
padmet-utils Documentation

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CHAPTER 1

Installation

Clone repository from github:

```
git clone https://github.com/AuReMe/padmet-utils.git
```


CHAPTER 2

Usage

To get started using padmet, install the library as described above. Once the library becomes available on the given system, it can be developed against. The developed scripts do not need to reside in any particular location on the system.

CHAPTER 3

Development

Anyone interested in contributing or tweaking the library is more then welcome to do so. To start, simply fork the [Git repository](#) on Github and start playing with it. Then, issue pull requests.

CHAPTER 4

API Documentation

4.1 scripts API

4.1.1 Scripts: Connection

Description:

#TODO

biggAPI_to_padmet

Description: Require internet access !

Allows to extract the bigg database from the API to create a padmet.

1./ Get all reactions universal id from <http://bigg.ucsd.edu/api/v2/universal/reactions>, escape reactions of biomass.

2./ Using async_list, extract all the informations for each reactions (compounds, stochio, name ...)

3./ Need to use sleep time to avoid to lose the server access.

4./ Because the direction fo the reaction is not set by default in bigg. We get all the models where the reaction is and the final direction will the one found in more than 75%

5./ Also extract xrefs

```
usage:  
  biggAPI_to_padmet.py --output=FILE [--pwy_file=FILE] [-v]  
  
options:  
  -h --help      Show help.  
  --output=FILE  path to output, the padmet file.
```

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```
--pwy_file=FILE    add kegg pathways from pathways file, line:'pwy_id, pwy_name, x,  
↳ rxn_id'.  
-v    print info.
```

padmet_utils.connection.biggAPI_to_padmet.**add_kegg_pwy** (pwy_file, padmetRef, verbose=False)

padmet_utils.connection.biggAPI_to_padmet.**biggAPI_to_padmet** (output, pwy_file=None, verbose=False)

Extract BIGG database using the api. Create a padmet file. Escape reactions of biomass. Require internet access !

Allows to extract the bigg database from the API to create a padmet.

1./ Get all reactions universal id from <http://bigg.ucsd.edu/api/v2/universal/reactions>, escape reactions of biomass. 2./ Using async_list, extract all the informations for each reactions (compounds, stochio, name ...) 3./ Need to use sleep time to avoid to lose the server access. 4./ Because the direction fo the reaction is not set by default in bigg. We get all the models where the reaction is and the final direction will be the one found in more than 75% 5./ Also extract xrefs

Parameters

- **output** (*str*) – path to output, the padmet file.
- **pwy_file** (*str*) – path to pathway file, add kegg pathways, line:'pwy_id, pwy_name, x, rxn_id'.
- **verbose** (*bool*) – if True print information

padmet_utils.connection.biggAPI_to_padmet.**main**()

check_orthology_input

enhanced_meneco_output

Description: The standard output of meneco return ids of reactions corresponding to the solution for gapfilling.

The ids are those from the sbml and so they are encoded.

This script extract the solution corresponding to the union of reactions “Computing union of reactions from all completion” Based on padmetRef return a file with more information for each reaction.

ex: RXN_45_5

RXN-5, common_name, ec-number, Formula (with id),Formula (with cname),Action,Comment Also, the output can be used as input of the script update_padmetSpec.py In the column Action: ‘add’ => To add the reaction, ‘’ => to do nothing

Comment: the reason of adding the reaction (ex: added for gap-filling by meneco)

```
usage:  
  enhanced_meneco_output.py --meneco_output=FILE --padmetRef=FILE --output=FILE [-v]  
  
options:  
  -h --help      Show help.  
  --meneco_output=FILE    pathname of a meneco run' result  
  --padmetRef=FILE    path to padmet file corresponding to the database of  
  ↳reference (the repair network)  
  --output=FILE    path to tsv output file
```

```
padmet_utils.connection.enhanced_meneco_output.enhanced_meneco_output(meneco_output_file,
    pad-
    me-
    tRef,
    out-
    put,
    ver-
    bose=False)
```

The standard output of meneco return ids of reactions corresponding to the solution for gapfilling. The ids are those from the sbml and so they are encoded. This script extract the solution corresponding to the union of reactions “Computing union of reactions from all completion” Based on padmetRef return a file with more information for each reaction.

ex: RXN_45_5 RXN-5, common_name, ec-number, Formula (with id),Formula (with cname),Action,Comment Also, the output can be used as input for manual_curation In the column Action: ‘add’ => To add the reaction, ‘’ => to do nothing Comment: the reason of adding the reaction (ex: added for gap-filling by meneco)

Parameters

- **meneco_output_file** (*str*) – pathname of a meneco run’ result
- **padmetRef** (*padmet.padmetRef*) – path to padmet file corresponding to the database of reference (the repair network)
- **output** (*str*) – path to tsv output file
- **verbose** (*bool*) – if True print information

```
padmet_utils.connection.enhanced_meneco_output.main()
```

extract_orthofinder

Description: After running orthofinder on n fasta file, read the output file ‘Orthogroups.csv’

Require a folder ‘orthology_based_folder’ with this archi:

- |– **model_a** – model_a.sbml
- |– **model_b** –model_b.sbml

And the name of the studied organism ‘study_id’

1. Read the orthogroups file, extract orthogroups in dict ‘all_orthogroups’, and all org names
2. In orthology folder search for sbml files ‘extension = .sbml’
3. For each models regroup all information in a dict dict_data:

```
{‘study_id’: study_id, ‘model_id’ : model_id, ‘sbml_template’: path to sbml of model’, ‘output’: path to the output sbml, ‘verbose’: bool, if true print information }
```

The output is by default: output_orthofinder_from_’model_id’.sbml

4. Store all previous dict_data in a list all_dict_data
5. iter on dict from all_dict_data and use function dict_data_to_sbml

Use a dict of data dict_data and dict of orthogroups dict_orthogroup to create sbml files.

dict_data and dict_orthogroup are obtained with fun orthofinder_to_sbml

6./ Read dict_orthogroups and check if model associated to dict_data and study org share orthologue

7./ Read sbml of model, parse all reactions and get genes associated to reaction.

8./ For each reactions:

Parse genes associated to sub part (ex: (gene-a and gene-b) or gene-c) = [(gene-a,gene-b), gene-c]

Check if study org have orthologue with at least one sub part (gene-a, gene-b) or gene-c

if yes: add the reaction to the new sbml and change genes ids by study org genes ids

Create the new sbml file.

```
usage:
    extract_orthofinder --sbml=FILE/DIR --orthologues=DIR --study_id=STR --output=DIR_
    ↵[--workflow=STR] [-v]
    extract_orthofinder --sbml=DIR --orthogroups=FILE --study_id=STR --output=DIR [--_
    ↵workflow=STR] [-v]

option:
    -h --help      Show help.
    --sbml=DIR     Folder with sub folder named as models name within sbml file name as_
    ↵model_name.sbml
        --orthogroups=FILE   Output file of Orthofinder run Orthogroups.tsv
        --orthologues=DIR   Output directory of Orthofinder run Orthologues
        --study_id=ID       name of the studied organism
        --workflow=ID       workflow id in ['aureme', 'aucome']. specific run architecture_
    ↵where to search sbml files
        --output=DIR        folder where to create all sbml output files
        -v      print info
```

Use a dict of data dict_data and dict of orthogroups dict_orthogroup to create sbml files. 1./ dict_data and dict_orthogroup are obtained with fun orthofinder_to_sbml 1./ Read dict_orthogroups and check if model associated to dict_data and study org share orthologue 2./ Read sbml of model, parse all reactions and get genes associated to reaction. 3./ For each reactions:

Parse genes associated to sub part (ex: (gene-a and gene-b) or gene-c) = [(gene-a,gene-b), gene-c]
Check if study org have orthologue with at least one sub part (gene-a, gene-b) or gene-c if yes: add the reaction to the new sbml and change genes ids by study org genes ids

4./ Create the new sbml file.

Parameters

- **dict_data** (*dict*) – {‘study_id’: study_id, ‘model_id’ : model_id, ‘sbml_template’: path to sbml of model’, ‘output’: path to the output sbml, ‘verbose’: bool, if true print information }
 - **dict_orthogroup** (*dict*) – k=orthogroup_id, v = {k = name, v = set of genes}
 - **verbose** (*bool*) – if True print information

```
padmet_utils.connection.extract_orthofinder.get_sbml_files(sbml, workflow=None, verbose=False)
```

```
padmet_utils.connection.extract_orthofinder.main()
```

```
padmet_utils.connection.extract_orthofinder.orthogroups_to_sbml(orthogroups_file,
                                                               all_model_sbml,
                                                               output_folder,
                                                               study_id, verbose=False)
```

After running orthofinder on n fasta file, read the output file ‘Orthogroups.csv’ Require a folder ‘orthology_based_folder’ with this archi: model_a

model_a.sbml

model_b model_b.sbml

And the name of the studied organism ‘study_id’ 1. Read the orthogroups file, extract orthogroups in dict ‘all_orthogroups’, and all org names 2. In orthology folder search for sbml files ‘extension = .sbml’ 3. For each models regroup all information in a dict dict_data:

```
{‘study_id’: study_id, ‘model_id’: model_id, ‘sbml_template’: path to sbml of model’, ‘output’: path to the output sbml, ‘verbose’: bool, if true print information } The output is by default: out-put_orthofinder_from_’model_id’.sbml
```

4. Store all previous dict_data in a list all_dict_data

5. iter on dict from all_dict_data and use function dict_data_to_sbml This function will create a sbml from each model and conserve only reactions associated to ortholog genes For more information read the doc of func dict_data_to_sbml

Parameters

- **orthogroups_file** (*str*) – path of Orthofinder output file ‘Orthogroups.csv’
- **orthology_based_folder** (*str*) – path of folder with model’s sbml
- **output** (*str*) – pathname of the output folder of all sbml extracted
- **study_id** (*str*) – name of the studied organism
- **verbose** (*bool*) – if True print information

```
padmet_utils.connection.extract_orthofinder.orthologue_to_sbml(orthologue_folder,
                                                               all_model_sbml,
                                                               output_folder,
                                                               study_id, verbose=False)
```

After running orthofinder on n fasta file, read the output file ‘Orthogroups.csv’ Require a folder ‘orthology_based_folder’ with this archi: model_a

model_a.sbml

model_b model_b.sbml

And the name of the studied organism ‘study_id’ 1. Read the orthogroups file, extract orthogroups in dict ‘all_orthogroups’, and all org names 2. In orthology folder search for sbml files ‘extension = .sbml’ 3. For each models regroup all information in a dict dict_data:

```
{‘study_id’: study_id, ‘model_id’: model_id, ‘sbml_template’: path to sbml of model’, ‘output’: path to the output sbml, ‘verbose’: bool, if true print information } The output is by default: out-put_orthofinder_from_’model_id’.sbml
```

4. Store all previous dict_data in a list all_dict_data

5. iter on dict from all_dict_data and use function dict_data_to_sbml This function will create a sbml from each model and conserve only reactions associated to ortholog genes For more information read the doc of func dict_data_to_sbml

Parameters

- **orthogroups_file** (*str*) – path of Orthofinder output file ‘Orthogroups.csv’
- **orthology_based_folder** (*str*) – path of folder with model’s sbml
- **output** (*str*) – pathname of the output folder of all sbml extracted
- **study_id** (*str*) – name of the studied organism
- **verbose** (*bool*) – if True print information

extract_rxn_with_gene_assoc

Description: From a given sbml file, create a sbml with only the reactions associated to a gene.

Need for a reaction, in section ‘note’, ‘GENE_ASSOCIATION’:

```
usage:  
    extract_rxn_with_gene_assoc.py --sbml=FILE --output=FILE [-v]  
  
options:  
    -h --help      Show help.  
    --sbml=FILE    path to the sbml file  
    --output=FILE   path to the sbml output (with only rxn with genes assoc)  
    -v     print info
```

```
padmet_utils.connection.extract_rxn_with_gene_assoc(sbml_document,  
                                                out-  
                                                put,  
                                                ver-  
                                                bose=False)
```

From a given sbml document, create a sbml with only the reactions associated to a gene. Need for a reaction, in section ‘note’, ‘GENE_ASSOCIATION’:

Parameters

- **sbml_file** (*libsbml.document*) – sbml document
- **output** (*str*) – pathname of the output sbml

```
padmet_utils.connection.extract_rxn_with_gene_assoc.main()
```

gbk_to_faa

gene_to_targets

Description: From a list of genes, get from the linked reactions the list of products.

R1 is linked to G1, R1 produces M1 and M2. output: M1,M2. Takes into account reversibility

```
usage:  
    gene_to_targets.py --padmetSpec=FILE --genes=FILE --output=FILE [-v]  
  
option:
```

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```
-h --help      Show help
--padmetSpec=FILE  path to the padmet file
--genes=FILE    path to the file containing gene ids, one id by line
--output=FILE   path to the output file containing all targets which can be reproduced by all reactions associated to the given genes
-v   print info
```

`padmet_utils.connection.gene_to_targets.gene_to_targets(padmet, genes_file, output, verbose=False)`

From a list of genes, get from the linked reactions the list of products. R1 is linked to G1, R1 produces M1 and M2. output: M1,M2. Takes into account reversibility

Parameters

- **padmet** (`padmet.classes.PadmetSpec`) – padmet to explore
- **genes_file** (`str`) – path of genes file, 1 gene id by line
- **output** (`str`) – pathname of the output file
- **verbose** (`bool`) – if True print information

`padmet_utils.connection.gene_to_targets.main()`

modelSeed_to_padmet

Description:

#TODO

```
usage:
    modelSeed_to_padmet.py --output=FILE --rxn_file=FILE --pwy_file=FILE [-v]

options:
    -h --help      Show help.
    --output=FILE  path of the padmet file to create
    --rxn_file=FILE path to json file of modelSeed reactions
    --pwy_file=FILE path to pathway reactions association from modelSeed
    -v   print info.
```

`padmet_utils.connection.modelSeed_to_padmet.add_kegg_pwy(pwy_file, padmetRef, verbose=False)`

`padmet_utils.connection.modelSeed_to_padmet.main()`

padmet_to_asp

Description: Convert PADMet to ASP following these predicates: common_name({reaction_id or enzyme_id or pathway_id or compound_id}, common_name) direction(reaction_id, reaction_direction). reaction_direction in[LEFT-TO-RIGHT,REVERSIBLE] ec_number(reaction_id, ec(x,x,x)). catalysed_by(reaction_id, enzyme_id). uniprotID(enzyme_id, uniprot_id). #if has has_xref and db = “UNIPROT” in_pathway(reaction_id, pathway_id). reactant(reaction_id, compound_id, stoehio_value). product(reaction_id, compound_id, stoehio_value). is_a(compound_id, class_id). is_a(pathway_id, pathway_id).

```
usage:
    padmet_to_asp.py --padmet=FILE --output=FILE [-v]

option:
    -h --help      Show help.
```

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```
--padmet=FILE      path to padmet file to convert.  
--output=FILE      path to output file in lp format.  
-v     print info.
```

padmet_utils.connection.padmet_to_asp.**asp_synt** (*pred*, *list_args*)

create a predcat for asp

example: asp_synt("direction",["R1","REVERSIBLE"]) => "direction('R1','reversible')."

Parameters

- **pred** (*str*) – the predcat
- **list_args** (*list*) – list of atoms to put in the predcat

Returns the predcat ‘pred(“list_args[0]”,‘list_args[1]’,...,‘list_args[n]’).’

Return type str

padmet_utils.connection.padmet_to_asp.**main**()

```
padmet_utils.connection.padmet_to_asp.padmet_to_asp (padmet_file,      output,      ver-  
          bosc=False)
```

Convert PADMet to ASP following these predicates: common_name({reaction_id or enzyme_id or pathway_id or compound_id}, common_name) direction(reaction_id, reaction_direction). reaction_direction in [LEFT-TO-RIGHT,REVERSIBLE] ec_number(reaction_id, ec(x,x,x)). catalysed_by(reaction_id, enzyme_id). uniprotoID(enzyme_id, uniprot_id). #if has has_xref and db = “UNIPROT” in_pathway(reaction_id, pathway_id). reactant(reaction_id, compound_id, stoechio_value). product(reaction_id, compound_id, stoechio_value). is_a(compound_id, class_id). is_a(pathway_id, pathway_id).

Parameters

- **padmet_file** (*str*) – the path to padmet file to convert
- **output** (*str*) – the path to the output to create
- **verbose** (*bool*) – print informations

padmet_to_matrix

Description: Create a stoichiometry matrix from a padmet file.

The columns represent the reactions and rows represent metabolites.

S[i,j] contains the quantity of metabolite ‘i’ produced (negative for consumed) by reaction ‘j’.

```
usage:  
    padmet_to_matrix.py --padmet=FILE --output=FILE  
  
option:  
    -h --help      Show help.  
    --padmet=FILE      path to the padmet file to convert.  
    --output=FILE      path to the output file, col: rxn, row: metabo, sep = " ".
```

padmet_utils.connection.padmet_to_matrix.**main**()

padmet_utils.connection.padmet_to_matrix.**padmet_to_matrix** (*padmet*, *output*)

Create a stoichiometry matrix from a padmet file. The columns represent the reactions and rows represent metabolites. S[i,j] contains the quantity of metabolite ‘i’ produced (negative for consumed) by reaction ‘j’.

Parameters

- **padmet** (*padmet.PadmetSpec*) – padmet instance
- **output** – path to the output file, col: rxn, row: metabo, sep = “ “

padmet_to_padmet

Description: Allows to merge 1-n padmet. 1./ Update the ‘init_padmet’ with the ‘to_add’ padmet(s). to_add can be a file or a folder with only padmet files to add.

padmetRef can be used to ensure data uniformization.

```
usage:
    padmet_to_padmet.py --to_add=FILE/DIR --output=FILE [--padmetRef=FILE] [-v]

options:
    -h --help      Show help.
    --to_add=FILE/DIR  path to the padmet file to add (sep: ;) or path to folder of
    ↵padmet files.
    --output=FILE   path to the new padmet file
    --padmetRef=FILE path to the padmet file representing to the database of
    ↵reference (ex: metacyc_18.5.padmet)
    -v  print info
```

padmet_utils.connection.padmet_to_padmet.main()

padmet_utils.connection.padmet_to_padmet.padmet_to_padmet(to_add, output, padmetRef=None, verbose=False)

padmet_to_tsv

Description: convert a padmet representing a database (padmetRef) and/or a padmet representing a model (padmet-Spec) to tsv files for askomics.

1./ Folder creation given the output directory. Create this directory if required and create a folder padmetRef filename and/or padmetSpec filename

2./

2.1/ For padmetRef:

2.1.a/ Nodes get all reactions nodes => extract data from misc with extract_nodes(rxn_nodes, “reaction”, “./rxn.tsv”)

get all compounds nodes => extract data from misc with extract_nodes(cpd_nodes, “compounds”, “./cpd.tsv”)

get all pathways nodes => extract data from misc with extract_nodes(pwy_nodes, “pathway”, “./pwy.tsv”)

get all xrefs nodes => extract data from misc with extract_nodes(xref_nodes, “xref”, “./xref.tsv”)

2.1.b/ Relations for each rxn in rxn nodes:

get all rlt consumes/produces => create list of data with extract_rxn_cpd(rxn_cpd_rlt)
fieldnames = “rxn_cpd”, “concerns@reaction”, “consumes@compound”, “produces@compound”, “stoichiometry”,

get all rlt is_in_pathway => create list of data with extract_rxn_pwy(rxn_pwy_rlt)
fieldnames = “rxn_pwy”, “concerns@reaction”, “in_pwy@pathway”

get all rlt has_xref => create list of data with extract_entity_xref(rxn_xref_rlt)

for each cpd in cpd nodes:

get all rlt has_xref => update previous list of data with extract_entity_xref(cpd_xref_rlt)
fieldnames = "entity_xref","concerns@reaction","concerns@compound","has_xref@xref"

usage:

```
padmet_to_tsv.py --padmetSpec=FILE [--padmetRef=FILE] --output_dir=DIR [-v]  
padmet_to_tsv.py --padmetRef=FILE [--padmetSpec=FILE] --output_dir=DIR [-v]
```

options:

```
-h --help      Show help.  
--padmetSpec=FILE    path of the padmet representing the network to convert  
--padmetRef=FILE    path of the padmet representing the database  
--output_dir=DIR  
-v
```

padmet_utils.connection.padmet_to_tsv.**entity_xref_file**(data, output)

padmet_utils.connection.padmet_to_tsv.**extract_entity_xref**(entity_xref_rlt, padmet)

padmet_utils.connection.padmet_to_tsv.**extract_nodes**(padmet, nodes, entity_id, output, opt_col={})

for n nodes in nodes. for each node.misc = {A:[‘x’],B:[‘y’,‘z’]} create a file with line = [node.id,A[0],B[0]], [node.id,”,B[1]] the order is defined in fieldnames. merge common name and synonyms in ‘name’

padmet_utils.connection.padmet_to_tsv.**extract_pwy**(padmet)

from padmet return a dict, k = pwy_id, v = set of rxn_id in pwy

padmet_utils.connection.padmet_to_tsv.**extract_rxn_cpd**(rxn_cpd_rlt)

for rlt in rxn_cpd_rlt, append in data: [rxn_id, cpd_id(consumed), ‘,stoich,compartment] and/or [rxn_id,’,cpd_id(produced),stoich,compartment]. The value in index 0 is a merge of all data to create a unique relation id

padmet_utils.connection.padmet_to_tsv.**extract_rxn_gene**(rxn_gene_rlt)

padmet_utils.connection.padmet_to_tsv.**extract_rxn_pwy**(rxn_pwy_rlt)

for rlt in rxn_pwy_rlt, append in data: [rxn_id,pwy_id]. The value in index 0 is a merge of all data to create a unique relation id

padmet_utils.connection.padmet_to_tsv.**extract_rxn_rec**(rxn_rec_rlt)

padmet_utils.connection.padmet_to_tsv.**main**()

padmet_utils.connection.padmet_to_tsv.**pwy_rate**(padmetRef, padmetSpec, metabolic_network, output)

pwy rate in padmetSpec is calculated based on padmetRef

padmet_utils.connection.padmet_to_tsv.**rxn_cpd_file**(data, output)

from data obtained with extract_rxn_cpd(), create file rxn_cpd

padmet_utils.connection.padmet_to_tsv.**rxn_gene_file**(data, output)

padmet_utils.connection.padmet_to_tsv.**rxn_pwy_file**(data, output)

padmet_utils.connection.padmet_to_tsv.**rxn_rec_file**(data, output)

pgdb_to_padmet

Description:

Read a PGDB folder (from BIOCYC/PATHWAYTOOLS) and create a padmet. 1./ To create a padmet without any genes information extracted use the first usage with:

pgdb: path to pgdb folder output: path to the padmet to create version: to specify the version of the pgdb (20.0, 22.0) db: to sepcify the name of the database (METACYC, ECOCYC, ...) enhance: to also read the file metabolic-reaction.xml and add the to the padmet

2./ To create a padmet and add only reactions from pgdb if they are in padmetRef specific. Copy information of the reaction not from the pgdb but from the padmetRef. This allow to uniform reaction to the same version of metacyc represented in the padmetRef For example, in some case 2 pgdb from different version can contain different information for a same reaction,pathway... In this case use:

padmetRef: path to the padmet of reference

3./ To create a padmet wth genes information extracted use: extract-gene

3.1/ To remove from the final padmet all reactions without genes associated use: no-orphan

4./ To read the metabolic-reaction.xml file, a sbml with some missing reactions in PGDB use:
enhance

For more information of the parsing process read information below.

classes.dat: For each class: create new node / class = class UNIQUE-ID (1) => node.id = UNIQUE-ID COMMON-NAME (0-n) => node.Misc['COMMON-NAME'] = COMMON-NAME TYPES (0-n) => for each, check or create new node class, create rlt (node is_a_class types) SYNONYMS (0-n) => for each, create new node name, create rlt (node has_name synonyms)

compounds.dat: for each compound: create new node / class = compound UNIQUE-ID (1) => node.id = UNIQUE-ID COMMON-NAME (0-n) => node.Misc['COMMON-NAME'] = COMMON-NAME INCHI-KEY (0-1) {InChIKey=XXX} => node.misc['INCHI_KEY': XXX] MOLECULAR-WEIGHT (0-1) => node.misc()['MOLECULAR_WEIGHT'] = MOLECULAR-WEIGHT SMILES (0-1) => node.misc()['SMILES'] = SMILES TYPES (0-n) => for each, check or create new node class, create rlt (node is_a_class types) SYNONYMS (0-n) => for each, create new node name, create rlt (node has_name name) DBLINKS (0-n) {(db "id" ...)} => for each, create new node xref, create rlt (node has_xref xref)

proteins.dat: for each protein: create new node / class = protein UNIQUE-ID (1) => node.id = UNIQUE-ID COMMON-NAME (0-n) => node.Misc['COMMON-NAME'] = COMMON-NAME INCHI-KEY (0-1) {InChIKey=XXX} => node.misc['INCHI_KEY': XXX] MOLECULAR-WEIGHT (0-1) => node.misc()['MOLECULAR_WEIGHT'] = MOLECULAR-WEIGHT SMILES (0-1) => node.misc()['SMILES'] = SMILES TYPES (0-n) => for each, check or create new node class, create rlt (node is_a_class types) SYNONYMS (0-n) => for each, create new node name, create rlt (node has_name name) DBLINKS (0-n) {(db "id" ...)} => for each, create new node xref, create rlt (node has_xref xref) SPECIES (0-1) => for each, check or create new node class, create rlt (node is_in_species class)

reactions.dat: for each reaction: create new node / class = reaction + node.misc()["DIRECTION"] = "UNKNOWN" by default UNIQUE-ID (1) => node.id = UNIQUE-ID COMMON-NAME (0-n) => node.Misc['COMMON-NAME'] = COMMON-NAME EC-NUMBER (0-n) => node.Misc['EC-NUMBER'] = EC-NUMBER REACTION-DIRECTION (0-1) => node.Misc['DIRECTION'] = reaction-direction, if REVERSIBLE, else: LEFT-TO-RIGHT RXN-LOCATIONS (0,n) => node.misc['COMPARTMENT'] = rxn-location TYPES (0-n) => check or create new node class, create rlt (node.id is_a_class types's_node.id) DBLINKS (0-n) {(db "id" ...)} => create new node xref, create rlt (node has_xref xref's_node.id) SYNONYMS (0-n) => create new node name, create rlt (node has_name name's_node.id) – for LEFT and RIGHT, also check 2 next lines if info about 'coefficient'

or ‘compartment’ default value: coefficient/stoichiometry = 1, compartment = unknown also check if the direction is ‘RIGHT-TO-LEFT’, if yes, inverse consumes and produces relations then change direction to ‘LEFT-TO-RIGHT’ LEFT (1-n) => create rlt (node.id consumes left’s_node.id) RIGHT (1-n) => create rlt (node.id produces right’s_node.id)

enzrxns.dat: for each association enzyme/reaction: create new rlt / type = catalyses ENZYME (1) => stock enzyme as ‘enzyme catalyses’ REACTION (1-n) => for each reaction after, create relation ‘enzyme catalyses reaction’

pathways.dat: for each pathway: create new node / class = pathway UNIQUE-ID (1) => node._id = UNIQUE-ID TYPES (0-n) => check or create new node class, create rlt (node is_a_class types) COMMON-NAME (0-n) => node.Misc[‘COMMON-NAME’] = COMMON-NAME DBLINKS (0-n) { (db “id” ...) } => create new node xref, create rlt (node has_xref xref) SYNONYMS (0-n) => create new node name, create rlt (node has_name name) IN-PATHWAY (0-n) => check or create new node pathway, create rlt (node is_in_pathway name) REACTION-LIST (0-n) => check or create new node pathway, create rlt (node is_in_pathway name)

```
usage:
    pgdb_to_padmet.py --pgdb=DIR --output=FILE [--version=V] [--db=ID] [--
    ↪padmetRef=FILE] [--source=STR] [-v] [--enhance]
    pgdb_to_padmet.py --pgdb=DIR --output=FILE --extract-gene [--no-orphan] [--
    ↪version=V] [--db=ID] [--padmetRef=FILE] [--source=STR] [-v] [--enhance]

options:
    -h --help      Show help.
    --version=V   Xcyc version [default: N.A].
    --db=ID       Biocyc database corresponding to the pgdb (metacyc, ecocyc, ...)
    ↪[default: N.A].
    --output=FILE  padmet file corresponding to the DB.
    --pgdb=DIR     directory containing all the .dat files of metacyc (data).
    --padmetRef=FILE  padmet of reference.
    --source=STR    Tag associated to the source of the reactions, used to ensure
    ↪traceability [default: GENOME].
    --enhance      use the metabolic-reactions.xml file to enhance the database.
    --extract-gene  use the genes_file (use if its a specie's pgdb, if metacyc, do
    ↪not use).
    --no-orphan     use the genes_file (use if its a specie's pgdb, if metacyc, do not
    ↪use).
    -v      print info.
```

`padmet_utils.connection.pgdb_to_padmet.classes_parser(filePath, padmet, verbose=False)`

from class.dat: get for each class, the UNIQUE-ID, COMMON-NAME, TYPES, SYNONYMS, DBLINKS Create a class node with node.id = UNIQUE-ID, node.misc = {COMMON-NAME:[COMMON-NAMES]} - For each types: A type is in fact a class. this information is stocked in padmet as: is_a_class relation btw a node and a class_node check if the type is already in the padmet if not create a new class_node (var: subClass) with subClass_node.id = type Create a relation current node is_a_class type - For each Synonyms: this information is stocked in padmet as: has_name relation btw a node and a name_node create a new name_node with name_node.id = class_id+_names” and name_node.misc = {LABEL:[synonyms]} Create a relation current node has_name name_node.id - For each DBLINKS: DBLINKS is parsed with regex_xref to get the db and the id this information is stocked in padmet as: has_xref relation btw a node and a xref_node create a new xref_node with xref_node.id = class_id+_xrefs” and xref_node.misc = {db:[id]} Create a relation current node has_xref xref_node.id

Parameters

- `filePath (str)` – path to classes.dat
- `padmet (padmet.PadmetRef)` – padmet instance

- **verbose** (`bool`) – if True print information

```
padmet_utils.connection.pgdb_to_padmet.compounds_parser(filePath, padmet, verbose=False)
```

Parameters

- **filePath** (`str`) – path to compounds.dat
- **padmet** (`padmet.PadmetRef`) – padmet instance
- **verbose** (`bool`) – if True print information

```
padmet_utils.connection.pgdb_to_padmet.enhance_db(metadata_reactions, padmet, with_genes, verbose=False)
```

Parse sbml metabolic_reactions and add reactions in padmet if with_genes: add also genes information

Parameters

- **metadata_reactions** (`str`) – path to sbml metabolic-reactions.xml
- **padmet** (`padmet.PadmetRef`) – padmet instance
- **with_genes** (`bool`) – if true also add genes information.

Returns padmet instance with pgdb within pgdb + metabolic-reactions.xml data

Return type `padmet.PadmetRef`

```
padmet_utils.connection.pgdb_to_padmet.enzrxns_parser(filePath, padmet, dict_protein_gene_id, source, verbose=False)
```

Parameters

- **filePath** (`str`) – path to enzrxns.dat
- **padmet** (`padmet.PadmetRef`) – padmet instance
- **verbose** (`bool`) – if True print information

```
padmet_utils.connection.pgdb_to_padmet.from_pgdb_to_padmet(pgdb_folder, db='NA', version='NA', source='GENOME', extract_gene=False, no_orphan=False, enhanced_db=False, padmetRef_file=None, verbose=False)
```

Parameters

- **pgdb_folder** (`str`) – path to pgdb
- **db** (`str`) – pgdb name, default is ‘NA’
- **version** (`str`) – pgdb version, default is ‘NA’
- **source** (`str`) – tag reactions for traceability, default is ‘GENOME’
- **extract_gene** (`bool`) – if true extract genes information
- **no_orphan** (`bool`) – if true, remove reactions without genes associated
- **enhanced_db** (`bool`) – if true, read metabolix-reactions.xml sbml file and add information in final padmet

- **padmetRef_file** (*str*) – path to padmetRef corresponding to metacyc in padmet format
- **verbose** (*bool*) – if True print information

Returns padmet instance with pgdb within pgdb data

Return type padmet.padmetRef

```
padmet_utils.connection.pgdb_to_padmet.genes_parser(filePath, padmet, verbose=False)
```

Parameters

- **filePath** (*str*) – path to genes.dat
- **padmet** (*padmet.PadmetRef*) – padmet instance
- **verbose** (*bool*) – if True print information

```
padmet_utils.connection.pgdb_to_padmet.main()
```

```
padmet_utils.connection.pgdb_to_padmet.map_gene_id(dict_protein_gene_id, map_gene_ids)
```

Map gene ID created by Pathway Tools with gene ID from the data. Automatically Pathway Tools uppercased all the letter in gene ID. So we need to do this mapping to retrieve the unuppercased gene ID.

```
padmet_utils.connection.pgdb_to_padmet.pathways_parser(filePath, padmet, verbose=False)
```

Parameters

- **filePath** (*str*) – path to pathways.dat
- **padmet** (*padmet.PadmetRef*) – padmet instance
- **verbose** (*bool*) – if True print information

```
padmet_utils.connection.pgdb_to_padmet.proteins_parser(filePath, padmet, verbose=False)
```

Parameters

- **filePath** (*str*) – path to proteins.dat
- **padmet** (*padmet.PadmetRef*) – padmet instance
- **verbose** (*bool*) – if True print information

```
padmet_utils.connection.pgdb_to_padmet.reactions_parser(filePath, padmet, extract_gene, source, verbose=False)
```

from reaction.dat: get for each reaction, the UNIQUE-ID, COMMON-NAME, TYPES, SYONYMS, DBLINKS Create a reaction node with node.id = UNIQUE-ID, node.misc = {COMMON-NAME:[COMMON-NAMES]} - For each types: A type is in fact a class. this information is stocked in padmet as: is_a_class relation btw a node and a class_node check if the type is already in the padmet if not create a new class_node (var: subClass) with subClass_node.id = type Create a relation current node is_a_class type - For each Synonyms: this information is stocked in padmet as: has_name relation btw a node and a name_node create a new name_node with name_node.id = reaction_id+"_names" name_node.misc = {LABEL:[synonyms]} Create a relation current node has_name name_node.id - For each DBLINKS: DBLINKS is parsed with regex_xref to get the db and the id this information is stocked in padmet as: has_xref relation btw a node and a xref_node create a new xref_node with xref_node.id = reaction_id+"_xrefs" and xref_node.misc = {db:[id]} Create a relation current node has_xref xref_node.id

Parameters

- **filePath** (*str*) – path to reactions.dat

- **padmet** (`padmet.PadmetRef`) – padmet instance
- **verbose** (`bool`) – if True print information

sbmlGenerator

Description: The module sbmlGenerator contains functions to generate sbml files from padmet and txt usign the libsbml package

```
usage:
    sbmlGenerator.py --padmet=FILE --output=FILE --sbml_lvl=STR [--model_id=STR] [--obj_fct=STR] [--mnx_chem_prop=FILE] [--mnx_chem_xref=FILE] [-v]
    sbmlGenerator.py --padmet=FILE --output=FILE [--init_source=STR] [-v]
    sbmlGenerator.py --compound=FILE --output=FILE [--padmetRef=FILE] [-v]
    sbmlGenerator.py --reaction=FILE --output=FILE --padmetRef=FILE [-v]

option:
    -h --help      Show help.
    --padmet=FILE   path of the padmet file to convert into sbml
    --output=FILE    path of the sbml file to generate.
    --mnx_chem_prop=FILE   path of the MNX chemical compounds properties.
    --mnx_chem_xref=FILE   path of the mnx dict of chemical compounds id mapping.
    --reaction=FILE    path of file of reactions ids, one by line to convert to sbml.
    --compound=FILE    path of file of compounds ids, one by line to convert to sbml.
    --init_source=STR   Select the reactions of padmet to convert on sbml based on
    ↪the source of the reactions, check relations rxn has_reconstructionData.
    --sbml_lvl=STR     sbml level of output.
    --obj_fct=STR      id of the reaction objective.
    -v     print info.
```

`padmet_utils.connection.sbmlGenerator.add_ga(rId_encoded, all_ga_subsets)`

if list_ga len == 1: only 1 list of gene: if len of this list is 1: just add gene, else create OR structure else: create OR structure, then for each list of gene for each ga in list_ga: if len == 1: if the only ga len == 1: just add gene, else create OR structure elif len > 1: create AND structure, then for each GA if len GA == 1: just add gene, else create OR structure if no suppdata, if linked_genes: if len linked_genes == 1: just add gene, else create OR structure

`padmet_utils.connection.sbmlGenerator.check(value, message)`

If ‘value’ is None, prints an error message constructed using ‘message’ and then exits with status code 1. If ‘value’ is an integer, it assumes it is a libSBML return status code. If the code value is LIBSBML_OPERATION_SUCCESS, returns without further action; if it is not, prints an error message constructed using ‘message’ along with text from libSBML explaining the meaning of the code, and exits with status code 1.

`padmet_utils.connection.sbmlGenerator.compound_to_sbml(species_compart, output, verbose=False)`

convert a list of compounds to sbml format if compart_name is not None, then the compounds id will by: M_originalID_compart_name if verbose and specified padmetRef and/or padmetSpec: will check if compounds are in one of the padmet files Ids are encoded for sbml using functions sbmlPlugin.convert_to_coded_id @param compounds_file: the pathname to the file containing the compounds ids and the compart, line = cpd-id compart. @param output: the pathname to the sbml file to create @param padmetRef_file: the pathname to the file padmet of reference @param padmetRef_file: the pathname to the file padmet of a species @param compart_name: the default compart to concatenate @param sbml_version: the sbml version @param verbose: print informations @type compounds_file, output, padmetRef_file, padmetSpec_file, verbose: str @type sbml_lvl, sbml_version: int @return: check return of writeSBMLToFile @rtype: int

```
padmet_utils.connection.sbmlGenerator.create_annotation(inchi, ref_id)
    dict_data, k = url, v = id

padmet_utils.connection.sbmlGenerator.create_note(dict_data)

padmet_utils.connection.sbmlGenerator.main()

padmet_utils.connection.sbmlGenerator.padmet_to_sbml(padmet,           output,
                                                       model_id=None,
                                                       obj_fct=None,      sbml_lvl=3,
                                                       mnx_chem_prop=None,
                                                       mnx_chem_xref=None,   verbose=False)
```

Convert padmet file to sbml file. Specificity: - ids are encoded for sbml using functions sbmlPlugin.convert_to_coded_id @param padmet_file: the pathname to the padmet file to convert @param output: the pathname to the sbml file to create @param obj_fct: the identifier of the objection function, the reaction to test in FBA @param sbml_lvl: the sbml level @param sbml_version: the sbml version @param verbose: print informations @type padmet_file, output, verbose: str @type sbml_lvl, sbml_version: int @return: check return of writeSBMLToFile @rtype: int

```
padmet_utils.connection.sbmlGenerator.parse_mnx_chem_prop(mnx_chem_prop)
padmet_utils.connection.sbmlGenerator.parse_mnx_chem_xref(mnx_chem_xref)
padmet_utils.connection.sbmlGenerator.reaction_to_sbml(reactions, output, padmetRef, verbose=False)
```

convert a list of reactions to sbml format based on a given padmet of reference. - ids are encoded for sbml using functions sbmlPlugin.convert_to_coded_id @param reactions: list of reactions ids @param padmetRef: padmet of reference @param output: the pathname to the sbml file to create @param sbml_lvl: the sbml level @param sbml_version: the sbml version @param verbose: print informations @type reactions: set @type output, verbose: str @type padmetRef: <Padmet> @type sbml_lvl, sbml_version: int @return: check return of writeSBMLToFile @rtype: int

sbml_to_curation_form

Description: extract all reactions from a sbml file to the form used in aureme for curation.

```
usage:
    sbml_to_curation_form.py --sbml=FILE --output=FILE --comment=STR [--rxn_id=ID]

options:
    -h --help      Show help.
    --sbml=FILE    path of the sbml.
    --output=FILE  form containing the reaction extracted, form used for manual_
    ↪curation in aureme.
    --rxn_id=FILE  id of one reaction or n reactions sep by ';', if None try to
    ↪extract the reaction with objective coefficient == 1.
    --comment=STR   comment associated to the reactions in the form. Used to track_
    ↪sources of curation in aureme.
```

sbml_to_padmet

Description: There are 3 cases of conversion sbml to padmet:

1./ Creation of a reference database in padmet format from sbml(s) (or updating one with new(s) sbml(s)) First usage, padmetRef is the padmetRef to create or to update. If it's an update case, the output can be used to create a new padmet, if output None, will overwrite the input padmetRef.

2./ Creation of a padmet representing an organism in padmet format from sbml(s) (or updating one with new(s) sbml(s))
 2.A/ Without a database of reference: Second usage, padmetSpec is the padmetSpec to create or update. If it's an update case, the output can be used to create a new padmet, if output None, will overwrite the input padmetSpec.

2.B/ With a database of reference: Third usage, padmetSpec is the padmetSpec to create or update. If it's an update case, the output can be used to create a new padmet, if output None, will overwrite the input padmetSpec. padmetRef is the padmet representing the database of reference.

It is possible to define a specific policy and info for the padmet. To learn more about policy and info check doc of lib.padmetRef/Spec. if the ids of reactions/compounds are not the same between padmetRef and the sbml, it is possible to use a dictionary of association (sbml_id padmetRef_id) with one line = 'id_sbml id_padmetRef' Finally if a reaction from sbml is not in padmetRef, it is possible to force the copy and creating a new reaction in padmetSpec with the arg -f

```
usage:
  sbml_to_padmet.py --sbml=FILE --padmetRef=FILE [--output=FILE] [--db=STR] [--version=STR] [-v]
  sbml_to_padmet.py --sbml=FILE --padmetSpec=FILE [--output=FILE] [--db=STR] [--version=STR] [-v]
  sbml_to_padmet.py --sbml=FILE --padmetSpec=FILE [--padmetRef=FILE] [--output=FILE] [--mapping=FILE] [--source_tool=STR] [--source_category=STR] [--source_id=STR] [-v] [-f]

options:
  -h --help      Show help.
  --padmetSpec=FILE    path to the padmet file to update with the sbml. If there's no padmetSpec, just specify the output
  --padmetRef=FILE    path to the padmet file representing to the database of reference (ex: metacyc_18.5.padmet)
  --sbml=FILE    1 sbml file to convert into padmetSpec (ex: my_network.xml/sbml) OR a directory with n SBML
  --output=FILE   pathanme to the new padmet file
  --mapping=FILE  dictionnary of association id_origin id_ref
  --db=STR        database name
  --version=STR   database version
  -v   print info
```

```
padmet_utils.connection.sbml_to_padmet.from_sbml_to_padmet(sbml, padmetSpec_file, source_tool, source_category, source_id, padmetRef_file, mapping, db='NA', version='NA', verbose=False)

padmet_utils.connection.sbml_to_padmet.main()
```

wikiGenerator

Description: Contains all necessary functions to generate wikiPages from a padmet file and update a wiki online.
 Require WikiManager module (with wikiMate, Vendor)

```
usage:
  wikiGenerator.py --padmet=FILE/DIR --output=DIR --wiki_id=STR [--database=STR] [--padmetRef=FILE] [--log_file=FILE] [-v]
```

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```
wikiGenerator.py --aureme_run=DIR --padmetSpec=ID -v

options:
  -h --help      Show help.
  --padmet=FILE  path to padmet file.
  --output=DIR   path to folder to create with all wikipages in subdir.
  --wiki_id=STR  id of the wiki.
  --padmetRef=FILE path to padmet of reference, ex: metacyc_xx.padmet, if given, able to calcul pathway rate completion.
  --log_file=FILE log file from an aureme run, use this file to create a wikipage with all the command used during the aureme run.
  --aureme_run=DIR can use an aureme run as input, will use from config file information for model_id and log_file and padmetRef.
  -v      print info.
```

```
padmet_utils.connection.wikiGenerator.add_collapsible(text_array, title=None)
#TODO

padmet_utils.connection.wikiGenerator.add_property(properties, prop_id, prop_values)
#TODO

padmet_utils.connection.wikiGenerator.copy_io_files()

padmet_utils.connection.wikiGenerator.createDirectory(output, verbose=False)
    create the folders genes, reactions, metabolites, pathways in the folder dirPath/ if already exist, it will replace old folders (and delete old files)

Parameters output (str) – path to output folder

padmet_utils.connection.wikiGenerator.create_biological_page(category, page_id,
                                                               page_dict_data,
                                                               total_padmet_data,
                                                               ext_link,         output_file,        pad-
                                                               metRef=None,       verbose=False)
#TODO

padmet_utils.connection.wikiGenerator.create_log_page(log_file, output_folder)
#TODO

padmet_utils.connection.wikiGenerator.create_main(wiki_id)
#TODO

padmet_utils.connection.wikiGenerator.create_navigation_page(total_padmet_data,
                                                               navigation_folder,
                                                               verbose=False)
#TODO

padmet_utils.connection.wikiGenerator.create_venn()
#TODO

padmet_utils.connection.wikiGenerator.draw_ellipse(fig, ax, x, y, w, h, a, fillcolor)
padmet_utils.connection.wikiGenerator.draw_text(fig, ax, x, y, text, color=[0, 0, 0, 1])
padmet_utils.connection.wikiGenerator.extract_padmet_data(padmetFile,          total_padmet_data,
                                                          global_pwy_rxn_dict=None,
                                                          padmetRef=None,    verbose=False)
```

```
total_padmet_data: k in ['reaction', 'gene', 'organism', 'pathway', ...] if k = 'reaction', v =
{'misc':{},'gene_assoc':{}}
```

For reaction in padmetFile:

```
if reaction_id not in total_padmet_data["reaction"].keys(): add total_padmet_data["reaction"][reaction_id][padmet_sou-
= dict()
```

else, add data only if different from first

```
padmet_utils.connection.wikiGenerator.get_cmd_label(cmd)
#TODO
```

```
padmet_utils.connection.wikiGenerator.get_labels(data, fill=['number'])
get a dict of labels for groups in data example: In [12]: get_labels([range(10), range(5,15), range(3,8)],
fill=['number']) Out[12]: {'001': '0', '010': '5', '011': '0', '100': '3', '101': '2', '110': '2', '111': '3'}
```

Parameters

- **data** (*list*) – data to get label for
- **fill** – ['number'|"logic"|"percent"]

Returns a dict of labels for different sets

Return type `dict`

```
padmet_utils.connection.wikiGenerator.main()
```

```
padmet_utils.connection.wikiGenerator.reduce_padmet_data(total_padmet_data, ver-
bose=False)
#TODO
```

```
padmet_utils.connection.wikiGenerator.update_basic_attrib(node, cur-
rent_node_dict, pad-
met_source)
#TODO
```

```
padmet_utils.connection.wikiGenerator.venn4(labels, names=['A', 'B', 'C', 'D'], **op-
tions)
plots a 4-set Venn diagram
```

Parameters

- **labels** (*dict*) – a label dict where keys are identified via binary codes ('0001', '0010', '0100', ...), hence a valid set could look like: {'0001': 'text 1', '0010': 'text 2', '0100': 'text 3', ...}. Unmentioned codes are considered as ''.
- **names** (*list*) – group names

Returns (Figure, AxesSubplot), pyplot Figure and AxesSubplot object

Return type `set`

```
padmet_utils.connection.wikiGenerator.wikiGenerator(padmet, output, wiki_id, pad-
metRef=None, database=None,
log_file=None, verbose=False)
```

```
padmet_utils.connection.wikiGenerator.xrefLink(dataInArray, db, ids)
#TODO
```

4.1.2 Scripts: Exploration

Description:

#TODO

compare_padmet

Description: #Compare 1-n padmet and create a folder output with files: genes.csv:

fieldnames = [gene, padmet_a, padmet_b, padmet_a_rxn_assoc, padmet_b_rxn_assoc] line = [gene-a, ‘present’ (if in padmet_a), ‘present’ (if in padmet_b), rxn-1;rxn-2 (names of reactions associated to gene-a in padmet_a), rxn-2]

reactions.csv: fieldnames = [reaction, padmet_a, padmet_b, padmet_a_genes_assoc, padmet_b_genes_assoc, padmet_a_formula, padmet_b_formula] line = [rxn-1, ‘present’ (if in padmet_a), ‘present’ (if in padmet_b), ‘gene-a;gene-b; gene-a, ‘cpd-1 + cpd-2 => cpd-3’, ‘cpd-1 + cpd-2 => cpd-3’]

pathways.csv: fieldnames = [pathway, padmet_a_completion_rate, padmet_b_completion_rate, padmet_a_rxn_assoc, padmet_b_rxn_assoc] line = [pwy-a, 0.80, 0.30, rxn-a;rxn-b; rxn-a]

compounds.csv: fieldnames = [‘metabolite’, padmet_a_rxn_consume, padmet_a_rxn_produce, padmet_b_rxn_consume, padmet_rxn_produce] line = [cpd-1, rxn-1,’,rxn-1,’]

```
usage:
    compare_padmet.py --padmet=FILES/DIR --output=DIR [--padmetRef=FILE] [-v]

option:
    -h --help      Show help.
    --padmet=FILES/DIR      pathname of the padmet files, sep all files by ',', ex: /
    ↪path/padmet1.padmet;/path/padmet2.padmet OR a folder
    --output=DIR      pathname of the output folder
    --padmetRef=FILE      pathanme of the database ref in padmet
```

padmet_utils.exploration.compare_padmet.**compare_padmet**(*padmet_path*, *output*,
padmetRef=None, *verbose=False*)

#Compare 1-n padmet and create a folder output with files: genes.csv:

fieldnames = [gene, padmet_a, padmet_b, padmet_a_rxn_assoc, padmet_b_rxn_assoc] line = [gene-a, ‘present’ (if in padmet_a), ‘present’ (if in padmet_b), rxn-1;rxn-2 (names of reactions associated to gene-a in padmet_a), rxn-2]

reactions.csv: fieldnames = [reaction, padmet_a, padmet_b, padmet_a_genes_assoc, padmet_b_genes_assoc, padmet_a_formula, padmet_b_formula] line = [rxn-1, ‘present’ (if in padmet_a), ‘present’ (if in padmet_b), ‘gene-a;gene-b; gene-a, ‘cpd-1 + cpd-2 => cpd-3’, ‘cpd-1 + cpd-2 => cpd-3’]

pathways.csv: fieldnames = [pathway, padmet_a_completion_rate, padmet_b_completion_rate, padmet_a_rxn_assoc, padmet_b_rxn_assoc] line = [pwy-a, 0.80, 0.30, rxn-a;rxn-b; rxn-a]

compounds.csv: fieldnames = [‘metabolite’, padmet_a_rxn_consume, padmet_a_rxn_produce, padmet_b_rxn_consume, padmet_rxn_produce] line = [cpd-1, rxn-1,’,rxn-1,’]

Parameters

- **padmet_path** (*str*) – pathname of the padmet files, sep all files by ‘,’, ex: /path/padmet1.padmet;/path/padmet2.padmet OR a folder
- **output** (*str*) – pathname of the output folder
- **padmetRef** (*padmet.classes.PadmetRef*) – padmet containing the database of reference, need to calculat pathway completion rate
- **verbose** (*bool*) – if True print information

```
padmet_utils.exploration.compare_padmet.main()
```

compare_sbml

Description: compare reactions in two sbml.

Returns if a reaction is missing

And if a reaction with the same id is using different species or different reversibility

```
usage:
    compare_sbml.py --sbml1=FILE --sbml2=FILE

option:
    -h --help      Show help.
    --sbml1=FILE   path of the first sbml file
    --sbml2=FILE   path of the second sbml file
```

compare_sbml_padmet

Description: compare reactions in sbml and padmet file

```
usage:
    compare_sbml_padmet.py --padmet=FILE --sbml=FILE

option:
    -h --help      Show help.
    --padmet=FILE  path of the padmet file
    --sbml=FILE    path of the sbml file
```

```
padmet_utils.exploration.compare_sbml_padmet.compare_sbml_padmet(sbml_document,
                                                                padmet)
compare reactions ids in sbml vs padmet, return nb of reactions in both and reactions id not in sbml or not in padmet
```

Parameters

- **padmet** (*padmet.classes.PadmetSpec*) – padmet to update
- **sbml_file** (*libsbml.document*) – sbml document

```
padmet_utils.exploration.compare_sbml_padmet.main()
```

convert_sbml_db

Description: This tool is use the MetaNetX database to check or convert a sbml. Flat files from MetaNetx are required to run this tool. They can be found in the aureme workflow or from the MetaNetx website. To use the tool set:

mnx_folder= the path to a folder containing MetaNetx flat files. the files must be named as ‘reac_xref.tsv’ and ‘chem_xref.tsv’ or set manually the different path of the flat files with:

mnx_reac= path to the flat file for reactions

mnx_chem= path to the flat file for chemical compounds (species)

To check the database used in a sbml:

to check all element of sbml (reaction and species) set: to-map=all

to check only reaction of sbml set: to-map=reaction

to check only species of sbml set: to-map=species

To map a sbml and obtain a file of mapping ids to a given database set:

to-map: as previously explained

db_out: the name of the database target: ['metacyc', 'bigg', 'kegg'] only

output: the path to the output file

For a given sbml using a specific database.

Return a dictionnary of mapping.

the output is a file with line = reaction_id/or species in sbml, reaction_id/species in db_out database

ex: For a sbml based on kegg database, db_out=metacyc: the output file will contains for ex:

R02283 ACETYLORNTRANSAM-RXN

```
usage:
    convert_sbml_db.py --mnx_reac=FILE --mnx_chem=FILE --sbml=FILE --to-map=STR [-v]
    convert_sbml_db.py --mnx_folder=DIR --sbml=FILE --to-map=STR [-v]
    convert_sbml_db.py --mnx_folder=DIR --sbml=FILE --output=FILE --db_out=ID --to-
    ↵map=STR [-v]
    convert_sbml_db.py --mnx_reac=FILE --mnx_chem=FILE --sbml=FILE --output=FILE --db_
    ↵out=ID --to-map=STR [-v]

options:
    -h --help      Show help.
    --to-map=STR    select the part of the sbml to check or convert, must be in ['all
    ↵', 'reaction', 'species']
    --mnx_reac=FILE    path to the MetaNetX file for reactions
    --mnx_chem=FILE    path to the MetaNetX file for compounds
    --sbml=FILE      path to the sbml file to convert
    --output=FILE     path to the file containing the mapping, sep = "      "
    --db_out=FILE     id of the output database in ["BIGG", "METACYC", "KEGG"]
    -v      verbose.
```

```
padmet_utils.exploration.convert_sbml_db.check_sbml_db(sbml_file,           to_map,
                                                       verbose=False,
                                                       mnx_reac_file=None,
                                                       mnx_chem_file=None,
                                                       mnx_folder=None)
```

Check sbml database of a given sbml.

Parameters

- **sbml_file** (*str*) – path to the sbml file to convert
- **to_map** (*str*) – select the part of the sbml to check must be in ['all', 'reaction', 'species']
- **verbose** (*bool*) – if true: more info during process
- **mnx_reac_file** (*str*) – path to the flat file for reactions (can be None if given mnx_folder)
- **mnx_chem_file** (*str*) – path to the flat file for chemical compounds (species) (can be None if given mnx_folder)
- **mnx_folder** (*str*) – the path to a folder containing MetaNetx flat files

Returns (name of the best matching database, dict of matching)

Return type tuple

```
padmet_utils.exploration.convert_sbml_db.get_from_mnx(mnx_dict,           element_id,
                                                       db_out)
padmet_utils.exploration.convert_sbml_db.intern_mapping(id_to_map, db_out, _type)
padmet_utils.exploration.convert_sbml_db.main()
padmet_utils.exploration.convert_sbml_db.map_sbml(sbml_file,      to_map,      db_out,
                                                 output,          verbose=False,
                                                 mnx_reac_file=None,
                                                 mnx_chem_file=None,
                                                 mnx_folder=None)
```

map a sbml and obtain a file of mapping ids to a given database.

Parameters

- **sbml_file** (*str*) – path to the sbml file to convert
- **to_map** (*str*) – select the part of the sbml to check must be in ['all', 'reaction', 'species']
- **db_out** (*str*) – the name of the database target: ['metacyc', 'bigg', 'kegg'] only
- **output** (*str*) – path to the file containing the mapping, sep = " "
- **verbose** (*bool*) – if true: more info during process
- **mnx_reac_file** (*str*) – path to the flat file for reactions (can be None if given mnx_folder)
- **mnx_chem_file** (*str*) – path to the flat file for chemical compounds (species) (can be None if given mnx_folder)
- **mnx_folder** (*str*) – the path to a folder containing MetaNetx flat files

Returns (name of the best matching database, dict of matching)

Return type tuple

```
padmet_utils.exploration.convert_sbml_db.mnx_reader(input_file, db_out)
```

dendrogram_reactions_distance

Description: Use reactions.csv file from compare_padmet.py to create a dendrogram using a Jaccard distance.

From the matrix absence/presence of reactions in different species computes a Jaccard distance between these species. Apply a hierarchical clustering on these data with a complete linkage. Then create a dendrogram. Apply also intervene to create an upset graph on the data.

```
usage:
    dendrogram_reactions_distance.py --reactions=FILE --output=FILE [--padmetRef=STR]
    ↪[--pvclust] [--upset=INT] [-v]

option:
    -h --help      Show help.
    -r --reactions=FILE    pathname of the file containing reactions in each species
    ↪of the comparison.
    -o --output=FOLDER    path to the output folder.
    --pvclust    launch pvclust dendrogram using R
    --padmetRef=STR    path to the padmet Ref file
    -u --upset=INT    number of cluster in the upset graph.
    -v      verbose mode.
```

flux_analysis

Description: Run flux balance analyse with cobra package. If the flux is >0. Run also FVA and return result in standard output

```
usage:
    flux_analysis.py --sbml=FILE
    flux_analysis.py --sbml=FILE --seeds=FILE --targets=FILE
    flux_analysis.py --sbml=FILE --all_species

option:
    -h --help      Show help.
    --sbml=FILE    pathname to the sbml file to test for fba and fva.
    --seeds=FILE   pathname to the sbml file containing the seeds (medium).
    --targets=FILE  pathname to the sbml file containing the targets.
    --all_species   allow to make FBA on all the metabolites of the given model.
```

get_pwy_from_rxn

Description: From a file containing a list of reaction, return the pathways where these reactions are involved. ex: if rxn-a in pwy-x => return, pwy-x; all rxn ids in pwy-x; all rxn ids in pwy-x FROM the list; ratio

```
usage:
    get_pwy_from_rxn.py --reaction_file=FILE --padmetRef=FILE --output=FILE

options:
    -h --help      Show help.
    --reaction_file=FILE    pathname of the file containing the reactions id, 1/line
    --padmetRef=FILE    pathname of the padmet representing the database.
    --output=FILE      pathname of the file with line = pathway id, all reactions id, ratio
    ↵reactions ids from reaction file, ratio. sep = " "
```

padmet_utils.exploration.get_pwy_from_rxn.**dict_pwys_to_file**(dict_pwy, output)

Create csv file from dict_pwy. dict_pwy is obtained with extract_pwys()

Parameters

- **dict_pwy** (*dict*) – dict, k=pathway_id, v=dict: k in [total_rxn, rxn_from_list, ratio ex: {pwy-x:{'total_rxn':[a,b,c], rxn_from_list:[a], ratio:1/3}}]
- **output** (*str*) – path to output file

padmet_utils.exploration.get_pwy_from_rxn.**extract_pwys**(padmet, reactions)

#extract from padmet pathways containing 1-n reactions from a set of reactions ‘reactions’ Return a dict of data. dict, k=pathway_id, v=dict: k in [total_rxn, rxn_from_list, ratio ex: {pwy-x:{'total_rxn':[a,b,c], rxn_from_list:[a], ratio:1/3}}]

Parameters

- **padmet** (*padmet.classes.PadmetSpec*) – padmet to update
- **reactions** (*set*) – set of reactions to match with pathways

Returns dict, k=pathway_id, v=dict: k in [total_rxn, rxn_from_list, ratio ex: {pwy-x:{'total_rxn':[a,b,c], rxn_from_list:[a], ratio:1/3}}]

Return type *dict*

padmet_utils.exploration.get_pwy_from_rxn.**main**()

padmet_stats

Description: From a file containing a list of reaction, return the pathways where these reactions are involved. ex: if rxn-a in pwy-x => return, pwy-x; all rxn ids in pwy-x; all rxn ids in pwy-x FROM the list; ratio

```
usage:
    get_pwy_from_rxn.py --reaction_file=FILE --padmetRef=FILE --output=FILE

options:
    -h --help      Show help.
    --reaction_file=FILE  pathname of the file containing the reactions id, 1/line
    --padmetRef=FILE   pathname of the padmet representing the database.
    --output=FILE     pathname of the file with line = pathway id, all reactions id,_
    ↪reactions ids from reaction file, ratio. sep = "      "
```

padmet_utils.exploration.get_pwy_from_rxn.**dict_pwys_to_file**(dict_pwy, output)
Create csv file from dict_pwy. dict_pwy is obtained with extract_pwys()

Parameters

- **dict_pwy** (*dict*) – dict, k=pathway_id, v=dict: k in [total_rxn, rxn_from_list, ratio ex: {pwy-x:{‘total_rxn’:[a,b,c], rxn_from_list:[a], ratio:1/3}}]
- **output** (*str*) – path to output file

padmet_utils.exploration.get_pwy_from_rxn.**extract_pwys**(padmet, reactions)
#extract from padmet pathways containing 1-n reactions from a set of reactions ‘reactions’ Return a dict of data. dict, k=pathway_id, v=dict: k in [total_rxn, rxn_from_list, ratio ex: {pwy-x:{‘total_rxn’:[a,b,c], rxn_from_list:[a], ratio:1/3}}]

Parameters

- **padmet** (*padmet.classes.PadmetSpec*) – padmet to update
- **reactions** (*set*) – set of reactions to match with pathways

Returns dict, k=pathway_id, v=dict: k in [total_rxn, rxn_from_list, ratio ex: {pwy-x:{‘total_rxn’:[a,b,c], rxn_from_list:[a], ratio:1/3}}]

Return type dict

padmet_utils.exploration.get_pwy_from_rxn.**main**()

report_network

Description: Create reports of a padmet file.

all_pathways.tsv: header = [“dbRef_id”, “Common name”, “Number of reaction found”, “Total number of reaction”, “Ratio (Reaction found / Total)”]

all_reactions.tsv: header = [“dbRef_id”, “Common name”, “formula (with id)”, “formula (with common name)”, “in pathways”, “associated genes”]

all_metabolites.tsv: header = [“dbRef_id”, “Common name”, “Produced (p), Consumed (c), Both (cp)”]

```
usage:
    report_network.py --padmetSpec=FILE --output_dir=dir [--padmetRef=FILE] [-v]

options:
    -h --help      Show help.
```

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```
--padmetSpec=FILE      pathname of the padmet file.  
--padmetRef=FILE      pathname of the padmet file used as database  
--output_dir=dir      directory for the results.  
-v      print info.
```

```
padmet_utils.exploration.report_network.main()
```

visu_path

4.1.3 Scripts: Management

Description:

```
#TODO
```

manual_curation

Description: Update a padmetSpec by filling specific forms.

1./ Create new reaction(s) to padmet file.

- Get the template form with --template_new_rxn
- Fill the template
- set --data as path to the filled template

2./ Add reaction(s) from padmetRef or remove reactions(s).

- Get the template form with --template_add_delete_rxn
- Fill the template
- set --date as path to the filled template

Update padmetSpec and create a new padmet (new_padmet) or overwrite the input

```
usage:  
    manual_curation.py --padmetSpec=FILE --data=FILE [--padmetRef=FILE] [--  
    ↪output=FILE] [--tool=STR] [--category=STR] [-v]  
    manual_curation.py --template_new_rxn=FILE  
    manual_curation.py --template_add_delete_rxn=FILE  
  
option:  
    -h --help      Show help.  
    --padmetSpec=FILE      path to the padmet to update  
    --padmetRef=FILE      path of the padmet representing the reference database  
    --data=FILE      path to the form with data for curation  
    --output=FILE      path to the output. if None. Overwriting padmetSpec  
    --tool=STR      specification of the tool used to allow this curation: ex a tool of  
    ↪gapfilling (meneco)  
    --category=STR      specification of the category of curation: ex if a reaction is  
    ↪added based on annotation info, use 'annotation'  
    --template_new_rxn=FILE      create a form used to create new reaction, use this  
    ↪form as input for 'data' option  
    --template_add_delete_rxn=FILE      create a form used to add or delete reaction,  
    ↪use this form as input for 'data' option  
    -v      print info
```

```
padmet_utils.management.manual_curation.add_delete_rxn(data_file,      padmetSpec,
                                                       output,    padmetRef=None,
                                                       source=None, tool=None,
                                                       category='MANUAL',
                                                       verbose=False)
```

Read a data_file (form created with template_add_delete and filed), for each reaction if column ‘Action’ == ‘add’:

add the reaction from padmetRef to padmetSpec.

elif column ‘Action’ == ‘delete’: remove the reaction

Can’t add a reaction without a padmetRef !

the source ensure the traceability of the reaction, its a simple tag ex ‘pathway_XX_update’ if not given the filename of data_file will be used. if a tool was used to infer the reaction, define tool=’name_of_the_tool’

Parameters

- **data_file** (*str*) – path to file based on template_new_rxn()
- **padmetSpec** (*padmet.classes.PadmetSpec*) – padmet to update
- **padmetRef** (*padmet.classes.PadmetRef*) – padmet containing the database of reference
- **output** (*str*) – path to the new padmet file
- **source** (*str*) – tag associated to the new reactions to create and add, used for traceability
- **tool** (*str*) – The eventual tool used to infer the reactions to create and add
- **category** (*str*) – The default category of the reaction added manually is ‘MANUAL’. Must not be changed.
- **verbose** (*bool*) – if True print information

```
padmet_utils.management.manual_curation.main()
```

```
padmet_utils.management.manual_curation.rxn_creator(data_file,      padmetSpec,   output,
                                                       padmetRef=None,
                                                       source=None,    tool=None,
                                                       category='MANUAL',   verbose=False)
```

Read a data_file (form created with template_new_rxn and filed), for each reaction to create, add the reaction in padmetSpec (only if the id of the reaction is not already in padmetSpec or in padmetRef if given) the source ensure the traceability of the reaction, its a simple tag ex ‘pathway_XX_update’ if not given the filename of data_file will be used. if a tool was used to infer the reaction, define tool=’name_of_the_tool’ the Padmet of reference padmetRef can be used to check that the reaction id is not already in the database and copy information from the database for existing compounds strongly recommended to give a padmetRef.

Parameters

- **data_file** (*str*) – path to file based on template_new_rxn()
- **padmetSpec** (*padmet.classes.PadmetSpec*) – padmet to update
- **output** (*str*) – path to the new padmet file
- **source** (*str*) – tag associated to the new reactions to create and add, used for traceability
- **tool** (*str*) – The eventual tool used to infer the reactions to create and add

- **category** (*str*) – The default category of the reaction added manually is ‘MANUAL’. Must not be changed.
- **padmetRef** (*padmet.classes.PadmetRef*) – padmet containing the database of reference
- **verbose** (*bool*) – if True print information

`padmet_utils.management.manual_curation.sniff_datafile(data_file)`

Read data_file and check which kind of data input it is. A reaction_creator file contains only 2 columns. Add reaction_add_delete more than 2. Basic, need to be improved.

Parameters `data_file` (*str*) – path to file of reaction_creator or reaction_add_delete.

Returns “rxn_creator” or “add_delete_rxn”

Return type *str*

`padmet_utils.management.manual_curation.template_add_delete(output)`

Generate template file used as input of add_delete_rxn function

Parameters `output` (*str*) – path for the template rxn_add_delete to create

`padmet_utils.management.manual_curation.template_new_rxn(output)`

Generate template file used as input of rxn_creator function

Parameters `output` (*str*) – path for the template new_rxn to create

padmet_compart

Description: For a given padmet file, check and update compartment.

1./ Get all compartment with 1st usage

2./ Remove a compartment with 2nd usage. Remove all reactions acting in the given compartment

3./ change compartment id with 3rd usage

```
usage:  
  padmet_compart.py --padmet=FILE  
  padmet_compart.py --padmet=FILE --remove=STR [--output=FILE] [-v]  
  padmet_compart.py --padmet=FILE --old=STR --new=STR [--output=FILE] [-v]  
  
options:  
  -h --help      Show help.  
  --padmet=FILE  pathname of the padmet file  
  --remove=STR   compartment id to remove  
  --old=STR      compartment id to change to new id  
  --new=STR      new compartment id  
  --output=FILE  new padmet pathname, if none, overwritting the original padmet  
  -v  print info
```

`padmet_utils.management.padmet_compart.remove_compart(padmet, to_remove, verbose=False)`

Remove all reaction associated to a compound in the compartment to remove.

Parameters

- **padmet** (*padmet.classes.PadmetSpec*) – padmet to update
- **to_remove** (*str*) – compartment id to remove, if many separate compartment id by ‘,’
- **verbose** (*bool*) – if True print information

Returns New padmet after removing compartment(s)

Return type padmet.classes.PadmetSpec

```
padmet_utils.management.padmet_compart.replace_compart(padmet, old_compart,
                                                       new_compart, verbose=False)
```

Replace compartment ‘old_compart’ by ‘new_compart’.

Parameters

- **padmet** (*padmet.classes.PadmetSpec*) – padmet to update
- **old_compart** (*str*) – compartment id to replace
- **new_compart** (*str*) – new compartment id
- **verbose** (*bool*) – if True print information

Returns New padmet after replacing compartment

Return type padmet.classes.PadmetSpec

padmet_medium

Description: For a given set of compounds representing the growth medium (or seeds). Create 2 reactions for each compounds to maintain consistency of the network for flux analysis. For each compounds create:

An exchange reaction: this reaction consumes the compound in the compartment ‘C-BOUNDARY’ and produces the compound in the compartment ‘e’ extracellular

A transport reaction: this reaction consumes the compound in the compartment ‘e’ extracellular and produces the compound in the compartment ‘c’ cytosol ex: for seed ‘cpd-a’

1/ check if cpd-a in padmetSpec, if not, copy from padmetRef.

2/ create exchange reaction: ExchangeSeed_cpd-a_b: 1 cpd-a (C-BOUNDARY) \leftrightarrow 1 cpd-a (e)

3/ create transport reaction: TransportSeed_cpd-a_e: 1 cpd-a (e) \Rightarrow 1 cpd-a (c)

4/ create a new file if output not None, or overwrite padmetSpec

```
usage:
    padmet_medium.py --padmetSpec=FILE
    padmet_medium.py --padmetSpec=FILE -r [--output=FILE] [-v]
    padmet_medium.py --padmetSpec=FILE --seeds=FILE [--padmetRef=FILE] [--output=FILE] [-v]

options:
    -h --help      Show help.
    --padmetSpec=FILE    path to the padmet file to update
    --padmetRef=FILE    path to the padmet file representing to the database of reference (ex: metacyc_18.5.padmet)
    --seeds=FILE     the path to the file containing the compounds ids and the compartment.
    -r               Use to remove all medium from padmet
    -v               print info
```

```
padmet_utils.management.padmet_medium.main()
```

```
padmet_utils.management.padmet_medium.manage_medium(padmet,  
                                  new_growth_medium=None,  
                                  padmetRef=None,               ver-  
                                  bose=False)
```

Manage medium of a padmet. If new_growth_medium give, use this list of compound to define the new medium and create transport and exchange reactions. if padmetRef given, use the information from padmetRef to create the missing compound. If no new_growth_medium given: remove the current medium in the padmet.

Parameters

- **padmet** (*padmet.classes.PadmetSpec*) – padmet to update
- **new_growth_medium** (*list*) – list of compound id representing the medium
- **padmetRef** (*padmet.classes.PadmetRef*) – padmet containing the database of reference
- **verbose** (*bool*) – if True print information

Returns New padmet after updating medium

Return type *padmet.classes.PadmetSpec*

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