

hpAnnot

**Marta R. Hidalgo^{*1}, Francisco Salavert², Alicia Amadoz³,
Çankut Cubuk⁴, José Carbonell-Caballero⁵, and Joaquín
Dopazo^{4,6,7}**

¹Unidad de Bioinformática y Bioestadística, Centro de Investigación Príncipe Felipe (CIPF), Valencia, 46012, Spain

²BioBam Bioinformatics S.L., Valencia, 46012, Spain

³Department of Bioinformatics, Igenomix S.A., Valencia, 46980, Spain

⁴Clinical Bioinformatics Area, Fundación Progreso y Salud (FPS), Hospital Virgen del Rocío, Sevilla, 41013, Spain

⁵Chromatin and Gene expression Lab, Gene Regulation, Stem Cells and Cancer Program, Centre de Regulació Genòmica (CRG), The Barcelona Institute of Science and Technology, PRBB, Barcelona, 08003, Spain

⁶Functional Genomics Node (INB), FPS, Hospital Virgen del Rocío, Sevilla, 41013, Spain.

⁷Bioinformatics in Rare Diseases (BiER), Centro de Investigación Biomédica en Red de Enfermedades Raras (CIBERER), FPS, Hospital Virgen del Rocío, Sevilla, 41013, Spain

*marta.hidalgo@outlook.es

2021-10-13

Abstract

HpAnnot is the annotation and data package of the *hipathia* package. *Hipathia* is a method for the computation of signal transduction along signaling pathways. The method is based on an iterative algorithm which is able to compute the signal intensity passing through the nodes of a network by taking into account the level of expression of each gene and the intensity of the signal arriving to it.

Contents

1	Usage	2
----------	--------------	----------

1 Usage

In order to access to the files stored in *AnnotationHub*, type:

```
library(AnnotationHub)
ah <- AnnotationHub()
hp <- query(ah, "hpAnnot")
hp
## AnnotationHub with 64 records
## # snapshotDate(): 2021-10-08
## # $datatypeprovider: BioMart, KEGG, GeneOntology
## # $species: Rattus norvegicus, Mus musculus, Homo sapiens, NA
## # $rdataclass: data.frame, list, igraph
## # additional mcols(): taxonomyid, genome, description,
## #   coordinate_1_based, maintainer, rdatadateadded, preparerclass, tags,
## #   rdatapath, sourceurl, sourcetype
## # retrieve records with, e.g., 'object[["AH60887"]]''
##
##           title
## AH60887 | annofuns_G0_hsa.rda
## AH60888 | annofuns_G0_mmu.rda
## AH60889 | annofuns_G0_rno.rda
## AH60890 | annofuns_uniprot_hsa.rda
## AH60891 | annofuns_uniprot_mmu.rda
## ...
## AH69121 | pmgi_rno_G0_v2.rda
## AH69122 | pmgi_rno_uniprot_v2.rda
## AH69123 | xref_hsa_v2.rda
## AH69124 | xref_mmu_v2.rda
## AH69125 | xref_rno_v2.rda
mcols(hp)[,c("title", "description")]
## DataFrame with 64 rows and 2 columns
##           title           description
##           <character>      <character>
## AH60887 annofuns_G0_hsa.rda Annotations from pat..
## AH60888 annofuns_G0_mmu.rda Annotations from pat..
## AH60889 annofuns_G0_rno.rda Annotations from pat..
## AH60890 annofuns_uniprot_hsa.. Annotations from pat..
## AH60891 annofuns_uniprot_mmu.. Annotations from pat..
## ...
## AH69121 pmgi_rno_G0_v2.rda Pseudo-pathways topo..
## AH69122 pmgi_rno_uniprot_v2... Pseudo-pathways topo..
## AH69123 xref_hsa_v2.rda XRef transformation ..
## AH69124 xref_mmu_v2.rda XRef transformation ..
## AH69125 xref_rno_v2.rda XRef transformation ..
xtabs(~datatypeprovider + species, mcols(hp))
##           species
## datatypeprovider Homo sapiens Mus musculus Rattus norvegicus
##       BioMart          12          12          12
##       KEGG             8             8             8
head(hp[["AH60887"]])
```

hpAnnot

```
##          effector.nodes      paths
## hsa03320.N-hsa03320-37 N-hsa03320-37 P-hsa03320-37
## hsa03320.N-hsa03320-611 N-hsa03320-61 P-hsa03320-61
## hsa03320.N-hsa03320-612 N-hsa03320-61 P-hsa03320-61
## hsa03320.N-hsa03320-613 N-hsa03320-61 P-hsa03320-61
## hsa03320.N-hsa03320-614 N-hsa03320-61 P-hsa03320-61
## hsa03320.N-hsa03320-615 N-hsa03320-61 P-hsa03320-61
##                                     funs
##                                     <NA>
## hsa03320.N-hsa03320-37
## hsa03320.N-hsa03320-611 negative regulation of very-low-density lipoprotein particle remodeling
## hsa03320.N-hsa03320-612 high-density lipoprotein particle remodeling
## hsa03320.N-hsa03320-613 phospholipid efflux
## hsa03320.N-hsa03320-614 regulation of Cdc42 protein signal transduction
## hsa03320.N-hsa03320-615 cholesterol efflux
hp$title
## [1] "annofuns_G0_hsa.rda"           "annofuns_G0_mmu.rda"
## [3] "annofuns_G0_rno.rda"           "annofuns_uniprot_hsa.rda"
## [5] "annofuns_uniprot_mmu.rda"       "annofuns_uniprot_rno.rda"
## [7] "annot_G0_hsa.rda"              "annot_G0_mmu.rda"
## [9] "annot_G0_rno.rda"              "annot_uniprot_hsa.rda"
## [11] "annot_uniprot_mmu.rda"         "annot_uniprot_rno.rda"
## [13] "entrez_hgnc_hsa.rda"           "entrez_hgnc_mmu.rda"
## [15] "entrez_hgnc_rno.rda"           "go_bp_frame.rda"
## [17] "go_bp_net.rda"                 "meta_graph_info_hsa.rda"
## [19] "meta_graph_info_mmu.rda"        "meta_graph_info_rno.rda"
## [21] "pmgi_hsa_genes.rda"            "pmgi_hsa_G0.rda"
## [23] "pmgi_hsa_uniprot.rda"          "pmgi_mmu_genes.rda"
## [25] "pmgi_mmu_G0.rda"               "pmgi_mmu_uniprot.rda"
## [27] "pmgi_rno_genes.rda"            "pmgi_rno_G0.rda"
## [29] "pmgi_rno_uniprot.rda"          "xref_hsa.rda"
## [31] "xref_mmu.rda"                  "xref_rno.rda"
## [33] "annofuns_G0_hsa_v2.rda"         "annofuns_G0_mmu_v2.rda"
## [35] "annofuns_G0_rno_v2.rda"         "annofuns_uniprot_hsa_v2.rda"
## [37] "annofuns_uniprot_mmu_v2.rda"     "annofuns_uniprot_rno_v2.rda"
## [39] "annot_G0_hsa_v2.rda"             "annot_G0_mmu_v2.rda"
## [41] "annot_G0_rno_v2.rda"             "annot_uniprot_hsa_v2.rda"
## [43] "annot_uniprot_mmu_v2.rda"        "annot_uniprot_rno_v2.rda"
## [45] "entrez_hgnc_hsa_v2.rda"          "entrez_hgnc_mmu_v2.rda"
## [47] "entrez_hgnc_rno_v2.rda"          "go_bp_frame_v2.rda"
## [49] "go_bp_net_v2.rda"                "meta_graph_info_hsa_v2.rda"
## [51] "meta_graph_info_mmu_v2.rda"       "meta_graph_info_rno_v2.rda"
## [53] "pmgi_hsa_genes_v2.rda"           "pmgi_hsa_G0_v2.rda"
## [55] "pmgi_hsa_uniprot_v2.rda"          "pmgi_mmu_genes_v2.rda"
## [57] "pmgi_mmu_G0_v2.rda"               "pmgi_mmu_uniprot_v2.rda"
## [59] "pmgi_rno_genes_v2.rda"             "pmgi_rno_G0_v2.rda"
## [61] "pmgi_rno_uniprot_v2.rda"          "xref_hsa_v2.rda"
## [63] "xref_mmu_v2.rda"                 "xref_rno_v2.rda"
```

For further information on this please refer to [AnnotationHub](#).