IPTK

Release 0.3.1

Hesham ElAbd

CONTENTS

1	Introduction:	3
2	Installation:	5
3	Funding: 3.1 Guide	7 7
4	Indices and tables	57
Ру	ython Module Index	59
In	dex	61



Analysis, Visualize, Compare and Integrate experimentally generated or in-silico predicted Immunopeptidomics data,

CONTENTS 1

2 CONTENTS

CHAPTER

ONE

INTRODUCTION:

IPTK is a Pythonic library specialized in the analysis of HLA-peptidomes identified through an Immunopeptiomics pipeline. The library provides a high level API for analyzing and visualizing the identified peptides, Integrating transcritomics and protein structure information for a rich analysis and for comparing different experiments and different runs.

СНАРТЕ	:R
TWO	0

INSTALLATION:

The library can installed using pip as follow

CHAPTER

THREE

FUNDING:

The project was funded by the German Research Foundation (DFG) (Research Training Group 1743, 'Genes, Environment and Inflammation')



3.1 Guide

3.1.1 License

Will be written here

3.1.2 Contact

for further question and communication please contact h.elabd@ikmb.uni-kiel.de

3.1.3 Get Started!

To get started with using the library check the Interactive Tutorials available at https://github.com/ikmb/iptoolkit/tree/master/Tutorials

3.1.4 IPTK

IPTK package

Subpackages

IPTK. Analysis package

Submodules

IPTK.Analysis.AnalysisFunction module

The module contain a collection of analysis function that can be used by the methods of the classes defined in the classes module.

```
IPTK.Analysis.AnalysisFunction.compute_binary_distance(peptides: List[str], dist\_func: Callable) \rightarrow numpy.ndarray
```

compare the distance between every pair of peptides in a collection of peptides. @param: peptides: a collection of peptides sequences. @param: dist_func: function to compute the distance between each pair of peptides. @note:

Parameters

- **peptides** (List[str]) a collection of peptides sequences.
- dist_func (Callable) a function to compute the distance between each pair of peptides.

Raises RuntimeError – make sure that the dist_function is suitable with the peptides which might have different lengths.

Returns the distance between each pair of peptides in the provided list of peptides

Return type np.ndarray

```
\label{lem:in_representation} IPTK. Analysis. AnalysisFunction. {\it compute\_change\_in\_protein\_representation} \ ({\it mapped\_prot\_cond1: numpy.ndarray}, \\ {\it mapped\_prot\_cond2: numpy.ndarray}) \\ {\it description of the protein\_to a protein\_representation} \ ({\it mapped\_prot\_cond1: numpy.ndarray}) \\ {\it description of the protein\_to a protein\_representation} \ ({\it mapped\_prot\_cond1: numpy.ndarray}) \\ {\it description of the protein\_to a p
```

Compute the change in the protein representation between two conditions, by computing the difference in the area under the curve, AUC.

Parameters

- mapped_prot_cond1 (np.ndarray) a mapped protein instance containing the protein coverage in the first condition
- mapped_prot_cond2 (np.ndarray) a mapped protein instance containing the protein coverage in the second condition

Raises ValueError – if the provided pair of proteins is of different length

Returns the difference in the area under the coverage curve between the two experiments.

Return type float

```
\begin{tabular}{ll} {\tt IPTK.Analysis.AnalysisFunction.} \textbf{compute\_difference\_in\_representation} (mapped\_prot\_cond1: & numpy.ndarray, & mapped\_prot\_cond2: & numpy.ndarray) \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & &
```

return the difference in the representation of a protein between two conditions by substracting the coverage of the first protein from the second proteins.

@param: mapped_prot_cond1: a mapped protein instance containing the protein coverage in the first condition @param: mapped_prot_cond2: a mapped protein instance containing the protein coverage in the second condition

Parameters

- mapped_prot_cond1 (np.ndarray) a mapped protein instance containing the protein coverage in the first condition
- mapped_prot_cond2 (np.ndarray) a mapped protein instance containing the protein coverage in the second condition

Returns an array that shows the difference in coverage between the two proteins at each amino acid position.

Return type np.ndarray

IPTK.Analysis.AnalysisFunction.compute_expression_correlation(expl:

IPTK.Classes.Experiment.Experiment, *exp2*: IPTK.Classes.Experiment.Experiment)

float

compute the correlation in the gene expression between two experiments by constructing a union of all the proteins expressed in the first and second experiments, extract the gene expression of these genes and then compute the correlation using SciPy stat module.

Parameters

- exp1 (Experiment) The first experimental object
- exp2 (Experiment) he second experimental object

Returns the correlation in gene expression of the proteins inferred in the provided pair of experiment

Return type float

```
IPTK.Analysis.AnalysisFunction.download_structure_file (pdb\_id: str) \rightarrow None Download PDB/mmCIF file containing the pbd_id from PDB using BioPython library
```

Parameters $pdb_id(str)$ – the protein id in protein databank

```
IPTK.Analysis.AnalysisFunction.get_binnary_peptide_overlap(expl:
```

IPTK.Classes.Experiment.Experiment, *exp2*:
IPTK.Classes.Experiment.Experiment)

PTK.Classes.Experiment.Experiment

→ List[str]

compare the peptide overlap between two experimental objects.

Parameters

- **exp1** (Experiment) an instance of class Experiment.
- exp2 (Experiment) an instance of class Experiment.

Returns a list of peptides that have been identified in both experiments.

Return type Peptides

```
IPTK.Analysis.AnalysisFunction.get_binnary_protein_overlap(expl:
```

IPTK.Classes.Experiment.Experiment, *exp2:*IPTK.Classes.Experiment.Experiment)
→ List[str]

compare the protein overlap between two experimental objects.

Parameters

- exp1 (Experiment) an instance of class Experiment.
- **exp2** (Experiment) an instance of class Experiment.

Returns a list of proteins that have been identified in both experiments.

Return type Proteins

```
IPTK.Analysis.AnalysisFunction.get_sequence_motif(peptides: List[str], temp_dir: str = './TEMP_DIR', verbose: bool = False, meme_params: Dict[str, str] = \{\}) \rightarrow None
```

compute the sequences motif from a collection of peptide sequences using meme software.

Parameters

- **peptides** (*Peptides*) a list of string containing the peptide sequences
- temp_dir(str, optional) he temp directory to write temp-files to it, defaults to "./TEMP_DIR"
- **verbose** (bool, optional) whether or not to print the output of the motif discovery tool to the stdout, defaults to False
- meme_params (Dict[str,str], optional) a dict object that contain meme controlling parameters, defaults to {}

Raises

- **FileNotFoundError** incase meme is not installed or could not be found in the path!
- **ValueError** incase the peptides have different length!

Module contents

IPTK.Classes package

Submodules

IPTK.Classes.Database module

This submodule define a collection of container classes that are used through the library

```
class IPTK.Classes.Database.CellularLocationDB (path2data: str, sep: str = \t')
Bases: object
```

The class provides an API to access the cellular location information from a database the follow the structure of the human Proteome Atlas sub-cellular location database. See https://www.proteinatlas.org/about/download for more details.

add_to_database ($genes_to_add$: IPTK.Classes.Database.CellularLocationDB) \rightarrow None add the the location of more proteins to the database.

Parameters genes_to_add (CellularLocationDB) - a CellularLocationDB instance containing the genes that shall be added to the database.

Raises

- ValueError if the genes to add to the database are already defined in the database
- **RuntimeError** Incase any other error has been encountered while merging the tables.
- **get_approved_location** ($gene_id: Optional[str] = None, gene_name=None) <math>\rightarrow$ List[str] return the location of the provided gene id or gene name

Parameters

- gene_id (str, optional) the id of the gene of interest, defaults to None
- **gene_name** ([type], optional) the name of the gene of interest, defaults to None

Raises

- ValueError if both gene_id and gene_name are None
- **KeyError** if gene_id is None and gene_name is not in the database
- **KeyError** if gene_name is None and gene_id is not in the database
- RuntimeError incase some error was encountered while running retriving the elements from the database

Returns the approved location where the protein the corresponds to the provided name or id is located.

Return type List[str]

```
get\_gene\_names() \rightarrow List[str]
```

return a list of all gene names in the dataset

Returns the names of all genes in the database

Return type List[str]

```
\texttt{get\_genes}() \rightarrow List[str]
```

return a list of all gene ids in the dataset

Returns all genes ids currently defined in the database

Return type List[str]

 get_go_names ($gene_id: Optional[str] = None, gene_name = None$) \rightarrow List[str] return the location of the provided gene id or gene name

Parameters

- gene_id(str, optional) the id of the gene of interest, defaults to None
- gene_name ([type], optional) the name of the gene of interest, defaults to None

Raises

- ValueError if both gene_id and gene_name are None
- **KeyError** if gene_id is None and gene_name is not in the database

- **KeyError** if gene_name is None and gene_id is not in the database
- RuntimeError incase some error was encountered while running retriving the elements from the database

Returns the gene ontology, GO, location where the protein the corresponds to the provided name or id is located.

Return type List[str]

get_main_location (gene_id: Optional[str] = None, corresponds=None) → List[str] return the main location(s) of the provided gene id or gene name. If both gene Id and gene name are provided, both gene_id has a higher precedence

Parameters

- gene_id (str, optional) the id of the gene of interest, defaults to None
- **gene_name** ([type], optional) the name of the gene of interest, defaults to None

Raises

- ValueError if both gene_id and gene_name are None
- **KeyError** if gene_id is None and gene_name is not in the database
- **KeyError** if gene_name is None and gene_id is not in the database
- RuntimeError incase some error was encountered while running retriving the elements from the database

Returns the main location where the protein the corresponds to the provided name or id is located.

Return type List[str]

 $\mathtt{get_table}$ () \rightarrow pandas.core.frame.DataFrame return the instance table

Returns the location table of the instance.

Return type pd.DataFrame

class IPTK.Classes.Database.GeneExpressionDB(path2data:

 $sep: str = \t^{\prime}$

Bases: object

provides an API to access gene expression data stored in table that follows the same structure as the Human proteome Atlas Normalized RNA Expression see https://www.proteinatlas.org/about/download for more details

```
get\_expression (gene\_name: Optional[str] = None, gene\_id: Optional[str] = None) \rightarrow pan-das.core.frame.DataFrame
```

Return a table summarizing the expression of the provided gene name or gene id accross different tissues.

Parameters

- gene_id (str, optional) the id of the gene of interest, defaults to None
- **gene_name** ([type], optional) the name of the gene of interest, defaults to None

Raises

• ValueError – if both gene_id and gene_name are None

- **KeyError** if gene_id is None and gene_name is not in the database
- **KeyError** if gene_name is None and gene_id is not in the database
- RuntimeError incase some error was encountered while running retriving the elements from the database

Returns A table summarizing the expression of the provided gene accross all tissues in the database

Return type pd.DataFrame

 $\texttt{get_expression_in_tissue}$ ($tissue_name: str$) \rightarrow pandas.core.frame.DataFrame return the expression profile of the provided tissue

Parameters tissue_name (str) – the name of the tissue

Raises

- **KeyError** incase the provided tissue is not provided in the database
- **RuntimeError** in case any error was encountered while generating the expression profile.

Returns a table summarizing the expression of all genes in the provided tissue.

Return type pd.DataFrame

```
get\_gene\_names() \rightarrow List[str]
```

return a list of the UNIQUE gene names currently in the database

Returns a list of the UNIQUE gene names currently in the database

Return type List[str]

```
get\_genes() \rightarrow List[str]
```

return a list of the UNIQUE gene ids currently in the database

Returns a list of the UNIQUE gene ids currently in the database

Return type List[str]

```
\texttt{get\_table}() \rightarrow pandas.core.frame.DataFrame
```

return a table containing the expression value of all the genes across all tissues in the current instance

Returns The expression of all genes across all tissues in the database.

Return type pd.DataFrame

```
get\_tissues() \rightarrow List[str]
```

return a list of the tissues in the current database

Returns a list containing the names of the UNIQUE tissues in the database.

Return type List[str]

```
class IPTK.Classes.Database.OrganismDB (path2Fasta: str)
```

Bases: object

Extract information about the source organsim of a collection of protein sequences from a fasta file and provides an API to query the results. The function expect the input fasta file to have header written in the UNIPROT format.

 $get_number_protein_per_organism() \rightarrow pandas.core.frame.DataFrame provides a table containing the number of proteins per organism.$

Returns a table containing the number of proteins per organism

Return type pd.DataFrame

```
get\_org(prot\_id: str) \rightarrow str
```

return the parent organism of the provided proteins

Parameters $prot_id(str)$ – the id of the protein of interest

Raises KeyError – incase the provided identifier is not in the database

Returns the name of the parent organism, i.e. the source organism.

Return type str

```
\texttt{get\_unique\_orgs}() \rightarrow List[str]
```

get the number of unique organisms in the database

Returns a list of all unique organisms in the current instance

Return type List[str]

```
class IPTK.Classes.Database.SeqDB (path2fasta: str)
```

Bases: object

load a fasta file and constructs a lock up dictionary where sequence ids are keys and sequences are values.

```
get\_seq(protein\_id: str) \rightarrow str
```

returns the corresponding sequence if the provided protein-id is defined in the database.

Parameters protein_id (str) - The protein id to retrive its sequence.

Raises KeyError – If the provided protein does not exist in the database

Returns the protein sequence

Return type str

 $has_sequence(sequence_id: str) \rightarrow bool$

check if the provided sequence id is an element of the database or not

Parameters sequence_name (str) - The id of the sequence

Returns True if the database has this id, False otherwise.

Return type bool

IPTK.Classes.Experiment module

This module provides an abstraction for an IP experiment.

```
class IPTK.Classes.Experiment.Experiment (proband: IPTK.Classes.Proband.Proband, hla_set: IPTK.Classes.HLASet.HLASet, tissue: IPTK.Classes.Tissue.Tissue, database: IPTK.Classes.Database.SeqDB, ident_table: pandas.core.frame.DataFrame)
```

Bases: object

A representation of an immunopeptidomic experiment.

```
add_org_info (prot2org: Dict[str, str]) \rightarrow None annotated the inferred proteins with their source organism
```

Parameters prot2org (*ProteinSource*) – a dict that contain the protein id as keys and its source organism as values and add this info to each protein inferred in the current experiment.

Raises RuntimeWarning – If the provided dictionary does cover all proteins in the experimental object.

 ${\tt annotate_proteins} \ (\textit{organisms_db:} \ IPTK.Classes.Database.OrganismDB) \ \rightarrow None$

Extract the parent organisms of each protein in the experiment from an organism database instance.

Parameters organisms_db (OrganismDB) – an OrgansimDB instance that will be used to annotate the proteins identified in the experiment.

```
drop\_peptide\_belong\_to\_org(org: str) \rightarrow None
```

Drop the all the peptides that belong to a user provided organism. Note that, this function will IRRE-VERSIBLY remove the peptide from the experimental object.

Parameters org (str) – the organims name

$\texttt{get_binarized_results}() \rightarrow List[numpy.ndarray]$

Return a list of NumPy arrays where each array represents a child peptide, parent protein mapped pair. Please note that, The function treat each peptide-protein pair individually, that is if two peptides originating from the same protein, it treat them independently and the same protein will be represented twice with the two different peptides. Incase an integrative mapping is needed, the function @get_integrated_binarized_results@ shall be used.

Returns a list of NumPy arrays containing the mapping between each peptide protein pair.

Return type MappedProtein

 $\texttt{get_c_terminal_flanked_seqs}$ (flank_length: int) \rightarrow List[IPTK.Classes.Peptide.Peptide] return the c-terminal flanking sequences

Parameters flank_length (*int*) – the length of the peptide downstream of the C-terminal of the peptide

Returns a list sequences contain the N-terminal flanking sequence for each peptide in the instance.

Return type Peptides

 $\mbox{\tt get_experiment_reference_tissue_expression}\ () \ \rightarrow \mbox{\tt pandas.core.frame.DataFrame} \ \ \mbox{\tt return the reference gene expression for the current tissue} \label{tissue}$

Returns A table that contain the expression value for ALL the genes in the instance Tissue **Return type** pd.DataFrame

return a table containing the expression of the proteins inferred in the current experiment from the current tissue. This method need internet connection as it need to access uniprot mapping API to map uniprot IDs to gene IDs.

Parameters non_mapped_dval (float, optional) - A default value to be added incase the parent protein is not define in the expression database, defaults to -1

Returns a table that contain the expression of the protein inferred in the database

Return type pd.DataFrame

$get_flanked_peptides (flank_length: int) \rightarrow List[str]$

returns a list of sequences containing the peptides identified in the experiment padded with the flanking regions from all the parents of each peptide.

Parameters flank length (int) - the length of the flanking region

Returns a list of the peptides + the flanking region.

Return type Sequences

@brief: @param: not_mapped_val: the default value to return incase the GO term of the protein can not be extracted. @note: This method need internet connection as it need to access uniprot mapping API to map uniprot IDs to gene IDs.

Parameters not_mapped_val (str, optional) – The default value to return incase the GO term of the protein can not be extracted, defaults to 'UNK'

Returns A table that contain the GO-location term for each protein in the current instance.

Return type pd.DataFrame

```
\mathtt{get\_hla\_allele}() \rightarrow List[str]
```

Returns the set of HLA alleles from which the instance peptides have been eluted

Return type List[str]

```
\mathtt{get\_hla\_class}() \rightarrow \mathtt{int}
```

Returns the HLA class

Return type int

das.core.frame.DataFrame retrun the main cellular location for the identified proteins. This method need internet connection as it need to access uniprot mapping API to map uniprot IDs to gene IDs.

Parameters not_mapped_val (str, optional) – The default value to return incase the location of a protein can not be extracted, defaults to 'UNK'

Returns A table that contain the main cellular compartment for each protein in the current instance.

Return type pd.DataFrame

```
get_mapped_protein (pro_id: str) → numpy.ndarray
```

return an NumPy array of shape 1 x protein length where each number in the array represents the total number of peptides identified in the experiment that have originated from the said position in the protein.

```
Parameters pro_id (str) - the protein id
```

Raises KeyError - if the provided protein id was inferred from the current experiment

Returns a NumPy array that contain the coverage of the protein.

Return type np.ndarray

```
get_mapped_proteins() → Dict[str, List[numpy.ndarray]]
```

return a dictionary of all the proteins identified in the current experiment with all inferred peptides mapped to them.

Returns a dictionary that contain the mapped proteins for all the proteins in the current instance.

Return type MappedProteins

```
get_mono_parent_peptides() → List[IPTK.Classes.Peptide.Peptide]
```

return a list of peptides that have only one parent protein

Returns list of peptide instance

Return type Peptides

 $get_n_terminal_flanked_seqs$ (flank_length: int) \rightarrow List[IPTK.Classes.Peptide.Peptide] return the n-terminal flanking sequences

Parameters flank_length (*int*) – the length of the flanking region upstream of the N-terminal of the peptide

Returns a list sequences contain the N-terminal flanking sequence for each peptide in the instance.

Return type Peptides

$get_negative_example (fold: int = 2) \rightarrow List[str]$

generate negative examples, i.e., non-bounding peptides from the proteins identified in the current experiment.

Parameters fold (*int*, *optional*) – the number of negative example to generate relative to the number of unique identified peptides, defaults to 2

Returns list of non-presented peptides from all inferred proteins.

Return type Sequences

 $\verb"get_num_peptide_expression_table" () \rightarrow pandas.core.frame.DataFrame$

Get a table that contain the id of all parent proteins, number of peptide per-proteins and the expression value of these parent transcripts. Please note, this method need internet connection as it need to access uniprot mapping API to map uniprot IDs to gene IDs.

Returns the number of peptides per protein table

Return type pd.DataFrame

get_num_peptide_per_go_term() → pandas.core.frame.DataFrame retrun the number of peptides per each GO-Term :return: A table that has two columns, namely, GO-Terms and Counts. :rtype: pd.DataFrame

get_num_peptide_per_location() → pandas.core.frame.DataFrame

retrun the number of peptides obtained from proteins localized to different sub-cellular compartments

Returns A table that has two columns, namely, Compartment and Counts.

Return type pd.DataFrame

$\verb"get_number_of_children" (\textit{pro_id: str}) \rightarrow \text{int}$

return the number of children, i.e. number of peptides belonging to a parent protein

Parameters $pro_id(str)$ – the id of the parent protein

Returns the number of peptides

Return type int

 $\mbox{\tt get_number_of_proteins_per_compartment}\ (\,) \ \rightarrow \mbox{\tt pandas.core.frame.DataFrame}$ get the number of proteins from each compartment

Returns A table that has two columns, namely, Compartment and Counts.

Return type pd.DataFrame

 $\mbox{\tt get_number_of_proteins_per_go_term} \ () \ \rightarrow \mbox{\tt pandas.core.frame.DataFrame} \ \mbox{\tt get the number of proteins from each GO-Term}$

Returns A table that has two columns, namely, GO-Terms and Counts.

Return type pd.DataFrame

get_orgs() → List[str]

return a list containing the UNIQUE organisms identified in the current experiment

Returns list of all UNIQUE organisms inferred from the inferred proteins.

Return type List[str]

 $get_peptide(pep_seq: str) \rightarrow IPTK.Classes.Peptide.Peptide$

return a peptide instance corresponding to the user provided peptide sequence.

Parameters $pep_seq(str)$ – the peptide sequence

Raises KeyError – if the peptide sequence has not been inferred from the current database.

Returns the peptide instance with the coresponding sequence

Return type Peptide

 $get_peptide_number_parent$ (ascending: bool = False) \rightarrow pandas.core.frame.DataFrame return a pandas dataframe with the peptide sequence in the first columns and the number of parent proteins in the second column.

Parameters ascending (bool, optional) – ascending sort the peptide by their number of parent proteins, defaults to False

Returns the number of parents for each peptide

Return type pd.DataFrame

get_peptides() → List[IPTK.Classes.Peptide.Peptide]

Returns a set of all the peptide stored in the experimental object

Return type Peptides

 $get_peptides_length() \rightarrow List[int]$

return a list containing the length of each unique peptide in the database.

Returns peptides' lengths

Return type List[int]

 $\texttt{get_peptides_per_organism}() \rightarrow pandas.core.frame.DataFrame$

return a pandas dataframe that contain the count of peptides belonging to each organism in the database

Returns a table with two columns, namely, Organisms and Counts

Return type pd.DataFrame

get_peptides_per_protein (ascending: bool = False) → pandas.core.frame.DataFrame return a pandas dataframe that contain the number of peptides belonging to each protein inferred in the experiment

Parameters ascending (bool, optional) – ascending sort the proteins by their number of parent number of child peptides, defaults to False

Returns a table with the following columns, Proteins and Number_of_Peptides

Return type pd.DataFrame

get_poly_parental_peptides() → List[IPTK.Classes.Peptide.Peptide]

return a list of peptides that have more than one parent proteins :return: [list of peptide instance :rtype: Peptides

 $\texttt{get_proband_name} () \rightarrow \mathsf{str}$

Returns the proband name

Return type str

 $\mathtt{get_proteins}() \rightarrow \mathtt{List}[\mathit{IPTK.Classes.Protein.Protein}]$

Returns a set of all the proteins in the experimental object

Return type Proteins

 $\mathtt{get_tissue}() \rightarrow \mathit{IPTK.Classes.Tissue.Tissue}$

Returns the tissue of the current experiment.

Return type *Tissue*

 $\texttt{get_tissue_name}() \rightarrow \mathsf{str}$

Returns the tissue name

Return type str

 $has_allele_group (gene_group: str) \rightarrow bool$

return whether or not the experiment contain peptides eluted from an HLA-alleles belonging to the provided allele group or not

Parameters gene_group (str) – the gene group to query the hla_set against

Returns True if the gene group has a member that is a member of the instance HLASet and False otherwise

Return type bool

 $has_gene(locus: str) \rightarrow bool$

return whether or not the experiment contain peptides eluted from an HLA-alleles belonging to the provided locus or not

Parameters locus (str) – the locus of the allele to query the hla_set against

Returns True if the locus has a member that is a member of the instance HLASet and False otherwise

Return type bool

 $has_hla_allele(individual: str) \rightarrow bool$

return whether or not the experiment contain an eluted peptides from the provided alleles

Parameters individual (str) – is the name of the allele as a string

Returns True if the allele is a member of the instance HLASet and False otherwise.

Return type bool

 $has_protein_group (protein_group: str) \rightarrow bool$

return whether or not the experiment contain peptides eluted from an HLA-alleles belonging to the provided protein group or not

Parameters protein_group (str) - The protein group to query the hla_set against

Returns True if the locus has a member that is a member of the instance HLASet and False otherwise

Return type bool

 $is_a_parent_protein(protein: str) \rightarrow bool$

Parameters protein – check if the protein is a member of the instance proteins or not.

Returns True if the protein has been identified in the current instance, False otherwise.

Return type bool

```
is\_member(peptide: str) \rightarrow bool
```

Parameters peptide (str) – check if the peptide is a member of the instance peptides or not.

Returns True if the peptide has been identified in the current instance, False otherwise.

Return type bool

IPTK.Classes.ExperimentalSet module

An Experimental set which is a collection of experiments. The class provides an API for integrating and comparing different experimental instances.

```
class IPTK.Classes.ExperimentalSet.ExperimentSet (**exp_id_pair)
    Bases: object
    an API for integrating and comparing different experimental instances
```

```
add_experiment (**exp_id_pair) \rightarrow None add an arbitrary number of experiments to the set
```

Parameters

- org (str) The name of the organism to query the database for it.
- abs_count (bool, optional) The absolute count, defaults to False

Returns the count of the peptides that belong to a specific organism in the database.

Return type pd.DataFrame

```
compare_peptide_counts() → pandas.core.frame.DataFrame
```

Returns A table that contain the total number of peptides and per-organism peptide counts among all experiments in the set

Return type pd.DataFrame

compute_average_distance_between_exps() → pandas.core.frame.DataFrame

compute the average distance between experiments by taking the average over the z-axis of the 3D tensor computed by the function compute_change_in_protein_representation.

Returns A 2D tensor with shape of (num-experiments, num-experiments)

Return type pd.DataFrame

$\label{local_compute_change_in_protein_representation} () \rightarrow numpy.ndarray$

Compute the change in protein representation among the proteins that are presented/ detect in all of the instance's experiments.

Returns a 3D tensor, T, with shape of (num-experiments, num-experiments, num-proteins), where T[i,j,k] is a the difference between experiment i & j with respect to the k th protein :rtype: np.ndarray

compute correlation in experssion() → pandas.core.frame.DataFrame

compute the correlation in parent protein gene-expression across all the experiments in the set. See the function **compute_binary_correlation** in the analysis module for information about the computational logic.

Returns return a 2D matrix containing the coorelation in gene expression between each pair of experiments inside the current instance collection of experiments.

Return type pd.DataFrame

 $compute_peptide_length_table() \rightarrow pandas.core.frame.DataFrame$

Returns A table that contain the length of each peptide in the experiment

Return type pd.DataFrame

compute_peptide_overlap_matrix() → numpy.ndarray

Returns a 2D matrix containing the number of peptide overlapping between each pair of experiments inside the current instance collection of experiment.

Return type np.ndarray

 ${\tt compute_peptide_representation_count}\;() \; \to Dict[str, int]$

Returns The number of times a peptide was observed accross experiments in the set

Return type Counts

 $compute_protein_coverage_over_the_set() \rightarrow Dict[str, numpy.ndarray]$

Returns the mapped representation for each protein in accross the entire set

Return type Dict[str, np.ndarray]

 $\verb|compute_protein_overlap_matrix|()| \rightarrow numpy.ndarray|$

Returns return a 2D matrix containing the number of proteins overlapping between each pair of experiments inside the current instance collection of experiment.

Return type np.ndarray

 $compute_protein_representation_count() \rightarrow Dict[str, int]$

Returns The number of times a protein was observed accross the experiment in the set

Return type Counts

 $drop_peptides_belong_to_org(org_name: str) \rightarrow None$

drop all the peptides that belong to the provided organisms from all experiments in the set.

Parameters org_name (str) – the name of the organism to drop

 $\texttt{get_allele_count}\;()\;\to Dict[str,int]$

Returns the number of experiments obtained from each allele in the instance.

Return type Counts

 $\texttt{get_experiment}(exp_name: str) \rightarrow IPTK.Classes.Experiment.Experiment$

return the experiment pointed to by the provided experimental name

Parameters exp_name (str) – the name of the experiment

Raises KeyError – if the provided experimental name is not in the dataset.

Returns the experiment corresponds to the provided name

Return type Experiment

 $\texttt{get_experimental_names}\:(\:)\:\to List[str]$

Returns a list with all the identifiers of the experiments in the set

Return type Names

```
get_experiments() → Dict[Experiment]
         Returns return a dict with all the experiments stored in the instance as value of ids as keys.
         Return type Dict[Experiment]
{\tt get\_num\_experiments\_in\_the\_set}\:(\:)\:\to int
         Returns The number of experiments currently in the set
         Return type int
\texttt{get\_peptides\_present\_in\_all}() \rightarrow List[IPTK.Classes.Peptide.Peptide]
         Returns the peptides that are observed in every experiments in the set.
         Return type Peptides
get_proband_count() → Dict[str, int]
         Returns The number of experiments obtained from each proband in the ExperimentalSet.
         Return type Counts
get\_proteins\_present\_in\_all() \rightarrow List[str]
         Returns the proteins that are inferred in all experiments of the set
         Return type Proteins
get tissue counts() → Dict[str, int]
         Returns The number of experiments obtained from each tissue in the current instance
         Return type Counts
\verb"get_total_peptide_per_org_count" () \rightarrow pandas.core.frame.DataFrame
         Returns The total count of peptides per organism accross the all experiments in the set.
         Return type pd.DataFrame
get\_unique\_orgs() \rightarrow List[str]
         Returns a list of the unique organisms in the set
         Return type List[str]
get\_unique\_peptides() \rightarrow List[IPTK.Classes.Peptide.Peptide]
         Returns The set of unique peptides in the experimentalSet
         Return type Peptides
get_unique_proteins() → List[str]
         Returns the set of unique proteins in the experimentalset
         Return type Proteins
group\_by\_proband() \rightarrow Dict[str, IPTK.Classes.ExperimentalSet.ExperimentSet]
         Returns a map between each proband and an Experimentalset object represent all the experi-
              ments objects belonging to this proband.
         Return type Dict[str, ExperimentSet]
group\_by\_tissue() \rightarrow Dict[str, IPTK.Classes.ExperimentalSet.ExperimentSet]
         Returns A map between each tissue and an ExperimentalSet object representing all experiments
```

belonging to that tissue.

```
Return type Dict[str, ExperimentSet]
      is_peptide_present_in_all(peptide: str) → bool
                Parameters peptide (str) - The peptide sequence to search its occurrences in every experi-
                     ment contained in the set
                Returns True if peptide is present in all experiments inside the instance, False otherwise
                Return type bool
      is\_protein\_present\_in\_all (protein: str) \rightarrow bool
                Parameters protein (str) – the protein id to search its occurrences in every experimental in
                     the set
                Returns True if peptide is present in all experiments inside the instance, False otherwise
                Return type bool
IPTK.Classes.HLAChain module
The implementation of an HLA molecules
class IPTK.Classes.HLAChain.HLAChain (name: str)
      Bases: object
      \texttt{get\_allele\_group}() \rightarrow \mathsf{str}
                Returns The allele group
                Return type str
      get\_chain\_class(gene\_name: str) \rightarrow int
                Parameters gene_name (str) - the name of the gene
                Returns 1 if the gene belongs to class one and 2 if it belong to class two
                Return type int
      \mathtt{get\_class}() \rightarrow \mathrm{int}
                Returns The HLA class
                Return type int
      \texttt{get\_gene}() \rightarrow \mathsf{str}
                Returns the gene name
                Return type str
      \mathtt{get}\_\mathtt{name}\left(\right) \to \mathrm{str}
                Returns The chain name
                Return type str
      \texttt{get\_protein\_group}() \rightarrow str
                Returns the protein name
```

3.1. Guide 23

Return type str

IPTK.Classes.HLAMolecules module

```
a representation of an HLA molecules
class IPTK.Classes.HLAMolecules.HLAMolecule(**hla_chains)
      Bases: object
      get_allele_group() \rightarrow List[str]
               Returns the allele group for the instance chain/pair of chains
               Return type AlleleGroup
      \mathtt{get\_class}() \rightarrow \mathrm{int}
               Returns The class of the HLA molecules
               Return type int
      get\_gene() \rightarrow List[str]
               Returns gene/pair of genes coding for the current HLA molecules
               Return type Genes
      get_name(sep: str = ':') \rightarrow str
               Parameters sep (str, optional) - the name of the allele by concatenating the names of
                    the individual chains using a separator, defaults to ':'
               Returns [description]
               Return type str
      get_protein_group() → List[str]
               Returns The protein group for the instance chain/pair of chains
               Return type ProteinGroup
IPTK.Classes.HLASet module
An abstraction for a collection of HLA alleles
class IPTK.Classes.HLASet.HLASet(hlas: List[str], gene_sep: str = ':')
      Bases: object
      get_alleles() \rightarrow List[str]
               Returns The class of the HLA-alleles in the current instance
               Return type int
      \texttt{get\_class}\,(\,)\,\to int
               Returns The class of the HLA-alleles in the current instance
               Return type int
      get_hla_count() \rightarrow int
               Returns the count of HLA molecules in the set
               Return type int
```

 $has_allele(allele: str) \rightarrow bool$

Parameters allele (str) – the name of the allele to check the instance for

Returns True, if the provided allele is in the current instance, False otherwise.

Return type bool

 $has_allele_group (allele_group: str) \rightarrow bool$

Parameters allele_group (str) – the allele group to search the set for

Returns True, if at least one allele in the set belongs to the provided allele group, False otherwise.

Return type bool

 $has_gene(gene_name: str) \rightarrow bool$

Parameters gene_name (str) - the gene name to search the set against.

Returns True, if at least one of the alleles in the set belongs to the provided gene. False otherwise

Return type bool

 $has_protein_group (protein_group: str) \rightarrow bool$

Parameters protein_group – The protein group to search the set for

Returns True, if at least one allele in the set belongs to the provided protein group

Return type bool

IPTK.Classes.Peptide module

A representation of the eluted peptides and its identified proteins.

```
class IPTK.Classes.Peptide.Peptide(pep_seq: str)
    Bases: object
```

An representation of an eluted peptide.

add_org_2_parent (prot_name: str, org: str) \rightarrow None add the source organism of one of the instance parent protein

Parameters

- $prot_name(str)$ The name of the protein, i.e. the identifier of the protein
- org(str) the name of the organism

Raises ValueError – incase the provided protein is not a parent of the provided peptide

add_parent_protein (parent_protein, start_index: int, end_index: int) → None

add a protein instance as a parent to the current peptide. The library use Python-based indexing where its 0-indexed and ranges are treated as [start, end). :param parent_protein: a Protein instance that act as a parent to the peptide. :type parent_protein: Protein :param start_index: the position in the parent protein where the peptide starts :type start_index: int :param end_index: the index of the amino acid that occurs after the last amino acid in the peptide, :type start_index: int

```
get_c_terminal_flank_seq(flank_len: int) → List[str]
```

:param flank_len:the length of the flanking regions :type flank_len: int :return: a list of string containing the sequences located downstream of the peptide in the parent protein. :rtype: [type]

 $\texttt{get_flanked_peptide} (\textit{flank_len: int}) \rightarrow List[str]$

Parameters flank len (int) – the length of the flanking regions

Returns A list of string containing the length of the peptide + the flanking region from both the N and C terminal of the instance peptide, from all proteins. **Return type** Sequences $\mathtt{get_length}\left(\right) \to \mathrm{int}$ **Returns** the length of the peptides **Return type** int $\texttt{get}_n_\texttt{terminal_flank_seq}(\mathit{flank_len:int}) \rightarrow List[str]$ **Parameters flank_len** (int) – the length of the flanking regions **Returns** a list of string containing the sequences located upstream of the peptide in the parent protein. **Return type** List[str] $\texttt{get_non_presented_peptides}$ (length: int) $\rightarrow List[str]$ **Parameters length** (int) – The length, i.e. number of amino acids, for the non-presented peptide **Returns** non-presented peptide from all the parent protein of the current peptide instance. Return type Sequences ${\tt get_number_of_parents}\:(\:)\:\to int$ **Returns** the number of instance parent proteins Return type int $\texttt{get_number_parent_protein} () \rightarrow int$ **Returns** the number of parent proteins this instance has Return type int get_parent (pro_id: str) **Parameters** pro_id (str) - The protein identifer **Returns** the parent protein that has an id matching the user defined pro id Return type *Protein* $\texttt{get_parent_proteins}() \rightarrow List[str]$ $\texttt{get_parents_org}\,(\,)\,\to List[str]$ **Returns** a list containing the name of each parent protein source organisms Return type Organisms $\texttt{get_peptide_seq}() \rightarrow \mathsf{str}$ **Returns** the sequence of the peptide. Return type str **get_pos_in_parent** (*pro_id: str*) → Tuple[int, int] **Parameters** pro_id (str) – the id of the parent protein Raises ValueError – If the identifier is not a parent of the instance

Returns the start and end position of the instance peptide in the parent pointed out by the provided identifier

Return type Range

```
is\_child\_of(pro\_id: str) \rightarrow bool
```

Parameters pro_id (str) - is the protein id

Returns True if a user provided protein-id is a parent for the instance peptide, False otherwise

Return type bool

```
map_to_parent_protein() → List[numpy.ndarray]
```

Mapped the instance peptide to the parent protein and returned a list of numpy arrays where each array has a size of 1 by protein length. within the protein the range representing the peptide is encoded as one while the rest is zero.

Returns A list of binary encoded arrays represent this mapping.

Return type MappedProtein

IPTK.Classes.Proband module

A description for an IP proband

```
class IPTK.Classes.Proband(**info)
    Bases: object
```

```
\texttt{get\_meta\_data}\,(\,)\,\to dict
```

Returns a dict that contain all the meta-data about the patient

Return type dict

```
\mathtt{get}\_\mathtt{name}\left(\right) \to \mathrm{str}
```

Returns the name of the patient

Return type str

```
update_info(**info) \rightarrow None
```

add new or update existing info about the patient using an arbitrary number of key-value pair to be added to added to the instance meta-info dict

IPTK.Classes.Protein module

A representation of a protein that has been inferred from an IP experiment.

representation of a protein that has been infered from an IP experiment.

```
\mathtt{get\_id}() \rightarrow \operatorname{str}
```

Returns return the protein identifier.

Return type str

get_non_presented_peptide ($exc_reg_s_idx$: int, $exc_reg_e_idx$: int, length: int) \rightarrow str sample a peptide from the protein sequences where the sampled peptides is not part of the experimentally identified regions.

Parameters

- exc_reg_s_idx (int) the start point in the reference protein sequence of the experimentally identified peptide.
- **exc_reg_e_idx** (*int*) the end point in the reference protein sequence of the experimentally identified peptide.
- **length** (*int*) length the non-presented peptides.

Raises

- **ValueError** if the length of the peptide is bigger than the protein length
- **ValueError** if the length of the peptide is smaller than or equal to zero

Returns a substring of the instance sequence

Return type str

```
\mathtt{get\_org}() \to \mathrm{str}
```

Returns the organism in which this instance protein belong.

Return type str

get_peptides_map (start_idxs: List[int], end_idxs: List[int]) → numpy.ndarray compute a coverage over the protein sequence

Parameters

- **start_idxs** (*Index*) a list of integers representing the start positions
- end_idxs a list of integers representing the end positions

Raises ValueError – if start_indxs and end_idxs MUST be of equal length are not of equal length

Returns A numpy array with shape of 1 by the length of the protein where every element in the array donates the number of times, It has been observed in the experiment.

Return type np.ndarray

```
get\_seq() \rightarrow str
```

Returns the protein sequence.

Return type str

```
set\_org(org: str) \rightarrow None
```

a post-instantitation mechanism to set the organism for which the protein belong.

Parameters org(str) – the name of the organism

IPTK.Classes.Tissue module

A representation of the Tissue used in an IP Experiment.

Bases: object

a representation of tissue reference expression value.

```
get\_gene\_id\_expression (gene\_id: str) \rightarrow float
                Parameters gene_id (str) – the gene id to retrive its expression value from the database
                Raises KeyError – if the provided id is not defined in the instance table
                Returns the expression value of the provided gene id.
                Return type float
      get\_gene\_name\_expression (gene\_name: str) \rightarrow float
                Parameters gene_name (str) - the gene name to retrive its expression value from the
                    database
                Raises KeyError – if the provided id is not defined in the instance table
                Returns the expression value of the provided gene name.
                Return type float
      \mathtt{get}\_\mathtt{name}\left(\right) \to \mathrm{str}
                Returns the name of the tissue which the instance profile its gene expression
                Return type str
      \texttt{get\_table} () \rightarrow pandas.core.frame.DataFrame
                Returns return a table that contain the expression of all the transcript in the current profile in-
                    cluding core and auxiliary proteins
                Return type pd.DataFrame
class IPTK.Classes.Tissue.Tissue(name:
                                                                                             main_exp_value:
                                                                            str,
                                                 IPTK.Classes.Database.GeneExpressionDB, main_location:
                                                 IPTK.Classes.Database.CellularLocationDB, aux_exp_value:
                                                 Optional/IPTK.Classes.Database.GeneExpressionDB/
                                                           None,
                                                                           aux location:
                                                 tional/IPTK.Classes.Database.CellularLocationDB/
                                                 None)
      Bases: object
      \texttt{get\_expression\_profile} () \rightarrow IPTK.Classes.Tissue.ExpressionProfile
                Returns the expresion profile of the current tissue
```

Return type ExpressionProfile

 $\texttt{get}_\texttt{name} \ (\) \ \to str$

Returns the name of the tissue

Return type str

get subCellular locations () \rightarrow IPTK. Classes. Database. Cellular Location DB

Returns the sub-cellular localization of all the proteins stored in current instance resources.

Return type *CellularLocationDB*

Module contents

IPTK.IO package

Submodules

IPTK.IO.InFunctions module

Parse different user inputs into a standard format/tables used by the library.

IPTK.IO.InFunctions.download_pdb_entry($prot_id: str$) \rightarrow str Download the structure of a protein from protein databank form as mmCIF file.

Parameters prot_id (str) - the protein id

Raises IOError – incase downloading and accessing the data failed

Returns the path to the downloaded file

Return type str

IPTK.IO.InFunctions.fasta2dict($path2fasta: str, filter_decoy: bool = True, decoy_string: str = 'DECOY'$) \rightarrow Dict[str, str]

loads a fasta file and construct a dict object where ids are keys and sequences are the value

Parameters

- path2fasta(str) The path to load the fasta file
- **filter_decoy** (bool, optional) A boolean of whether or not to filter the decoy sequences from the database, defaults to True
- **decoy_string** (*str*, *optional*) The decoy database prefix, only valid incase filter_decoy is set to true, defaults to 'DECOY'

Raises IOError – [description]

Returns a dict where the protein ids are the keys and the protein sequences are the value

Return type Dict[str,str]

IPTK.IO.InFunctions.load_identification_table ($input_path: str, sep: str$) \rightarrow pandas.core.frame.DataFrame load & process an identification table

Parameters

- **input_path** (*str*) the path two the identification table. with the following columns: peptides which hold the identified peptide sequence, protein which hold the identified protein sequence, start_index, and end_index where the last two columns define the position of the peptide in the parent protein.
- **sep** (str) The separator to parse the provided table.

Raises

- IOError [description]
- ValueError [description]

Returns [description]

Return type pd.DataFrame

```
IPTK.IO.InFunctions.parse_mzTab_to_identification_table (path2mzTab: str, path2fastaDB: str, fasta\_reader\_param: Dict[str, str] = \{'decoy\_string': 'DECOY', 'filter\_decoy': True\}) \rightarrow pandas.core.frame.DataFrame
```

parse a user provided mzTab to an identification table

Parameters

- path2mzTab (str) the path to the input mzTab file
- path2fastaDB (str) the path to a fasta sequence database to obtain the protein sequences
- **fasta_reader_param** (Dict[str,str], optional) a dict of parameters for controlling the behavior of the fasta reader, defaults to {'filter_decoy':True, 'decoy string':'DECOY' }

Raises

- IOError if the mztab file could not be open and loaded or if the fasta database could not be read
- **KeyError** if a protein id defined in the mzTab file could not be extracted from a matched sequence database
- ValueError if the peptide can not be mapped to the identified protein

Returns the identification table

Return type pd.DataFrame

```
IPTK.IO.InFunctions.parse_text_table (path2file: str, path2fastaDB: str, sep=',', fasta_reader_param: Dict[str, str] = {'decoy_string': 'DECOY', 'filter_decoy': True}, seq_column: str = 'Sequence', accession_column: str = 'Protein Accessions', protein_group_sep: str = ';', remove_protein_version: bool = True, remove_if_not_matched: bool = True) \rightarrow pandas.core.frame.DataFrame
```

Parse a user defined table to extract the columns containing the identification table

Parameters

- path2file (str) The path to load the CSV file holding the results
- path2fastaDB (str) The path to a fasta sequence database to obtain the protein sequences
- **sep** (str, optional) The table separators, defaults to ','
- **fasta_reader_param** (Dict[str,str], optional) a dict of parameters for controlling the behavior of the fasta reader, defaults to {'filter_decoy':True, 'decoy_string':'DECOY' }
- **seq_column** (*str*, *optional*) The name of the columns containing the peptide sequence, defaults to 'Sequence'
- accession_column (str, optional) The name of the column containing the protein accession, defaults to 'Protein Accessions'
- protein_group_sep (str, optional) The separator for the protein group,, defaults to ';'

- remove_protein_version (bool, optional) A bool if true strip the version number from the protein, defaults to True
- remove_if_not_matched (bool, optional) remove the peptide if it could not be matched to the parent protein, defaults to True

Raises

- IOError Incase either the sequences database or the identification table can not be open and loaded
- **KeyError** In case the provided column names not in the provided identification table.
- **KeyError** Incase the protein sequence can not be extract from the sequence database
- ValueError incase the peptide could not be located in the protein sequence

Returns an identification table

Return type pd.DataFrame

```
IPTK.IO.InFunctions.parse_xml_based_format_to_identification_table (path2XML_file:
```

```
str.
path2fastaDB:
str,
       de-
coy_prefix:
str
'DECOY',
is idXML:
bool
False.
fasta_reader_param:
Dict[str,
str]
      {'de-
coy_string':
'DECOY',
'fil-
ter_decoy':
True}, re-
move_if_not_matched:
bool
True)
      pan-
\rightarrow
das.core.frame.DataFrame
```

parse either a pepXML or an idXML file to generate an identification table,

Parameters

- path2XML_file (str) The path to the input pepXML files
- path2fastaDB (str) The path to a fasta sequence database to obtain the protein sequences
- **decoy_prefix** (*str*, *optional*) the prefix of the decoy sequences, default is DE-COY, defaults to 'DECOY'
- **is_idXML** (bool, optional) Whether or not the provided file is an idXML, default is false which assume the provided file is a pepXML file, defaults to False

32 Chapter 3. Funding:

- **fasta_reader_param** (Dict[str,str], optional) a dict of parameters for controlling the behavior of the fasta reader, defaults to {'filter_decoy':True, 'decoy_string':'DECOY'}
- remove_if_not_matched (bool, optional) remove the peptide if it could not be matched to the parent protein, defaults to True

Raises

- **IOError** if the fasta database could not be open
- **ValueError** if the XML file can not be open
- **KeyError** if a protein id defined in the mzTab file could not be extracted from a matched sequence database
- **ValueError** if the peptide can not be mapped to the identified protein

Returns the identification table

Return type pd.DataFrame

IPTK.IO.MEMEInterface module

The module contains functions to to call meme software via a system call.

```
IPTK.IO.MEMEInterface.call_meme (input_fasta_file: str, output_dir: str, verbose: bool = True, objunc: str = 'classic', test: str = 'mhg', use_llr: bool = False, shuf: int = 2, hsfrac: float = 0.5, cefrac: float = 0.25, search-size: int = -1, maxsize: int = -1, norand: bool = False, csites: int = -1, seed: int = -1, mod: str = 'oops', nmotifs: int = -1, evt: float = -1.0, time: int = -1, nsite: int = -1, minsites: int = -1, maxsite: int = -1, nomatrim: bool = False, wg: int = -1, ws: int = -1, noendgaps: bool = False, maxiter: int = -1, prior: str = 'dirichlet', b: int = -1, p: int = -1) \rightarrow None
```

warper for making a system call to meme software for sequence motif finding for the reset of the function parameters use the function **get_meme_help** defined in the module IO, submodule MEMEInterface.

Parameters

- input_fasta_file (str) The path to input FASTA files.
- output_dir (str) the output dir to write the results, IT WILL OVERWRITE EX-ISTING DIRECTORY
- **verbose** (bool) whether or not to print the output of calling meme to the screen, default is True.

IPTK.IO.MEMEInterface.get_meme_help() \to None print the command line help interface for the meme tool

Raises FileNotFoundError – if meme is not callable

<code>IPTK.IO.MEMEInterface.is_meme_callable()</code> $\rightarrow bool$

Returns True if meme is callable, False otherwise.

Return type bool

IPTK.IO.OutFunctions module

Write the results generated by the library into a wide variety of formats.

```
IPTK.IO.OutFunctions.write_annotated_sequences(peptides: List[str], labels: List[int], path2write: str, sep: str = ',', shuffle: bool = True) \rightarrow None
```

take a list of peptides along with it sequences and write the results to a CSV file.

Parameters

- **peptides** (List[str]) a list of peptide sequences
- labels (List[int]) a list of numerical labels associated with the peptides
- path2write (str) the path to write the generated file
- **sep** (str, optional) The separator in the resulting table, defaults to ','
- shuffle (bool, optional) Whether or not to shuffle the table, defaults to True

Raises

- **ValueError** incase the length of the tables and labels is not matching
- **IOError** In case writing the output table failed

```
IPTK.IO.OutFunctions.write_auto_named_peptide_to_fasta (peptides:
```

```
List[IPTK.Classes.Peptide.Peptide],
```

 $output_file: str) \rightarrow None$

Takes a list of peptides, generate automatic names for the peptides and write the results to the disk as FASTA files

Parameters

- **peptides** (*Peptides*) a list of peptide sequences
- output_file (str) the name of the output file to write the results to, it will OVER-WRITE existing files

```
IPTK.IO.OutFunctions.write_mapped_tensor_to_h5py (tensor: numpy.ndarray, path2write: str, dataSet_name: str = 'MAPPED_TENSOR') \rightarrow None
```

Write a mapped tensor to an hdf5 file

Parameters

- **tensor** (np. ndarray) The provided tensor to write it to the hdf5 file.
- path2write (str) The path of the output file
- dataSet_name(str, optional) The name of the dataset inside the mapped tensor, defaults to 'MAPPED_TENSOR'

Raises IOError – In case opening the file for writing failed

Takes a list of names and peptide sequences and writes them as an output file to the disk as fasta files

Parameters

- names (Names) A list of sequences names
- peptides (Peptides) A list of peptide sequences

output_file (str) – The name of the output file to write the results to, it will OVER-WRITE existing files

Raises

- ValueError Incase the length of the tables and labels is not matching
- **IOError** In case writing the output file failed

Takes a file and write the peptides to .pep file which is a text file that contain one peptide per line

Parameters

- **peptides** (*Peptides*) a list of peptide sequences
- output_file (str) the name of the output file to write the results to, it will OVER-WRITE existing files

Raises IOError – In case writing the output file failed

Module contents

IPTK.Utils package

Submodules

IPTK.Utils.DevFunctions module

The module contain functions that can be used for developing & testing other functions of the library

```
IPTK.Utils.DevFunctions.generate_random_peptide_seq(peptide_length: int, num\_peptides: int) \rightarrow List[str] generate a list of random peptides for testing and developing purposes.
```

Parameters

- peptide_length (int) The peptide length
- num_peptides (int) The number of peptides in the generate list

Returns a list of random peptides

simulate an IP identification table from a fasta file. Please Note, if the reminder of num_pep over num_prot does not equal to zero, the floor of this ratio will be used to sample peptides from each proteins

Parameters

• path2load (str) - The path to load the Fasta files

- num_pep (int) The number of peptides in the tables
- num_prot (int) The number of UNIQUE proteins in the table

Raises ValueError – if number of proteins or number of peptide is zero

Returns an identification table

Return type pd.DataFrame

IPTK.Utils.DevFunctions.simulate_an_expression_table ($num_transcripts: int = 100$) \rightarrow pandas.core.frame.DataFrame create a dummy expression table to be used for testing and developing Tissue based classes

Parameters num_transcripts (int, optional) – The number of transcripts that shall be contained in the transcript, defaults to 100

Raises ValueError – incase number of transcripts is 0

Returns [description]

Return type pd.DataFrame

IPTK.Utils.DevFunctions.simulate_mapped_array_list ($min_len: int = 20, max_len: int = 100, num_elem: int = 100) <math>\rightarrow$ List[numpy.ndarray]

Simulate a list of mapped arrays proteins to be used for developing purposes

Parameters

- min_len (int, optional) the minmum length of the protein, defaults to 20
- $max_len(int, optional)$ the maximum length for the protein , defaults to 100
- num_elem (int, optional) the number of arrays in the protein, defaults to 100

Returns a list of simulated NumPy array that represent protein peptide coverage

Return type List[np.ndarray]

```
IPTK.Utils.DevFunctions.simulate_random_experiment (alleles: List[str], path2fasta: str, tissue_name: str = {}^{'}TEST\_TISSUE', num_pep: int = 10, num_prot: int = 5, proband_name: str = None) \rightarrow IPTK.Classes.Experiment.Experiment
```

Simulate a random experiment objects

Parameters

- alleles (List[str]) a list of alleles names.
- path2fasta (str) The path to load the database objects
- tissue_name (str, optional) the name of the tissue, defaults to 'TEST_TISSUE'
- $num_pep(int, optional)$ the number of peptides in the table, defaults to 10
- num prot (int, optional) number of proteins, defaults to 5
- proband_name (str, optional) The name of the Proband, defaults to None

Returns a simulated experimental object

Return type Experiment

IPTK.Utils.Mapping module

A submodule that contain function to map different database keys

```
IPTK.Utils.Mapping.map_from_uniprot_gene (uniprots: List[str]) \rightarrow pandas.core.frame.DataFrame map from uniprot id to ensemble gene ids
```

Parameters uniprots (List[str]) – a list of uniprot IDs

Returns A table that contain the mapping between each uniprot and its corresponding Gene ID/IDs

Return type pd.DataFrame

```
IPTK.Utils.Mapping.map_from_uniprot_pdb (uniprots: List[str]) \rightarrow pandas.core.frame.DataFrame map from uniprot id to protein data bank identifiers
```

Parameters uniprots (List[str]) – a list of uniprot IDs

Returns A table that contain the mapping between each uniprot and its corresponding PDB ID/IDs

Return type pd.DataFrame

IPTK.Utils.Types module

Contain a definition of commonly used types through the library

IPTK.Utils.UtilityFunction module

Utility functions that are used through the library

```
IPTK.Utils.UtilityFunction.append_to_calling_string(param: str, def_value, cur_val, calling_string: str, is_flag: bool = False) \rightarrow str
```

help function that take a calling string, a parameter, a default value and current value if the parameter does not equal its default value the function append the parameter with its current value to the calling string adding a space before the calling_string.

Parameters

- param (str) The name of the parameter that will be append to the calling string
- **def_value** ([type]) The default value for the parameter
- **cur_val** ([type]) The current value for the parameter
- calling_string (str) The calling string in which the parameter and the current value might be appended to it
- **is_flag** (bool, optional) If the parameter is a control flag, i.e. a boolean switch, it append the parameter to the calling string without associating a value to it, defaults to False

Returns the updated version of the calling string

Return type str

```
\label{lem:construct} \begin{tabular}{ll} \end{tabular} IPTK. Utils. UtilityFunction. build_sequence_table (sequence_dict: Dict[str, str]) $\rightarrow$ pandas.core.frame. DataFrame construct a sequences database from sequences dict object \\ \end{tabular}
```

Parameters sequence_dict (Dict[str, str]) – a dict that contain the protein ids as keys and sequences as values.

Returns pandas dataframe that contain the protein ID and the associated protein sequence

Return type pd.DataFrame

IPTK.Utils.UtilityFunction.check_peptide_made_of_std_20_aa (peptide: str) → str Check if the peptide is made of the standard 20 amino acids, if this is the case, it return the peptide sequence, otherwise it return an empty string

Parameters peptide (str) – a peptide sequence to check its composition

Returns True, if the peptide is made of the standard 20 amino acids, False otherwise.

Return type str

 $\label{lem:color_scale} IPTK. Utils. UtilityFunction. \textbf{generate_color_scale} (color_ranges: int) \rightarrow \text{mat-plotlib.colors.} LinearSegmentedColormap generate a color gradient with number of steps equal to color_ranges -1$

Parameters color_ranges (int) - the number of colors in the range

Returns a color gradient palette

Return type matplotlib.colors.LinearSegmentedColormap

IPTK.Utils.UtilityFunction.generate_random_name(name_length: int) → str

Parameters name_length (int) - Generate a random ASCII based string

Returns [description]

Return type str

IPTK.Utils.UtilityFunction.generate_random_protein_mapping(protein_len: int, $max_coverage$: int) \rightarrow numpy.ndarray

Generate a NumPy array with shape of 1 by protein_len where the elements in the array is a random integer between zero & max_coverage.

Parameters

- **protein_len** (*int*) The length of the protein
- $max_coverage(int)$ The maximum peptide coverage at each position

Returns a NumPy array contain a simulate protein coverage

Return type np.ndarray

IPTK.Utils.UtilityFunction.get_idx_peptide_in_sequence_table (sequence_table:

pan- das.core.frame.DataFrame, $peptide: str) \rightarrow$ List[str]

check the sequences table if the provided peptide is locate in one of its sequences and returns a list of protein identifiers containing the identifier of the hit proteins.

Parameters

- **sequence_table** (pd.DataFrame) pandas dataframe that contain the protein ID and the associated protein sequence
- **peptide** (str) the peptide sequence to query the protein with

Returns a list of protein identifiers containing the identifier of the hit proteins

Return type List[str]

```
IPTK.Utils.UtilityFunction.load_3d_figure (file_path: str) → matplotlib.figure.Figure
```

Parameters file_path (str) - Load a pickled 3D figure from thr provided path

Raises IOError – The path of the pickled figure.

Returns a matplotlib figure

Return type plt. Figure

```
IPTK.Utils.UtilityFunction.pad_mapped_proteins (list_array: List[numpy.ndarray], pre\_pad: bool = True, padding\_char: int = -1) \rightarrow numpy.ndarray
```

Pad the provided list of array into a 2D tensor of shape number of arrays by maxlength.

Parameters

- list_array (List[np.ndarray]) A list of NumPy arrays where each array is a mapped_protein array, the expected shape of these arrays is 1 by protein length.
- **pre_pad** (bool, optional) pre or post padding of shorter array in the library.Default is pre-padding, defaults to True
- padding_char (int, optional) The padding char, defaults to -1

Returns A 2D tensor of shape number of arrays by maxlength.

Return type np.ndarray

IPTK.Utils.UtilityFunction.save_3d_figure (outpath: str, fig2save: matplotlib.figure.Figure)

None
write a pickled version of the a 3D figure so it can be loaded later for more interactive analysis

Parameters

- **outpath** (str) The output path of the writer function
- **fig2save** (plt.Figure) The figure to save to the output file

Raises IOError – In case writing the file failed

Parameters

- num_conditions (int) The number of conditions to simulate
- protein_length (int) The Length of the protein

Returns A 2D matrix of shape protein_length by number of conditions, where each element can be either zero.

Return type np.ndarray

```
IPTK.Utils.UtilityFunction.simulate_protein_representation (num\_conditions: int, protein\_len: int, protein\_coverage: int) \rightarrow Dict[str, numpy.ndarray]
```

Simulate protein peptide coverage under-different conditions

Parameters

- num_conditions ([type]) The number of condition to simulate
- protein_len ([type]) The length of the protein
- protein_coverage ([type]) The maximum protein coverage

Returns a dict of length num_conditions contains that the condition index and a simulated protein array

Return type Dict[str, np.ndarray]

Module contents

IPTK.Visualization package

Submodules

IPTK. Visualization. vizTools module

The module contain visualization functions the can be used to plot the results obtained from the methods of the classes defined in the Class module or from the analysis functions defined in the Analysis Module.

```
IPTK.Visualization.vizTools.imposed_coverage_on_3D_structure(path2mmCIF: str, mapped_protein: numpy.ndarray, background_color: str = 'black', low: str = 'red', high: str = 'blue') \rightarrow None
```

A function to impose the peptide coverage on top of a protein 3D structure where the color of each position is marked by a color gradient that reflect the number of peptides aligned to this position. Note: Use the function with Jupyter-note book as it return an NGLWidget object that your can explore and navigate from the browser.

Parameters

- path2mmCIF (str) The path to load the mmCIF file containing the protein structure
- mapped_protein (np.ndarray) a Numpy array of containg the number of peptides originated from each position in the array
- background_color(str, optional) the color of the background, default is black , defaults to 'black'
- low (str, optional) the color of low covered position, default is red. , defaults to 'red'
- high (str, optional) the color of high covered position, default is violet., defaults to 'blue'

Raises

- **ValueError** incase the provided path to the structure file does not exists
- **IOError** if the structure file can not be loaded or if more than one file are located in the provided path

```
IPTK. Visualization. vizTools.plot_change_in_presentation_between_experiment (change_in_presentation numpy.ndarray,
```

index_first: int, index second: int, plotting_kwargs: Dict[str, str] {}, title='Change inprotein presentation', xlabel='Proteins', ylabel = 'magnitudeofchange inprotein count') \rightarrow mat-

plotlib.figure.Figure

plot the change in protein presentation between two experiment

Parameters

- **change_in_presentation_array** (np.ndarray) a 3D tensor of shape number of experiments by number of experiment by number of identified proteins.
- index_first (int) [description]
- index_second (int) the index of the first experiment in the tensor
- plotting_kwargs (Dict[str, str], optional) a dict object containing parameters for the sns.scatterplot function, defaults to {}
- title (str, optional) The title of the figure, defaults to 'Change in protein presentation'
- xlabel (str, optional) The axis on the x-axis, defaults to "Proteins"
- ylabel (str, optional) The axis on the y-axis, defaults to "magnitude of change in protein count"

Raises

- **ValueError** if the provided tensor is not of rank 3
- **IndexError** if the provided indices are out of bound

```
IPTK.Visualization.vizTools.plot_experiment_set_counts (counts_table: pandas.core.frame.DataFrame, log_scale: bool = False, plotting_kwargs: Dict[str, str] = \{\}) \rightarrow matplotlib.pyplot.figure
```

visualize the number of peptides and number of peptides-per-organism per experiment.

Parameters

- counts_table (pd.DataFrame) a pandas dataframe that contain the count, organism name and the count
- log_scale (bool, optional) Normalize the peptide counts one log 10, defaults to False
- plotting_kwargs (Dict[str,str], optional) a dict object containing parameters for the sns.catplot function, defaults to {}

```
IPTK. Visualization.vizTools.plot_gene_expression_vs_num_peptides (exp_count_table:
```

```
pan-
das.core.frame.DataFrame,
tissue name:
str,
def value:
float =
1.
        plot-
ting_kwargs:
Dict[str,
str] = {},
xlabel:
    'Number
of peptides',
ylabel:
     'Expres-
sion value',
title: str =
'Peptides
per
      protein
     Expres-
Vs.
sion Level')
        mat-
plotlib.figure.Figure
```

Plot the correlation between the gene expression and the num of peptids per protein using seaborn library

Parameters

- **exp_count_table** (pd.DataFrame) A table that contain the number of peptides and the expresion value for each protein in the database
- tissue_name (str) The name of the tissue
- **def_value** (*float*, *optional*) The default value for proteins that could not be mapped to the expression database, defaults to -1

- plotting_kwargs (Dict[str,str], optional) a dict object containing parameters for the sns.scatter function, defaults to {}
- xlabel (str, optional) the label on the x-axis, defaults to 'Number of peptides'
- ylabel (str, optional) the label on the y-axis, defaults to 'Expression value'
- **title** (str, optional) The title of the figure, defaults to 'Peptides per protein Vs. Expression Level'

```
IPTK. Visualization. vizTools.plot_motif (pwm\_df: pandas.core.frame.DataFrame, plot-ting\_kwargs: Dict[str, str] = \{'fade\_probabilities': True\}\} \rightarrow matplotlib.figure.Figure
```

A generic motif plotter that forward its argument to logomaker for making plots

Parameters

- pwm_df (pd. DataFrame) A pandas dataframe containing the position weighted matrix
- **plotting_kwargs** (*PlottingKeywards*, *optional*) A dictionary of parameter to control the behavior of the logo plotter, defaults to {'fade_probabilities':True}

Returns a matplotlib figure instance containing the ploted motif

Return type plt. Figure

```
IPTK. Visualization.vizTools.plot_num_peptide_per_go_term (pep2goTerm: pan-das.core.frame.DataFrame, plotting\_kwargs: Dict[str, str] = \{\}, drop\_unknown: bool = False, xlabel: str = 'Number of peptides', ylabel: <math>str = 'GO-Term', title: str = 'Number of peptides per GO Term') \rightarrow matplotlib.figure. Figure
```

plot the number of peptides obtained per Go-Term using matplotlib library.

Parameters

- pep2goTerm (pd.DataFrame) A table that contain the count of peptides from each GO-Term
- plotting_kwargs (Dict[str,str], optional) a dict object containing parameters for the sns.barplot function, defaults to {}
- **drop_unknown** (bool, optional) whether or not to drop peptide with unknown GO-term, defaults to False
- xlabel (str, optional) the label on the x-axis, defaults to 'Number of peptides'
- ylabel (str, optional) the label on the y-axis, defaults to 'GO-Term'
- title (str, optional) The title of the figure, defaults to 'Number of peptides per GO Term'

Returns [description]

Return type plt. Figure

```
\begin{tabular}{ll} \label{table:per_location} IPTK. Visualization. vizTools.plot_num_peptides_per_location (pep2loc: pan-das.core.frame.DataFrame, plotting_kwargs: \\ Dict[str, str] = \{\}, \\ drop\_unknown: bool \\ = False, xlabel: str = 'Number of peptides', ylabel: str = 'Compartment', title: str = 'Number of peptides per sub-cellular compartment') <math>\rightarrow matplotlib.figure. Figure \begin{tabular}{ll} Pan-location (pep2loc: pan-das.core.frame.DataFrame, plotlib.figure.Figure) & pan-das.core.frame.DataFrame, plotlib.figure.Figure & pan-das.core.frame.DataFrame, plotlib.figure.Figure & plotlib.figure.Figure & pan-das.core.frame.DataFrame, plotlib.figure.frame.DataFrame, plotlib.figure.Figure & pan-das.core.frame.DataFrame, plotlib.figure.frame.DataFrame, plotlib.figure.frame.DataFrame, plotlib.figure.frame.DataFrame, plotlib.figure.frame.DataFrame, plotlib.frame.DataFrame, plotlib.frame.DataFrame, plotli
```

plot the number of peptides obtained from each compartment using seaborn library.

Parameters

- pep21oc (pd.DataFrame) A table that contain the count of peptides from each location
- plotting_kwargs (Dict[str, str], optional) a dict object containing parameters for the sns.barplot function, defaults to {}
- **drop_unknown** (bool, optional) whether or not to drop protein with unknown location, defaults to False
- **xlabel** (str, optional) The label on the x-axis, defaults to 'Number of peptides'
- ylabel (str, optional) The label on the y-axis, defaults to 'Compartment'
- **title** (*str*, *optional*) The title of the figure, defaults to 'Number of peptides per sub-cellular compartment'

```
IPTK. Visualization.vizTools.plot_num_peptides_per_organism(pep_per_org:
                                                                                                pan-
                                                                                das.core.frame.DataFrame,
                                                                                log_scale:
                                                                                                bool
                                                                                     False,
                                                                                                plot-
                                                                                ting_kwargs: Dict[str,
                                                                                str] = {}, xlabel:
                                                                                str = 'Number of
                                                                               peptides',
                                                                                              ylabel:
                                                                                          'Organism',
                                                                                title: str = 'Num-
                                                                                ber of peptides per
                                                                                organism') \rightarrow mat-
                                                                               plotlib.figure.Figure
     plot the number of peptides per each organism inferred from the experiment using seaborn and matlotlib.
```

Parameters

- **pep_per_org** (pd.DataFrame) A table that contain the number of peptides belonging to each organism
- log_scale (bool, optional) Whether or not to scale the number of peptides using a log scale, default is False, defaults to False
- plotting_kwargs (Dict[str, str], optional) a dict object containing parameters for the sns.barplot function, defaults to {}
- **xlabel** (str, optional) the label on the x-axis, defaults to 'Number of peptides'

- ylabel (str, optional) The label on the y-axis, defaults to 'Organism'
- **title** (*str*, *optional*) The title of the figure, defaults to 'Number of peptides per organism'

```
IPTK.Visualization.vizTools.plot_num_peptides_per_parent (nums_table: pan-das.core.frame.DataFrame, num_prot: int = -1, plot-ting_kwargs: Dict[str, str] = {}, x_label: str = 'Number of peptides', y_label: str = 'Protein ID', title: str = 'Number of peptides per protein')
```

Visualize a histogram of the eluted peptide length.

Parameters

- nums_table (pd.DataFrame) a pandas dataframe containing number of peptides identified from each protein.
- num_prot (int, optional) the number of protein to show relative to the first element, for example, the first 10, 20 etc. If the default value of -1 is used then all protein will be plotted, however, this might lead to a crowded figure, defaults to -1.
- plotting_kwargs a dict object containing parameters for the function seaborn::distplot, defaults to {}
- **x_label** (str, optional) the label of the x-axis, defaults to 'Number of peptides'
- y_label (str, optional) the label of the y-axis, defaults to 'Protein ID'
- **title** (*str*, *optional*) The title of the figure, defaults to 'Number of peptides per protein'

Raises ValueError - if the num_prot is bigger than the number of elements in the provided table

```
IPTK. Visualization.vizTools.plot_num_protein_per_go_term (protein2goTerm:
                                                                           das.core.frame.DataFrame,
                                                                           tissue_name: str, plot-
                                                                           ting_kwargs:
                                                                                           Dict[str,
                                                                           str] = {}, drop\_unknown:
                                                                           bool = False, xlabel:
                                                                           str = 'Number of Pro-
                                                                           teins', ylabel:
                                                                                            str =
                                                                            'Compartment',
                                                                                              title:
                                                                           str = 'Number of pro-
                                                                           teins per sub-cellular
                                                                           compartment') \rightarrow mat-
                                                                           plotlib.figure.Figure
```

plot the number of proteins per each GO Term

Parameters

- protein2goTerm (pd.DataFrame) A table that contain the count of proteins from each GO-Term
- **tissue_name** (str) a dict object containing parameters for the sns.barplot function.
- plotting_kwargs (Dict[str, str], optional) a dict object containing parameters for the sns.barplot function, defaults to {}

- drop_unknown (bool, optional) whether or not to drop protein with unknown location, defaults to False
- xlabel (str, optional) the label on the x-axis, defaults to 'Number of Proteins'
- ylabel (str, optional) the label on the y-axis, defaults to 'Compartment'
- **title** (*str*, *optional*) the title of the figure, defaults to 'Number of proteins per sub-cellular compartment'

```
IPTK. Visualization. vizTools.plot_num_protein_per_location (protein_loc:
                                                                                           pan-
                                                                          das.core.frame.DataFrame,
                                                                          plotting_kwargs:
                                                                          Dict[str, str] = \{\},
                                                                          drop_unknown:
                                                                                           bool
                                                                          = False, xlabel: str
                                                                          = 'Number of Pro-
                                                                          teins', ylabel: str =
                                                                           'Compartment',
                                                                                           title:
                                                                          str = 'Number of pro-
                                                                          teins per sub-cellular
                                                                          compartment')
                                                                          matplotlib.figure.Figure
```

plot the number of proteins per each sub-cellular compartment

Parameters

- protein_loc (pd.DataFrame) A table that contain the count of protein from each location.
- plotting_kwargs (Dict[str, str], optional) a dict object containing parameters for the sns.barplot function, defaults to {}
- **drop_unknown** (bool, optional) whether or not to drop protein with unknown location, defaults to False
- xlabel (str, optional) the label on the x-axis, defaults to 'Number of Proteins'
- ylabel (str, optional) the label on the y-axis, defaults to 'Compartment'
- **title** (*str*, *optional*) the title of the figure, defaults to 'Number of proteins per sub-cellular compartment'

```
IPTK.Visualization.vizTools.plot_overlap_heatmap (results_df: pandas.core.frame.DataFrame, plotting_kwargs: Dict[str, str] = \{\}) \rightarrow seaborn.matrix.ClusterGrid
```

Plot a user provided dataframe as a cluster heatmap using seaborn library

Parameters

- results_df (pd.DataFrame) A pandas dataframe table that hold the overlapping number
- plotting_kwargs (PlottingKeywards, optional) forward parameter to the clustermap function, defaults to {}

Returns the generated clustermap

Return type sns.matrix.ClusterGrid

```
IPTK. Visualization.vizTools.plot_overlay_representation (proteins: Dict[str, Dict[str, numpy.ndarray]], alpha: float = 0.5, title: str = None, legend_pos: int = 2) \rightarrow matplotlib.figure.Figure
```

plot an overlayed representation for the SAME protein or proteins OF EQUAL LENGTH in different conditions in which the mapped representation of each protein are stacked on top of each other to generate a representation for the protein representability under different conditions.

Parameters

- **proteins** (Dict[str, np.ndarray]]) a nested dict object containing for each protein a child dict that contain the mapping array and the color in the figure.
- alpha (float, optional) the transparency between proteins, defaults to 0.5
- title (str, optional) The title of the figure, defaults to None
- legend_pos (int, optional) the position of the legend, defaults to 2

Raises ValueError – if the provided protein have different lengths

Returns a matplotlib figure containing the mapped representation

Return type plt. Figure

```
IPTK.Visualization.vizTools.plot_paired_represention (protein_one_repr: Dict[str, numpy.ndarray], protein\_two\_repr: Dict[str, numpy.ndarray], color\_first: str = 'red', color\_second: str = 'blue', alpha: float = 0.9, title=' Parired protein representation') <math>\rightarrow matplotlib.figure.Figure
```

visualize the difference in representation for the same protein between two experiments using matplotlib library.

Parameters

- **protein_one_repr** (Dict[str, np.ndarray]) a dict object containing the legand of the first condition along with its mapped array
- protein_two_repr (Dict[str, np.ndarray]) a dict object containing the legand of the second condition along with its mapped array
- alpha (the transparency of the figure.) the transparency of the figure.

Param color_first: the color of representation for the first condition

Param color_second: the color of the second condition

Param title: the title of the figure.

Returns A matplotlib Figure containing the representation

Return type plt. Figure

IPTK. Visualization.vizTools.plot_parent_protein_expression_in_tissue (expression_table:

```
pan-
das.core.frame.DataFrame,
ref_expression:
pan-
das.core.frame.DataFrame,
tis-
sue_name:
str,
sam-
pling_num:
int
10.
plot-
ting_kwargs:
Dict[str,
str] =
{'ori-
ent':
v',
def_value:
float
= -1,
ylabel:
str
'Nor-
mal-
ized
Expres-
sion')
\rightarrow mat-
plotlib.figure.Figure
```

Plot the parent protein expression in tissue relative a sampled collection of non-presented data using seaborn library.

Parameters

- **expression_table** (pd.DataFrame) The protein expression table which contains the expression value for each parent protein
- ref_expression (pd.DataFrame) The reference expression of the tissue under investigation.
- **tissue_name** (str) The name of the tissue.
- **sampling_num** (*int*, *optional*) The number of times to sample from the non-prsenter., defaults to 10
- plotting_kwargs (Dict[str, str], optional) a dict object containing parameters for the sns.violinplot function., defaults to {'orient':'v'}
- **def_value** (*float*, *optional*) The default value for proteins that could not be mapped to the expression database, defaults to -1
- ylabel (str, optional) the label on the y-axis., defaults to 'Normalized Expression'

Raises ValueError - if the reference gene expression table is smaller than the number of parents

```
IPTK. Visualization. vizTools.plot_peptide_length_dist (pep\_length: List[int], plot-ting_kwargs: Dict[str, str] = {}, x_label: str = 'Peptide Length', y_label: str = 'Frequency', title: str = 'Peptide Length distribution')
```

Visualize a histogram of the eluted peptide length using seaborn library.

Parameters

- pep_length (List[int]) [description]
- plotting_kwargs (Dict[str, str], optional) a dict object containing parameters for the function seaborn::distplot, defaults to {}
- x_label (str, optional) the label of the x-axis, defaults to 'Peptide Length'
- **y_label** (str, optional) the label of the y-axis, defaults to 'Frequency'
- title (str, optional) the title of the figure, defaults to 'Peptide Length distribution'

```
IPTK. Visualization.vizTools.plot_peptide_length_per_experiment (counts_table:
```

```
pan-
das.core.frame.DataFrame,
plot-
ting_kwargs:
Dict[str, str]
= {}) → mat-
plotlib.pyplot.figure
```

visualize the peptide length distribution among the experiments defined in the set

Parameters

- **counts_table** (pd.DataFrame) a pandas dataframe that contain the length of each peptide defined in the experiment along with the experiment name
- plotting_kwargs (Dict[str,str], optional) a dict object containing parameters for the sns.catplot function, defaults to {}

```
IPTK. Visualization. vizTools.plot_protein_coverage (mapped_protein: numpy.ndarray, col: str = 'r', prot_name: str = None) \rightarrow matplotlib.figure. Figure
```

plot the peptide coverage for a given protein.

Parameters

- mapped_protein (np.ndarray) a NumPy array with shape of 1 by protein length or shape protein-length
- col (str, optional) the color of the coverage respresentation, defaults to 'r'
- prot_name (str, optional) The default protein name, defaults to None

Return type plt. Figure

IPTK. Visualization. vizTools.plot_protein_presentation_3D (proteins:

```
Dict[str, Dict[str, numpy.ndarray]], plot-
ting_args={'color': 'red'}, title: str
= None) \rightarrow mat-
plotlib.figure.Figure
```

plot a 3D surface representation for the SAME protein or proteins OF EQAUL LENGTH in different conditions.

Parameters

- **proteins** ([type]) a nested dict object containing for each protein a child dict that contain the mapping array and the color in the figure.
- plotting_args (dict, optional) a dict that contain further parameter to the plot surface functions, defaults to {'color':'red'}
- title (str, optional) the title of the figure, defaults to None

Raises ValueError – if the provided proteins are of different length

Return type plt. Figure

IPTK. Visualization. vizTools.plotly_gene_expression_vs_num_peptides (exp_count_table:

```
das.core.frame.DataFrame,
tis-
sue_name:
str.
def_value:
float = -1,
xlabel: str
   'Num-
ber
         of
peptides',
ylabel:
str = 'Ex-
pression
value',
title: str =
'Peptides
per pro-
        Vs.
tein
Protein
Expres-
sion
Level')
\rightarrow
      mat-
plotlib.figure.Figure
```

Plot the correlation between the gene expression and the number of peptids per protein using plotly library.

Parameters

- **exp_count_table** (pd. DataFrame) A table that contain the number of peptides and the expression value for each protein in the database
- **tissue_name** (str) The name of the tissue
- **def_value** (*float*, *optional*) The default value for proteins that could not be mapped to the expression database, defaults to -1
- xlabel (str, optional) The label on the x-axis, defaults to 'Number of peptides'
- ylabel (str, optional) The label on the y-axis., defaults to 'Expression value'
- **title** (*str*, *optional*) The title of the figure, defaults to 'Peptides per protein Vs. Protein Expression Level'

```
IPTK. Visualization. vizTools.plotly_multi_traced_coverage_representation (proteins:
```

```
Dict[str,
numpy.ndarray]],
ti-
tle:
str
=
'Pro-
tein
Cov-
er-
age
Across
')
→
plotly.graph_objs_figure.F.
```

Dict[str.

plot a multi-traced representation for the same protein across using plotly library

Parameters

- **proteins** ([type]) A dict object containing for each protein the corresponding mapped array.
- title (str, optional) the title of the figure, defaults to "Protein Coverage Across "

Returns a multitraced traced figure showing the coverage of proteins accross different conditions **Return type** Figure

```
IPTK.Visualization.vizTools.plotly_num_peptide_per_go_term (pep2goTerm: pan-das.core.frame.DataFrame, drop\_unknown: bool
= False, xlabel: str
= 'Number of peptides', ylabel: str = 'GO-Term', title: str = 'Number of peptides' per GO Term') <math>\rightarrow plotly.graph_objs._figure.Figure
```

plot the number of peptides obtained per Go-Term using plotly library.

Parameters

- pep2goTerm (pd.DataFrame) A table that contain the count of peptides from each GO-Term
- **drop_unknown** (bool, optional) whether or not to drop peptide with unknown GO-term, defaults to False
- xlabel (str, optional) the label on the x-axis, defaults to 'Number of peptides'
- ylabel (str, optional) the label on the y-axis, defaults to 'GO-Term'
- title (str, optional) the title of the figure, defaults to 'Number of peptides per GO Term'

```
IPTK.Visualization.vizTools.plotly_num_peptides_per_location (pep2loc: pan-das.core.frame.DataFrame, drop_unknown: bool = False, xlabel: str = 'Number of peptides', ylabel: str = 'Compartment', title: str = 'Number of peptides per sub-cellular compartment') \rightarrow matplotlib.figure.Figure
```

plot the number of peptides obtained from each compartment using plotly library

Parameters

- pep21oc (pd.DataFrame) A table that contain the count of peptides from each location
- **drop_unknown** (bool, optional) whether or not to drop protein with unknown location, defaults to False
- xlabel (str, optional) The label on the x-axis, defaults to 'Number of peptides'
- ylabel (str, optional) The label on the y-axis, defaults to 'Compartment'
- **title** (*str*, *optional*) The title of the figure, defaults to 'Number of peptides per sub-cellular compartment'

```
{\tt IPTK.Visualization.vizTools.plotly\_num\_peptides\_per\_organism} \ (\textit{pep\_per\_org: pan-peptides\_per\_organism}) \ \ \textit{pep\_per\_org: pan-peptides\_per\_organism} \ \ \textit{pep\_per\_org: pan-peptides\_per\_organism} \ \ \textit{pep\_per\_org: pan-peptides\_per\_org: pan-peptides\_per\_organism} \ \ \textit{pep\_per\_org: pan-peptides\_per\_org: pan-peptides\_per
```

```
das.core.frame.DataFrame,
log_scale: bool
= False, xlabel:
str = 'Number of
Peptides', ylabel:
str = 'Organism',
title: str = 'Num-
ber of peptides
per organism') →
plotly.graph_objs._figure.Figure
```

plot the number of peptides per each organism inferred from the experiment using plotly library.

Parameters

- **pep_per_org** (pd. DataFrame) A table that contain the count of peptides from each organism
- log_scale (bool, optional) Whether or not to scale the number of peptide using a log scale, defaults to False
- xlabel (str, optional) The label on the x-axis, defaults to 'Number of Peptides'
- ylabel (str, optional) The label on the y-axis, defaults to 'Organism'
- **title** (*str*, *optional*) the title of the figure , defaults to 'Number of peptides per organism'

Visualize a histogram of the the number of peptides per each inferred protein.

Parameters

- nums_table (pd.DataFrame) a pandas dataframe containing number of peptides identified from each protein.
- num_prot the number of protein to show relative to the first element, for example, the first 10, 20 etc. If the default value of -1 is used then all protein will be plotted, however, this might lead to a crowded figure., defaults to -1 :type num_prot: int, optional
- **x_label** (str, optional) the label of the x-axis, defaults to 'Number of peptides'
- y_label (str, optional) the label of the y-axis, defaults to 'Protein ID'
- title (str, optional) title, defaults to 'Number of peptides per protein'

Raises ValueError - if the num_prot is bigger than the number of elements in the provided table

```
IPTK.Visualization.vizTools.plotly_num_protein_per_go_term (protein2goTerm: pan-das.core.frame.DataFrame, drop_unknown: bool = False, \ xlabel: \ str \\ = \ 'Number \ of \ Proteins', \ ylabel: \ str = \ 'GO-Term', \ title: \ str \\ = \ 'Number \ of \ proteins \\ per \ GO-Term') \ \rightarrow \\ \text{plotly.graph_objs._figure.Figure}
```

plot the number of proteins per each GO Term using plotly library

Parameters

- protein2goTerm (pd.DataFrame) A table that contain the count of proteins from each GO-Term
- **drop_unknown** (bool, optional) whether or not to drop protein with unknown location, defaults to False
- xlabel (str, optional) the label on the x-axis, defaults to 'Number of Proteins'
- ylabel (str, optional) the label on the y-axis, defaults to 'GO-Term'
- title (str, optional) the title of the figure, defaults to 'Number of proteins per GO-Term'

Returns [description]

Return type Figure

```
 \begin{tabular}{ll} {\tt IPTK.Visualization.vizTools.plotly\_num\_protein\_per\_location} & (protein\_loc: pan-das.core.frame.DataFrame, drop\_unknown: bool \\ &= False, xlabel: str \\ &= 'Number \ of \ Proteins', ylabel: str = 'Compartment', title: str = 'Number \ of \ proteins \ per \ sub-cellular \ compartment') \ \rightarrow \\ &= pan-das.core.frame.DataFrame, drop\_unknown: bool \\ &= False, xlabel: str \\ &= 'Number \ of \ Proteins', ylabel: str = 'Compartment', title: \ str = 'Number \ of \ proteins \ per \ sub-cellular \ compartment') \ \rightarrow \\ &= plotly.graph\_objs.\_figure.Figure \end{tabular}
```

plot the number of proteins per each sub-cellular compartment

Parameters

- protein_loc (pd.DataFrame) A table that contain the count of protein from each location
- **drop_unknown** (bool, optional) whether or not to drop protein with unknown location, defaults to False
- xlabel (str, optional) the label on the x-axis, defaults to 'Number of Proteins'
- ylabel (str, optional) the label on the y-axis, defaults to 'Compartment'
- **title** (*str*, *optional*) [description], defaults to 'Number of proteins per sub-cellular compartment'

```
\begin{tabular}{ll} \begin{tabular}{ll} IPTK. Visualization. vizTools. \textbf{plotly_overlap_heatmap} (\it{results\_df: pan-das.core.frame.DataFrame}) & \rightarrow & & & & & & \\ & & & & & & & & \\ & & & & & & & \\ & & & & & & & \\ & & & & & & & \\ & & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ &
```

Plot a user provided dataframe as a heatmap using plotly library

Parameters results_df (pd.DataFrame) – A pandas dataframe table that hold the overlapping number.

Returns a plotly Figure containing the heatmap

Return type Figure

```
IPTK.Visualization.vizTools.plotly_paired_representation(protein_one_repr:

Dict[str, numpy.ndarray],
protein_two_repr:
Dict[str, numpy.ndarray],
title: str = 'Parired protein representation') \rightarrow
```

plotly.graph_objs._figure.Figure compare the peptide coverage for the same protein under different conditions using the same protein using plotly library.

Parameters

- protein_one_repr (Dict[str, np.ndarray]) a dict object containing the legand of the first condition along with its mapped array
- protein_two_repr (Dict[str, np.ndarray]) a dict object containing the legand of the second condition along with its mapped array

Returns A plotly Figure containing the representation

Return type Figure

```
IPTK. Visualization. vizTools.plotly_parent_protein_expression_in_tissue (expression_table:
```

```
pan-
das.core.frame.DataFrame,
ref_expression:
pan-
das.core.frame.DataFrame,
tis-
sue_name:
str,
sam-
pling_num:
int
10,
def_value:
float
= -
1.
yla-
bel:
str
'Nor-
mal-
ized
Ex-
pres-
sion')
plotly.graph_objs._figure.Fig
```

plot the parent protein expression in tissue relative a sampled collection of non-presented genes using plotly library.

Parameters

- **expression_table** (pd.DataFrame) The protein expression table which contains the expression value for each parent proteins.
- **ref_expression** (pd.DataFrame) The reference expression of the tissue under investigation.
- **tissue_name** (str) The name of the tissue.
- **sampling_num** (*int*, *optional*) the number of times to sample from the non-prsenter, defaults to 10
- **def_value** (*float*, *optional*) The default value for proteins that could not be mapped to the expression database, defaults to -1
- ylabel (The label on the y-axis, optional) [description], defaults to 'Normalized Expression'

Raises ValueError – If the reference gene expression table is smaller than the number of parents

```
IPTK.Visualization.vizTools.plotly_peptide_length_dist (pep_length: List[int], x_label: str = 'Peptide Length', y_label: str = 'Counts', title: str = 'Peptide Length' distribution')
```

visualize a histogram of the eluted peptide length using plotly library

Parameters

- pep_length (List[int]) a list of integer containing the peptides' lengths
- **x_label** (str, optional) the label of the x-axis, defaults to 'Peptide Length'
- y_label (str, optional) the label of the y-axis, defaults to 'Counts'
- title (str, optional) the figure title, defaults to 'Peptide Length distribution'

```
IPTK. Visualization.vizTools.plotly_protein_coverage (mapped_protein:
```

```
numpy.ndarray, prot_name:

str = None) \rightarrow

plotly.graph_objs._figure.Figure
```

Plot the peptide coverage for a given protein

Parameters

- mapped_protein (np.ndarray) A NumPy array with shape of 1 by protein length or shape protein-length
- prot_name (str, optional) The default protein name, defaults to None

Return type Figure

Module contents

Module contents

CHAPTER

FOUR

INDICES AND TABLES

- genindex
- modindex
- search

PYTHON MODULE INDEX

```
IPTK, 56
IPTK. Analysis, 10
IPTK. Analysis. Analysis Function, 8
IPTK.Classes, 30
IPTK.Classes.Database, 10
IPTK.Classes.Experiment, 14
IPTK.Classes.ExperimentalSet, 20
IPTK.Classes.HLAChain, 23
IPTK.Classes.HLAMolecules, 24
IPTK.Classes.HLASet, 24
IPTK.Classes.Peptide, 25
IPTK.Classes.Proband, 27
IPTK.Classes.Protein, 27
IPTK.Classes.Tissue, 28
IPTK. 10, 35
IPTK.IO.InFunctions, 30
IPTK.IO.MEMEInterface, 33
IPTK.IO.OutFunctions, 34
IPTK.Utils, 40
IPTK.Utils.DevFunctions, 35
IPTK.Utils.Mapping, 37
IPTK.Utils.Types, 37
IPTK.Utils.UtilityFunction, 37
IPTK. Visualization, 56
IPTK. Visualization. vizTools, 40
```

60 Python Module Index

INDEX

A	<pre>compute_change_in_protein_representation(</pre>
add_experiment()(IPTK.Classes.ExperimentalSet.Ex	sperimentSein module IPTK.Analysis.AnalysisFunction), 8
method), 20	compute_cnange_in_protein_representation(
add_org_2_parent()	(IPTK.Classes.ExperimentalSet.ExperimentSet
(IPTK.Classes.Peptide.Peptide method),	method), 20
25	compute_correlation_in_experssion()
add_org_info() (IPTK.Classes.Experiment.Experiment.method), 14	method), 20
<pre>add_parent_protein()</pre>	<pre>compute_difference_in_representation()</pre>
(IPTK.Classes.Peptide.Peptide method),	(in module IPTK.Analysis.AnalysisFunction), 8
25	compute_expression_correlation() (in mod-
add_to_database()	ule IPTK.Analysis.AnalysisFunction), 9
(IPTK. Classes. Database. Cellular Location DB	compute_peptide_length_table()
method), 10	(IPTK.Classes.ExperimentalSet.ExperimentSet
annotate_proteins()	method), 21
(IPTK.Classes.Experiment.Experiment	compute_peptide_overlap_matrix()
method), 15	(IPTK.Classes.ExperimentalSet.ExperimentSet
<pre>append_to_calling_string() (in module</pre>	method), 21
IPTK.Utils.UtilityFunction), 37	compute_peptide_representation_count()
D	(IPTK.Classes.ExperimentalSet.ExperimentSet
В	method), 21
ouild_sequence_table() (in module IPTK.Utils.UtilityFunction), 37	<pre>compute_protein_coverage_over_the_set() (IPTK.Classes.ExperimentalSet.ExperimentSet method), 21</pre>
C	<pre>compute_protein_overlap_matrix()</pre>
	(IPTK.Classes.ExperimentalSet.ExperimentSet
call_meme() (in module IPTK.IO.MEMEInterface), 33	<pre>method), 21 compute_protein_representation_count()</pre>
CellularLocationDB (class in	(IPTK.Classes.ExperimentalSet.ExperimentSet
IPTK.Classes.Database), 10	method), 21
<pre>check_peptide_made_of_std_20_aa() (in</pre>	memou), 21
module IPTK.Utils.UtilityFunction), 38	D
<pre>compare_org_count_among_exps()</pre>	download_pdb_entry() (in module
(IPTK. Classes. Experimental Set. Experiment Set	IPTK.IO.InFunctions), 30
method), 20	download_structure_file() (in module
<pre>compare_peptide_counts()</pre>	IPTK.Analysis.AnalysisFunction), 9
(IPTK.Classes.ExperimentalSet.ExperimentSet	drop_peptide_belong_to_org()
method), 20	(IPTK.Classes.Experiment.Experiment
<pre>compute_average_distance_between_exps()</pre>	method), 15
(IPTK.Classes.ExperimentalSet.ExperimentSet	drop_peptides_belong_to_org()
method), 20	(IPTK.Classes.ExperimentalSet.ExperimentSet
compute_binary_distance() (in module	method), 21
IPTK. Analysis. Analysis Function), 8	

E	<pre>get_class() (IPTK.Classes.HLAMolecules.HLAMolecule</pre>
Experiment (class in IPTK.Classes.Experiment), 14	method), 24 get_class() (IPTK.Classes.HLASet.HLASet
ExperimentSet (class in IPTK.Classes.ExperimentalSet), 20	<pre>get_class() (IPTK.Classes.HLASet.HLASet method), 24</pre>
ExpressionProfile (class in IPTK.Classes.Tissue), 28	<pre>get_experiment() (IPTK.Classes.ExperimentalSet.ExperimentSet</pre>
=	<pre>get_experiment_reference_tissue_expression()</pre>
Foot of digt () (in module IPTV IO In Functions) 20	(IPTK.Classes.Experiment.Experiment method), 15
<pre>fasta2dict() (in module IPTK.IO.InFunctions), 30</pre>	<pre>get_experimental_names()</pre>
G	(IPTK.Classes.ExperimentalSet.ExperimentSet method), 21
GeneExpressionDB (class in IPTK.Classes.Database), 12	get_experiments() (IPTK.Classes.ExperimentalSet.ExperimentSet
generate_color_scale() (in module IPTK.Utils.UtilityFunction), 38	method), 21
generate_random_name() (in module IPTK.Utils.UtilityFunction), 38	<pre>get_expression() (IPTK.Classes.Database.GeneExpressionDB method), 12</pre>
generate_random_peptide_seq() (in module IPTK.Utils.DevFunctions), 35	get_expression_in_tissue() (IPTK.Classes.Database.GeneExpressionDB
generate_random_protein_mapping() (in	<pre>method), 13 get_expression_of_parent_proteins()</pre>
<pre>module IPTK.Utils.UtilityFunction), 38 get_allele_count()</pre>	(IPTK.Classes.Experiment.Experiment
(IPTK.Classes.ExperimentalSet.ExperimentSet method), 21	<pre>method), 15 get_expression_profile()</pre>
get_allele_group()	(IPTK.Classes.Tissue.Tissue method), 29 get_flanked_peptide()
(IPTK.Classes.HLAChain.HLAChain method), 23	(IPTK.Classes.Peptide.Peptide method), 25
<pre>get_allele_group() (IPTK.Classes.HLAMolecules.HLAMolecule</pre>	get_flanked_peptides()
method), 24	(IPTK.Classes.Experiment.Experiment method), 15
<pre>get_alleles() (IPTK.Classes.HLASet.HLASet</pre>	get_gene() (IPTK.Classes.HLAChain.HLAChain
get_approved_location()	method), 23 get_gene() (IPTK.Classes.HLAMolecules.HLAMolecule
(IPTK.Classes.Database.CellularLocationDB method), 11	method), 24
get_binarized_results()	<pre>get_gene_id_expression() (IPTK.Classes.Tissue.ExpressionProfile</pre>
(IPTK.Classes.Experiment.Experiment method), 15	method), 28
get_binnary_peptide_overlap() (in module IPTK.Analysis.AnalysisFunction), 9	<pre>get_gene_name_expression() (IPTK.Classes.Tissue.ExpressionProfile</pre>
get_binnary_protein_overlap() (in module	<pre>method), 29 get_gene_names() (IPTK.Classes.Database.CellularLocationDB</pre>
<pre>IPTK.Analysis.AnalysisFunction), 10 get_c_terminal_flank_seq()</pre>	<pre>method), 11 get_gene_names() (IPTK.Classes.Database.GeneExpressionDB</pre>
(IPTK.Classes.Peptide.Peptide method), 25	method), 13
get_c_terminal_flanked_seqs()	get_genes() (IPTK.Classes.Database.CellularLocationDB method), 11
(IPTK.Classes.Experiment.Experiment method), 15	get_genes() (IPTK.Classes.Database.GeneExpressionDB
memoa), 13 get_chain_class()	method), 13
(IPTK.Classes.HLAChain.HLAChain method), 23	<pre>get_go_location_id_parent_proteins() (IPTK.Classes.Experiment.Experiment</pre>
get_class() (IPTK.Classes.HLAChain.HLAChain	<pre>method), 16 get_go_names() (IPTK.Classes.Database.CellularLocationDB</pre>
method), 23	method) 11

<pre>get_hla_allele() (IPTK.Classes.Experiment.Experim</pre>	
method), 16	(IPTK. Classes. Experimental Set. Experiment Set
<pre>get_hla_class() (IPTK.Classes.Experiment.Experime</pre>	
method), 16	<pre>get_num_peptide_expression_table()</pre>
get_hla_count() (IPTK.Classes.HLASet.HLASet	(IPTK.Classes.Experiment.Experiment
method), 24	method), 17
<pre>get_id() (IPTK.Classes.Protein.Protein method), 27</pre>	<pre>get_num_peptide_per_go_term()</pre>
<pre>get_idx_peptide_in_sequence_table() (in</pre>	(IPTK.Classes.Experiment.Experiment
module IPTK. Utils. UtilityFunction), 38	method), 17
get_length() (IPTK.Classes.Peptide.Peptide	<pre>get_num_peptide_per_location()</pre>
method), 26	(IPTK.Classes.Experiment.Experiment
get_main_location()	method), 17
(IPTK. Classes. Database. Cellular Location DB	<pre>get_number_of_children()</pre>
method), 12	(IPTK.Classes.Experiment.Experiment
get_main_sub_cellular_location_of_parent	_protei <i>m&kod</i>), 17
(IPTK.Classes.Experiment.Experiment	<pre>get_number_of_parents()</pre>
method), 16	(IPTK.Classes.Peptide.Peptide method),
<pre>get_mapped_protein()</pre>	26
(IPTK.Classes.Experiment.Experiment	<pre>get_number_of_proteins_per_compartment()</pre>
method), 16	(IPTK.Classes.Experiment.Experiment
get_mapped_proteins()	method), 17
(IPTK.Classes.Experiment.Experiment	<pre>get_number_of_proteins_per_go_term()</pre>
method), 16	(IPTK.Classes.Experiment.Experiment
<pre>get_meme_help()</pre>	method), 17
IPTK.IO.MEMEInterface), 33	<pre>get_number_parent_protein()</pre>
<pre>get_meta_data() (IPTK.Classes.Proband.Proband</pre>	(IPTK.Classes.Peptide.Peptide method),
method), 27	26
get_mono_parent_peptides()	<pre>get_number_protein_per_organism()</pre>
(IPTK.Classes.Experiment.Experiment	(IPTK.Classes.Database.OrganismDB
method), 16	method), 13
get_n_terminal_flank_seq()	<pre>get_org() (IPTK.Classes.Database.OrganismDB</pre>
(IPTK.Classes.Peptide.Peptide method),	method), 14
26	<pre>get_org() (IPTK.Classes.Protein.Protein method), 28</pre>
get_n_terminal_flanked_seqs()	<pre>get_orgs() (IPTK.Classes.Experiment.Experiment</pre>
(IPTK.Classes.Experiment.Experiment	method), 18
method), 17	<pre>get_parent() (IPTK.Classes.Peptide.Peptide</pre>
get_name() (IPTK.Classes.HLAChain.HLAChain	method), 26
method), 23	<pre>get_parent_proteins()</pre>
get_name()(IPTK.Classes.HLAMolecules.HLAMolecule	
method), 24	26
get_name() (IPTK.Classes.Proband.Proband	<pre>get_parents_org() (IPTK.Classes.Peptide.Peptide</pre>
method), 27	method), 26
<pre>get_name() (IPTK.Classes.Tissue.ExpressionProfile</pre>	<pre>get_peptide() (IPTK.Classes.Experiment.Experiment</pre>
method), 29	method), 18
get_name() (IPTK.Classes.Tissue.Tissue method), 29	<pre>get_peptide_number_parent()</pre>
get_negative_example()	(IPTK.Classes.Experiment.Experiment
(IPTK.Classes.Experiment.Experiment	method), 18
method), 17	<pre>get_peptide_seq() (IPTK.Classes.Peptide.Peptide</pre>
get_non_presented_peptide()	method), 26
(IPTK.Classes.Protein.Protein method),	<pre>get_peptides()(IPTK.Classes.Experiment.Experiment</pre>
27	method), 18
get_non_presented_peptides()	<pre>get_peptides_length()</pre>
(IPTK.Classes.Peptide.Peptide method),	(IPTK.Classes.Experiment.Experiment
26	method), 18

<pre>get_peptides_map()</pre>	(IPTK.Classes.Experiment.Experiment
(IPTK.Classes.Protein.Protein method),	method), 19
28	<pre>get_tissues() (IPTK.Classes.Database.GeneExpressionDB</pre>
<pre>get_peptides_per_organism()</pre>	method), 13
(IPTK.Classes.Experiment.Experiment	<pre>get_total_peptide_per_org_count()</pre>
method), 18	(IPTK.Classes.ExperimentalSet.ExperimentSet
<pre>get_peptides_per_protein()</pre>	method), 22
(IPTK.Classes.Experiment.Experiment	get_unique_orgs()
method), 18	(IPTK.Classes.Database.OrganismDB
<pre>get_peptides_present_in_all()</pre>	method), 14
(IPTK.Classes.ExperimentalSet.ExperimentSet	get_unique_orgs()
method), 22	(IPTK.Classes.ExperimentalSet.ExperimentSet
get_poly_parental_peptides()	method), 22
(IPTK.Classes.Experiment.Experiment	get_unique_peptides()
method), 18	(IPTK.Classes.ExperimentalSet.ExperimentSet
get_pos_in_parent()	method), 22
(IPTK.Classes.Peptide.Peptide method),	<pre>get_unique_proteins()</pre>
26	(IPTK.Classes.ExperimentalSet.ExperimentSet
get_proband_count()	method), 22
(IPTK.Classes.ExperimentalSet.ExperimentSet	group_by_proband()
method), 22	(IPTK. Classes. Experimental Set. Experiment Set
<pre>get_proband_name()</pre>	method), 22
(IPTK.Classes.Experiment.Experiment	<pre>group_by_tissue()</pre>
method), 18	(IPTK. Classes. Experimental Set. Experiment Set
<pre>get_protein_group()</pre>	method), 22
(IPTK.Classes.HLAChain.HLAChain method),	
23	H
<pre>get_protein_group()</pre>	has_allele() (IPTK.Classes.HLASet.HLASet
(IPTK.Classes.HLAMolecules.HLAMolecule	method), 24
method), 24	has_allele_group()
<pre>get_proteins() (IPTK.Classes.Experiment.Experiment</pre>	
method), 19	method), 19
<pre>get_proteins_present_in_all()</pre>	has_allele_group()
(IPTK.Classes.ExperimentalSet.ExperimentSet	(IPTK.Classes.HLASet method),
method), 22	(II TR. Classes. III Asei. III Asei method),
get_seq() (IPTK.Classes.Database.SeqDB method),	has_gene() (IPTK.Classes.Experiment.Experiment
14	
	method), 19
<pre>get_seq() (IPTK.Classes.Protein.Protein method), 28 get_sequence motif() (in module</pre>	
	25
IPTK.Analysis.AnalysisFunction), 10	has_hla_allele() (IPTK.Classes.Experiment.Experiment
get_subCellular_locations()	method), 19
(IPTK.Classes.Tissue.Tissue method), 29	has_protein_group()
<pre>get_table() (IPTK.Classes.Database.CellularLocation</pre>	(
method), 12	method), 19
<pre>get_table() (IPTK.Classes.Database.GeneExpressionI</pre>	
method), 13	(IPTK.Classes.HLASet.HLASet method),
<pre>get_table() (IPTK.Classes.Tissue.ExpressionProfile</pre>	25
method), 29	has_sequence() (IPTK.Classes.Database.SeqDB
<pre>get_tissue() (IPTK.Classes.Experiment.Experiment</pre>	method), 14
method), 19	HLAChain (class in IPTK. Classes. HLAChain), 23
<pre>get_tissue_counts()</pre>	HLAMolecule (class in IPTK. Classes. HLAMolecules),
(IPTK.Classes.ExperimentalSet.ExperimentSet	24
method), 22	HLASet (class in IPTK. Classes. HLASet), 24
get tissue name()	(

		is_a_parent_protein()
<pre>imposed_coverage_on_3D_structure() module IPTK.Visualization.vizTools), 40</pre>	(in	(IPTK.Classes.Experiment.Experiment method), 19
IPTK		is_child_of() (IPTK.Classes.Peptide.Peptide
module, 56		method), 27
IPTK.Analysis		<pre>is_member() (IPTK.Classes.Experiment.Experiment</pre>
module, 10		method), 20
IPTK.Analysis.AnalysisFunction		is_meme_callable() (in module
module, 8		IPTK.IO.MEMEInterface), 33
IPTK.Classes		is_peptide_present_in_all()
module, 30		(IPTK. Classes. Experimental Set. Experiment Set
IPTK.Classes.Database		method), 23
module, 10		is_protein_present_in_all()
IPTK.Classes.Experiment		(IPTK. Classes. Experimental Set. Experiment Set
module, 14		method), 23
IPTK.Classes.ExperimentalSet		1
module, 20		L
IPTK.Classes.HLAChain		<pre>load_3d_figure()</pre>
module, 23		IPTK.Utils.UtilityFunction), 39
IPTK.Classes.HLAMolecules		<pre>load_identification_table() (in module</pre>
module, 24		IPTK.IO.InFunctions), 30
IPTK.Classes.HLASet		
module, 24		M
IPTK.Classes.Peptide		<pre>map_from_uniprot_gene() (in module</pre>
module, 25		IPTK.Utils.Mapping), 37
IPTK.Classes.Proband		<pre>map_from_uniprot_pdb() (in module</pre>
module, 27		IPTK.Utils.Mapping), 37
IPTK.Classes.Protein		<pre>map_to_parent_protein()</pre>
module, 27		(IPTK.Classes.Peptide.Peptide method),
IPTK.Classes.Tissue		27
module, 28		module
IPTK.IO		IPTK, 56
module, 35		IPTK.Analysis, 10
IPTK.IO.InFunctions		IPTK.Analysis.AnalysisFunction, 8
module, 30		IPTK.Classes, 30
IPTK.IO.MEMEInterface		IPTK.Classes.Database, 10
module, 33		IPTK.Classes.Experiment, 14
IPTK.IO.OutFunctions		${ t IPTK.Classes.ExperimentalSet,20}$
module, 34		IPTK.Classes.HLAChain, 23
IPTK.Utils		IPTK.Classes.HLAMolecules, 24
module, 40		IPTK.Classes.HLASet, 24
IPTK.Utils.DevFunctions		IPTK.Classes.Peptide, 25
module, 35		IPTK.Classes.Proband, 27
IPTK.Utils.Mapping		IPTK.Classes.Protein,27
module, 37		IPTK.Classes.Tissue, 28
IPTK.Utils.Types		IPTK.10,35
module, 37		IPTK.IO.InFunctions, 30
IPTK.Utils.UtilityFunction		IPTK.IO.MEMEInterface, 33
module, 37		IPTK.IO.OutFunctions, 34
IPTK. Visualization		IPTK.Utils,40
module, 56		IPTK.Utils.DevFunctions, 35
IPTK.Visualization.vizTools		IPTK.Utils.Mapping, 37
module,40		IPTK.Utils.Types,37
		IPTK.Utils.UtilityFunction,37

IPTK.Visualization, 56 IPTK.Visualization.vizTools, 40	<pre>plotly_multi_traced_coverage_representation()</pre>
0	<pre>plotly_num_peptide_per_go_term() (in mod- ule IPTK.Visualization.vizTools), 51</pre>
OrganismDB (class in IPTK.Classes.Database), 13	<pre>plotly_num_peptides_per_location() (in</pre>
P	<pre>plotly_num_peptides_per_organism() (in module IPTK.Visualization.vizTools), 52</pre>
pad_mapped_proteins() (in module IPTK.Utils.UtilityFunction), 39	plotly_num_peptides_per_parent() (in mod- ule IPTK.Visualization.vizTools), 52
<pre>parse_mzTab_to_identification_table()</pre>	plotly_num_protein_per_go_term() (in mod- ule IPTK.Visualization.vizTools), 53
parse_text_table() (in module IPTK.IO.InFunctions), 31	plotly_num_protein_per_location() (in
parse_xml_based_format_to_identification (in module IPTK.IO.InFunctions), 32	n_table (Notatie II TK. Visualization. Viz1001s), 53 plotly_overlap_heatmap() (in module
Peptide (class in IPTK.Classes.Peptide), 25	11 1K. visuaitzation.viz.100is), 54
plot_change_in_presentation_between_experiments. plot_change_in_presentation_between_experiments. plot in module IPTK. Visualization. vizTools), 40	11 1 K. Visuatization. Viz. 10013), 54
plot_experiment_set_counts() (in module IPTK.Visualization.vizTools), 42	plotly_parent_protein_expression_in_tissue() (in module IPTK.Visualization.vizTools), 54
<pre>plot_gene_expression_vs_num_peptides()</pre>	plotly_peptide_length_dist() (in module IPTK.Visualization.vizTools), 55
plot_motif() (in module IPTK.Visualization.vizTools), 43	plotly_protein_coverage() (in module IPTK.Visualization.vizTools), 56
plot_num_peptide_per_go_term() (in module IPTK.Visualization.vizTools), 43	Proband (class in IPTK.Classes.Proband), 27 Protein (class in IPTK.Classes.Protein), 27
<pre>plot_num_peptides_per_location() (in mod- ule IPTK.Visualization.vizTools), 43</pre>	S
<pre>plot_num_peptides_per_organism() (in mod- ule IPTK.Visualization.vizTools), 44</pre>	<pre>save_3d_figure() (in module</pre>
<pre>plot_num_peptides_per_parent() (in module</pre>	SeqDB (class in IPTK.Classes.Database), 14
IPTK. Visualization. vizTools), 45	set_org() (IPTK.Classes.Protein.Protein method), 28
plot_num_protein_per_go_term() (in module IPTK.Visualization.vizTools), 45	<pre>simulate_an_experimental_ident_table_from_fasta()</pre>
<pre>plot_num_protein_per_location() (in mod- ule IPTK.Visualization.vizTools), 46</pre>	simulate_an_expression_table() (in module IPTK.Utils.DevFunctions), 36
plot_overlap_heatmap() (in module IPTK.Visualization.vizTools), 46	<pre>simulate_mapped_array_list() (in module</pre>
plot_overlay_representation() (in module IPTK.Visualization.vizTools), 46	<pre>simulate_protein_binary_represention() (in module IPTK.Utils.UtilityFunction), 39</pre>
plot_paired_represention() (in module IPTK.Visualization.vizTools), 47	simulate_protein_representation() (in module IPTK.Utils.UtilityFunction), 39
plot_parent_protein_expression_in_tissue (in module IPTK.Visualization.vizTools), 47	eqimulate_random_experiment() (in module IPTK.Utils.DevFunctions), 36
plot_peptide_length_dist() (in module IPTK.Visualization.vizTools), 48	Т
<pre>plot_peptide_length_per_experiment() (in</pre>	Tissue (class in IPTK.Classes.Tissue), 29
plot_protein_coverage() (in module	U
IPTK.Visualization.vizTools), 49	update_info() (IPTK.Classes.Proband.Proband
plot_protein_presentation_3D() (in module IPTK.Visualization.vizTools), 49	method), 27
plotly_gene_expression_us_num_peptides()) v v
(in module IPTK.Visualization.vizTools), 50	write_annotated_sequences() (in module

IPTK.IO.OutFunctions), 34
write_auto_named_peptide_to_fasta() (in
 module IPTK.IO.OutFunctions), 34
write_mapped_tensor_to_h5py() (in module
 IPTK.IO.OutFunctions), 34
write_named_peptides_to_fasta() (in module IPTK.IO.OutFunctions), 34
write_pep_file() (in module IPTK.IO.OutFunctions), 35