
MethylNet Documentation

Release 0.1

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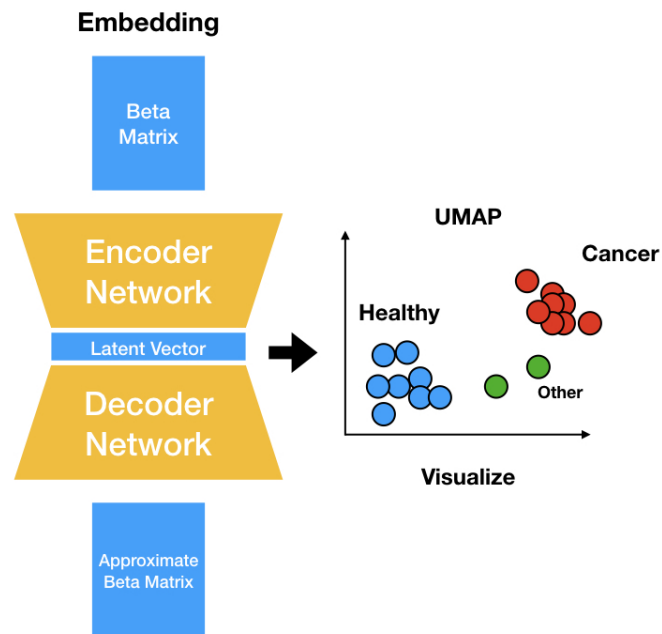
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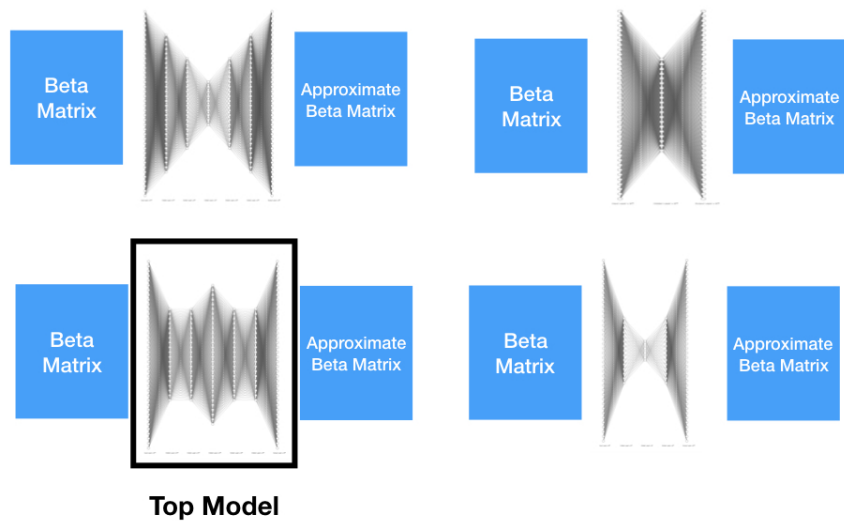
<https://github.com/Christensen-Lab-Dartmouth/MethylNet>

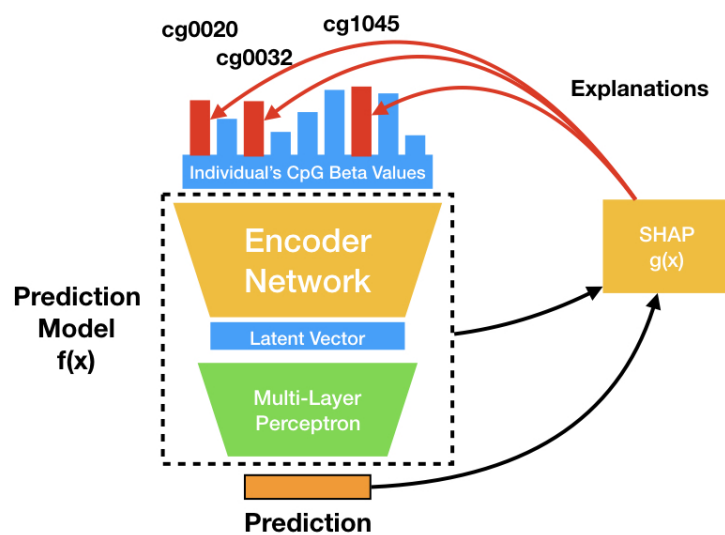
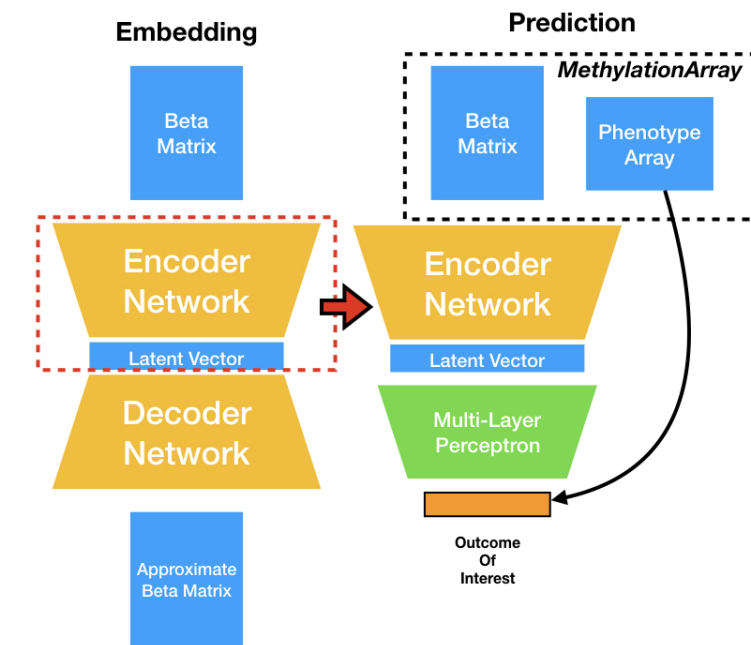
See README.md in Github repository for install directions and for example scripts for running the pipeline (not all datasets may be available on GEO at this time).

There is both an API and CLI available for use. Examples for CLI usage can be found in ./example_scripts.



Hyper-Parameter Scan





DATASETS.PY

Contains datatypes core to loading MethylationArrays into Torch tensors.

```
class methylnet.datasets.MethylationDataSet (methylation_array, transform, outcome_col="", categorical=False, categorical_encoder=False)
```

Pytorch Dataset that contains instances of methylation array samples to be loaded.

Parameters

- methylation_array** [MethylationArray] Methylation Array input.
- transform** [Transformer] Transforms data into torch tensor.
- outcome_col** [str] Pheno column(s) to train on.
- categorical** [bool] Whether predicting categorical/classification.
- categorical_encoder** [encoder] Encoder used to binarize categorical column.

Attributes

- samples** [np.array] Samples of MethylationArray
- features** [np.array] List CpGs.
- encoder** : Encoder used to binarize categorical column.
- new_shape** : Shape of torch tensors.
- length** : Number CpGs.
- methylation_array**
- outcome_col**
- transform**

Methods

<code>to_methyl_array(self)</code>	Convert torch dataset back into methylation array, useful because turning into torch dataset can cause the original MethylationArray beta matrix to turn into numpy array, when needs turn back into pandas dataframe.
------------------------------------	--

```
to_methyl_array (self)  
    Convert torch dataset back into methylation array, useful because turning into torch dataset can cause
```

the original MethylationArray beta matrix to turn into numpy array, when needs turn back into pandas dataframe.

Returns

MethylationArray

```
class methylnet.datasets.MethylationPredictionDataSet (methylation_array, transform, outcome_col="", categorical=False, categorical_encoder=False)
```

MethylationArray Torch Dataset that contains instances of methylation array samples to be loaded, specifically targetted for prediction.

Parameters

methylation_array [MethylationArray] Methylation Array input.

transform [Transformer] Transforms data into torch tensor.

outcome_col [str] Pheno column(s) to train on.

categorical [bool] Whether predicting categorical/classification.

categorical_encoder [encoder] Encoder used to binarize categorical column.

Attributes

samples [np.array] Samples of MethylationArray

features [np.array] List CpGs.

encoder : Encoder used to binarize categorical column.

new_shape : Shape of torch tensors.

length : Number CpGs.

methylation_array

outcome_col

transform

Methods

<code>to_methyl_array(self)</code>	Convert torch dataset back into methylation array, useful because turning into torch dataset can cause the original MethylationArray beta matrix to turn into numpy array, when needs turn back into pandas dataframe.
------------------------------------	--

```
class methylnet.datasets.RawBetaArrayDataSet (beta_array, transform)
```

Torch Dataset just for beta matrix in numpy format.

Parameters

beta_array [numpy.array] Beta value matrix in numpy form.

transform : Transforms data to torch tensor.

Attributes

length : Number CpGs.

beta_array**transform****class** methylnet.datasets.**Transformer** (*convolutional=False, cpg_per_row=30000, l=None*)

Push methyl array sample to pytorch tensor, possibly turning into image.

Parameters**convolutional** [bool] Whether running convolutions on torch dataset.**cpg_per_row** [int] Number of CpGs per row of image if convolutional.**l** [tuple] Attributes that describe image.**Attributes****shape** [tuple] Shape of methylation array image for sample**convolutional****cpg_per_row****l****Methods***generate*(self)

Generate function for transform.

generate (*self*)

Generate function for transform.

Returns**function**methylnet.datasets.**cat** ()Concatenates the given sequence of `seq` tensors in the given dimension. All tensors must either have the same shape (except in the concatenating dimension) or be empty.`torch.cat()` can be seen as an inverse operation for `torch.split()` and `torch.chunk()`.`torch.cat()` can be best understood via examples.**Args:****tensors (sequence of Tensors): any python sequence of tensors of the same type.** Non-empty tensors provided must have the same shape, except in the cat dimension.**dim** (int, optional): the dimension over which the tensors are concatenated out (Tensor, optional): the output tensor**Example:**

```

>>> x = torch.randn(2, 3)
>>> x
tensor([[ 0.6580, -1.0969, -0.4614],
        [-0.1034, -0.5790,  0.1497]])
>>> torch.cat((x, x, x), 0)
tensor([[ 0.6580, -1.0969, -0.4614],
        [-0.1034, -0.5790,  0.1497],
        [ 0.6580, -1.0969, -0.4614],

```

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```
        [-0.1034, -0.5790,  0.1497],
        [ 0.6580, -1.0969, -0.4614],
        [-0.1034, -0.5790,  0.1497]])
>>> torch.cat((x, x, x), 1)
tensor([[ 0.6580, -1.0969, -0.4614,  0.6580, -1.0969, -0.4614,  0.6580,
        -1.0969, -0.4614],
        [-0.1034, -0.5790,  0.1497, -0.1034, -0.5790,  0.1497, -0.1034,
        -0.5790,  0.1497]])
```

```
methylnet.datasets.get_methylation_dataset(methylation_array, outcome_col, con-
                                          volutional=False, cpg_per_row=1200,
                                          predict=False, categorical=False, categori-
                                          cal_encoder=False)
```

Turn methylation array into pytorch dataset.

Parameters

methylation_array [MethylationArray] Input MethylationArray.

outcome_col [str] Pheno column to train on.

convolutional [bool] Whether running CNN on methylation data

cpg_per_row [int] If convolutional, number of cpgs per image row.

predict [bool] Running prediction algorithm vs VAE.

categorical [bool] Whether training on categorical vs continuous variables.

categorical_encoder : Scikit learn encoder.

Returns

Pytorch Dataset

```
methylnet.datasets.stack()
```

Concatenates sequence of tensors along a new dimension.

All tensors need to be of the same size.

Arguments: seq (sequence of Tensors): sequence of tensors to concatenate dim (int): dimension to insert. Has to be between 0 and the number

of dimensions of concatenated tensors (inclusive)

out (Tensor, optional): the output tensor

HYPERPARAMETER_SCANS.PY

Run randomized grid search to find ideal model hyperparameters, with possible deployments to batch system for scalability.

```
methylnet.hyperparameter_scans.coarse_scan(hyperparameter_input_csv, hyperparameter_output_log, generate_input, job_chunk_size, stratify_column, reset_all, torque, gpu, gpu_node, nohup, mlp=False, custom_jobs=[], model_complexity_factor=0.9, set_beta=1.0, n_jobs=4, categorical=True, add_softmax=False, additional_command="", cuda=True, new_grid={})
```

Perform randomized hyperparameter grid search

Parameters

hyperparameter_input_csv [type] CSV file containing hyperparameter inputs.

hyperparameter_output_log [type] CSV file containing prior runs.

generate_input [type] Generate hyperparameter input csv.

job_chunk_size [type] Number of jobs to be launched at same time.

stratify_column [type] Performing classification?

reset_all [type] Rerun all jobs previously scanned.

torque [type] Run jobs using torque.

gpu [type] What GPU to use, set to -1 to be agnostic to GPU selection.

gpu_node [type] What GPU to use, set to -1 to be agnostic to GPU selection, for torque submission.

nohup [type] Launch jobs using nohup.

mlp [type] If running prediction job (classification/regression) after VAE.

custom_jobs [type] Supply custom job parameters to be run.

model_complexity_factor [type] Degree of neural network model complexity for hyperparameter search. Search for less wide networks with a lower complexity value, bounded between 0 and infinity.

set_beta [type] Don't hyperparameter scan over beta (KL divergence weight), and set it to value.

n_jobs [type] Number of jobs to generate.

categorical [type] Classification task?

add_softmax [type] Add softmax layer at end of neural network.

cuda [type] Whether to use GPU?

`methylnet.hyperparameter_scans.find_top_jobs` (*hyperparameter_input_csv*,
hyperparameter_output_log,
n_top_jobs, *crossover_p=0*,
val_loss_column='min_val_loss')
Finds top performing jobs from hyper parameter scan to rerun and cross-over parameters.

Parameters

hyperparameter_input_csv [str] CSV file containing hyperparameter inputs.

hyperparameter_output_log [str] CSV file containing prior runs.

n_top_jobs [int] Number of top jobs to select

crossover_p [float] Rate of cross over of reused hyperparameters

val_loss_column [str] Loss column used to select top jobs

Returns

list List of list of new parameters for jobs to run.

`methylnet.hyperparameter_scans.generate_topology` (*topology_grid*, *probability_decay_factor=0.9*)
Generates list denoting neural network topology, list of hidden layer sizes.

Parameters

topology_grid [list] List of different hidden layer sizes (number neurons) to choose from.

probability_decay_factor [float] Degree of neural network model complexity for hyperparameter search. Search for less wide networks with a lower complexity value, bounded between 0 and infinity.

Returns

list List of hidden layer sizes.

`methylnet.hyperparameter_scans.replace_grid` (*old_grid*, *new_grid*, *topology_grid*)
Replaces old hyperparameter search grid with one supplied by YAML.

TORQUE_JOBS.PY

Wraps and runs your commands through torque.

`methylnet.torque_jobs.assemble_replace_dict` (*command, use_gpu, additions, queue, time, ngpu*)
Create dictionary to update BASH submission script for torque.

Parameters

command [type] Command to executer through torque.
use_gpu [type] GPUs needed?
additions [type] Additional commands to add (eg. module loads).
queue [type] Queue to place job in.
time [type] How many hours to run job for.
ngpu [type] Number of GPU to use.

Returns

Dict Dictionary used to update Torque Script.

`methylnet.torque_jobs.assemble_run_torque` (*command, use_gpu, additions, queue, time, ngpu, additional_options=""*)
Runs torque job after passing commands to setup bash file.

Parameters

command [type] Command to executer through torque.
use_gpu [type] GPUs needed?
additions [type] Additional commands to add (eg. module loads).
queue [type] Queue to place job in.
time [type] How many hours to run job for.
ngpu [type] Number of GPU to use.
additional_options [type] Additional options to pass to Torque scheduler.

Returns

job Custom job name.

`methylnet.torque_jobs.run_torque_job` (*replace_dict, additional_options=""*)
Run torque job after creating submission script.

Parameters

replace_dict [type] Dictionary used to replace information in bash script to run torque job.

additional_options [type] Additional options to pass scheduler.

Returns

str Custom torque job name.

INTERPRETATION_CLASSES.PY

Contains core classes and functions to extracting explanations for the predictions at single samples, and then interrogates important CpGs for biological plausibility.

class `methylnet.interpretation_classes.BioInterpreter` (*dict_top_cpgs*)
Interrogate CpGs found to be important from SHAP through enrichment and overlap tests.

Parameters

dict_top_cpgs [type] Dictionary of topCpGs output from ShapleyExplorer object

Attributes

top_cpgs [type] Dictionary of top cpgs to be interrogated.

Methods

<code>get_nearby_cpg_shapleys</code> (self, all_cpgs, max_gap)	Query for CpGs that are within a max_gap of the topCpGs in the genome, helps find cpGs that could be highly correlated and thus also important.
<code>gometh</code> (self[, collection, allcpGs, ...])	Run GO, KEGG, GSEA analyses or return nearby genes.
<code>return_overlap_score</code> (self[, set_cpGs, ...])	Perform overlap test, overlapping top CpGs with IDOL CpGs, age related cpGs, etc.
<code>run_lola</code> (self[, all_cpGs, lola_db, cores, ...])	Run LOLA enrichment test.

get_nearby_cpg_shapleys (*self, all_cpGs, max_gap*)

Query for CpGs that are within a max_gap of the topCpGs in the genome, helps find cpGs that could be highly correlated and thus also important.

Parameters

all_cpGs [type] List of all cpGs in methylationarray.

max_gap [type] Max radius to search for nearby CpGs.

Returns

Dict Results for each class/indiv's top CpGs, in the form of nearby CpGs to each of these CpGs sets.

gometh (*self, collection='GO', allcpGs=[], length_output=20, gsea_analyses=[], gsea_pickle=""*)

Run GO, KEGG, GSEA analyses or return nearby genes.

Parameters

collection [type] GO, KEGG, GSEA, GENE overlaps?

allcpgs [type] All CpGs defines CpG universe, empty if use all CpGs, smaller group of CpGs may yield more significant results.

length_output [type] How many lines should the output be, maximum number of results to print.

gsea_analyses [type] GSEA analyses/collections to target.

gsea_pickle [type] Location of gsea pickle containing gene sets, may need to run download_help_data to acquire.

Returns

Dict Dictionary containing results from each test run on all of the keys in top_cpgs.

return_overlap_score (*self*, *set_cpgs*='IDOL', *platform*='450k', *all_cpgs*=[], *output_csv*='output_bio_intersect.csv', *extract_library*=False)
Perform overlap test, overlapping top CpGs with IDOL CpGs, age related cpgs, etc.

Parameters

set_cpgs [type] Set of reference cpgs.

platform [type] 450K or 850K

all_cpgs [type] All CpGs to build a universe.

output_csv [type] Output CSV for results of overlaps between classes, how much do they share these CpGs overlapped.

extract_library [type] Can extract a list of the CpGs that ended up being overlapped with.

Returns

List Optional output of overlapped library of CpGs.

run_lola (*self*, *all_cpgs*=[], *lola_db*="", *cores*=8, *collections*=[], *depletion*=False)
Run LOLA enrichment test.

Parameters

all_cpgs [type] CpG universe for LOLA.

lola_db [type] Location of LOLA database, can be downloaded using methylnet-interpret. Set to extended or core, and collections correspond to these.

cores [type] Number of cores to use.

collections [type] LOLA collections to run, leave empty to run all.

depletion [type] Set true to look for depleted regions over enriched.

Returns

type Description of returned object.

class methylnet.interpretation_classes.**CpGExplainer** (*prediction_function*=None, *cuda*=False)

Produces SHAPley explanation scores for individual predictions, approximating a complex model with a basic linear one, one coefficient per CpG per individual.

Parameters

prediction_function [type] Model or function that makes predictions on the data, can be sklearn .predict method or pytorch forward pass, etc. . .

cuda [type] Run on GPUs?

Attributes

explainer [type] Type of SHAPley explanation method; kernel (uses LIME model agnostic explainer), deep (uses DeepLift) or Gradient (integrated gradients and backprop for explanations)?

prediction_function

cuda

Methods

<code>build_explainer(self, train_methyl_array[, ...])</code>	Builds SHAP explainer using background samples.
<code>classifier_assign_scores_to_shap_data(self, ...)</code>	Assigns the SHAP scores to a SHAPley data object, populating nested dictionaries (containing class-individual information) with SHAPley information and “top CpGs”.
<code>feature_select(self, methyl_array, ...)</code>	Perform feature selection based on the best overall SHAP scores across all samples.
<code>from_explainer(method, cuda)</code>	Load custom SHAPley explainer
<code>regressor_assign_scores_to_shap_data(self, ...)</code>	Assigns the SHAP scores to a SHAPley data object, populating nested dictionaries (containing multioutput and single output regression-individual information) with SHAPley information and “top CpGs”.
<code>return_shapley_predictions(self, ..., encoder])</code>	Method in development or may be deprecated.
<code>return_shapley_scores(self, ..., ...])</code>	Generate explanations for individual predictions, in the form of SHAPley scores for supplied test data.

build_explainer (*self*, *train_methyl_array*, *method*=*'kernel'*, *batch_size*=*100*)

Builds SHAP explainer using background samples.

Parameters

train_methyl_array [type] Train Methylation Array from which to populate background samples to make estimated explanations.

method [type] SHAP explanation method?

batch_size [type] Break up prediction explanation creation into smaller batch sizes for lower memory consumption.

classifier_assign_scores_to_shap_data (*self*, *test_methyl_array*, *n_top_features*, *interest_col*=*'disease'*, *prediction_classes*=*None*)

Assigns the SHAP scores to a SHAPley data object, populating nested dictionaries (containing class-individual information) with SHAPley information and “top CpGs”.

Parameters

test_methyl_array [type] Testing MethylationArray.

n_top_features [type] Number of top SHAP scores to use for a subdict called “top CpGs”

interest_col [type] Column in pheno sheet from which class names reside.

prediction_classes [type] User supplied prediction classes.

feature_select (*self*, *methyl_array*, *n_top_features*)

Perform feature selection based on the best overall SHAP scores across all samples.

Parameters

methyl_array [type] MethylationArray to run feature selection on.

n_top_features [type] Number of CpGs to select.

Returns

MethylationArray Subsetted by top overall CpGs, overall positive contributions to prediction. May need to update.

classmethod from_explainer (*method*, *cuda*)

Load custom SHAPley explainer

regressor_assign_scores_to_shap_data (*self*, *test_methyl_array*, *n_top_features*, *cell_names=[]*)

Assigns the SHAP scores to a SHAPley data object, populating nested dictionaries (containing multioutput and single output regression-individual information) with SHAPley information and “top CpGs”.

Parameters

test_methyl_array [type] Testing MethylationArray.

n_top_features [type] Number of top SHAP scores to use for a subdict called “top CpGs”

cell_names [type] If multi-output regression, create separate regression classes to access for all individuals.

return_shapley_predictions (*self*, *test_methyl_array*, *sample_name*, *interest_col*, *encoder=None*)

Method in development or may be deprecated.

return_shapley_scores (*self*, *test_methyl_array*, *n_samples*, *n_outputs*, *shap_sample_batch_size=None*, *top_outputs=None*)

Generate explanations for individual predictions, in the form of SHAPley scores for supplied test data. One SHAP score per individual prediction per CpG, and more if multiclass/output. For multiclass/multivariate outcomes, scores are generated for each outcome class.

Parameters

test_methyl_array [type] Testing MethylationArray.

n_samples [type] Number of SHAP score samples to produce. More SHAP samples provides convergence to correct score.

n_outputs [type] Number of outcome classes.

shap_sample_batch_size [type] If not None, break up SHAP score sampling into batches of this size to be averaged.

top_outputs [type] For deep explainer, limit number of output classes due to memory consumption.

Returns

type Description of returned object.

class `methylnet.interpretation_classes.DistanceMatrixCompute` (*methyl_array*, *pheno_col*)

From any embeddings, calculate pairwise distances between classes that label embeddings.

Parameters

methyl_array [MethylationArray] Methylation array storing beta and pheno data.

pheno_col [str] Column name from which to extract sample names from and group by class.

Attributes

col [pd.DataFrame] Column of pheno array.

classes [np.array] Unique classes of pheno column.

embeddings [pd.DataFrame] Embeddings of beta values.

Methods

<code>calculate_p_values(self)</code>	Compute pairwise p-values between different clusters using manova.
<code>compute_distances(self[, metric, trim])</code>	Compute distance matrix between classes by average distances between points of classes.
<code>return_distances(self)</code>	Return the distance matrix
<code>return_p_values(self)</code>	Return the distance matrix

calculate_p_values (*self*)

Compute pairwise p-values between different clusters using manova.

compute_distances (*self*, *metric*='cosine', *trim*=0.0)

Compute distance matrix between classes by average distances between points of classes.

Parameters

metric [str] Scikit-learn distance metric.

return_distances (*self*)

Return the distance matrix

Returns

pd.DataFrame Distance matrix between classes.

return_p_values (*self*)

Return the distance matrix

Returns

pd.DataFrame MANOVA values between classes.

class methylnet.interpretation_classes.**PlotCircos**

Plot Circos Diagram using ggbio (regular circos software may be better)

Attributes

generate_base_plot [type] Function to generate base plot

add_plot_data [type] Function to add more plot data

generate_final_plot [type] Function to plot data using Circos

Methods

<code>plot_cpgs(self, top_cpgs[, output_dir])</code>	Plot top CpGs location in genome.
--	-----------------------------------

plot_cpgs (*self*, *top_cpgs*, *output_dir*='./')
Plot top CpGs location in genome.

Parameters

top_cpgs [type] Input dataframe of top CpG name and SHAP scores. Future: Plot SHAP scores in locations?

output_dir [type] Where to output plots.

class `methylnet.interpretation_classes.ShapleyData`

Store SHAP results that stores feature importances of CpGs on varying degrees granularity.

Attributes

top_cpgs [type] Quick accessible CpGs that have the n highest SHAP scores.

shapley_values [type] Storing all SHAPley values for CpGs for varying degrees granularity. For classification problems, this only includes SHAP scores particular to the actual class.

Methods

<code>add_class(self, class_name, shap_df, cpgs, ...)</code>	Store SHAP scores for particular class to Shapley_Data class.
<code>add_global_importance(self, ...)</code>	Add overall feature importances, globally across all samples.
<code>from_pickle(input_pkl[, from_dict])</code>	Load SHAPley data from pickle.
<code>to_pickle(self, output_pkl[, output_dict])</code>	Export Shapley data to pickle.

add_class (*self*, *class_name*, *shap_df*, *cpgs*, *n_top_cpgs*, *add_top_negative=False*, *add_top_abs=False*)

Store SHAP scores for particular class to Shapley_Data class. Save feature importances on granular level, explanations for individual and aggregate predictions.

Parameters

class_name [type] Particular class name to be stored in dictionary structure.

shap_df [type] SHAP data to pull from.

cpgs [type] All CpGs.

n_top_cpgs [type] Number of top CpGs for saving n top CpGs.

add_top_negative [type] Only consider top negative scores.

add_global_importance (*self*, *global_importance_shaps*, *cpgs*, *n_top_cpgs*)

Add overall feature importances, globally across all samples.

Parameters

global_importance_shaps [type] Overall SHAP values to be saved, aggregated across all samples.

cpgs [type] All CpGs.

n_top_cpgs [type] Number of top CpGs to rank and save as well.

classmethod `from_pickle(input_pkl, from_dict=False)`

Load SHAPley data from pickle.

Parameters

input_pkl [type] Input pickle.

to_pickle (*self*, *output_pkl*, *output_dict=False*)
Export Shapley data to pickle.

Parameters

output_pkl [type] Output file to save SHAPley data.

class `methylnet.interpretation_classes.ShapleyDataExplorer` (*shapley_data*)
Datatype used to explore saved ShapleyData.

Parameters

shapley_data [type] ShapleyData instance to be explored.

Attributes

indiv2class [type] Maps individuals to their classes used for quick look-up.

shapley_data

Methods

<code>add_abs_value_classes(self)</code>	WIP.
<code>add_bin_continuous_classes(self)</code>	Bin continuous outcome variable and extract top positive and negative CpGs for each new class.
<code>extract_class(self, class_name[, ...])</code>	Extract the top cpgs from a class
<code>extract_individual(self, individual[, ...])</code>	Extract the top cpgs from an individual
<code>extract_methylation_array(self, methyl_arr)</code>	Subset MethylationArray Beta values by some top SHAP CpGs from classes or overall.
<code>jaccard_similarity_top_cpgs(self, class_names)</code>	Calculate Jaccard Similarity matrix between classes and individuals within those classes based on how they share sets of CpGs.
<code>limit_number_top_cpgs(self, n_top_cpgs)</code>	Reduce the number of top CpGs.
<code>list_classes(self)</code>	List classes in ShapleyData object.
<code>list_individuals(self[, return_list])</code>	List the individuals in the ShapleyData object.
<code>regenerate_individual_shap_values(self, ...)</code>	Use original SHAP scores to make nested dictionary of top CpGs based on shapley score, can do this for ABS SHAP or Negative SHAP scores as well.
<code>return_binned_shapley_data(self, ..., ...)</code>	Converts existing shap data based on continuous variable predictions into categorical variable.
<code>return_cpg_sets(self)</code>	Return top sets of CpGs from classes and individuals, and the complementary set.
<code>return_global_importance_cpgs(self)</code>	Return overall globally important CpGs.
<code>return_shapley_data_by_methylation_status(self)</code>	Return dictionary containing two SHAPley datasets, each split by low/high levels of methylation.
<code>return_top_cpgs(self[, classes, ...])</code>	Given list of classes and individuals, export a dictionary containing data frames of top CpGs and their SHAP scores.
<code>view_methylation(self, individual, methyl_arr)</code>	Use MethylationArray to output top SHAP values for individual with methylation data appended to output data frame.

make_shap_scores_abs	
-----------------------------	--

add_abs_value_classes (*self*)
WIP.

add_bin_continuous_classes (*self*)
Bin continuous outcome variable and extract top positive and negative CpGs for each new class.

extract_class (*self*, *class_name*, *class_intersect=False*, *get_shap_values=False*)
Extract the top cpGs from a class

Parameters

class_name [type] Class to extract from?

class_intersect [type] Bool to extract from aggregation of SHAP values from individuals of current class, should have been done already.

Returns

DataFrame CpGs and SHAP Values

extract_individual (*self*, *individual*, *get_shap_values=False*)
Extract the top cpGs from an individual

Parameters

individual [type] Individual to extract from?

Returns

Tuple Class name of individual, DataFrame of CpGs and SHAP Values

extract_methylation_array (*self*, *methylation_arr*, *classes_only=True*, *global_vals=False*,
n_extract=1000, *class_name=""*)
Subset MethylationArray Beta values by some top SHAP CpGs from classes or overall.

Parameters

methylation_arr [type] MethylationArray.

classes_only [type] Setting this to False will include the overall top CpGs per class and the top CpGs for individuals in that class.

global_vals [type] Use top CpGs overall, across the entire dataset.

n_extract [type] Number of top CpGs to subset.

class_name [type] Which class to subset top CpGs from? Blank to use union of all top CpGs.

Returns

MethylationArray Stores methylation beta and pheno data, reduced here by queried CpGs.

jaccard_similarity_top_cpGs (*self*, *class_names*, *individuals=False*, *overall=False*, *cooccurrence=False*)

Calculate Jaccard Similarity matrix between classes and individuals within those classes based on how they share sets of CpGs.

Parameters

class_names [type] Classes to include.

individuals [type] Individuals to include.

overall [type] Whether to use overall class top CpGs versus aggregate.

cooccurrence [type] Output cooccurrence instead of jaccard.

Returns

pd.DataFrame Similarity matrix.

limit_number_top_cpgs (*self*, *n_top_cpgs*)

Reduce the number of top CpGs.

Parameters

n_top_cpgs [type] Number top CpGs to retain.

Returns

ShapleyData Returns shapley data with fewer top CpGs.

list_classes (*self*)

List classes in ShapleyData object.

Returns

List List of classes

list_individuals (*self*, *return_list=False*)

List the individuals in the ShapleyData object.

Parameters

return_list [type] Return list of individual names rather than a dictionary of class:individual key-value pairs.

Returns

List or Dictionary Class: Individual dictionary or individuals are elements of list

regenerate_individual_shap_values (*self*, *n_top_cpgs*, *abs_val=False*, *neg_val=False*)

Use original SHAP scores to make nested dictionary of top CpGs based on shapley score, can do this for ABS SHAP or Negative SHAP scores as well.

Parameters

n_top_cpgs [type] Description of parameter *n_top_cpgs*.

abs_val [type] Description of parameter *abs_val*.

neg_val [type] Description of parameter *neg_val*.

Returns

type Description of returned object.

return_binned_shapley_data (*self*, *original_class_name*, *outcome_col*,
add_top_negative=False)

Converts existing shap data based on continuous variable predictions into categorical variable.

Parameters

original_class_name [type] Regression results were split into ClassName_pos and ClassName_neg, what is the ClassName?

outcome_col [type] Feed in from pheno sheet one the column to bin samples on.

add_top_negative [type] Looking to include negative SHAPs?

Returns

ShapleyData With new class labels, built from regression results.

return_cpg_sets (*self*)

Return top sets of CpGs from classes and individuals, and the complementary set. Returns dictionary of sets of CpGs and the union of all the other sets minus the current set.

Returns

cpg_sets Dictionary of individuals and classes; CpGs contained in set for particular individual/class

cpg_exclusion_sets Dictionary of individuals and classes; CpGs not contained in set for particular individual/class

return_global_importance_cpgs (*self*)

Return overall globally important CpGs.

Returns

list

return_shapley_data_by_methylation_status (*self*, *methylation_array*, *threshold*)

Return dictionary containing two SHAPley datasets, each split by low/high levels of methylation. Todo: Define this using median methylation value vs 0.5.

Parameters

methylation_array [type] MethylationArray instance.

Returns

dictionary Contains shapley data by methylation status.

return_top_cpgs (*self*, *classes*=[], *individuals*=[], *class_intersect*=False, *cpg_exclusion_sets*=None, *cpg_sets*=None)

Given list of classes and individuals, export a dictionary containing data frames of top CpGs and their SHAP scores.

Parameters

classes [type] Higher level classes to extract CpGs from, list of classes to extract top CpGs from.

individuals [type] Individual samples to extract top CpGs from.

class_intersect [type] Whether the top CpGs should be chosen by aggregating the remaining individual scores.

cpg_exclusion_sets [type] Dict of sets of CpGs, where these ones contain CpGs not particular to particular class.

cpg_sets [type] Contains top CpGs found for each class.

Returns

Dict Top CpGs accessed and returned for further processing.

view_methylation (*self*, *individual*, *methylation_arr*)

Use MethylationArray to output top SHAP values for individual with methylation data appended to output data frame.

Parameters

individual [type] Individual to query from ShapleyData

methylation_arr [type] Input MethylationArray

Returns

class_name

individual

top_shap_df Top Shapley DataFrame with top cpgs for individual, methylation data appended.

`methylnet.interpretation_classes.cooccurrence_fn(list1, list2)`

Cooccurrence of elements between two lists.

Parameters

list1

list2

Returns

float Cooccurrence between elements in list.

`methylnet.interpretation_classes.jaccard_similarity(list1, list2)`

Jaccard score between two lists.

Parameters

list1

list2

Returns

float Jaccard similarity between elements in list.

`methylnet.interpretation_classes.main_prediction_function(n_workers, batch_size, model, cuda)`

Combine dataloader and prediction function into final prediction function that takes in tensor data, for Kernel Explanations.

Parameters

n_workers [type] Number of workers.

batch_size [type] Batch size for input data.

model [type] Model/predidction function.

cuda [type] Running on GPUs?

Returns

function Final prediction function.

`methylnet.interpretation_classes.plot_lola_output_(lola_csv, plot_output_dir, description_col, cell_types)`

Plots any LOLA output in the form of a forest plot.

Parameters

lola_csv [type] CSV containing lola results.

plot_output_dir [type] Plot output directory.

description_col [type] Column that will label the points on the plot.

cell_types [type] Column containing cell-types, colors plots and are rows to be compared.

`methylnet.interpretation_classes.return_dataloader_construct(n_workers, batch_size)`

Decorator to build dataloader compatible with KernelExplainer for SHAP.

Parameters

n_workers [type] Number CPU.

batch_size [type] Batch size to load data into pytorch.

Returns

DataLoader Pytorch dataloader.

`methylnet.interpretation_classes.return_predict_function(model, cuda)`

Decorator to build the supplied prediction function, important for kernel explanations.

Parameters

model [type] Prediction function with predict method.

cuda [type] Run using cuda.

Returns

function Predict function.

`methylnet.interpretation_classes.return_shap_values(test_arr, explainer, method,
n_samples, additional_opts)`

Return SHAP values, sampled a number of times, used in CpGExplainer class.

Parameters

test_arr [type] Testing MethylationArray.

explainer [type] SHAP explainer object.

method [type] Method of explaining.

n_samples [type] Number of samples to estimate SHAP scores.

additional_opts [type] Additional options to be passed into non-kernel/gradient methods.

Returns

np.array/list Shapley scores.

`methylnet.interpretation_classes.to_tensor(arr)`

Turn np.array into tensor.

MODELS.PY

Contains core PyTorch Models for running VAE and VAE-MLP.

```
class methylnet.models.AutoEncoder (autoencoder_model,  n_epochs,  loss_fn,  optimizer,  
                                     cuda=True,  kl_warm_up=0,  beta=1.0,  scheduler_opts={})
```

Wraps Pytorch VAE module into Scikit-learn like interface for ease of training, validation and testing.

Parameters

- autoencoder_model** [type] Pytorch VAE Model to supply.
- n_epochs** [type] Number of epochs to train for.
- loss_fn** [type] Pytorch loss function for reconstruction error.
- optimizer** [type] Pytorch Optimizer.
- cuda** [type] GPU?
- kl_warm_up** [type] Number of epochs until fully utilizing KLLoss, begin saving models here.
- beta** [type] Weighting for KLLoss.
- scheduler_opts** [type] Options to feed learning rate scheduler, which modulates learning rate of optimizer.

Attributes

- model** [type] Pytorch VAE model.
- scheduler** [type] Learning rate scheduler object.
- vae_animation_fname** [type] Save VAE embeddings evolving over epochs to this file name. Defunct for now.
- loss_plt_fname** [type] Where to save loss curves. This has been superceded by plot_training_curves in methylnet-visualize command.
- plot_interval** [type] How often to plot data; defunct.
- embed_interval** [type] How often to embed; defunct.
- validation_set** [type] MethylationArray DataLoader, produced from Pytorch Methylation-Dataset of Validation MethylationArray.
- n_epochs**
- loss_fn**
- optimizer**
- cuda**

kl_warm_up

beta

Methods

<code>add_validation_set(self, validation_data)</code>	Add validation data in the form of Validation DataLoader.
<code>fit(self, train_data)</code>	Fit VAE model to training data, best model returned with lowest validation loss over epochs.
<code>fit_transform(self, train_data)</code>	Fit VAE model and transform Methylation Array using VAE model.
<code>transform(self, train_data)</code>	

Parameters	
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add_validation_set (*self*, *validation_data*)

Add validation data in the form of Validation DataLoader. Adding this will use validation data for early termination / generalization of model to unseen data.

Parameters

validation_data [type] Pytorch DataLoader housing validation MethylationDataset.

fit (*self*, *train_data*)

Fit VAE model to training data, best model returned with lowest validation loss over epochs.

Parameters

train_data [DataLoader] Training DataLoader that is loading MethylationDataset in batches.

Returns

self Autoencoder object with updated VAE model.

fit_transform (*self*, *train_data*)

Fit VAE model and transform Methylation Array using VAE model.

Parameters

train_data [type] Pytorch DataLoader housing training MethylationDataset.

Returns

np.array Latent Embeddings.

np.array Sample names from MethylationArray

np.array Outcomes from column of methylarray.

transform (*self*, *train_data*)

Parameters

train_data [type] Pytorch DataLoader housing training MethylationDataset.

Returns

np.array Latent Embeddings.

np.array Sample names from MethylationArray

np.array Outcomes from column of methylarray.

class methylnet.models.**MLPFinetuneVAE** (*mlp_model, n_epochs=None, loss_fn=None, optimizer_vae=None, optimizer_mlp=None, cuda=True, categorical=False, scheduler_opts={}, output_latent=True, train_decoder=False*)

Wraps VAE_MLP pytorch module into scikit-learn interface with fit, predict and fit_predict methods for ease-of-use model training/evaluation.

Parameters

mlp_model [type] VAE_MLP model.

n_epochs [type] Number epochs train for.

loss_fn [type] Loss function, pytorch, CrossEntropy, BCE, MSE depending on outcome.

optimizer_vae [type] Optimizer for VAE layers for finetuning original pretrained network.

optimizer_mlp [type] Optimizer for new appended MLP layers.

cuda [type] GPU?

categorical [type] Classification or regression outcome?

scheduler_opts [type] Options for learning rate scheduler, modulates learning rates for VAE and MLP.

output_latent [type] Whether to output latent embeddings during evaluation.

train_decoder [type] Retrain decoder to adjust for finetuning of VAE?

Attributes

model [type] VAE_MLP.

scheduler_vae [type] Learning rate modulator for VAE optimizer.

scheduler_mlp [type] Learning rate modulator for MLP optimizer.

loss_plt_fname [type] File where to plot loss over time; defunct.

embed_interval [type] How often to return embeddings; defunct.

validation_set [type] Validation set used for hyperparameter tuning and early stopping criteria for generalization.

return_latent [type] Return embedding during evaluation?

n_epochs

loss_fn

optimizer_vae

optimizer_mlp

cuda

categorical

output_latent

train_decoder

Methods

<code>add_validation_set(self, validation_data)</code>	Add validation data to reduce overfitting.
<code>fit(self, train_data)</code>	Fit MLP to training data to make predictions.
<code>predict(self, test_data)</code>	Short summary.

add_validation_set (*self*, *validation_data*)

Add validation data to reduce overfitting.

Parameters

validation_data [type] Validation Dataloader MethylationDataset.

fit (*self*, *train_data*)

Fit MLP to training data to make predictions.

Parameters

train_data [type] DataLoader with Training MethylationDataset.

Returns

self MLPFinetuneVAE with updated parameters.

predict (*self*, *test_data*)

Short summary.

Parameters

test_data [type] Test DataLoader MethylationDataset.

Returns

np.array Predictions

np.array Ground truth

np.array Latent Embeddings

np.array Sample names.

```
class methylnet.models.TybaltTitusVAE (n_input, n_latent, hid-  
                                         den_layer_encoder_topology=[100, 100, 100],  
                                         cuda=False)
```

Pytorch NN Module housing VAE with fully connected layers and customizable topology.

Parameters

n_input [type] Number of input CpGs.

n_latent [type] Size of latent embeddings.

hidden_layer_encoder_topology [type] List, length of list contains number of hidden layers for encoder, and each element is number of neurons, mirrored for decoder.

cuda [type] GPU?

Attributes

cuda_on [type] GPU?

pre_latent_topology [type] Hidden layer topology for encoder.

post_latent_topology [type] Mirrored hidden layer topology for decoder.

encoder_layers [list] Encoder pytorch layers.

encoder [type] Encoder layers wrapped into pytorch module.

z_mean [type] Linear layer from last encoder layer to mean layer.

z_var [type] Linear layer from last encoder layer to var layer.

z_develop [type] Linear layer connecting sampled latent embedding to first layer decoder.

decoder_layers [type] Decoder layers wrapped into pytorch module.

output_layer [type] Linear layer connecting last decoder layer to output layer, which is same size as input..

decoder [type] Wraps decoder_layers and output_layers into Sequential module.

n_input

n_latent

Methods

<code>__call__(self, *input, **kwargs)</code>	Call self as a function.
<code>add_module(self, name, module)</code>	Adds a child module to the current module.
<code>apply(self, fn)</code>	Applies fn recursively to every submodule (as returned by <code>.children()</code>) as well as self.
<code>buffers(self[, recurse])</code>	Returns an iterator over module buffers.
<code>children(self)</code>	Returns an iterator over immediate children modules.
<code>cpu(self)</code>	Moves all model parameters and buffers to the CPU.
<code>cuda(self[, device])</code>	Moves all model parameters and buffers to the GPU.
<code>decode(self, z)</code>	Decode latent embeddings back into reconstructed input.
<code>double(self)</code>	Casts all floating point parameters and buffers to double datatype.
<code>encode(self, x)</code>	Encode input into latent representation.
<code>eval(self)</code>	Sets the module in evaluation mode.
<code>extra_repr(self)</code>	Set the extra representation of the module
<code>float(self)</code>	Casts all floating point parameters and buffers to float datatype.
<code>forward(self, x)</code>	Return reconstructed output, mean and variance of embeddings.
<code>forward_predict(self, x)</code>	Forward pass from input to reconstructed input.
<code>get_latent_z(self, x)</code>	Encode X into reparameterized latent representation.
<code>half(self)</code>	Casts all floating point parameters and buffers to half datatype.
<code>load_state_dict(self, state_dict[, strict])</code>	Copies parameters and buffers from state_dict into this module and its descendants.
<code>modules(self)</code>	Returns an iterator over all modules in the network.
<code>named_buffers(self[, prefix, recurse])</code>	Returns an iterator over module buffers, yielding both the name of the buffer as well as the buffer itself.
<code>named_children(self)</code>	Returns an iterator over immediate children modules, yielding both the name of the module as well as the module itself.

Continued on next page

Table 3 – continued from previous page

<code>named_modules(self[, memo, prefix])</code>	Returns an iterator over all modules in the network, yielding both the name of the module as well as the module itself.
<code>named_parameters(self[, prefix, recurse])</code>	Returns an iterator over module parameters, yielding both the name of the parameter as well as the parameter itself.
<code>parameters(self[, recurse])</code>	Returns an iterator over module parameters.
<code>register_backward_hook(self, hook)</code>	Registers a backward hook on the module.
<code>register_buffer(self, name, tensor)</code>	Adds a persistent buffer to the module.
<code>register_forward_hook(self, hook)</code>	Registers a forward hook on the module.
<code>register_forward_pre_hook(self, hook)</code>	Registers a forward pre-hook on the module.
<code>register_parameter(self, name, param)</code>	Adds a parameter to the module.
<code>sample_z(self, mean, logvar)</code>	Sample latent embeddings, reparameterize by adding noise to embedding.
<code>state_dict(self[, destination, prefix, ...])</code>	Returns a dictionary containing a whole state of the module.
<code>to(self, *args, **kwargs)</code>	Moves and/or casts the parameters and buffers.
<code>train(self[, mode])</code>	Sets the module in training mode.
<code>type(self, dst_type)</code>	Casts all parameters and buffers to <code>dst_type</code> .
<code>zero_grad(self)</code>	Sets gradients of all model parameters to zero.

share_memory	
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decode (*self*, *z*)

Decode latent embeddings back into reconstructed input.

Parameters**z** [type] Reparameterized latent embedding.**Returns****torch.tensor** Reconstructed input.**encode** (*self*, *x*)

Encode input into latent representation.

Parameters**x** [type] Input methylation data.**Returns****torch.tensor** Learned mean vector of embeddings.**torch.tensor** Learned variance of learned mean embeddings.**forward** (*self*, *x*)

Return reconstructed output, mean and variance of embeddings.

forward_predict (*self*, *x*)

Forward pass from input to reconstructed input.

get_latent_z (*self*, *x*)

Encode X into reparameterized latent representation.

Parameters**x** [type] Input methylation data.

Returns

torch.tensor Latent embeddings.

sample_z (*self, mean, logvar*)

Sample latent embeddings, reparameterize by adding noise to embedding.

Parameters

mean [type] Learned mean vector of embeddings.

logvar [type] Learned variance of learned mean embeddings.

Returns

torch.tensor Mean + noise, reparameterization trick.

```
class methylnet.models.VAE_MLP (vae_model,      n_output,      categorical=False,      hid-
                                den_layer_topology=[100, 100, 100], dropout_p=0.2,
                                add_softmax=False)
```

VAE_MLP, pytorch module used to both finetune VAE embeddings and simultaneously train downstream MLP layers for classification/regression tasks.

Parameters

vae_model [type] VAE pytorch model for methylation data.

n_output [type] Number of outputs at end of model.

categorical [type] Classification or regression problem?

hidden_layer_topology [type] Hidden Layer topology, list of size number of hidden layers for MLP and each element contains number of neurons per layer.

dropout_p [type] Apply dropout regularization to reduce overfitting.

add_softmax [type] Softmax the output before evaluation.

Attributes

vae [type] Pytorch VAE module.

topology [type] List with hidden layer topology of MLP.

mlp_layers [type] All MLP layers (# layers and neurons per layer)

output_layer [type] nn.Linear connecting last MLP layer and output nodes.

mlp [type] nn.Sequential wraps all layers into sequential ordered pytorch module.

output_z [type] Whether to output latent embeddings.

n_output

categorical

add_softmax

dropout_p

Methods

<code>__call__(self, *input, **kwargs)</code>	Call self as a function.
<code>add_module(self, name, module)</code>	Adds a child module to the current module.

Continued on next page

Table 4 – continued from previous page

<code>apply(self, fn)</code>	Applies <code>fn</code> recursively to every submodule (as returned by <code>.children()</code>) as well as self.
<code>buffers(self[, recurse])</code>	Returns an iterator over module buffers.
<code>children(self)</code>	Returns an iterator over immediate children modules.
<code>cpu(self)</code>	Moves all model parameters and buffers to the CPU.
<code>cuda(self[, device])</code>	Moves all model parameters and buffers to the GPU.
<code>decode(self, z)</code>	Run VAE decoder on embeddings.
<code>double(self)</code>	Casts all floating point parameters and buffers to <code>double</code> datatype.
<code>eval(self)</code>	Sets the module in evaluation mode.
<code>extra_repr(self)</code>	Set the extra representation of the module
<code>float(self)</code>	Casts all floating point parameters and buffers to <code>float</code> datatype.
<code>forward(self, x)</code>	Pass data in to return predictions and embeddings.
<code>forward_embed(self, x)</code>	Return predictions, latent embeddings and reconstructed input.
<code>forward_predict(self, x)</code>	Make predictions, based on <code>output_z</code> , either output predictions or output embeddings.
<code>half(self)</code>	Casts all floating point parameters and buffers to <code>half</code> datatype.
<code>load_state_dict(self, state_dict[, strict])</code>	Copies parameters and buffers from <code>state_dict</code> into this module and its descendants.
<code>modules(self)</code>	Returns an iterator over all modules in the network.
<code>named_buffers(self[, prefix, recurse])</code>	Returns an iterator over module buffers, yielding both the name of the buffer as well as the buffer itself.
<code>named_children(self)</code>	Returns an iterator over immediate children modules, yielding both the name of the module as well as the module itself.
<code>named_modules(self[, memo, prefix])</code>	Returns an iterator over all modules in the network, yielding both the name of the module as well as the module itself.
<code>named_parameters(self[, prefix, recurse])</code>	Returns an iterator over module parameters, yielding both the name of the parameter as well as the parameter itself.
<code>parameters(self[, recurse])</code>	Returns an iterator over module parameters.
<code>register_backward_hook(self, hook)</code>	Registers a backward hook on the module.
<code>register_buffer(self, name, tensor)</code>	Adds a persistent buffer to the module.
<code>register_forward_hook(self, hook)</code>	Registers a forward hook on the module.
<code>register_forward_pre_hook(self, hook)</code>	Registers a forward pre-hook on the module.
<code>register_parameter(self, name, param)</code>	Adds a parameter to the module.
<code>state_dict(self[, destination, prefix, ...])</code>	Returns a dictionary containing a whole state of the module.
<code>to(self, *args, **kwargs)</code>	Moves and/or casts the parameters and buffers.
<code>toggle_latent_z(self)</code>	Toggle whether to output latent embeddings during forward pass.
<code>train(self[, mode])</code>	Sets the module in training mode.
<code>type(self, dst_type)</code>	Casts all parameters and buffers to <code>dst_type</code> .
<code>zero_grad(self)</code>	Sets gradients of all model parameters to zero.

share_memory	
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decode (*self*, *z*)

Run VAE decoder on embeddings.

Parameters

z [type] Embeddings.

Returns

torch.tensor Reconstructed Input.

forward (*self*, *x*)

Pass data in to return predictions and embeddings.

Parameters

x [type] Input data.

Returns

torch.tensor Predictions

torch.tensor Embeddings

forward_embed (*self*, *x*)

Return predictions, latent embeddings and reconstructed input.

Parameters

x [type] Input data

Returns

torch.tensor Predictions

torch.tensor Embeddings

torch.tensor Reconstructed input.

forward_predict (*self*, *x*)

Make predictions, based on output_z, either output predictions or output embeddings.

Parameters

x [type] Input Data.

Returns

torch.tensor Predictions or embeddings.

toggle_latent_z (*self*)

Toggle whether to output latent embeddings during forward pass.

`methylnet.models.project_vae` (*model*, *loader*, *cuda=True*)

Return Latent Embeddings of any data supplied to it.

Parameters

model [type] VAE Pytorch Model.

loader [type] Loads data one batch at a time.

cuda [type] GPU?

Returns

np.array Latent Embeddings.

np.array Sample names from MethylationArray

np.array Outcomes from column of methylarray.

`methylnet.models.test_mlp(model, loader, categorical, cuda=True, output_latent=True)`
Evaluate MLP on testing set, output predictions.

Parameters

model [type] VAE_MLP model.

loader [type] DataLoader with MethylationDataSet

categorical [type] Categorical or continuous predictions.

cuda [type] GPU?

output_latent [type] Output latent embeddings in addition to predictions?

Returns

np.array Predictions

np.array Ground truth

np.array Latent Embeddings

np.array Sample names.

`methylnet.models.train_decoder_(model, x, z)`

Run if retraining decoder to adjust for adjusted latent embeddings during finetuning of embedding layers for VAE_MLP.

Parameters

model [type] VAE_MLP model.

x [type] Input methylation data.

z [type] Latent Embeddings

Returns

nn.Module VAE_MLP module with updated decoder parameters.

float Reconstruction loss over all batches.

`methylnet.models.train_mlp(model, loader, loss_func, optimizer_vae, optimizer_mlp, cuda=True, categorical=False, train_decoder=False)`

Train Multi-layer perceptron appended to latent embeddings of VAE via transfer learning. Do this for one iteration.

Parameters

model [type] VAE_MLP model.

loader [type] DataLoader with MethylationDataset.

loss_func [type] Loss function (BCE, CrossEntropy, MSE).

optimizer_vae [type] Optimizer for pytorch VAE.

optimizer_mlp [type] Optimizer for outcome MLP layers.

cuda [type] GPU?

categorical [type] Predicting categorical or continuous outcomes.

train_decoder [type] Retrain decoder during training loop to adjust for fine-tuned embeddings.

Returns

nn.Module Training VAE_MLP model with updated parameters.

float Training loss over all batches

```
methylnet.models.train_vae(model, loader, loss_func, optimizer, cuda=True, epoch=0,  
                           kl_warm_up=0, beta=1.0)
```

Function for parameter update during VAE training for one iteration.

Parameters

model [type] VAE torch model

loader [type] Data loader, generator that calls batches of data.

loss_func [type] Loss function for reconstruction error, nn.BCELoss or MSELoss

optimizer [type] SGD or Adam pytorch optimizer.

cuda [type] GPU?

epoch [type] Epoch of training, passed in from outer loop.

kl_warm_up [type] How many epochs until model is fully utilizes KL Loss.

beta [type] Weight given to KL Loss.

Returns

nn.Module Pytorch VAE model

float Total Training Loss across all batches

float Total Training reconstruction loss across all batches

float Total KL Loss across all batches

```
methylnet.models.vae_loss(output, input, mean, logvar, loss_func, epoch, kl_warm_up=0,  
                          beta=1.0)
```

Function to calculate VAE Loss, Reconstruction Loss + Beta KLLoss.

Parameters

output [torch.tensor] Reconstructed output from autoencoder.

input [torch.tensor] Original input data.

mean [type] Learned mean tensor for each sample point.

logvar [type] Variation around that mean sample point, learned from reparameterization.

loss_func [type] Loss function for reconstruction loss, MSE or BCE.

epoch [type] Epoch of training.

kl_warm_up [type] Number of epochs until fully utilizing KLLoss, begin saving models here.

beta [type] Weighting for KLLoss.

Returns

torch.tensor Total loss

torch.tensor Recon loss

torch.tensor KL loss

```
methylnet.models.val_decoder_(model, x, z)
```

Validation Loss over decoder.

Parameters

model [type] VAE_MLP model.

x [type] Input methylation data.

z [type] Latent Embeddings

Returns

float Reconstruction loss over all batches.

`methylnet.models.val_mlp(model, loader, loss_func, cuda=True, categorical=False, train_decoder=False)`
Find validation loss of VAE_MLP over one Epoch.

Parameters

model [type] VAE_MLP model.

loader [type] DataLoader with MethylationDataset.

loss_func [type] Loss function (BCE, CrossEntropy, MSE).

cuda [type] GPU?

categorical [type] Predicting categorical or continuous outcomes.

train_decoder [type] Retrain decoder during training loop to adjust for fine-tuned embeddings.

Returns

nn.Module VAE_MLP model.

float Validation loss over all batches

`methylnet.models.val_vae(model, loader, loss_func, optimizer, cuda=True, epoch=0, kl_warm_up=0, beta=1.0)`
Function for validation loss computation during VAE training for one epoch.

Parameters

model [type] VAE torch model

loader [type] Validation Data loader, generator that calls batches of data.

loss_func [type] Loss function for reconstruction error, nn.BCELoss or MSELoss

optimizer [type] SGD or Adam pytorch optimizer.

cuda [type] GPU?

epoch [type] Epoch of training, passed in from outer loop.

kl_warm_up [type] How many epochs until model is fully utilizes KL Loss.

beta [type] Weight given to KL Loss.

Returns

nn.Module Pytorch VAE model

float Total Validation Loss across all batches

float Total Validation reconstruction loss across all batches

float Total Validation KL Loss across all batches

PLOTTER.PY

Plotting mechanisms for training that are now defunct.

class `methylnet.plotter.Plot` (*title, xlab='vae1', ylab='vae2', data=[]*)
Stores plotting information; defunct, superceded, see `methylnet-visualize`.

Methods

return_img	
-------------------	--

class `methylnet.plotter.PlotTransformer` (*data, color_list*)
Plotting Transformer to help with plotting embeddings changing over epochs; defunct.

Methods

transform	
------------------	--

class `methylnet.plotter.Plotter` (*plots, animation=True*)
Plot embeddings and training curve from Plot objects; defunct, superceded, see `methylnet-visualize`.

Methods

animate	
write_plots	

SCHEDULERS.PY

Learning rate schedulers that help enable better and more generalizable models.

```
class methylnet.schedulers.CosineAnnealingWithRestartsLR(optimizer, T_max,
                                                         eta_min=0, last_epoch=-
                                                         1, T_mult=1.0, al-
                                                         pha_decay=1.0)
```

Borrowed from: https://github.com/mpyrozok/adamwr/blob/master/cyclic_scheduler.py Needs to be updated to reflect newest changes. From original docstring: Set the learning rate of each parameter group using a cosine annealing schedule, where η_{max} is set to the initial lr and T_{cur} is the number of epochs since the last restart in SGDR:

$$\eta_t = \eta_{min} + \frac{1}{2}(\eta_{max} - \eta_{min})(1 + \cos(\frac{T_{cur}}{T_{max}}\pi))$$

When last_epoch=-1, sets initial lr as lr. It has been proposed in

SGDR: Stochastic Gradient Descent with Warm Restarts. This implements the cosine annealing part of SGDR, the restarts and number of iterations multiplier.

Args: optimizer (Optimizer): Wrapped optimizer. T_max (int): Maximum number of iterations. T_mult (float): Multiply T_max by this number after each restart. Default: 1. eta_min (float): Minimum learning rate. Default: 0. last_epoch (int): The index of last epoch. Default: -1.

Attributes

step_n

Methods

load_state_dict(self, state_dict)	Loads the schedulers state.
state_dict(self)	Returns the state of the scheduler as a dict.

cosine	
get_lr	
restart	
step	

```
class methylnet.schedulers.Scheduler(optimizer=None, opts={'T_max': 10, 'T_mult': 2,
                                                         'eta_min': 5e-08, 'lr_scheduler_decay': 0.5, 'sched-
                                                         uler': 'null'})
```

Scheduler class that modulates learning rate of torch optimizers over epochs.

Parameters

optimizer [type] torch.Optimizer object
opts [type] Options of setting the learning rate scheduler, see default.

Attributes

schedulers [type] Different types of schedulers to choose from.
scheduler_step_fn [type] How scheduler updates learning rate.
initial_lr [type] Initial set learning rate.
scheduler_choice [type] What scheduler type was chosen.
scheduler [type] Scheduler object chosen that will more directly update optimizer LR.

Methods

<code>get_lr(self)</code>	Return current learning rate.
<code>step(self)</code>	Update optimizer learning rate

get_lr (*self*)
Return current learning rate.

Returns

float Current learning rate.

step (*self*)
Update optimizer learning rate

METHYLNET-EMBED

```
methylnet-embed [OPTIONS] COMMAND [ARGS]...
```

Options

--version
Show the version and exit.

8.1 launch_hyperparameter_scan

Launch randomized grid search of neural network hyperparameters.

```
methylnet-embed launch_hyperparameter_scan [OPTIONS]
```

Options

-hcsv, --hyperparameter_input_csv <hyperparameter_input_csv>
CSV file containing hyperparameter inputs. [default: embeddings/embed_hyperparameters_scan_input.csv]

-hl, --hyperparameter_output_log <hyperparameter_output_log>
CSV file containing prior runs. [default: embeddings/embed_hyperparameters_log.csv]

-g, --generate_input
Generate hyperparameter input csv.

-c, --job_chunk_size <job_chunk_size>
If not series, chunk up and run these number of commands at once..

-sc, --stratify_column <stratify_column>
Column to stratify samples on. [default: disease]

-r, --reset_all
Run all jobs again.

-t, --torque
Submit jobs on torque.

-gpu, --gpu <gpu>
If torque submit, which gpu to use. [default: -1]

-gn, --gpu_node <gpu_node>
If torque submit, which gpu node to use. [default: -1]

- nh, --nohup**
Nohup launch jobs.
- mc, --model_complexity_factor** <model_complexity_factor>
Degree of neural network model complexity for hyperparameter search. Search for less wide and less deep networks with a lower complexity value, bounded between 0 and infinity. [default: 1.0]
- b, --set_beta** <set_beta>
Set beta value, bounded between 0 and infinity. Set to -1 [default: 1.0]
- j, --n_jobs** <n_jobs>
Number of jobs to generate.
- n, --n_jobs_relaunch** <n_jobs_relaunch>
Relaunch n top jobs from previous run. [default: 0]
- cp, --crossover_p** <crossover_p>
Rate of crossover between hyperparameters. [default: 0.0]
- v, --val_loss_column** <val_loss_column>
Validation loss column.
- a, --additional_command** <additional_command>
Additional command to input for torque run.
- cu, --cuda**
Use GPUs.
- grid, --hyperparameter_yaml** <hyperparameter_yaml>
YAML file with custom subset of hyperparameter grid.

8.2 perform_embedding

Perform variational autoencoding on methylation dataset.

`methylnet-embed perform_embedding [OPTIONS]`

Options

- i, --train_pkl** <train_pkl>
Input database for beta and phenotype data. [default: ./train_val_test_sets/train_methyl_array.pkl]
- o, --output_dir** <output_dir>
Output directory for embeddings. [default: ./embeddings/]
- c, --cuda**
Use GPUs.
- n, --n_latent** <n_latent>
Number of latent dimensions. [default: 64]
- lr, --learning_rate** <learning_rate>
Learning rate. [default: 0.001]
- wd, --weight_decay** <weight_decay>
Weight decay of adam optimizer. [default: 0.0001]
- e, --n_epochs** <n_epochs>
Number of epochs to train over. [default: 50]

-hlt, --hidden_layer_encoder_topology <hidden_layer_encoder_topology>
Topology of hidden layers, comma delimited, leave empty for one layer encoder, eg. 100,100 is example of 5-hidden layer topology. [default:]

-kl, --kl_warm_up <kl_warm_up>
Number of epochs before introducing kl_loss. [default: 0]

-b, --beta <beta>
Weighting of kl divergence. [default: 1.0]

-s, --scheduler <scheduler>
Type of learning rate scheduler. [default: null]

-d, --decay <decay>
Learning rate scheduler decay for exp selection. [default: 0.5]

-t, --t_max <t_max>
Number of epochs before cosine learning rate restart. [default: 10]

-eta, --eta_min <eta_min>
Minimum cosine LR. [default: 1e-06]

-m, --t_mult <t_mult>
Multiply current restart period times this number given number of restarts. [default: 2.0]

-bce, --bce_loss
Use bce loss instead of MSE.

-bs, --batch_size <batch_size>
Batch size. [default: 50]

-vp, --val_pkl <val_pkl>
Validation Set Methylation Array Location. [default: ./train_val_test_sets/val_methyl_array.pkl]

-w, --n_workers <n_workers>
Number of workers. [default: 9]

-conv, --convolutional
Use convolutional VAE.

-hs, --height_kernel_sizes <height_kernel_sizes>
Heights of convolutional kernels.

-ws, --width_kernel_sizes <width_kernel_sizes>
Widths of convolutional kernels.

-v, --add_validation_set
Evaluate validation set.

-l, --loss_reduction <loss_reduction>
Type of reduction on loss function. [default: sum]

-hl, --hyperparameter_log <hyperparameter_log>
CSV file containing prior runs. [default: embeddings/embed_hyperparameters_log.csv]

-sc, --stratify_column <stratify_column>
Column to stratify samples on. [default: disease]

-j, --job_name <job_name>
Embedding job name. [default: embed_job]

METHYLNET-PREDICT

```
methylnet-predict [OPTIONS] COMMAND [ARGS]...
```

Options

--version
Show the version and exit.

9.1 classification_report

Generate classification report that gives results from classification tasks.

```
methylnet-predict classification_report [OPTIONS]
```

Options

-r, --results_pickle <results_pickle>
Results from training, validation, and testing. [default: predictions/results.p]

-o, --output_dir <output_dir>
Output directory. [default: results/]

-e, --categorical_encoder <categorical_encoder>
One hot encoder if categorical model. If path exists, then return top positive controbutions per samples of that class. Encoded values must be of sample class as interest_col. [default: ./predictions/one_hot_encoder.p]

-a, --average_mechanism <average_mechanism>
Output directory. [default: micro]

9.2 launch_hyperparameter_scan

Run randomized hyperparameter scan of neural network hyperparameters.

```
methylnet-predict launch_hyperparameter_scan [OPTIONS]
```

Options

- hcsv, --hyperparameter_input_csv** <hyperparameter_input_csv>
CSV file containing hyperparameter inputs. [default: predictions/predict_hyperparameters_scan_input.csv]
- hl, --hyperparameter_output_log** <hyperparameter_output_log>
CSV file containing prior runs. [default: predictions/predict_hyperparameters_log.csv]
- g, --generate_input**
Generate hyperparameter input csv.
- c, --job_chunk_size** <job_chunk_size>
If not series, chunk up and run these number of commands at once..
- ic, --interest_cols** <interest_cols>
Column to stratify samples on.
- cat, --categorical**
Whether to run categorical analysis or not.
- r, --reset_all**
Run all jobs again.
- t, --torque**
Submit jobs on torque.
- gpu, --gpu** <gpu>
If torque submit, which gpu to use. [default: -1]
- gn, --gpu_node** <gpu_node>
If torque submit, which gpu node to use. [default: -1]
- nh, --nohup**
Nohup launch jobs.
- n, --n_jobs_relaunch** <n_jobs_relaunch>
Relaunch n top jobs from previous run. [default: 0]
- cp, --crossover_p** <crossover_p>
Rate of crossover between hyperparameters. [default: 0.0]
- mc, --model_complexity_factor** <model_complexity_factor>
Degree of neural network model complexity for hyperparameter search. Search for less wide networks with a lower complexity value, bounded between 0 and infinity. [default: 1.0]
- j, --n_jobs** <n_jobs>
Number of jobs to generate.
- v, --val_loss_column** <val_loss_column>
Validation loss column.
- sft, --add_softmax**
Add softmax for predicting probability distributions.
- a, --additional_command** <additional_command>
Additional command to input for torque run.
- cu, --cuda**
Use GPUs.
- grid, --hyperparameter_yaml** <hyperparameter_yaml>
YAML file with custom subset of hyperparameter grid.

9.3 make_new_predictions

Run prediction model again to further assess outcome. Only evaluate prediction model.

```
methylnet-predict make_new_predictions [OPTIONS]
```

Options

- tp, --test_pkl** <test_pkl>
Test database for beta and phenotype data. [default: ./train_val_test_sets/test_methyl_array.pkl]
- m, --model_pickle** <model_pickle>
Pytorch model containing forward_predict method. [default: ./predictions/output_model.p]
- bs, --batch_size** <batch_size>
Batch size. [default: 50]
- w, --n_workers** <n_workers>
Number of workers. [default: 9]
- ic, --interest_cols** <interest_cols>
Specify columns looking to make predictions on. [default: disease]
- cat, --categorical**
Multi-class prediction. [default: False]
- c, --cuda**
Use GPUs.
- e, --categorical_encoder** <categorical_encoder>
One hot encoder if categorical model. If path exists, then return top positive controbutions per samples of that class. Encoded values must be of sample class as interest_col. [default: ./predictions/one_hot_encoder.p]
- o, --output_dir** <output_dir>
Output directory for predictions. [default: ./new_predictions/]

9.4 make_prediction

Train prediction model by fine-tuning VAE and appending/training MLP to make classification/regression predictions on MethylationArrays.

```
methylnet-predict make_prediction [OPTIONS]
```

Options

- i, --train_pkl** <train_pkl>
Input database for beta and phenotype data. [default: ./train_val_test_sets/train_methyl_array.pkl]
- tp, --test_pkl** <test_pkl>
Test database for beta and phenotype data. [default: ./train_val_test_sets/test_methyl_array.pkl]
- vae, --input_vae_pkl** <input_vae_pkl>
Trained VAE. [default: ./embeddings/output_model.p]

-o, --output_dir <output_dir>
Output directory for predictions. [default: ./predictions/]

-c, --cuda
Use GPUs.

-ic, --interest_cols <interest_cols>
Specify columns looking to make predictions on. [default: disease]

-cat, --categorical
Multi-class prediction. [default: False]

-do, --disease_only
Only look at disease, or text before subtype_delimiter.

-hlt, --hidden_layer_topology <hidden_layer_topology>
Topology of hidden layers, comma delimited, leave empty for one layer encoder, eg. 100,100 is example of 5-hidden layer topology. [default:]

-lr_vae, --learning_rate_vae <learning_rate_vae>
Learning rate VAE. [default: 1e-05]

-lr_mlp, --learning_rate_mlp <learning_rate_mlp>
Learning rate MLP. [default: 0.001]

-wd, --weight_decay <weight_decay>
Weight decay of adam optimizer. [default: 0.0001]

-dp, --dropout_p <dropout_p>
Dropout Percentage. [default: 0.2]

-e, --n_epochs <n_epochs>
Number of epochs to train over. [default: 50]

-s, --scheduler <scheduler>
Type of learning rate scheduler. [default: null]

-d, --decay <decay>
Learning rate scheduler decay for exp selection. [default: 0.5]

-t, --t_max <t_max>
Number of epochs before cosine learning rate restart. [default: 10]

-eta, --eta_min <eta_min>
Minimum cosine LR. [default: 1e-06]

-m, --t_mult <t_mult>
Multiply current restart period times this number given number of restarts. [default: 2.0]

-bs, --batch_size <batch_size>
Batch size. [default: 50]

-vp, --val_pkl <val_pkl>
Validation Set Methylation Array Location. [default: ./train_val_test_sets/val_methyl_array.pkl]

-w, --n_workers <n_workers>
Number of workers. [default: 9]

-v, --add_validation_set
Evaluate validation set.

-l, --loss_reduction <loss_reduction>
Type of reduction on loss function. [default: sum]

- hl, --hyperparameter_log** <hyperparameter_log>
CSV file containing prior runs. [default: predictions/predict_hyperparameters_log.csv]
- j, --job_name** <job_name>
Embedding job name. [default: predict_job]
- sft, --add_softmax**
Add softmax for predicting probability distributions. Experimental.

9.5 regression_report

Generate regression report that gives concise results from regression tasks.

```
methylnet-predict regression_report [OPTIONS]
```

Options

- r, --results_pickle** <results_pickle>
Results from training, validation, and testing. [default: predictions/results.p]
- o, --output_dir** <output_dir>
Output directory. [default: results/]

METHYLNET-INTERPRET

```
methylnet-interpret [OPTIONS] COMMAND [ARGS]...
```

Options

--version
Show the version and exit.

10.1 bin_regression_shaps

Take aggregate individual scores from regression results, that normally are not categorized, and aggregate across new category created from continuous data.

```
methylnet-interpret bin_regression_shaps [OPTIONS]
```

Options

-s, --shapley_data <shapley_data>
Pickle containing top CpGs. [default: ./interpretations/shapley_explanations/shapley_data.p]

-t, --test_pkl <test_pkl>
Pickle containing testing set. [default: ./train_val_test_sets/test_methyl_array.pkl]

-c, --col <col>
Column to turn into bins. [default: age]

-n, --n_bins <n_bins>
Number of bins. [default: 10]

-ot, --output_test_pkl <output_test_pkl>
Binned shap pickle for further testing. [default: ./train_val_test_sets/test_methyl_array_shap_binned.pkl]

-os, --output_shap_pkl <output_shap_pkl>
Pickle containing top CpGs, binned phenotype. [default: ./interpretations/shapley_explanations/shapley_binned.p]

10.2 extract_methylation_array

Subset and write methylation array using top SHAP cpGs.

```
methylnet-interpret extract_methylation_array [OPTIONS]
```

Options

- s, --shapley_data** <shapley_data>
Pickle containing top CpGs. [default: ./interpretations/shapley_explanations/shapley_data.p]
- co, --classes_only**
Only take top CpGs from each class. [default: False]
- o, --output_dir** <output_dir>
Output directory for methylation array. [default: ./interpretations/shapley_explanations/top_cpgs_extracted_methylarr/]
- t, --test_pkl** <test_pkl>
Pickle containing testing set. [default: ./train_val_test_sets/test_methyl_array.pkl]
- c, --col** <col>
Column to color for output csvs. [default:]
- g, --global_vals**
Only take top CpGs globally. [default: False]
- n, --n_extract** <n_extract>
Number cpGs to extract. [default: 1000]

10.3 grab_lola_db_cache

Download core and extended LOLA databases for enrichment tests.

```
methylnet-interpret grab_lola_db_cache [OPTIONS]
```

Options

- o, --output_dir** <output_dir>
Output directory for lola dbs. [default: ./lola_db/]

10.4 interpret_biology

Interrogate CpGs with high SHAPley scores for individuals, classes, or overall, for enrichments, genes, GO, KEGG, LOLA, overlap with popular cpG sets, GSEA, find nearby cpGs to top.

```
methylnet-interpret interpret_biology [OPTIONS]
```

Options

- a, --all_cpGs_pickle** <all_cpGs_pickle>
List of all cpGs used in shapley analysis. [default: ./interpretations/shapley_explanations/all_cpGs.p]
- s, --shapley_data_list** <shapley_data_list>
Pickle containing top CpGs. [default: ./interpretations/shapley_explanations/shapley_data.p]

```

-o, --output_dir <output_dir>
    Output directory for interpretations. [default: ./interpretations/biological_explanations/]

-w, --analysis <analysis>
    Choose biological analysis. [default: GO]

-n, --n_workers <n_workers>
    Number workers. [default: 8]

-l, --lola_db <lola_db>
    LOLA region db. [default: ./lola_db/core/nm/t1/resources/regions/LOLACore/hg19/]

-i, --individuals <individuals>
    Individuals to evaluate. [default: ]

-c, --classes <classes>
    Classes to evaluate. [default: ]

-m, --max_gap <max_gap>
    Genomic distance to search for nearby CpGs than found top cpgs shapleys. [default: 1000]

-ci, --class_intersect
    Compute class shapleys by intersection of individuals. [default: False]

-lo, --length_output <length_output>
    Number enriched terms to print. [default: 20]

-ss, --set_subtraction
    Only consider CpGs relevant to particular class. [default: False]

-int, --intersection
    CpGs common to all classes. [default: False]

-col, --collections <collections>
    Lola collections. [default: ]

-gsea, --gsea_analyses <gsea_analyses>
    Gene set enrichment analysis to choose from, if chosen will override other analysis options. http://software.broadinstitute.org/gsea/msigdb/collections.jsp [default: ]

-gsp, --gsea_pickle <gsea_pickle>
    Gene set enrichment analysis to choose from. [default: ./data/gsea_collections.p]

-d, --depletion
    Run depletion LOLA analysis instead of enrichment. [default: False]

-ov, --overlap_test
    Run overlap test with IDOL library instead of all other tests. [default: False]

-cgs, --cpg_set <cpg_set>
    Test for clock or IDOL enrichments. [default: IDOL]

-g, --top_global
    Look at global top cpgs, overwrites classes and individuals. [default: False]

-ex, --extract_library
    Extract a library of cpgs to subset future array for inspection or prediction tests. [default: False]

```

10.5 interpret_embedding_classes

Compare average distance between classes in embedding space.

```
methylnet-interpret interpret_embedding_classes [OPTIONS]
```

Options

- i, --embedding_methyl_array_pkl** <embedding_methyl_array_pkl>
Use ./predictions/vae_mlp_methyl_arr.pkl or ./embeddings/vae_methyl_arr.pkl for vae interpretations. [default: ./embeddings/vae_methyl_arr.pkl]
- c, --pheno_col** <pheno_col>
Column to separate on. [default: disease]
- m, --metric** <metric>
Distance metric to compare classes. [default: cosine]
- t, --trim** <trim>
Trim outlier distances. Number 0-0.5. [default: 0.0]
- o, --output_csv** <output_csv>
Distances between classes. [default: ./results/class_embedding_differences.csv]
- op, --output_pval_csv** <output_pval_csv>
If specify a CSV file, outputs pairwise manova tests between embeddings for clusters. [default:]

10.6 list_classes

List classes/multioutput regression cell-types that have SHAPley data to be interrogated.

```
methylnet-interpret list_classes [OPTIONS]
```

Options

- s, --shapley_data** <shapley_data>
Pickle containing top CpGs. [default: ./interpretations/shapley_explanations/shapley_data.p]

10.7 list_individuals

List individuals that have ShapleyData. Not all made the cut from the test dataset.

```
methylnet-interpret list_individuals [OPTIONS]
```

Options

- s, --shapley_data** <shapley_data>
Pickle containing top CpGs. [default: ./interpretations/shapley_explanations/shapley_data.p]

10.8 order_results_by_col

Order results that produces some CSV by phenotype column, alphabetical ordering to show maybe that results group together. Plot using pymethyl-visualize.

```
methylnet-interpret order_results_by_col [OPTIONS]
```

Options

- i, --input_csv** <input_csv>
Output directory for cpg jaccard_stats. [default: ./interpretations/shapley_explanations/top_cpgs_jaccard/all_jaccard.csv]
- t, --test_pkl** <test_pkl>
Pickle containing testing set. [default: ./train_val_test_sets/test_methyl_array.pkl]
- c, --col** <col>
Column to sort on. [default: disease]
- o, --output_csv** <output_csv>
Output directory for cpg jaccard_stats. [default: ./interpretations/shapley_explanations/top_cpgs_jaccard/all_jaccard_sorted.csv]
- sym, --symmetric**
Is symmetric? [default: False]

10.9 plot_all_lola_outputs

Iterate through all LOLA csv results and plot forest plots of all enrichment scores.

```
methylnet-interpret plot_all_lola_outputs [OPTIONS]
```

Options

- l, --head_lola_dir** <head_lola_dir>
Location of lola output csvs. [default: interpretations/biological_explanations/]
- o, --plot_output_dir** <plot_output_dir>
Output directory for interpretations. [default: ./interpretations/biological_explanations/lola_plots/]
- d, --description_col** <description_col>
Description column for categorical variables [default: description]
- c, --cell_types** <cell_types>
Cell types. [default:]
- ov, --overwrite**
Overwrite existing plots. [default: False]

10.10 plot_lola_output

Plot LOLA results via forest plot for one csv file.

```
methylnet-interpret plot_lola_output [OPTIONS]
```

Options

- l, --lola_csv** <lola_csv>
Location of lola output csv. [default:]
- o, --plot_output_dir** <plot_output_dir>
Output directory for interpretations. [default: ./interpretations/biological_explanations/lola_plots/]
- d, --description_col** <description_col>
Description column for categorical variables [default: description]
- c, --cell_types** <cell_types>
Cell types. [default:]

10.11 plot_top_cpgs

Plot the top cpgs locations in the genome using circos.

```
methylnet-interpret plot_top_cpgs [OPTIONS]
```

Options

- s, --shapley_data** <shapley_data>
Pickle containing top CpGs. [default: ./interpretations/shapley_explanations/shapley_data.p]
- o, --output_dir** <output_dir>
Output directory for output plots. [default: ./interpretations/output_plots/]
- i, --individuals** <individuals>
Individuals to evaluate. [default:]
- c, --classes** <classes>
Classes to evaluate. [default:]

10.12 produce_shapley_data

Explanations (coefficients for CpGs) for every individual prediction. Produce SHAPley scores for each CpG for each individualized prediction, and then aggregate them across coarser classes. Store CpGs with top SHAPley scores as well for quick access. Store in Shapley data object.

```
methylnet-interpret produce_shapley_data [OPTIONS]
```

Options

- i, --train_pkl** <train_pkl>
Input database for beta and phenotype data. Use ./predictions/vae_mlp_methyl_arr.pkl or ./embeddings/vae_methyl_arr.pkl for vae interpretations. [default: ./train_val_test_sets/train_methyl_array.pkl]

```

-v, --val_pkl <val_pkl>
    Val database for beta and phenotype data. Use ./predictions/vae_mlp_methyl_arr.pkl or ./embed-
    dings/vae_methyl_arr.pkl for vae interpretations. [default: ./train_val_test_sets/val_methyl_array.pkl]

-t, --test_pkl <test_pkl>
    Pickle containing testing set. [default: ./train_val_test_sets/test_methyl_array.pkl]

-m, --model_pickle <model_pickle>
    Pytorch model containing forward_predict method. [default: ./predictions/output_model.p]

-w, --n_workers <n_workers>
    Number of workers. [default: 9]

-bs, --batch_size <batch_size>
    Batch size. [default: 512]

-c, --cuda
    Use GPUs.

-ns, --n_samples <n_samples>
    Number of samples for SHAP output. [default: 200]

-nf, --n_top_features <n_top_features>
    Top features to select for shap outputs. If feature selection, instead use this number to choose number features
    that make up top global score. [default: 20]

-o, --output_dir <output_dir>
    Output directory for interpretations. [default: ./interpretations/shapley_explanations/]

-meth, --method <method>
    Explainer type. [default: kernel]

-ssbs, --shap_sample_batch_size <shap_sample_batch_size>
    Break up shapley computations into batches. Set to 0 for only 1 batch. [default: 0]

-r, --n_random_representative <n_random_representative>
    Number of representative samples to choose for background selection. Is this number for regression models, or
    this number per class for classification problems. [default: 0]

-col, --interest_col <interest_col>
    Column of interest for sample selection for explainer training. [default: disease]

-rt, --n_random_representative_test <n_random_representative_test>
    Number of representative samples to choose for test set. Is this number for regression models, or this number
    per class for classification problems. [default: 0]

-e, --categorical_encoder <categorical_encoder>
    One hot encoder if categorical model. If path exists, then return top positive controbutions per samples of that
    class. Encoded values must be of sample class as interest_col. [default: ./predictions/one_hot_encoder.p]

-cc, --cross_class
    Find importance of features for prediction across classes.

-fs, --feature_selection
    Perform feature selection using top global SHAP scores. [default: False]

-top, --top_outputs <top_outputs>
    Get shapley values for fewer outputs if feature selection. [default: 0]

-vae, --vae_interpret
    Use model to get shapley values for VAE latent dimensions, only works if proper datasets are set. [default:
    False]

```

-cl, --pred_class <pred_class>
Prediction class top cpgs. [default:]

-r, --results_csv <results_csv>
Remove all misclassifications. [default: ./predictions/results.csv]

-ind, --individual <individual>
One individual top cpgs. [default:]

-rc, --residual_cutoff <residual_cutoff>
Activate for regression interpretations. Standard deviation in residuals before removing. [default: 0.0]

-cn, --cell_names <cell_names>
Multioutput names for multi-output regression. [default:]

-dsd, --dump_shap_data
Dump raw SHAP data to numpy file, warning, may be large file. [default: False]

10.13 reduce_top_cpgs

Reduce set of top cpgs.

```
methylnet-interpret reduce_top_cpgs [OPTIONS]
```

Options

-s, --shapley_data <shapley_data>
Pickle containing top CpGs. [default: ./interpretations/shapley_explanations/shapley_data.p]

-nf, --n_top_features <n_top_features>
Top features to select for shap outputs. [default: 500]

-o, --output_pkl <output_pkl>
Pickle containing top CpGs, reduced number. [default: ./interpretations/shapley_explanations/shapley_reduced_data.p]

10.14 regenerate_top_cpgs

Increase size of Top CpGs using the original SHAP scores.

```
methylnet-interpret regenerate_top_cpgs [OPTIONS]
```

Options

-s, --shapley_data <shapley_data>
Pickle containing top CpGs. [default: ./interpretations/shapley_explanations/shapley_data.p]

-nf, --n_top_features <n_top_features>
Top features to select for shap outputs. [default: 500]

-o, --output_pkl <output_pkl>
Pickle containing top CpGs, reduced number. [default: ./interpretations/shapley_explanations/shapley_reduced_data.p]

- a, --abs_val**
Top CpGs found using absolute value.
- n, --neg_val**
Return top CpGs that are making negative contributions.

10.15 return_shap_values

Return matrix of shapley values per class, with option to, classes/individuals are columns, CpGs are rows, option to plot multiple histograms/density plots.

```
methylnet-interpret return_shap_values [OPTIONS]
```

Options

- s, --shapley_data <shapley_data>**
Pickle containing top CpGs. [default: ./interpretations/shapley_explanations/shapley_data.p]
- o, --output_dir <output_dir>**
Output directory for output plots. [default: ./interpretations/shap_outputs/]
- i, --individuals <individuals>**
Individuals to evaluate. [default:]
- c, --classes <classes>**
Classes to evaluate. [default:]
- hist, --output_histogram**
Whether to output a histogram for each class/individual of their SHAP scores.
- abs, --absolute**
Use sums of absolute values in making computations.
- log, --log_transform**
Log transform final results for plotting.

10.16 shapley_jaccard

Plot Shapley Jaccard Similarity Matrix to demonstrate sharing of Top CpGs between classes/groupings of individuals.

```
methylnet-interpret shapley_jaccard [OPTIONS]
```

Options

- s, --shapley_data <shapley_data>**
Pickle containing top CpGs. [default: ./interpretations/shapley_explanations/shapley_data.p]
- c, --class_names <class_names>**
Class names. [default:]
- o, --output_dir <output_dir>**
Output directory for cpg jaccard_stats. [default: ./interpretations/shapley_explanations/top_cpgs_jaccard/]

- ov, --overall**
Output overall similarity. [default: False]
- i, --include_individuals**
Output individuals. [default: False]
- co, --cooccurrence**
Output cooccurrence instead jaccard. [default: False]
- opt, --optimize_n_cpgs**
Search for number of top CpGs to use. [default: False]

10.17 split_hyper_hypo_methylation

Split SHAPleyData object by methylation type (needs to change; but low methylation is 0.5 beta value and below) and output to file.

```
methylnet-interpret split_hyper_hypo_methylation [OPTIONS]
```

Options

- s, --shapley_data <shapley_data>**
Pickle containing top CpGs. [default: ./interpretations/shapley_explanations/shapley_data.p]
- o, --output_dir <output_dir>**
Output directory for hypo/hyper shap data. [default: ./interpretations/shapley_explanations/shapley_data_by_methylation/]
- t, --test_pkl <test_pkl>**
Pickle containing testing set. [default: ./train_val_test_sets/test_methyl_array.pkl]
- thr, --threshold <threshold>**
Pickle containing testing set. [default: original]

10.18 view_methylation_top_cpgs

Write the Top CpGs for each class/individuals to file along with methylation information.

```
methylnet-interpret view_methylation_top_cpgs [OPTIONS]
```

Options

- s, --shapley_data <shapley_data>**
Pickle containing top CpGs. [default: ./interpretations/shapley_explanations/shapley_data.p]
- i, --individuals <individuals>**
Individuals. [default:]
- o, --output_dir <output_dir>**
Output directory for cpg methylation shap. [default: ./interpretations/shapley_explanations/top_cpgs_methylation/]
- t, --test_pkl <test_pkl>**
Pickle containing testing set. [default: ./train_val_test_sets/test_methyl_array.pkl]

METHYLNET-VISUALIZE

```
methylnet-visualize [OPTIONS] COMMAND [ARGS]...
```

Options

--version
Show the version and exit.

11.1 plot_roc_curve

Plot ROC Curves from classification tasks; requires classification_report be run first.

```
methylnet-visualize plot_roc_curve [OPTIONS]
```

Options

-r, --roc_curve_csv <roc_curve_csv>
Weighted ROC Curve. [default: results/Weighted_ROC.csv]

-o, --outputfilename <outputfilename>
Output image. [default: results/roc_curve.png]

11.2 plot_training_curve

Plot training curves as output from either the VAE training or VAE MLP.

```
methylnet-visualize plot_training_curve [OPTIONS]
```

Options

-t, --training_curve_file <training_curve_file>
Training and validation loss and learning curves. [default: predictions/training_val_curve.p]

-o, --outputfilename <outputfilename>
Output image. [default: results/training_curve.png]

-vae, --vae_train

Plot VAE Training curves.

-thr, --threshold <threshold>

Loss values get rid of greater than this.

METHYLNETH-TORQUE

```
methylnet-torque [OPTIONS] COMMAND [ARGS]...
```

Options

--version
Show the version and exit.

12.1 run_torque_job

Run torque job.

```
methylnet-torque run_torque_job [OPTIONS]
```

Options

-c, --command <command>
Command to execute through torque. [default:]

-gpu, --use_gpu
Specify whether to use GPUs. [default: False]

-a, --additions <additions>
Additional commands to add, one liner for now. [default:]

-q, --queue <queue>
Queue for torque submission, gpuq also a good one if using GPUs. [default: default]

-t, --time <time>
Walltime in hours for job. [default: 1]

-n, --ngpu <ngpu>
Number of gpus to request. [default: 0]

METHYLNTEST

```
methylnet-test [OPTIONS] COMMAND [ARGS]...
```

Options

--version

Show the version and exit.

13.1 test_pipeline

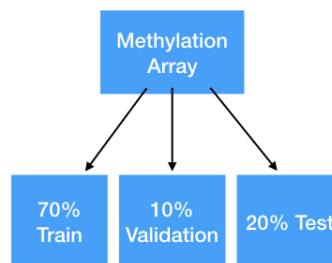
```
methylnet-test test_pipeline [OPTIONS]
```


EXAMPLE USAGE

Pipeline

- Download
- Format
- Preprocess
- Visualize
- Embedding
- Predict
- Explain

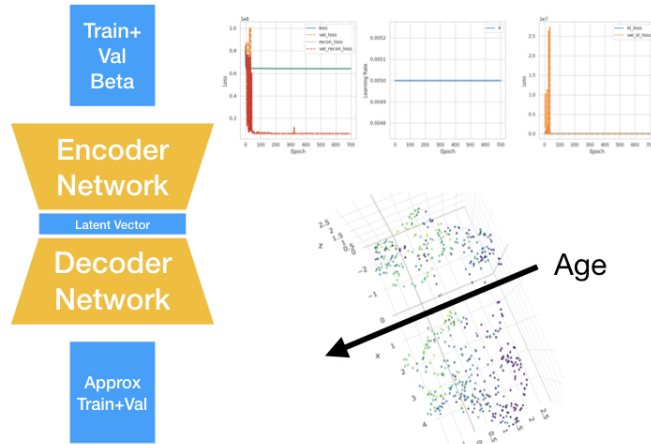
```
pymethyl-utils train_test_val_split -tp .8 -vp .125
```



Pipeline

```
python embedding.py launch_hyperparameter_scan -sc Age -t -mc 0.84 -b 1. -g -j 20
python embedding.py launch_hyperparameter_scan -sc Age -t -g -n 1 -b 1.
pymethyl-visualize transform_plot -i embeddings/vae_methyl_arr.pkl -nn 8 -c Age
```

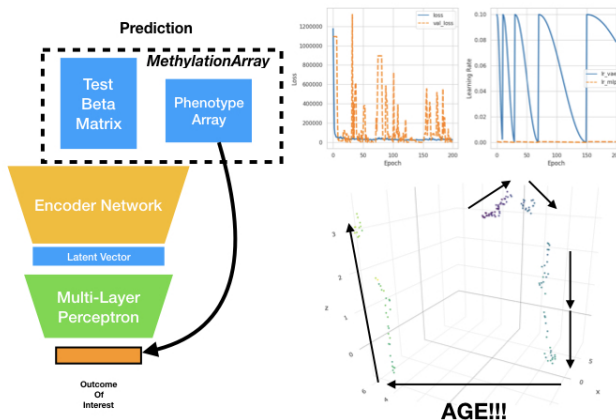
- Download
- Format
- Preprocess
- Visualize
- **Embedding**
- Predict
- Explain



Pipeline

```
python predictions.py launch_hyperparameter_scan -ic Age -t -mc 0.84 -g -j 200
python predictions.py launch_hyperparameter_scan -ic Age -t -g -n 1
pymethyl-visualize transform_plot -i predictions/vae_mlp_methyl_arr.pkl -nn 8 -c Age
```

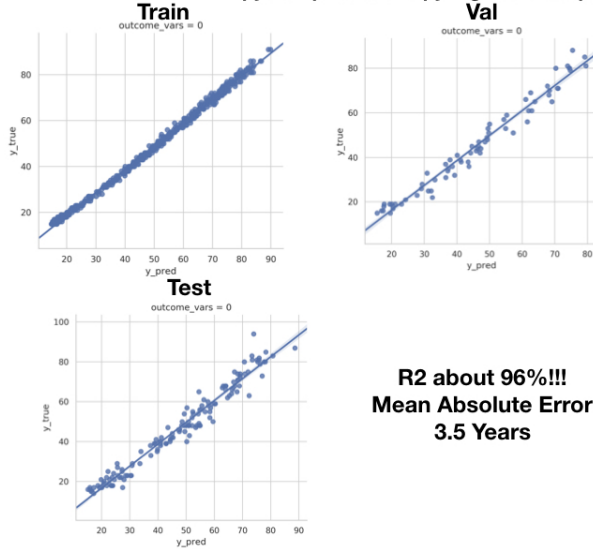
- Download
- Format
- Preprocess
- Visualize
- Embedding
- **Predict**
- Explain



Pipeline

python predictions.py regression_report

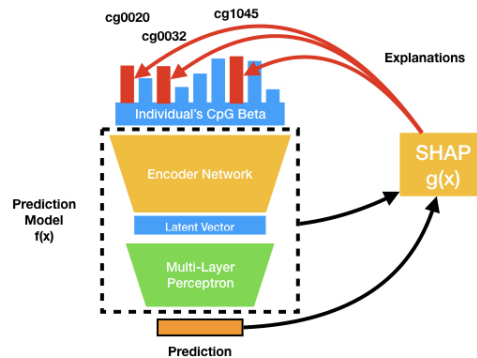
- Download
- Format
- Preprocess
- Visualize
- Embedding
- **Predict**
- Explain



Pipeline

python model_interpretability.py produce_shapley_data -mth gradient -ssbs 30 -ns 300 -bs 100 -rc 4. -r 0 -rt 0 -cn Age -nf 4000 -c

- Download
- Format
- Preprocess
- Visualize
- Embedding
- Predict
- **Explain**



Pipeline

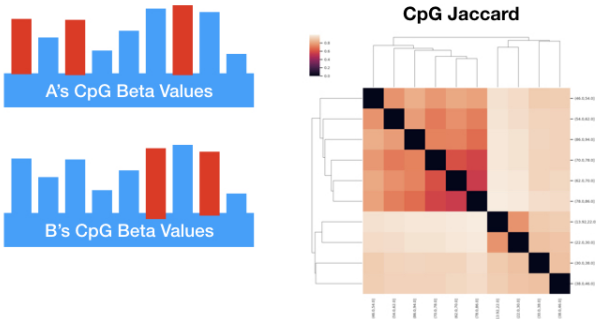
- Download
- Format
- Preprocess
- Visualize
- Embedding
- Predict
- Explain

```
python model_interpretability.py split_hyper_hypo_methylation

python model_interpretability.py bin_regression_shaps -c Age

python model_interpretability.py shapley_jaccard -c all -s ./interpretations/
shapley_explanations/shapley_binned.p -o ./interpretations/
shapley_explanations/top_cpgs_jaccard/ -ov

pymethyl-visualize plot_heatmap -m similarity -fs .7 -i ./interpretations/
shapley_explanations/top_cpgs_jaccard/all_jaccard.sorted.csv -o ./
interpretations/shapley_explanations/top_cpgs_jaccard/all_jaccard.png -
x -y -c
```

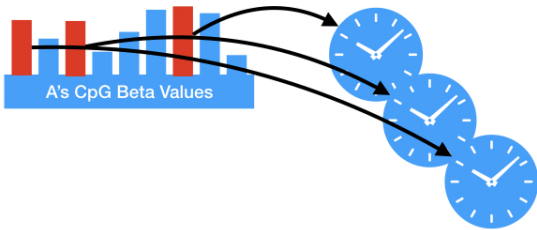


Pipeline

- Download
- Format
- Preprocess
- Visualize
- Embedding
- Predict
- Explain

```
python model_interpretability.py interpret_biology -ov -c all -s
interpretations/shapley_explanations/shapley_binned.p -cgs
clock

(54.0,62.0] top cpgs overlap with 9.63% of clock cpgs
(70.0,78.0] top cpgs overlap with 11.33% of clock cpgs
(62.0,70.0] top cpgs overlap with 12.46% of clock cpgs
(22.0,30.0] top cpgs overlap with 0.57% of clock cpgs
(78.0,86.0] top cpgs overlap with 10.2% of clock cpgs
(38.0,46.0] top cpgs overlap with 1.98% of clock cpgs
(13.92,22.0] top cpgs overlap with 0.85% of clock cpgs
(30.0,38.0] top cpgs overlap with 1.42% of clock cpgs
(46.0,54.0] top cpgs overlap with 7.93% of clock cpgs
(86.0,94.0] top cpgs overlap with 8.22% of clock cpgs
```



Pipeline

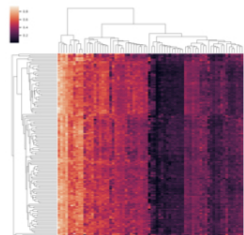
- Download
- Format
- Preprocess
- Visualize
- Embedding
- Predict
- Explain

```
pymethyl-utils subset_array -i train_val_test_sets/  
test_methyl_array.pkl -c ./interpretations/biological_explanations/  
cpg_library.pkl
```

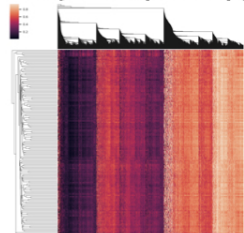
```
pymethyl-utils pkl_to_csv -i subset/methyl_array.pkl -o subset/  
pymethyl-visualize plot_heatmap -fs .7 -i ./subset/beta.csv -o ./  
subset/beta.png -c &
```

```
python model_interpretability.py  
extract_methylation_array -s  
interpretations/shapley_explanations/  
shapley_binned.p -c
```

Horvath CpGs



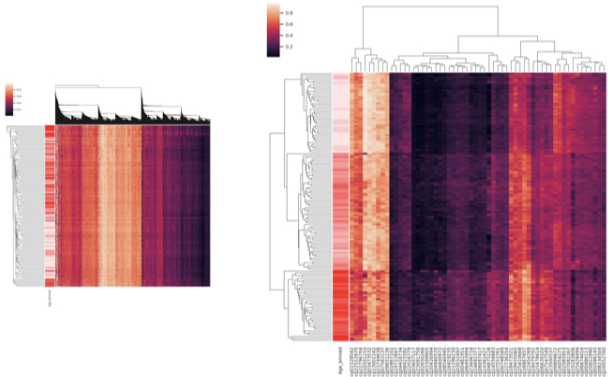
MethylNet Top 10k CpGs



Pipeline

- Download
- Format
- Preprocess
- Visualize
- Embedding
- Predict
- Explain

(13.92,22.0] top cpgs overlap with 0.0% of hannum cpgs
(22.0,30.0] top cpgs overlap with 0.0% of hannum cpgs
(30.0,38.0] top cpgs overlap with 1.45% of hannum cpgs
(38.0,46.0] top cpgs overlap with 4.35% of hannum cpgs
(46.0,54.0] top cpgs overlap with 39.13% of hannum cpgs
(54.0,62.0] top cpgs overlap with 78.26% of hannum cpgs
(62.0,70.0] top cpgs overlap with 79.71% of hannum cpgs
(70.0,78.0] top cpgs overlap with 82.61% of hannum cpgs
(78.0,86.0] top cpgs overlap with 65.22% of hannum cpgs
(86.0,94.0] top cpgs overlap with 65.22% of hannum cpgs



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