

# Package ‘keggorthology’

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**Title** graph support for KO, KEGG Orthology

**Version** 2.61.0

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**Description** graphical representation of the Feb 2010 KEGG Orthology.  
The KEGG orthology is a set of pathway IDs that are not to be  
confused with the KEGG ortholog IDs.

**Depends** R (>= 2.5.0), hgu95av2.db, graph

**Imports** AnnotationDbi, DBI, grDevices, methods, tools, utils

**Suggests** RBGL, ALL

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**License** Artistic-2.0

**biocViews** Pathways, GraphAndNetwork, Visualization, KEGG

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|             |                                                       |
|-------------|-------------------------------------------------------|
| getK0probes | <i>obtain probe set IDs associated with a KO term</i> |
|-------------|-------------------------------------------------------|

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## Description

obtain probe set IDs associated with a KO term

## Usage

```
getK0probes(str, useAcc=TRUE, plat="hgu95av2", na.action=na.omit)
```

## Arguments

|           |                                                                                |
|-----------|--------------------------------------------------------------------------------|
| str       | string giving a KEGG orthology term                                            |
| useAcc    | logical – use all accessible terms?                                            |
| plat      | platform corresponding to a bioconductor annotation package, e.g., hgu95av2.db |
| na.action | function for dealing with NA                                                   |

## Details

Based on keggorthology read of KEGG orthology, March 2 2010. Specifically, we run wget on <ftp://ftp.genome.jp/pub/kegg/brite/ko/ko00001.keg> and use parsing and modeling code given in inst/keggHTML to generate a data frame respecting the hierarchy, and then [keggDF2graph](#) to construct the graph.

Looks up the requested term and gives back the unique probe set ids on the platform.

## Value

character vector, typically processed by na.omit

## Author(s)

Vince Carey <stvjc@channing.harvard.edu>

## Examples

```
getK0tags("insulin")
es = graph::acc(K0graph, "Endocrine System")
nm = names(es[[1]])
nm
esp = lapply(nm, getK0probes)
names(esp) = nm
sapply(esp, length)
```

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`indRender`*indented textual rendering of nodes of a hierarchical graph*

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**Description**

indented textual rendering of nodes of a hierarchical graph

**Usage**

```
indRender(klike, from=nodes(klike)[1], indent="  ")
```

**Arguments**

|                     |                                                                                               |
|---------------------|-----------------------------------------------------------------------------------------------|
| <code>klike</code>  | a graph, with tree structure similar to <a href="#">K0graph</a>                               |
| <code>from</code>   | a node name from which the rendering should proceed to all leaves                             |
| <code>indent</code> | token to use for indentation – will be replicated to depth of node to be rendered to its left |

**Details**

Based on keggorthology read of KEGG orthology, March 2 2010. Specifically, we run wget on <ftp://ftp.genome.jp/pub/kegg/brite/ko/ko00001.keg> and use parsing and modeling code given in `inst/keggHTML` to generate a data frame respecting the hierarchy, and then [keggDF2graph](#) to construct the graph.

**Value**

NULL

**Author(s)**

Vince Carey <stvjc@channing.harvard.edu>

**Examples**

```
data(K0graph)
indRender(K0graph, "Human Diseases")
```

---

`keggDF2graph`*create a graph from a specific data frame format for KEGG orthology*

---

**Description**

create a graph from a specific data frame format for KEGG orthology

**Usage**

```
keggDF2graph(df, root="K0.June07root")
data(K0graph)
```

**Arguments**

|      |                      |
|------|----------------------|
| df   | the data frame       |
| root | a name for root node |

**Details**

the obvious directed graph structure from root to leaf nodes (pathway names) is instantiated for the orthology, nodeData attribute tag is loaded with the numerical tag for the term in KEGG, and nodeData attribute depth is loaded with depth from root

Based on keggorthology read of KEGG orthology, March 2 2010. Specifically, we run wget on <ftp://ftp.genome.jp/pub/kegg/brite/ko/ko00001.keg> and use parsing and modeling code given in inst/keggHTML to generate a data frame respecting the hierarchy, and then [keggDF2graph](#) to construct the graph.

**Value**

a [graphNEL-class](#) instance

**Note**

This is only a support function. The graph is serialized in the package data directory.

**Author(s)**

Vince Carey <stvjc@channing.harvard.edu>

**Examples**

```
data(keggOrthDF)
keggOrthDF[1:5,]
data(K0graph)
graph::nodes(K0graph)[1:4]
graph::nodeData(K0graph,,"tag")[1:5]
graph::nodeData(K0graph,,"depth")[1:5]
```

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