

# Package ‘msigdb’

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**Title** An ExperimentHub Package for the Molecular Signatures Database (MSigDB)

**Version** 1.5.0

**Description** This package provides the Molecular Signatures Database (MSigDB) in a R accessible objects. Signatures are stored in GeneSet class objects from the GSEABase package and the entire database is stored in a GeneSetCollection object. These data are then hosted on the ExperimentHub. Data used in this package was obtained from the MSigDB of the Broad Institute. Metadata for each gene set is stored along with the gene set in the GeneSet class object.

**biocViews** ExperimentHub, Homo\_sapiens\_Data, Mus\_musculus\_Data

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appendKEGG	<i>Add KEGG pathway derived gene sets to a collection</i>
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### Description

This function adds gene sets derived from KEGG pathways to the MSigDB data stored in this package. Direct user-end download from the MSigDB is required to ensure KEGG licenses are adhered to.

### Usage

```
appendKEGG(gsc, version = getMsigdbVersions())
```

### Arguments

<code>gsc</code>	a GeneSetCollection object, containing MSigDB genesets in the form of GeneSet objects.
<code>version</code>	a character, stating the version of MSigDB to be retrieved (should be $\geq 7.2$ ). See <code>getMsigdbVersions()</code> .

### Value

a GeneSetCollection object, storing gene sets from the MSigDB including the downloaded KEGG gene sets.

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getIMEX	<i>Retrieve IMEx PPI hosted on the hub</i>
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**Description**

Download International Molecular Exchange (IMEx) protein-protein interaction (PPI) hosted on the ExperimentHub or retrieve pre-downloaded version from cache. This package currently hosts versions for human and mouse with both symbol and Entrez identifiers.

**Usage**

```
getIMEX(org = c("hs", "mm"), inferred = FALSE, version = getIMEXVersions())
```

**Arguments**

org	a character, representing the organism whose PPI database needs to be retrieved ("hs" for human and "mm" for mouse).
inferred	a logical, indicating whether inference from other organisms should be included in the PPI.
version	a character, stating the version of IMEX to be retrieved. See getMsigdbVersions().

**Value**

a data.frame, containing the IMEx PPI.

**Examples**

```
imex = getIMEX('hs')
```

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getIMEXVersions	<i>Get IMEX versions included in the msigdb package</i>
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**Description**

Get IMEX versions included in the msigdb package

**Usage**

```
getIMEXVersions()
```

**Value**

a character, stating the IMEX versions available in this package.

## Examples

```
getIMEXVersions()
```

---

getMsigdb

*Retrieve MSigDB data hosted on the hub*

---

## Description

Download molecular signatures database (MSigDB) hosted on the ExperimentHub or retrieve pre-downloaded version from cache. This package currently hosts versions greater than 7.2 for human and mouse with both symbol and Entrez identifiers.

## Usage

```
getMsigdb(  
  org = c("hs", "mm"),  
  id = c("SYM", "EZID"),  
  version = getMsigdbVersions()  
)
```

## Arguments

org	a character, representing the organism whose signature database needs to be retrieved ("hs" for human and "mm" for mouse).
id	a character, representing the ID type to use ("SYM" for gene symbols and "EZID" for Entrez IDs).
version	a character, stating the version of MSigDB to be retrieved (should be $\geq 7.2$ ). See <code>getMsigdbVersions()</code> .

## Value

a `GeneSetCollection`, containing `GeneSet` objects from the specified version of the molecular signatures database (MSigDB).

## Examples

```
gsc = getMsigdb('hs', 'SYM')
```

---

getMsigdbVersions	<i>Get MSigDB versions included in the msigdb package</i>
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---

**Description**

Get MSigDB versions included in the msigdb package

**Usage**

```
getMsigdbVersions()
```

**Value**

a character, stating the MSigDB versions available in this package.

**Examples**

```
getMsigdbVersions()
```

---

getMsigIdType	<i>Infer gene identifier type for the gene set collection</i>
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**Description**

The gene identifier (Symbol or Entrez ID) of a gene set collection is inferred from the IDs present in the data. A collection should ideally store gene sets using a single identifier type. This function returns the identifier type (either SymbolIdentifier or EntrezIdentifier) of the collection. It returns an error if the identifier is neither of these.

**Usage**

```
getMsigIdType(gsc)
```

**Arguments**

gsc	a GeneSetCollection object, containing MSigDB genesets in the form of GeneSet objects.
-----	--

**Value**

a GSEABase::SymbolIdentifier or GSEABASE::EntrezIdentifier object, specifying the gene identifier type (gene symbols or Entrez IDs respectively).

**Examples**

```
gsc <- getMsigdb()  
id <- getMsigIdType(gsc)
```

---

getMsigOrganism	<i>Infer organism type for the gene set collection</i>
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### Description

Since both Human and Mouse MSigDB collections are hosted in this package, this function infers the type of organism represented in a gene set collection based on the gene IDs present. If not all gene IDs belong to the same organism, the organism with more than 50% gene IDs present in the collection is returned. In any other case, the function returns an error.

### Usage

```
getMsigOrganism(gsc, idType)
```

### Arguments

gsc	a GeneSetCollection object, containing MSigDB genesets in the form of GeneSet objects.
idType	a GSEABase::SymbolIdentifier or GSEABASE::EntrezIdentifier object, representing the ID type inferred from the <code>getMsigIdType()</code> function. Avoid providing this manually.

### Value

a character, either "mm" (representing *Mus musculus* - mouse) or "hs" (representing *Homo sapiens* - human).

### Examples

```
gsc <- getMsigdb()
id <- getMsigIdType(gsc)
getMsigOrganism(gsc, id)
```

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listCollections	<i>List all collection types within a MSigDB gene set collection</i>
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### Description

This function lists all the collection types present in a MSigDB gene set collection. Descriptions of collections can be found at the MSigDB website.

### Usage

```
listCollections(gsc)
```

**Value**

a character vector, containing character codes for all collections present in the GeneSetCollection object.

**Examples**

```
gsc = getMsigdb('hs', 'SYM')
listCollections(gsc)
```

---

listSubCollections	<i>List all sub-collection types within a MSigDB gene set collection</i>
--------------------	--

---

**Description**

This function lists all the sub-collection types present in a MSigDB gene set collection. Descriptions of sub-collections can be found at the MSigDB website.

**Usage**

```
listSubCollections(gsc)
```

**Value**

a character vector, containing character codes for all sub-collections present in the GeneSetCollection object.

**Examples**

```
gsc = getMsigdb('hs', 'SYM')
listSubCollections(gsc)
```

---

msigdb	<i>The Molecular Signatures Database (MSigDB)</i>
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**Description**

This ExperimentHub package contains gene expression signatures from the molecular signatures database (MSigDB) for versions  $\geq 7.2$ . Collections for human and mouse are currently supported in this package. The mouse version was developed in conjunction with Gordon K Smyth and Alexandra Garnham, and reflects the collections available from WEHI (<https://bioinf.wehi.edu.au/MSigDB/>).

## Format

A GeneSetCollection object composed of GeneSet objects representing all non-empty gene expression signatures from the molecular signatures database (MSigDB).

## Details

The molecular signatures database (MSigDB) is a collection of over 25000 gene expression signatures that are grouped into collections and sub-collections. Metadata associated with signatures is collected and stored in the data in this package.

All data in this package are stored in a GeneSetCollection object from the GSEABase package. Each gene expression signature in the collection is stored in a GeneSet object from the GSEABase package. This data does not include KEGG gene sets due to copyrights. Users can download this data using functions provided in the package (see Details).

Data in this package does not include gene sets from the KEGG database due to licensing limitations. Users can use the `appendKEGG()` function in this package to download KEGG gene sets directly from the MSigDB and append to existing data objects.

The mouse MSigDB is created by translating human genes to mouse homologs using annotations from the Mouse Genome Informatics (MGI) database for most gene sets. Gene sets in the collections c1 (positional gene sets) and c5 (ontologies) are recreated as information in these gene sets is organism specific. Positional gene sets are created using gene information from NCBI. Gene sets representing gene ontologies are derived from the mouse R/Bioconductor organism database (`org.Mm.eg.db`).

## Acknowledgement

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## References

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Liberzon, A., Subramanian, A., Pinchback, R., Thorvaldsdóttir, H., Tamayo, P., & Mesirov, J. P. (2011). Molecular signatures database (MSigDB) 3.0. *Bioinformatics*, 27(12), 1739-1740.

Liberzon, A., Birger, C., Thorvaldsdóttir, H., Ghandi, M., Mesirov, J. P., & Tamayo, P. (2015). The molecular signatures database hallmark gene set collection. *Cell systems*, 1(6), 417-425.



## Examples

```
library(ExperimentHub)
eh <- ExperimentHub()
msigdb_datasets <- query(eh, "msigdb")

#load data using different approaches
msigdb <- getMsigdb('hs', 'SYM')
msigdb <- eh[["EH5421"]]
```

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subsetCollection	<i>Subset collections and sub-collections of MSigDB</i>
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## Description

The molecular signatures database (MSigDB) is composed of collections and sub-collection. Many analyses (e.g. gene-set enrichment using `limma::fry`) are best carried out within specific collections rather than across the entire database of signatures. This function allows subsetting of MSigDB data objects within this package using collection and sub-collection types.

## Usage

```
subsetCollection(gsc, collection = c(), subcollection = c())
```

## Arguments

<code>gsc</code>	a <code>GeneSetCollection</code> object, containing MSigDB genesets in the form of <code>GeneSet</code> objects.
<code>collection</code>	a character, stating the collection(s) to be retrieved. The collection(s) must be one from the <code>listCollections()</code> function.
<code>subcollection</code>	a character, stating the sub-collection(s) to be retrieved. The sub-collection(s) must be one from the <code>listSubCollections()</code> function.

## Value

a `GeneSetCollection` object, containing gene sets belonging to the queries collection and/or sub-collection.

## Examples

```
gsc = getMsigdb('hs', 'SYM')
subsetCollection(gsc, collection = "h")
```

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