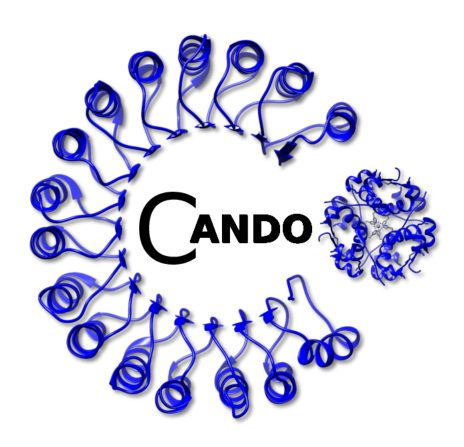
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1 CANDO

Computational Analysis of Novel Drug Opportunities

CANDO is a unique computational drug discovery, design, and repurposing platform.

2 Install

You may download the source code via the releases or cloning the git repository. However, we suggest using anaconda to install the CANDO package, as this is the easiest and quickest way to start using our platform!

The CANDO package relies on multiple "conda-forge" dependencies. Therefore, we require that you add "conda-forge" to your anaconda channels:

```
conda config --add channels conda-forge
```

Then you can install CANDO using the following command:

```
conda install -c ram-compbio cando
```

3 Test

You can test your install by running our script:

```
run_test.py
```

4 Authors

- · William Mangione
- Zackary Falls
- · James Schuler
- Matt Hudson
- · Liana Bruggemann
- · Ram Samudrala

For general questions, please contact Ram Samudrala (ram@compbio.org). For direct questions about source code for cando.py, please contact William Mangione (wmangion@buffalo.edu) or Zackary Falls (zmfalls@buffalo.edu).

5 LICENSE 3

5 LICENSE

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6 Namespace Index

6.1 Packages

Here are the packages with brief descriptions (if available):

cando 5

7 Hierarchical Index

7.1 Class Hierarchy

This inheritance list is sorted roughly, but not completely, alphabetically:

object

cando.ADR	11
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cando.Compound_pair	49
cando.Indication	52
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8 Class Index	
o oldos macx	
8.1 Class List	
Here are the classes, structs, unions and interfaces with brief descriptions:	
cando.ADR An object to represent an adverse reaction	11
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An object to represent all aspects of CANDO (compounds, indications, etc.)	matrix,
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An object to represent a protein	60

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9 File Index

9.1 File List

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10 Namespace Documentation

10.1 cando Namespace Reference

Classes

class ADR

An object to represent an adverse reaction.

• class CANDO

An object to represent all aspects of CANDO (compounds, indications, matrix, etc.)

class Compound

An object to represent a compound/drug.

class Compound_pair

An object to represent a compound/drug-pair.

· class Indication

An object to represent an indication (disease)

class Matrix

An object to represent a matrix.

class Pathway

An object to represent a pathway.

class Protein

An object to represent a protein.

Functions

def generate_matrix (v="v2.2", fp="rd_ecfp4", vect="int", dist="dice", org="nrpdb", bs="coach", c_cutoff=0.0, p_cutoff=0.0, percentile_cutoff=0.0, i_score="P", out_file=", out_path=".", nr_ligs=True, approved_only=False, lig_name=False, lib_path=", prot_← path=", ncpus=1)

Generate a matrix using our in-house protocol BANDOCK.

- def calc_scores (c, c_fps, l_fps, p_dict, dist, pscore_cutoff=0.0, cscore_cutoff=0.0, percentile_cutoff=0.0, i_score='P', nr_ligs=[], lig_name=False)
- def generate_signature (cmpd_file, fp="rd_ecfp4", vect="int", dist="dice", org="nrpdb", bs="coach", c_cutoff=0.0, p_cutoff=0.0, percentile_cutoff=0.0, i_score="P", out_file=", out_path=".", nr_ligs=True, prot_path=")

Generate an interaction signature for a query compound using our in-house protocol BANDO← CK.

def add_cmpds (cmpd_list, file_type='smi', fp="rd_ecfp4", vect="int", cmpd_dir=".", v=None)

Add new compounds to an existing CANDO Compound library, or create a new Compoun library using our in-house protocol BANDOCK.

- def cosine_dist (A)
- def tanimoto_sparse (str1, str2)

Calculate the tanimoto coefficient for a pair of sparse vectors.

def tanimoto_dense (list1, list2)

Calculate the tanimoto coefficient for a pair of dense vectors.

def get_fp_lig (fp)

Download precompiled binding site ligand fingerprints using the given fingerprint method.

def get_data (v="v2.2", org='nrpdb', fp='rd_ecfp4', vect='int')

Download CANDO v2.2+ data.

def clear_cache ()

Clear files in "data/" directory.

def get_tutorial ()

Download data for tutorial.

• def get test ()

Download data for test script.

def dl_dir (url, out, l)

Function to recursively download a directory.

• def dl file (url, out file)

Function to download a file.

def load_version (v='v2.3', protlib='nrpdb', i_score='CxP', approved_only=False, compute_distance=False, dist_metric='cosine', protein_set=", ncpus=1)

Directly load a pre-compiled version of CANDO.

10.1.1 Function Documentation

10.1.1.1 add_cmpds()

Add new compounds to an existing CANDO Compound library, or create a new Compoun library using our in-house protocol BANDOCK.

Parameters

cmpd_list	str: filepath to all input compounds
fp	str: the chemical fingerprint to use (rd_ecfp4, rd_ecfp10, etc)
vect	str: integer "int" or binary "bit" vector for fingerprint
cmpd_dir	str: ??
V	str: ??

Returns

Returns None

10.1.1.2 calc_scores()

```
10.1.1.3 clear_cache()
```

```
def cando.clear_cache ( )
```

Clear files in "data/" directory.

Returns

Returns None

10.1.1.4 cosine_dist()

```
def cando.cosine\_dist (
A )
```

10.1.1.5 dl_dir()

```
def cando.dl_dir (
    url,
    out,
    l )
```

Function to recursively download a directory.

Prints the name of the directory and a progress bar.

Parameters

url	str: URL of the dir to be downloaded	
out	str: Path to where the dir will be downloaded	
1	list: List of files in dir to be downloaded	

Returns

Returns None

10.1.1.6 dl_file()

Function to download a file.

Prints the name of the file and a progress bar.

Parameters

url	str: URL of the file to be downloaded
out_file	str: File path to where the file will be downloaded

Returns

Returns None

10.1.1.7 generate_matrix()

```
def cando.generate_matrix (
            v = "v2.2",
            fp = "rd_ecfp4",
            vect = "int",
            dist = "dice",
            org = "nrpdb",
            bs = "coach",
            c\_cutoff = 0.0,
            p\_cutoff = 0.0,
            percentile_cutoff = 0.0,
            i_score = "P",
            out_file = '',
            out_path = ".",
            nr_ligs = True,
            approved_only = False,
            lig_name = False,
            lib_path = '',
            prot_path = '',
            ncpus = 1)
```

Generate a matrix using our in-house protocol BANDOCK.

V	str: version to use (supports v2.2 - v2.5)
fp	str: the chemical fingerprint to use (rd_ecfp4, rd_ecfp10, etc)
vect	str: integer "int" or binary "bit" vector for fingerprint
dist	str: use Sorenson-Dice "dice" for vect="int" and Tanimoto "tani" for vect="bit"
org	str: protein library to use ('nrpdb' or 'homo_sapien')
bs	str: the method to use, just use "coach"
c_cutoff	float: minimum Cscore (Tanimoto/Dice similarity score) to consider for scoring
p_cutoff	float: minimum Pscore (binding site score from COACH) to consider for scoring
percentile_cutoff	float: ile cutoff for fingerprint similarity scores in 'dC' scoring protocols
i_score	str: the scoring protocol to use ('P', 'C', 'dC', 'CxP', dCxP')
out_file	str: filename of the output matrix
out_path	str: path to the output matrix
nr_ligs	bool: use only the non-redundant set of ligands for 'dC' scoring protocols (recommended)
approved_only	bool: use only approved drugs to create the matrix
lig_name	bool: output the ligand chosen for the compound-protein interaction score instead of the score
lib_path	str: specify a local compound fingerprint set for custom analyses
prot_path	str: specify a local protein library for custom analyses
ncpus	int: number of cores to run on

Returns

Returns None

10.1.1.8 generate_signature()

```
i_score = "P",
out_file = '',
out_path = ".",
nr_ligs = True,
prot_path = '')
```

Generate an interaction signature for a query compound using our in-house protocol BANDOCK.

Note: the parameters for this function MUST MATCH the parameters used to generate the matrix in use. Otherwise, the scores will be incompatible.

Parameters

cmpd_file	str: filepath to an input mol file
fp	str: the chemical fingerprint to use (rd_ecfp4, rd_ecfp10, etc)
vect	str: integer "int" or binary "bit" vector for fingerprint
dist	str: use Sorenson-Dice "dice" for vect="int" and Tanimoto "tani" for vect="bit"
org	str: protein library to use ('nrpdb' or 'homo_sapien')
bs	str: the method to use, just use "coach"
c_cutoff	float: minimum Cscore (Tanimoto/Dice similarity score) to consider for
	scoring
p_cutoff	float: minimum Pscore (binding site score from COACH) to consider for
	scoring
percentile_cutoff	float: ile cutoff for fingerprint similarity scores in 'dC' scoring protocols
i_score	str: the scoring protocol to use ('P', 'C', 'dC', 'CxP', dCxP')
out_file	str: filename of the output signature
out_path	str: path to the output signature
nr_ligs	bool: use only the non-redundant set of ligands for 'dC' scoring protocols
	(recommended)
prot_path	str: specify a local protein library for custom analyses

Returns

Returns None

10.1.1.9 get_data()

```
def cando.get_data ( v = "v2.2", \\ org = 'nrpdb', \\ fp = 'rd_ecfp4', \\ vect = 'int')
```

Download CANDO v2.2+ data.

V	str: version to use (supports v2.2 - v2.5)
org	str: protein library to use ('nrpdb' or 'homo_sapien')
fp	str: the chemical fingerprint to use (rd_ecfp4, rd_ecfp10, etc)
vect	str: integer "int" or binary "bit" vector for fingerprint

Returns

Returns None

10.1.1.10 get_fp_lig()

```
def cando.get_fp_lig (
          fp )
```

Download precompiled binding site ligand fingerprints using the given fingerprint method.

Parameters

fp str: Fingerprinting method used to compile each binding site ligand fingerprint

Returns

Returns None

10.1.1.11 get_test()

```
def cando.get_test ( )
```

Download data for test script.

Returns

Returns None

10.1.1.12 get_tutorial()

```
def cando.get_tutorial ( )
```

Download data for tutorial.

Returns

Returns None

10.1.1.13 load_version()

```
def cando.load_version (
    v = 'v2.3',
    protlib = 'nrpdb',
    i_score = 'CxP',
    approved_only = False,
    compute_distance = False,
    dist_metric = 'cosine',
    protein_set = '',
    ncpus = 1 )
```

Directly load a pre-compiled version of CANDO.

Parameters

V	str: version to use (supports v2.2 - v2.5)
protlib	str: protein library to use ('nrpdb' or 'homo_sapien')
i_score	str: the scoring protocol to use ('P', 'C', 'dC', 'CxP', dCxP')
approved_only	bool: use only approved drugs to create the matrix
compute_distance	bool: compute distance between compounds for specified matrix
dist_metric	str: the distance metric to use if compute_distance=True ('cosine',
	'rmsd', etc)
protein_set	str: path to a file containing a subset of proteins of interest
ncpus	int: number of cores to run on

Returns

Returns **CANDO** object

10.1.1.14 tanimoto_dense()

Calculate the tanimoto coefficient for a pair of dense vectors.

Parameters

list1	list: List of positions that have a 1 in first compound fingerprint
list2	list: List of positions that have a 1 in second compound fingerprint

Returns

Returns float

10.1.1.15 tanimoto_sparse()

```
def cando.tanimoto_sparse ( str1, str2 )
```

Calculate the tanimoto coefficient for a pair of sparse vectors.

Parameters

str1	str: String of 1s and 0s representing the first compound fingerprint
str2	str: String of 1s and 0s representing the second compound fingerprint

Returns

Returns float

11 Class Documentation

11.1 cando.ADR Class Reference

An object to represent an adverse reaction.

Inheritance diagram for cando.ADR:



Public Member Functions

def __init__ (self, id_, name)

Public Attributes

id

str: Identification for the given ADR

• name

str: Name of the given ADR

• compounds

list: Compound objects associated with the given ADR

• compound_pairs

11.1.1 Detailed Description

An object to represent an adverse reaction.

11.1.2 Constructor & Destructor Documentation

11.1.3 Member Data Documentation

11.1.3.1 compound_pairs

cando.ADR.compound_pairs

11.1.3.2 compounds

cando.ADR.compounds

list: Compound objects associated with the given ADR

List: Compound object pairs (tuples) associated with the given ADR.

11.1.3.3 id_

cando.ADR.id_

str: Identification for the given ADR

11.1.3.4 name

cando.ADR.name

str: Name of the given ADR

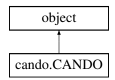
The documentation for this class was generated from the following file:

cando.py

11.2 cando.CANDO Class Reference

An object to represent all aspects of CANDO (compounds, indications, matrix, etc.)

Inheritance diagram for cando.CANDO:



Public Member Functions

- def __init__ (self, c_map, i_map, matrix=", compound_set='all', compute_distance=False, save_dists=", read_dists=", pathways=", pathway_quantifier='max', indication_\circ
 pathways=", indication_proteins=", similarity=False, dist_metric='rmsd', protein_set=",
 rm_zeros=False, rm_compounds=", ddi_compounds=", ddi_adrs=", adr_map=", protein\circ
 _distance=False, protein_map=", ncpus=1)
- def search_compound (self, name, n=5)

Print closest Compound names/IDs for input search str.

def get_compound (self, cmpd_id, quiet=False)

Get Compound object from Compound id or fuzzy match to Compound name.

def get_compound_pair (self, ids)

Get Compound_pair object from Compound_pair id.

def get_protein (self, protein_id)

Get Protein object from Protein id.

def get_indication (self, ind_id)

Get Indication object from Indication id.

def get_pathway (self, id_)

Get Pathway object from Pathway id.

def get adr (self, id)

Get ADR (adverse drug reaction) from ADR id.

def search_indication (self, name, n=5)

Print closest MeSH IDs for Indication name.

def top_targets (self, cmpd, n=10, negative=False, save_file=")

Get the top scoring protein targets for a given compound.

def common_targets (self, cmpds_file, n=10, negative=False, save_file=")

Get the consensus top scoring protein targets for a set of compounds.

- def virtual_screen (self, protein, n=10, negative=False, compound_set='all', save_file=")
 Get the top scoring compounds for a given protein.
- def uniprot set index (self, prots)

Gather proteins from input matrix that map to UniProt IDs from 'protein_set=' param.

def generate_similar_sigs (self, cmpd, sort=False, proteins=[], aux=False)

For a given compound, generate the similar compounds using distance of sigs.

• def generate_similar_sigs_cp (self, cmpd_pair, sort=False, proteins=[], aux=False)

For a given compound pair, generate the similar compound pairs using distance of sigs.

def generate_some_similar_sigs (self, cmpds, sort=False, proteins=[], aux=False)

For a given list of compounds, generate the similar compounds based on dist of sigs This is pathways/genes for all intents and purposes.

def quantify_pathways (self, indication=None)

Uses the pathway quantifier defined in the CANDO instantiation to make a pathway signature for all pathways in the input file (NOTE: does not compute distances)

def results_analysed (self, f, metrics, effect_type)

Creates the results analysed named file for the benchmarking and computes final avg indication accuracies.

def canbenchmark (self, file_name, indications=[], continuous=False, bottom=False, rank-ing='standard', adrs=False)

Benchmarks the platform based on compound similarity of those approved for the same diseases.

def canbenchmark_associated (self, file_name, indications=[], continuous=False, rank-ing='standard')

Benchmark only the compounds in the indication mapping, aka get rid of "noisy" compounds.

def canbenchmark_bottom (self, file_name, indications=[], ranking='standard')

Benchmark the reverse ranking of similar compounds as a control.

def canbenchmark_ndcg (self, file_name)

Benchmark using the normalized discounted cumulative gain metric.

def canbenchmark_cluster (self, n_clusters=5)

Benchmark using k-means clustering.

- def compounds analysed (self, f, metrics)
- def canbenchmark_compounds (self, file_name, adrs=[], continuous=False, bot-tom=False, ranking='standard')

Benchmarks the platform based on compound similarity of those known to interact with other compounds.

def canbenchmark_ddi (self, file_name, adrs=[], continuous=False, bottom=False, rank-ing='standard')

Benchmarks the platform based on compound pairs known to cause ADRs.

 def ml (self, method='rf', effect=None, benchmark=False, adrs=False, predict=[], threshold=0.5, negative='random', seed=42, out=")

Create an ML classifier for a specified indication to make drug-disease predictions or all inds for benchmarking.

- def raw results roc (self, rr files, labels, save='roc-raw results.pdf')
- def canpredict_denovo (self, method='count', threshold=0.0, topX=10, ind_id=None, proteins=None, cmpd_set='all', save=")

This function is used for predicting putative therapeutics for an indication of interest by summing/counting the number of interactions above a certain input interaction threshold for all proteins or a specified subset of proteins.

def canpredict_compounds (self, ind_id, n=10, topX=10, keep_associated=False, cmpd
 set='all', save=")

This function is used for predicting putative therapeutics for an indication of interest using a homology-based approach.

def canpredict_indications (self, cmpd, n=10, topX=10, save=")

This function is the inverse of canpredict compounds.

def canpredict_adr (self, cmpd, n=10, topX=10, save=")

This function is the inverse of canpredict_compounds.

def canpredict_ddi_cmpds (self, cmpd, n=10, topX=10, save=")

Input a compound of interest cando_cmpd and the most similar compounds to it will be computed and outputted as potential drug-drug-interactions.

def canpredict_ddi_adrs (self, cmpd_pair, n=10, topX=10, save=")

Similarly to canpredict_adrs(), input a compound pair of interest (cmpd_pair) and the most similar compound pairs to it will be computed.

• def similar_compounds (self, cmpd, n=10)

Computes and prints the top n most similar compounds to an input Compound object cando cmpd or input novel signature new sig.

def add_cmpd (self, new_sig, new_name=")

Add a new Compound object to the platform.

• def sigs (self, rm)

Return a list of all signatures, rm is a list of compound ids you do not want in the list.

def save_dists_to_file (self, f)

Write calculated distances of all compounds to all compounds to file.

def fusion (self, cando objs, out file=", method='sum')

This function re-ranks the compounds according to the desired comparison specified by 'method' -> currently supports 'min', 'avg', 'mult', and 'sum'.

def normalize (self)

Normalize the distance scores to between [0,1].

def __str__ (self)

Print stats about the CANDO object.

Public Attributes

c_map

str: File path to the compound mapping file (relative or absolute)

i_map

str: File path to the indication mapping file (relative or absolute)

· matrix

str: File path to the cando matrix file (relative or absolute)

· compound_set

str or List str: what compounds to use, such as all, approved, experimental, etc

protein_set

str: File path to protein subset file (relative or absolute)

pathways

str: File path to pathway file

- accuracies
- compute distance

bool: Calculate the distance for each Compound against all other Compounds using chosen distance metric

· protein_distance

bool: Calculate the distance for each Protein against all other Proteins using chosen distance metric

- · clusters
- rm_zeros

bool: Remove Compounds with all-zero signatures from CANDO object

• rm_compounds

list: Compounds to remove from the CANDO object

- rm_cmpds
- · save dists

bool: Write the calculated distances to file after computation (set compute_distances=True)

read_dists

str: File path to pre-computed distance matrix

similarity

bool: Use similarity instead of distance

· dist metric

str: Distance metric to be used for computing Compound-Compound distances

ncpus

int: Number of CPUs used for parallelization

· pathway_quantifier

str: Method used to quantify a all Pathways

indication pathways

str: File path to Indication-Pathway association file

indication_proteins

str: File path to Indication-Protein association file

adr_map

str: File path to ADR mapping file

protein_map

str: File path to Protein metadata mapping file

ddi_compounds

str: File path to Drug-drug mapping file

- · ddi_adrs
- proteins

List: Protein objects in the platform.

- protein_id_to_index
- compounds

List: Compound objects in the platform.

- compound_ids
- compound_pairs

List: Compound_pair objects in the platform.

- · compound pair ids
- indications

List: Indication objects in the platform.

• indication_ids

· adrs

List: ADR objects in the platform.

- adr_ids
- · short_matrix_path
- · short_read_dists
- · short protein set
- cmpd_set
- data_name

11.2.2.1 __init__()

11.2.1 Detailed Description

An object to represent all aspects of CANDO (compounds, indications, matrix, etc.)

To instantiate you need the compound mapping (c_map), an indication mapping file (i_map), and typically and a compound-protein matrix (matrix=) or or precomputed compound-compound distance matrix (read_rmsds=), but those are optional.

11.2.2 Constructor & Destructor Documentation

```
def cando.CANDO.__init__ (
            self,
            c_{map}
            i_map,
            matrix = '',
            compound_set = 'all',
            compute_distance = False,
            save_dists = '',
            read_dists = '',
            pathways = '',
            pathway_quantifier = 'max',
            indication_pathways = '',
            indication_proteins = '',
            similarity = False,
            dist_metric = 'rmsd',
            protein_set = '',
            rm_zeros = False,
            rm_compounds = '',
            ddi compounds = '',
            ddi_adrs = '',
            adr_map = '',
            protein_distance = False,
```

protein_map = '',

ncpus = 1)

11.2.3 Member Function Documentation

Print stats about the CANDO object.

11.2.3.2 add_cmpd()

Add a new Compound object to the platform.

Parameters

new_sig	str: Path to the tab-separated interaction scores
new_name	str: Name for the new Compound

Returns

Returns None

11.2.3.3 canbenchmark()

Benchmarks the platform based on compound similarity of those approved for the same diseases.

file_name	str: Name to be used for the various results files (e.g. file_name=test ->
	summary_test.tsv)
indications	list or str: List of Indication ids to be benchmarked, otherwise all will be used.
continuous	bool: Use the percentile of distances from the similarity matrix as the
	benchmarking cutoffs
bottom	bool: Reverse the ranking (descending) for the benchmark
ranking	str: What ranking method to use for the compounds. This really only affects
	ties. (standard, modified, and ordinal)
adrs	bool: ADRs are used as the Compounds' phenotypic effects instead of
	Indications

Returns

Returns None

11.2.3.4 canbenchmark_associated()

Benchmark only the compounds in the indication mapping, aka get rid of "noisy" compounds.

This function returns the filtered CANDO object in the event that you want to explore further.

Parameters

file_name	str: Name to be used for the variosu results files (e.g. file_name=test -> summary_test.tsv)
indications	list: List of Indication ids to be used for this benchmark, otherwise all will be used.
continuous	bool: Use the percentile of distances from the similarity matrix as the benchmarking cutoffs
ranking	str: What ranking method to use for the compounds. This really only affects ties. (standard, modified, and ordinal)

Returns

Returns None

11.2.3.5 canbenchmark_bottom()

Benchmark the reverse ranking of similar compounds as a control.

Parameters

file_name	str: Name to be used for the variosu results files (e.g. file_name=test -> summary_test.tsv)
indications	list: List of Indication ids to be used for this benchmark, otherwise all will be used.
ranking	str: What ranking method to use for the compounds. This really only affects ties. (standard, modified, and ordinal)

Returns

Returns None

11.2.3.6 canbenchmark_cluster()

```
def cando.CANDO.canbenchmark_cluster ( self, n\_clusters = 5 )
```

Benchmark using k-means clustering.

Parameters

n_clusters int: Number of clusters for k-means

Returns

Returns None

11.2.3.7 canbenchmark_compounds()

Benchmarks the platform based on compound similarity of those known to interact with other compounds.

Parameters

file_name	str: Name to be used for the various results files (e.g. file_name=test -> summary test.tsv)
	,
adrs	list: List of ADR ids to be used for this benchmark, otherwise all will be used.
continuous	bool: Use the percentile of distances from the similarity matrix as the cutoffs for
	benchmarking
bottom	bool: Reverse the ranking (descending) for the benchmark
ranking	str: What ranking method to use for the compounds. This really only affects
	ties. (standard, modified, and ordinal)

Returns

Returns None

11.2.3.8 canbenchmark_ddi()

Benchmarks the platform based on compound pairs known to cause ADRs.

file_name	str: Name to be used for the results files (file_name=test ->
	summary_test-ddi_adr.tsv)
continuous	bool: Use the percentile of distances from the similarity matrix as the cutoffs for
	benchmarking
bottom	bool: Reverse the ranking (descending) for the benchmark
ranking	str: What ranking method to use for the compounds. This really only affects
	ties. (standard, modified, and ordinal)

Returns

Returns None

11.2.3.9 canbenchmark_ndcg()

Benchmark using the normalized discounted cumulative gain metric.

Parameters

file_name	str: Name to be used for the results files (file_name=test ->	
	summary_ndcg-test.tsv)	

Returns

Returns None

11.2.3.10 canpredict_adr()

This function is the inverse of canpredict_compounds.

Input a compound of interest cando_cmpd (or a novel protein signature of interest new_sig) and the most similar compounds to it will be computed. The ADRs associated with the top n most similar compounds to the query compound will be examined to see if any are repeatedly enriched.

Parameters

cmpd	Compound: Compound object to be used
n	int: top number of similar Compounds to be used for prediction
topX	int: top number of predicted Indications to be printed

Returns

Returns None

11.2.3.11 canpredict compounds()

This function is used for predicting putative therapeutics for an indication of interest using a homology-based approach.

Input an ind_id id and for each of the associated compounds, it will generate the similar compounds (based on distance) and add them to a dictionary with a value of how many times it shows up (enrichment). If a compound not approved for the indication of interest keeps showing up, that means it is similar in signature to the drugs that are ALREADY approved for the indication, so it may be a target for repurposing. Control how many similar compounds to consider with the argument 'n'. In the output, 'score1' refers to the number of times the compound shows up in the top 'n' drugs associated with the indication and 'score2' is the average of the ranks for 'score1' (note: 'score2' <= 'n').

Parameters

ind_id	str: Indication id
n	int: top number of similar Compounds to be used for each Compound
	associated with the given Indication

topX	int: top number of predicted Compounds to be printed
keep_associated	bool: Print Compounds that are already approved/associated for the
	Indication
cmpd_set	str: specify the compound set to use ('all', 'approved', or 'other')
save	str: name of a file to save results

Returns

Returns None

11.2.3.12 canpredict_ddi_adrs()

Similarly to canpredict_adrs(), input a compound pair of interest (cmpd_pair) and the most similar compound pairs to it will be computed.

The ADRs associated with the top n most similar compound pairs to the query pair will be examined to see if any are repeatedly enriched.

Parameters

cmpd_pair	Compound_pair: Compound_pair object to be used
n	int: top number of similar Compounds to be used for prediction
topX	int: top number of predicted Indications to be printed

Returns

Returns None

11.2.3.13 canpredict_ddi_cmpds()

Input a compound of interest cando_cmpd and the most similar compounds to it will be computed and outputted as potential drug-drug-interactions.

Parameters

cmpd	Compound: Compound object to be used
n	int: top number of similar Compounds to be used for prediction
topX	int: top number of predicted Drug-drug Interactions to be printed

Returns

Returns None

11.2.3.14 canpredict_denovo()

```
def cando.CANDO.canpredict_denovo (
    self,
    method = 'count',
    threshold = 0.0,
    topX = 10,
    ind_id = None,
    proteins = None,
    cmpd_set = 'all',
    save = '')
```

This function is used for predicting putative therapeutics for an indication of interest by summing/counting the number of interactions above a certain input interaction threshold for all proteins or a specified subset of proteins.

An indication can be specified to mark drugs associated with that indication in the output. The threshold will vary based on the values of the input matrix. Method can either be 'count' (score1), which ranks compounds based on the number of interactions above the threshold, or 'sum' (score2), which ranks the compounds based on the highest total sum for interaction scores above the threshold (these two are highly correlated but can differ for larger sets of proteins or

lower thresholds). A third option is 'targets', which inspects and outputs the top protein interactions on an individual basis without summing/counting per drug (the output format differs from the other two options). If indication_proteins flag is used for the CANDO object instantiation, the proteins associated with the input indication will automatically be used. Otherwise, the 'proteins=' input can be used. The output can be saved to a file specified by 'save='. If ind_id is used, compounds associated with the indication will be included and marked in the output for comparison.

Parameters

method	str: 'sum', 'count', or 'targets'	
threshold	thold float: a interaction score cutoff to use (ignores values for sum/count less than threshold)	
topX	int: top number of predicted Compounds to be printed/saved	
ind_id	str: an indication id for marking drug output/ specifying protein set	
proteins	List str: list of protein IDs from the matrix to use for the sum/count	
cmpd_set	str: specify the compound set to use ('all', 'approved', or 'other')	
save	str: name of a file to save results	

Returns

Returns None

11.2.3.15 canpredict_indications()

This function is the inverse of canpredict_compounds.

Input a compound of interest cando_cmpd (or a novel protein signature of interest new_sig) and the most similar compounds to it will be computed. The indications associated with the top n most similar compounds to the query compound will be examined to see if any are repeatedly enriched.

Parameters

cmpd	Compound: Compound object to be used
n	int: top number of similar Compounds to be used for prediction
topX	int: top number of predicted Indications to be printed

Returns

Returns None

11.2.3.16 common_targets()

Get the consensus top scoring protein targets for a set of compounds.

Parameters

cmpds_file str: File containing a list of Compound IDs for which to search common targe		
n	int: number of top targets to print/return	
negative	int: if the interaction scores are negative (stronger) energies	
save_file	str: save results to file name	

Returns

Returns list: list of tuples (protein id_, score)

11.2.3.17 compounds_analysed()

11.2.3.18 fusion()

This function re-ranks the compounds according to the desired comparison specified by 'method' -> currently supports 'min', 'avg', 'mult', and 'sum'.

cando_objs	list: List of CANDO objects
out_file	str: Path to where the result will be written
method	str: Method of fusion to be used (e.g., sum, mult, etc.)

Returns

Returns **CANDO** object

11.2.3.19 generate_similar_sigs()

For a given compound, generate the similar compounds using distance of sigs.

Parameters

cmpd	object: Compound object
sort	bool: Sort the list of similar compounds
proteins	list: Protein objects to identify a subset of the Compound signature
aux	bool: Use an auxiliary signature (default: False)

Returns

Returns list: Similar Compounds to the given Compound

11.2.3.20 generate_similar_sigs_cp()

For a given compound pair, generate the similar compound pairs using distance of sigs.

cmpd_pair	object: Compound_pair object
sort	bool: Sort the list of similar compounds
proteins	list: Protein objects to identify a subset of the Compound signature
aux bool: Use an auxiliary signature (default: False)	

Returns

Returns list: Similar Compounds to the given Compound

11.2.3.21 generate_some_similar_sigs()

For a given list of compounds, generate the similar compounds based on dist of sigs This is pathways/genes for all intents and purposes.

Parameters

cmpds	list: Compound objects
sort	bool: Sort similar compounds for each Compound
proteins	list: Protein objects to identify a subset of the Compound signature
aux	bool: Use an auxiliary signature (default: False)

Returns

Returns list: Similar Compounds to the given Compound

11.2.3.22 get_adr()

Get ADR (adverse drug reaction) from ADR id.

```
id↔ str: ADR id
```

Returns

Returns object: ADR object

11.2.3.23 get_compound()

Get Compound object from Compound id or fuzzy match to Compound name.

Parameters

cmpd⇔	int or str: Compound id or Compound name
_id	

Returns

Returns object: Compound object or None if no exact match is found

11.2.3.24 get_compound_pair()

Get Compound_pair object from Compound_pair id.

Parameters

```
id← int: Compound_pair id _
```

Returns

Returns object: Compound_pair object

11.2.3.25 get_indication()

Get Indication object from Indication id.

Parameters

ind⇔	str: Indication id
_id	

Returns

Returns object: Indication object

11.2.3.26 get_pathway()

```
def cando.CANDO.get_pathway ( self, \\ id\_ \ )
```

Get Pathway object from Pathway id.

Parameters

```
id← str: Pathway id _←
```

Returns

Returns object: Pathway object

11.2.3.27 get_protein()

Get Protein object from Protein id.

Parameters

```
protein← str: Protein name
```

Returns

Returns object: Protein object

11.2.3.28 ml()

```
def cando.CANDO.ml (
    self,
    method = 'rf',
    effect = None,
    benchmark = False,
    adrs = False,
    predict = [],
    threshold = 0.5,
    negative = 'random',
    seed = 42,
    out = '')
```

Create an ML classifier for a specified indication to make drug-disease predictions or all inds for benchmarking.

Parameters

method str: type of machine learning algorithm to use ('rf' or 'log')

Parameters

effect	Indication or ADR: provide a specific Indication or ADR object to train a
	classifer
benchmark	bool: benchmark the ML pipeline by training a classifier with LOOCV for each
	Indication or ADR
adrs	bool: if the models are trained with ADRs instead of Indications
predict	list: provide a list of Compound objects to classify with the model (only used in
	combination with effect=Indication/ADR object)
threshold	float: decision threshold for positive vs negative classification
negative	str: choose random negative samples (default) or 'inverse' for most opposite
	signatures
seed	int: choose a seed for reproducibility
out	str: file name extension for the output of benchmark (note: must have
	benchmark=True)

Returns

Returns None

11.2.3.29 normalize()

```
\begin{tabular}{ll} $\operatorname{def}$ cando. CANDO. normalize ( \\ $\operatorname{\it self}$ ) \end{tabular}
```

Normalize the distance scores to between [0,1].

Simply divides all scores by the largest distance between any two compounds.

Returns

Returns None

11.2.3.30 quantify_pathways()

Uses the pathway quantifier defined in the CANDO instantiation to make a pathway signature for all pathways in the input file (NOTE: does not compute distances)

Parameters

ion object: Indication obje

Returns

Returns None

11.2.3.31 raw_results_roc()

11.2.3.32 results_analysed()

Creates the results analysed named file for the benchmarking and computes final avg indication accuracies.

Parameters

f	str: File path for results analysed named	
metrics	list: Cutoffs used for the benchmarking protocol	
effect_type	str: Defines the effect as either an Indication (disease) or ADR (adverse reaction)	

Returns

Returns dct: dict of accuracies at each cutoff

11.2.3.33 save_dists_to_file()

Write calculated distances of all compounds to all compounds to file.

Parameters

```
f | File name to save distances
```

11.2.3.34 search_compound()

Print closest Compound names/IDs for input search str.

Parameters

name	str: Compound name
n	int: Number of outputted compounds

Returns

Returns None

11.2.3.35 search_indication()

```
def cando.CANDO.search_indication (
self,
name,
n = 5)
```

Print closest MeSH IDs for Indication name.

Parameters

name	str: Indication name
n	int: Number of outputted indications

Returns

Returns None

11.2.3.36 sigs()

Return a list of all signatures, rm is a list of compound ids you do not want in the list.

Parameters

rm list: List of compound ids to remove from list of signatures

Returns

list: List of all signatures

11.2.3.37 similar_compounds()

```
def cando.CANDO.similar_compounds ( self, cmpd, n = 10 )
```

Computes and prints the top n most similar compounds to an input Compound object cando_← cmpd or input novel signature new_sig.

Parameters

cmpd	Compound: Compound object
n	int: top number of similar Compounds to be used for prediction

Returns

Returns None

11.2.3.38 top_targets()

Get the top scoring protein targets for a given compound.

Parameters

cmpd	Compound or int: Compound object or int id_ for which to print targets
n	int: number of top targets to print/return
negative	int: if the interaction scores are negative (stronger) energies
save_file	str: output file for results

Returns

Returns list: list of tuples (protein id_, score)

11.2.3.39 uniprot_set_index()

Gather proteins from input matrix that map to UniProt IDs from 'protein_set=' param.

Parameters

prots list: UniProt IDs (str)

Returns

Returns list: Protein chains (str) matching input UniProt IDs

11.2.3.40 virtual_screen()

Get the top scoring compounds for a given protein.

Parameters

protein	Protein int or str: Protein (object, int index, or str id_) of which to screen for top scores
n	int: number of top compounds to print/return
negative	int: if the interaction scores are negative (stronger) energies
compound_set	str: use all Compounds ('all') or only approved Compounds ('approved')
save_file	str: save results to file name

Returns

Returns None

11.2.4 Member Data Documentation

11.2.4.1 accuracies

```
cando.CANDO.accuracies
```

11.2.4.2 adr_ids

cando.CANDO.adr_ids

11.2.4.3 adr_map

cando.CANDO.adr_map

str: File path to ADR mapping file

11.2.4.4 adrs

cando.CANDO.adrs

List: ADR objects in the platform.

11.2.4.5 c_map

cando.CANDO.c_map

str: File path to the compound mapping file (relative or absolute)

11.2.4.6 clusters

cando.CANDO.clusters

11.2.4.7 cmpd_set

cando.CANDO.cmpd_set

11.2.4.8 compound_ids

cando.CANDO.compound_ids

11.2.4.9 compound_pair_ids

cando.CANDO.compound_pair_ids

11.2.4.10 compound_pairs

cando.CANDO.compound_pairs

List: Compound_pair objects in the platform.

11.2.4.11 compound_set

cando.CANDO.compound_set

str or List str: what compounds to use, such as all, approved, experimental, etc

11.2.4.12 compounds

cando.CANDO.compounds

List: Compound objects in the platform.

11.2.4.13 compute_distance

cando.CANDO.compute_distance

bool: Calculate the distance for each Compound against all other Compounds using chosen distance metric

11.2.4.14 data_name

cando.CANDO.data_name

11.2.4.15 ddi_adrs

cando.CANDO.ddi_adrs

11.2.4.16 ddi_compounds

cando.CANDO.ddi_compounds

str: File path to Drug-drug mapping file

str: File path to Drug-Drug-ADE mapping file

11.2.4.17 dist_metric

cando.CANDO.dist_metric

str: Distance metric to be used for computing Compound-Compound distances

11.2.4.18 i_map

cando.CANDO.i_map

str: File path to the indication mapping file (relative or absolute)

11.2.4.19 indication_ids

cando.CANDO.indication_ids

11.2.4.20 indication_pathways

cando.CANDO.indication_pathways

str: File path to Indication-Pathway association file

11.2.4.21 indication_proteins

cando.CANDO.indication_proteins

str: File path to Indication-Protein association file

11.2.4.22 indications

cando.CANDO.indications

List: Indication objects in the platform.

11.2.4.23 matrix

cando.CANDO.matrix

str: File path to the cando matrix file (relative or absolute)

11.2.4.24 ncpus

cando.CANDO.ncpus

int: Number of CPUs used for parallelization

11.2.4.25 pathway_quantifier

 $\verb| cando.CANDO.pathway_quantifier| \\$

str: Method used to quantify a all Pathways

11.2.4.26 pathways

cando.CANDO.pathways

str: File path to pathway file

11.2.4.27 protein_distance

```
cando.CANDO.protein_distance
```

bool: Calculate the distance for each Protein against all other Proteins using chosen distance metric

11.2.4.28 protein_id_to_index

```
cando.CANDO.protein_id_to_index
```

11.2.4.29 protein_map

```
cando.CANDO.protein_map
```

str: File path to Protein metadata mapping file

11.2.4.30 protein_set

```
cando.CANDO.protein_set
```

str: File path to protein subset file (relative or absolute)

11.2.4.31 proteins

```
cando.CANDO.proteins
```

List: Protein objects in the platform.

11.2.4.32 read_dists

```
cando.CANDO.read_dists
```

str: File path to pre-computed distance matrix

11.2.4.33 rm_cmpds

cando.CANDO.rm_cmpds

11.2.4.34 rm_compounds

 $\verb|cando.CANDO.rm_compounds||$

list: Compounds to remove from the CANDO object

11.2.4.35 rm_zeros

cando.CANDO.rm_zeros

bool: Remove Compounds with all-zero signatures from CANDO object

11.2.4.36 save_dists

cando.CANDO.save_dists

bool: Write the calculated distances to file after computation (set compute_distances=True)

11.2.4.37 short_matrix_path

cando.CANDO.short_matrix_path

11.2.4.38 short_protein_set

cando.CANDO.short_protein_set

11.2.4.39 short_read_dists

```
cando.CANDO.short_read_dists
```

11.2.4.40 similarity

```
cando.CANDO.similarity
```

bool: Use similarity instead of distance

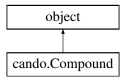
The documentation for this class was generated from the following file:

· cando.py

11.3 cando.Compound Class Reference

An object to represent a compound/drug.

Inheritance diagram for cando. Compound:



Public Member Functions

- def __init__ (self, name, id_, index, status='N/A')
- def add_indication (self, ind)

Add an Indication to the list of Indications associated to this Compound.

Public Attributes

name

str: Name of the Compound (e.g., 'caffeine')

id

int: CANDO id from mapping file (e.g., 1, 10, 100, ...)

index

int: The order in which the Compound appears in the mapping file (e.g, 1, 2, 3, ...)

status

str: The clinical trial status of the compound from DrugBank ('approved' or 'other')

sig

list: Signature is essentially a column of the Matrix

• aux_sig

list: Potentially temporary signature for things like pathways, where "c.sig" needs to be preserved

· indications

list: This is every indication the Compound is associated with from the mapping file

similar

list: This is the ranked list of compounds with the most similar interaction signatures

similar_computed

bool: Have the distances of all Compounds to the given Compound been computed?

· similar_sorted

bool: Have the most similar Compounds to the given Compound been sorted?

cluster_id

int: The cluster id this Compound was assigned from clustering method

• adrs

list: List of ADRs associated with this Compound

alt_ids

dict: dict of other ids inputted with compound mapping

metabolites

list: List of all metabolites from the compound

is_metabolite

bool: bool if the drug is a metabolite itself

parent

Compound: Compound object to which this compound is a metabolite.

compounds

List Compound: Compound objects to which this compound is associated.

11.3.1 Detailed Description

An object to represent a compound/drug.

11.3.2 Constructor & Destructor Documentation

11.3.3 Member Function Documentation

11.3.3.1 add_indication()

Add an Indication to the list of Indications associated to this Compound.

Parameters

ind object: Indication object to add

11.3.4 Member Data Documentation

11.3.4.1 adrs

cando.Compound.adrs

list: List of ADRs associated with this Compound

```
11.3.4.2 alt_ids
```

```
{\tt cando.Compound.alt\_ids}
```

dict: dict of other ids inputted with compound mapping

```
11.3.4.3 aux_sig
```

```
cando.Compound.aux_sig
```

list: Potentially temporary signature for things like pathways, where "c.sig" needs to be preserved

11.3.4.4 cluster_id

```
cando.Compound.cluster_id
```

int: The cluster id this Compound was assigned from clustering method

11.3.4.5 compounds

cando.Compound.compounds

List Compound: Compound objects to which this compound is associated.

11.3.4.6 id_

cando.Compound.id_

int: CANDO id from mapping file (e.g., 1, 10, 100, ...)

11.3.4.7 index

cando.Compound.index

int: The order in which the Compound appears in the mapping file (e.g, 1, 2, 3, ...)

11.3.4.8 indications

cando.Compound.indications

list: This is every indication the Compound is associated with from the mapping file

11.3.4.9 is_metabolite

cando.Compound.is_metabolite

bool: bool if the drug is a metabolite itself

11.3.4.10 metabolites

cando.Compound.metabolites

list: List of all metabolites from the compound

11.3.4.11 name

cando.Compound.name

str: Name of the Compound (e.g., 'caffeine')

11.3.4.12 parent

cando.Compound.parent

Compound: Compound object to which this compound is a metabolite.

11.3.4.13 sig

cando.Compound.sig

list: Signature is essentially a column of the Matrix

11.3.4.14 similar

cando.Compound.similar

list: This is the ranked list of compounds with the most similar interaction signatures

11.3.4.15 similar_computed

cando.Compound.similar_computed

bool: Have the distances of all Compounds to the given Compound been computed?

11.3.4.16 similar_sorted

cando.Compound.similar_sorted

bool: Have the most similar Compounds to the given Compound been sorted?

11.3.4.17 status

cando.Compound.status

str: The clinical trial status of the compound from DrugBank ('approved' or 'other')

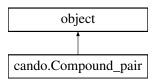
The documentation for this class was generated from the following file:

cando.py

11.4 cando.Compound_pair Class Reference

An object to represent a compound/drug-pair.

Inheritance diagram for cando.Compound_pair:



Public Member Functions

- def __init__ (self, name, id_, index)
- def add_adr (self, adr)

Add an ADR to the list of Indications associated to this Compound.

Public Attributes

name

str: Name of the Compound (e.g., 'caffeine')

id

int: CANDO id from mapping file (e.g., 1, 10, 100, ...)

index

int: The order in which the Compound appears in the mapping file (e.g, 1, 2, 3, ...)

• sig

list: Signature is essentially a column of the Matrix

aux sig

list: Potentially temporary signature for things like pathways, where "c.sig" needs to be preserved

• similar

list: This is the ranked list of compounds with the most similar interaction signatures

· similar computed

bool: Have the distances of all Compounds to the given Compound been computed?

similar_sorted

bool: Have the most similar Compounds to the given Compound been sorted?

• adrs

list: List of ADRs associated with this Compound

11.4.1 Detailed Description

An object to represent a compound/drug-pair.

11.4.2 Constructor & Destructor Documentation

```
11.4.2.1 __init__()
```

11.4.3 Member Function Documentation

```
11.4.3.1 add_adr()
```

Add an ADR to the list of Indications associated to this Compound.

Parameters

ind object: Indication object to add

11.4.4 Member Data Documentation

11.4.4.1 adrs

cando.Compound_pair.adrs

list: List of ADRs associated with this Compound

11.4.4.2 aux_sig

```
cando.Compound_pair.aux_sig
```

list: Potentially temporary signature for things like pathways, where "c.sig" needs to be preserved

```
11.4.4.3 id_
cando.Compound_pair.id_
int: CANDO id from mapping file (e.g., 1, 10, 100, ...)
11.4.4.4 index
cando.Compound_pair.index
int: The order in which the Compound appears in the mapping file (e.g, 1, 2, 3, ...)
11.4.4.5 name
cando.Compound_pair.name
str: Name of the Compound (e.g., 'caffeine')
11.4.4.6 sig
cando.Compound_pair.sig
list: Signature is essentially a column of the Matrix
11.4.4.7 similar
cando.Compound_pair.similar
list: This is the ranked list of compounds with the most similar interaction signatures
11.4.4.8 similar_computed
cando.Compound_pair.similar_computed
```

bool: Have the distances of all Compounds to the given Compound been computed?

11.4.4.9 similar_sorted

```
cando.Compound_pair.similar_sorted
```

bool: Have the most similar Compounds to the given Compound been sorted?

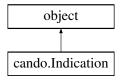
The documentation for this class was generated from the following file:

· cando.py

11.5 cando.Indication Class Reference

An object to represent an indication (disease)

Inheritance diagram for cando. Indication:



Public Member Functions

def __init__ (self, ind_id, name)

Public Attributes

• id_

str: MeSH or OMIM ID for the indication from the mapping file

name

str: Name for the indication from the mapping file

· compounds

list: Every associated compound object from the mapping file

pathways

list: Every pathway associated to the indication from the mapping file

proteins

list: Every protein associated to the indication form the mapping file

pathogen

bool: Whether or not this indication is caused by a pathogen

11.5.1 Detailed Description

An object to represent an indication (disease)

11.5.2 Constructor & Destructor Documentation

11.5.3 Member Data Documentation

11.5.3.1 compounds

```
cando.Indication.compounds
```

list: Every associated compound object from the mapping file

```
11.5.3.2 id_
```

```
cando.Indication.id_
```

str: MeSH or OMIM ID for the indication from the mapping file

11.5.3.3 name

```
cando.Indication.name
```

str: Name for the indication from the mapping file

11.5.3.4 pathogen

cando.Indication.pathogen

bool: Whether or not this indication is caused by a pathogen

11.5.3.5 pathways

```
cando. Indication. pathways
```

list: Every pathway associated to the indication from the mapping file

11.5.3.6 proteins

```
cando. Indication. proteins
```

list: Every protein associated to the indication form the mapping file

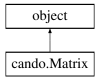
The documentation for this class was generated from the following file:

· cando.py

11.6 cando.Matrix Class Reference

An object to represent a matrix.

Inheritance diagram for cando.Matrix:



Public Member Functions

- def __init__ (self, matrix_file, dist=False, convert_to_tsv=False)
- def convert (self, out_file)

Convert similarity matrix to distance matrix or vice versa.

def normalize (self, outfile, dimension='drugs', method='avg')

Normalize the interaction scores across drugs (default) or proteins (not implemented yet).

Public Attributes

matrix_file

str: Path to file with interaction scores

dist

bool: if the matrix_file is an dist file

· convert_to_tsv

bool: Convert old matrix format (.fpt) to .tsv

proteins

list: Proteins in the Matrix

values

list: Values in the Matrix

11.6.1 Detailed Description

An object to represent a matrix.

Intended for easier handling of matrices. Convert between fpt and tsv, as well as distance to similarity (and vice versa)

11.6.2 Constructor & Destructor Documentation

11.6.3 Member Function Documentation

11.6.3.1 convert()

Convert similarity matrix to distance matrix or vice versa.

The first value in the matrix will determine the type of conversion (0.0 means distance to similarity, 1.0 means similarity to distance).

Parameters

Returns

Returns None

11.6.3.2 normalize()

Normalize the interaction scores across drugs (default) or proteins (not implemented yet).

Parameters

outfile	str: File path to which is written the converted matrix.		
dimension	str: which vector to normalize - either 'drugs' to normalize all scores within the		
	proteomic vector or 'proteins' to normalize for a protein against all drug scores.		
method	str: normalize by the average or max within the vectors		

Returns

Returns None

11.6.4 Member Data Documentation

11.6.4.1 convert_to_tsv

```
cando.Matrix.convert_to_tsv
```

bool: Convert old matrix format (.fpt) to .tsv

11.6.4.2 dist

cando.Matrix.dist

bool: if the matrix_file is an dist file

11.6.4.3 matrix_file

cando.Matrix.matrix_file

str: Path to file with interaction scores

11.6.4.4 proteins

cando.Matrix.proteins

list: Proteins in the Matrix

11.6.4.5 values

cando.Matrix.values

list: Values in the Matrix

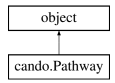
The documentation for this class was generated from the following file:

• cando.py

11.7 cando.Pathway Class Reference

An object to represent a pathway.

Inheritance diagram for cando. Pathway:



Public Member Functions

def __init__ (self, id_)

Public Attributes

• proteins

list: Protein objects associated with the given Pathway

id__

str: Identification for the given Pathway

• indications

list: Indication objects associated with the given Pathway

11.7.1 Detailed Description

An object to represent a pathway.

11.7.2 Constructor & Destructor Documentation

11.7.3 Member Data Documentation

```
11.7.3.1 id_
cando.Pathway.id_
```

str: Identification for the given Pathway

11.7.3.2 indications

cando.Pathway.indications

list: Indication objects associated with the given Pathway

11.7.3.3 proteins

cando.Pathway.proteins

list: Protein objects associated with the given Pathway

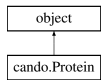
The documentation for this class was generated from the following file:

· cando.py

11.8 cando.Protein Class Reference

An object to represent a protein.

Inheritance diagram for cando. Protein:



Public Member Functions

def __init__ (self, id_, sig)

Public Attributes

id__

PDB or UniProt ID for the given protein.

· alt id

Used when a second identifier mapping is available (such as SIFTs project)

• sig

List of scores representing each drug interaction with the given protein.

pathways

List of Pathway objects in which the given protein is involved.

indications

List of Indication objects to which the protein is associated.

• name

str: the common name of the protein (not currently used)

• gene

str: the gene name from which the protein is produced

11.8.1 Detailed Description

An object to represent a protein.

11.8.2 Constructor & Destructor Documentation

11.8.3 Member Data Documentation

11.8.3.1 alt_id

```
cando.Protein.alt_id
```

Used when a second identifier mapping is available (such as SIFTs project)

11.8.3.2 gene

```
cando.Protein.gene
```

str: the gene name from which the protein is produced

11.8.3.3 id

```
cando.Protein.id_
```

PDB or UniProt ID for the given protein.

11.8.3.4 indications

```
cando.Protein.indications
```

List of Indication objects to which the protein is associated.

11.8.3.5 name

```
cando.Protein.name
```

str: the common name of the protein (not currently used)

11.8.3.6 pathways

```
cando.Protein.pathways
```

List of Pathway objects in which the given protein is involved.

11.8.3.7 sig

```
cando.Protein.sig
```

List of scores representing each drug interaction with the given protein.

The documentation for this class was generated from the following file:

· cando.py

12 File Documentation

12.1 AUTHORS.md File Reference

12.2 cando.py File Reference

Classes

· class cando.Protein

An object to represent a protein.

· class cando.Compound

An object to represent a compound/drug.

class cando.Compound_pair

An object to represent a compound/drug-pair.

· class cando.Indication

An object to represent an indication (disease)

· class cando.Pathway

An object to represent a pathway.

· class cando.ADR

An object to represent an adverse reaction.

class cando.CANDO

An object to represent all aspects of CANDO (compounds, indications, matrix, etc.)

class cando.Matrix

An object to represent a matrix.

Namespaces

cando

Functions

def cando.generate_matrix (v="v2.2", fp="rd_ecfp4", vect="int", dist="dice", org="nrpdb", bs="coach", c_cutoff=0.0, p_cutoff=0.0, percentile_cutoff=0.0, i_score="P", out_file=", out_path=".", nr_ligs=True, approved_only=False, lig_name=False, lib_path=", prot_← path=", ncpus=1)

Generate a matrix using our in-house protocol BANDOCK.

- def cando.calc_scores (c, c_fps, l_fps, p_dict, dist, pscore_cutoff=0.0, cscore_cutoff=0.0, percentile_cutoff=0.0, i_score='P', nr_ligs=[], lig_name=False)
- def cando.generate_signature (cmpd_file, fp="rd_ecfp4", vect="int", dist="dice", org="nrpdb", bs="coach", c_cutoff=0.0, p_cutoff=0.0, percentile_cutoff=0.0, i_score="P", out_file=", out_path=".", nr_ligs=True, prot_path=")

Generate an interaction signature for a query compound using our in-house protocol BANDO← CK.

 def cando.add_cmpds (cmpd_list, file_type='smi', fp="rd_ecfp4", vect="int", cmpd_dir=".", v=None)

Add new compounds to an existing CANDO Compound library, or create a new Compoun library using our in-house protocol BANDOCK.

- def cando.cosine_dist (A)
- def cando.tanimoto_sparse (str1, str2)

Calculate the tanimoto coefficient for a pair of sparse vectors.

def cando.tanimoto_dense (list1, list2)

Calculate the tanimoto coefficient for a pair of dense vectors.

def cando.get_fp_lig (fp)

Download precompiled binding site ligand fingerprints using the given fingerprint method.

• def cando.get_data (v="v2.2", org='nrpdb', fp='rd_ecfp4', vect='int')

Download CANDO v2.2+ data.

def cando.clear cache ()

Clear files in "data/" directory.

def cando.get_tutorial ()

Download data for tutorial.

def cando.get test ()

Download data for test script.

def cando.dl_dir (url, out, l)

Function to recursively download a directory.

def cando.dl_file (url, out_file)

Function to download a file.

• def cando.load_version (v='v2.3', protlib='nrpdb', i_score='CxP', approved_only=False, compute_distance=False, dist_metric='cosine', protein_set="', ncpus=1)

Directly load a pre-compiled version of CANDO.

12.3 LICENSE.md File Reference

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