?subtab=view>

**Examples**

```
data(PL_FlyTF_site_specific_TFs)
```

runQuery	<i>Run InterMineR queries</i>
----------	-------------------------------

## Description

Returns results from a query against data held inside the mine. These queries are similar to SQL queries, in that they request certain defined output columns of output, filtering the results through a series of "constraints".

## Usage

```
runQuery(im, qry, timeout = 60)
```

## Arguments

- |         |  |
|---------|--|
| im      | a list, containing the base URL and API token.   |
| qry     | an InterMineR or a list object, representing the query to the database.                |
| timeout | an integer, representing the number of seconds to wait for the web service to respond. |

## Details

Use [setQuery](#) function to create an InterMineR object. For setting a single constraint with multiple values, the function [setConstraints](#) can be used.

Alternatively, the user can define manually the constraints, the selection of data to be returned and the value by which they are ordered, as a list object.

For more information checkout the vignette of the package.

## Value

a data.frame containing the data which were retrieved from the InterMine instance.

## Methods

```
# S4 method for class 'InterMineR'
```

The method accepts an object of the class InterMineR and uses its information to perform the query on the defined InterMine instance.

[InterMineR-class](#) objects can contain a single constraint with multiple values.

```
# S4 method for class 'list'
```

`runQuery(im, qry = "InterMineR")`  
`runQuery(im, qry = "list")` The method accepts an object of the class list and uses its information to perform the query on the defined InterMine instance.  
 Queries defined as lists can possess only constraints with one value.

## Author(s)

InterMine Team

## See Also

[setConstraints](#), [setQuery](#), [InterMineR-class](#), [newQuery](#)

## Examples

```
# 1. Adapt 'GO_Gene' template query from FlyMine to 'InterMineR'

# get FlyMine instance
im.fly = initInterMine(listMines()["FlyMine"])

# get GO_Gene template query
qGO_Gene = getTemplateQuery(im.fly, "GO_Gene")

# constraint with GO value
qGO_Gene$where[[3]]

# modify GO_Gene template query to have more than one GO values
go.constraints = setConstraints(
    values = list(c("DNA repair", "cellular response to DNA damage stimulus")),
    modifyQueryConstraints = qGO_Gene,
    m.index = 3
)

go.constraints[[3]]

# create 'InterMineR' object
go.query = setQuery(
    inheritQuery = qGO_Gene,
    where = go.constraints
)

go.query

# run InterMineR query
go.results = runQuery(
    im = im.fly,
    qry = go.query
)

head(go.results)

# 2. Create similar query manually for Homo sapiens, using HumanMine

# get HumanMine instance
im.human = initInterMine(listMines()["HumanMine"])

# create constraints using GO terms and organism as values
hsa.go.constraints = setConstraints(
    paths = c("Gene.goAnnotation.ontologyTerm.parents.name",
              "Gene.organism.name"),
    operators = rep("=", 2),
    values = list(c("DNA repair", "cellular response to DNA damage stimulus"),
                  "Homo sapiens")
)

hsa.go.constraints

# create 'InterMineR' object
hsa.go.query = setQuery(
    select = c("Gene.secondaryIdentifier",
```

```

    "Gene.symbol",
    "Gene.goAnnotation.ontologyTerm.parents.name",
    "Gene.goAnnotation.ontologyTerm.parents.identifier",
    "Gene.goAnnotation.ontologyTerm.name",
    "Gene.goAnnotation.ontologyTerm.identifier"),
  orderBy = list(c(Gene.secondaryIdentifier = "ASC")),
  where = hsa.go.constraints
)

hsa.go.query

# run InterMineR query
hsa.go.results = runQuery(
  im = im.human,
  qry = hsa.go.query
)

head(hsa.go.results)

```

**setConstraints**

*setConstraints* function is used to create a new or modify an existing list of constraints

**Description**

*setConstraints* function is used to create or modify a list containing the constraints for an InterMine query. These constraints can be later passed on to a query formed by [setQuery](#) function.

**Usage**

```

setConstraints(
  paths,
  operators,
  values,
  modifyQueryConstraints,
  m.index
)

```

**Arguments**

<b>paths</b>	a character vector defining in which bioterm type the value(s) of each constraint belong. One assigned to each constraint.
<b>operators</b>	a character vector describing the operators for each constraint.
<b>values</b>	a list containing the values assigned to each constraints. The path of each argument defines the type of value by which the results of the query will be filtered.
<b>modifyQueryConstraints</b>	an existing list query (e.g. template query) whose constraints will be modified.
<b>m.index</b>	a vector of integer values indicating which constraints of the list assigned to <i>modifyQueryConstraints</i> will be modified.

## Details

It is important to note that:

1. setConstraints assigns each argument to a constraint based on the order in which they are assigned and one at a time. This means that paths, operators and values arguments must be of the same length and carefully assigned.
2. setConstraints can create a list of new constraints or modify an existing one with the arguments modifyQueryConstraints and m.index
3. only one of the constraints can be assigned with multiple values, aiming to retrieve results for multiple bioterms (genes, etc.)
4. to assign multiple bioterms in a constraint, values argument is assigned with a vector at the appropriate order: e.g.

to assign c("a", "b") to the second constraint one must use either:

```
values = list(first_element, c("a", "b"), ...)
```

or when modifying an existing list of constraints:

```
values = list(c("a", "b"))
```

```
m.index = 2
```

## Value

a list containing the constraints for an InterMineR query. It can be passed to [setQuery](#) function.

## Author(s)

InterMine Team

## See Also

[setQuery](#), [runQuery](#), [InterMineR-class](#)

## Examples

```
# get mine instance
im.fly = initInterMine(listMines()["FlyMine"])

# get GO_Gene template query
qGO_Gene = getTemplateQuery(im.fly, "GO_Gene")

# constraint with GO value
qGO_Gene$where[[3]]

# modify GO_Gene template query to have more than one GO values
go.constraints = setConstraints(
    values = list(c("DNA repair", "cellular response to DNA damage stimulus")),
    modifyQueryConstraints = qGO_Gene,
    m.index = 3
)
go.constraints[[3]]

# 2. Create similar query manually for Homo sapiens, using HumanMine

# get HumanMine instance
```

```

im.human = initInterMine(listMines()["HumanMine"])

# create constraints using GO terms and organism as values
hsa.go.constraints = setConstraints(
  paths = c("Gene.goAnnotation.ontologyTerm.parents.name",
            "Gene.organism.name"),
  operators = rep("=", 2),
  values = list(c("DNA repair", "cellular response to DNA damage stimulus"),
                "Homo sapiens")
)
hsa.go.constraints

```

**setQuery***Initialize a new InterMineR query or modify an existing list query***Description**

setQuery function is used to create a query for an InterMine instance. It can also accept and modify pre-defined queries of class list, like the template queries which can be retrieved with the function [getTemplateQuery](#). It returns a defined query stored as an object of class 'InterMineR'.

A query needs to have at least view, constraints and constraintLogic. The view is a vector containing the columns of the query output. The constraint is a matrix containing the following columns: path (the path of the constraint), op (the constraint operator, one of '=', '!=', 'LOOKUP', 'ONE OF', 'NONE OF', '>', '<', '>=', '<='), value (the constraint value), code (the name of the constraint), extraValue (optional, required for LOOKUP constraints). The constraintLogic by default is "AND" operation, e.g., "A and B", where A and B are the codes in the constraints.

**Usage**

```

setQuery(
  select,
  orderBy,
  where,
  name = "",
  description = "",
  inheritQuery
)

```

**Arguments**

<b>select</b>	a character vector defining the type of data to be returned.
<b>orderBy</b>	a list the name of the column and the type of sorting which will be used to order the retrieved data.frame.
<b>where</b>	a list containing the constraints used to restrict the query. Each constraint constitutes a list as well. If one wishes for a single constraint to possess more than one values, the <a href="#">setConstraints</a> function can be used.
<b>name</b>	Assign with a character string giving a name to the query. Pre-fixed with "".
<b>description</b>	Assign with a character string describing the purpose and the type of data retrieved by the query. Pre-fixed with "".

inheritQuery    Assign with a pre-defined query list, the input values of which can be modified and inherited in the new query object of class 'InterMineR'.

## Details

setQuery function can be used to create queries with a single constraint containing multiple values. The constraints can be first defined with the [setConstraints](#) function. The resulting 'InterMineR' object is passed to the [runQuery](#) function to perform the query.

## Value

An InterMineR object

## Author(s)

InterMine Team

## See Also

[setConstraints](#), [runQuery](#), [InterMineR-class](#)

## Examples

```
# get mine instance
im.fly = initInterMine(listMines()["FlyMine"])

# get GO_Gene template query
qGO_Gene = getTemplateQuery(im.fly, "GO_Gene")

# constraint with GO value
qGO_Gene$where[[3]]

# modify GO_Gene template query to have more than one GO values
go.constraints = setConstraints(
  values = list(c("DNA repair", "cellular response to DNA damage stimulus")),
  modifyQueryConstraints = qGO_Gene,
  m.index = 3
)
go.constraints[[3]]

# create 'InterMineR' object
go.query = setQuery(
  inheritQuery = qGO_Gene,
  where = go.constraints
)

class(go.query)
go.query

# 2. Create similar query manually for Homo sapiens, using HumanMine

# get HumanMine instance
im.human = initInterMine(listMines()["HumanMine"])

# create constraints using GO terms and organism as values
hsa.go.constraints = setConstraints(
```

```

paths = c("Gene.goAnnotation.ontologyTerm.parents.name",
        "Gene.organism.name"),
operators = rep("=", 2),
values = list(c("DNA repair", "cellular response to DNA damage stimulus"),
              "Homo sapiens")
)

hsa.go.constraints

# create 'InterMineR' object
hsa.go.query = setQuery(
  select = c("Gene.secondaryIdentifier",
            "Gene.symbol",
            "Gene.goAnnotation.ontologyTerm.parents.name",
            "Gene.goAnnotation.ontologyTerm.parents.identifier",
            "Gene.goAnnotation.ontologyTerm.name",
            "Gene.goAnnotation.ontologyTerm.identifier"),
  orderBy = list(c(Gene.secondaryIdentifier = "ASC")),
  where = hsa.go.constraints
)

class(hsa.go.query)
hsa.go.query

```

<b>simplifyResult</b>	<i>Convert multiple values of a column into Comma-separated character strings</i>
-----------------------	---

## Description

This function converts the values of a column variable within a dataset into comma-separated, character strings. The process is achieved by using another column of the same dataset as index.

## Usage

```

simplifyResult(
  dataset,
  index_column,
  values_column,
  returnInDataframe = FALSE
)

```

## Arguments

<b>dataset</b>	a data.frame containing the data
<b>index_column</b>	a character string or a numeric/integer value indicating the column which will be used as index.
<b>values_column</b>	a character string or a numeric/integer value indicating the column whose values will be converted into comma-separated, character strings.
<b>returnInDataframe</b>	a logical value indicating whether to return only the converted columns or to append the character strings to the dataset.

**Value**

If returnInDataframe argument is set to FALSE then a data.frame is returned with the unique values of the index column and the character strings of the values\_column that correspond to each.

If returnInDataframe argument is set to TRUE then the original dataset is returned containing an extra column in which the character strings of the values\_column have been appended.

**Author(s)**

InterMine Team

**Examples**

```
# get HumanMine
im.human = initInterMine(listMines()["HumanMine"])

# get template query for retrieving GOTerms for specific genes
qGene_GO = getTemplateQuery(im.human, "Gene.GO")

# retrieve GOTerms for four different genes
rGene_GO = list(NULL)
for(i in seq(length(c("PPARG", "RPL5", "RPL11", "TP53")))){
  g = c("PPARG", "RPL5", "RPL11", "TP53")[i]
  qGene_GO[[i]]$value = g
  rGene_GO[[i]] = runQuery(im.human, qGene_GO)

}

# rbind results to data.frame
rGene_GO = do.call(rbind, rGene_GO)

# return simplified GOTerms results for each Gene
simplify_GOTerms = simplifyResult(
  dataset = rGene_GO,
  index_column = "Gene.symbol",
  values_column = "Gene.goAnnotation.ontologyTerm.identifier",
  returnInDataframe = FALSE
)

# return simplified GOTerms results for each Gene, within the original data.frame
simplify_GOTerms.2 = simplifyResult(
  dataset = rGene_GO,
  index_column = "Gene.symbol",
  values_column = "Gene.goAnnotation.ontologyTerm.identifier",
  returnInDataframe = TRUE
)
```

**Description**

Summarize the information about the constraints contained by an object of the class InterMineR.

**Usage**

```
summary(object, ...)
```

**Arguments**

- |        |  |
|--------|--|
| object | an object of the class InterMineR.                   |
| ...    | additional arguments affecting the summary produced. |

**Value**

a data.frame containing the constraints of the InterMineR object as rows. Each constraint is constituted by a path, an operator and one or more values. Multiple values are returned as a comma-separated character string.

**Author(s)**

InterMine Team

**See Also**

[InterMineR-class](#), [setQuery](#)

**Examples**

```
# get FlyMine instance
im.fly = initInterMine(listMines()["FlyMine"])

# get GO_Gene template query
qGO_Gene = getTemplateQuery(im.fly, "GO_Gene")

# modify GO_Gene template query to have more than one GO values
go.constraints = setConstraints(
    values = list(c("DNA repair", "cellular response to DNA damage stimulus")),
    modifyQueryConstraints = qGO_Gene,
    m.index = 3
)

# create 'InterMineR' object
go.query = setQuery(
    inheritQuery = qGO_Gene,
    where = go.constraints
)

# get InterMineR constraint summary
summary(go.query)
```

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