# **PyMethylProcess Documentation**

Release 0.1

Joshua Levy

1	PreProcessDataTypes.py	5
2	MethylationDataTypes.py	9
3	meffil_functions.py	13
4	general_machine_learning.py	15
5	pymethyl-install 5.1 change_gcc_path 5.2 install_bioconductor 5.3 install_custom 5.4 install_meffil 5.5 install_minfi_others 5.6 install_r_packages 5.7 install_some_deps 5.8 install_tcga_biolinks	17 17 17 18 18 18 18
6	pymethyl-visualize6.1plot_cell_type_results6.2plot_heatmap6.3transform_plot	19 19 19 20
7	pymethyl-preprocess 7.1 batch_deploy_preprocess 7.2 combine_methylation_arrays 7.3 concat_sample_sheets 7.4 create_sample_sheet 7.5 download_clinical 7.6 download_geo 7.7 download_tcga 7.8 feature_select 7.9 get_categorical_distribution 7.10 imputation_pipeline 7.11 meffil_encode 7.12 merge_sample_sheets 7.13 na_report 7.14 preprocess_pipeline 7.15 remove_diseases 7.16 split_preprocess_input_by_subtype	23 24 24 24 25 26 26 26 27 28 28 29 30 30

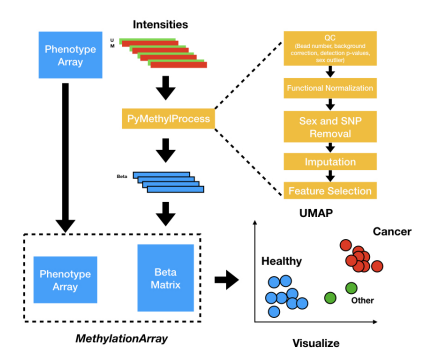
8	pyme	thyl-utils contained to the second contained to the se	33
	8.1	backup_pkl	33
	8.2	bin_column	33
	8.3	concat_csv	34
	8.4	counts	34
	8.5	create_external_validation_set	34
	8.6	est_age	35
	8.7	feature_select_train_val_test	35
	8.8	fix_key	36
	8.9	modify_pheno_data	36
	8.10	move_jpg	37
	8.11	overwrite_pheno_data	37
	8.12	pkl_to_csv	37
	8.13	print_number_sex_cpgs	38
	8.14	print_shape	38
	8.15	rate_regression	38
	8.16	ref_estimate_cell_counts	38
	8.17	ref_free_cell_deconv	39
	8.18	remove_sex	39
	8.19	remove_snps	40
	8.20	set_part_array_background	40
	8.21	stratify	40
	8.22	subset_array	41
	8.23	train_test_val_split	41
	8.24	write_cpgs	42
9	Indic	es and tables	43
Рy	thon N	Andule Index	45
Ind	dev		47

## https://github.com/Christensen-Lab-Dartmouth/PyMethylProcess

To get started, download pymethylprocess using Docker (joshualevy44/pymethylprocess) or PIP (pymethylprocess) and run pymethyl-install\_r\_dependencies.

See README.md in Github repository for more 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.



# **Pipeline**

pymethyl-preprocess download\_geo -g GSE87571

- Download

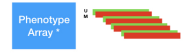


- Format
- Preprocess
- Visualize

# **Pipeline**

pymethyl-preprocess create\_sample\_sheet -is ./geo\_idats/ GSE87571\_clinical\_info.csv -s geo -i geo\_idats/ -os geo\_idats/samplesheet.csv -d "disease state:ch1" -c include\_col.txt

- Download
- Format
- Preprocess
- Visualize



# **Pipeline**

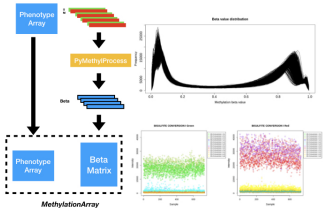
pymethyl-preprocess preprocess\_pipeline -i geo\_idats/ -p minfi -noob

pymethyl-utils remove\_sex -i preprocess\_outputs/methyl\_array.pkl

pymethyl-preprocess imputation\_pipeline -i ./autosomal/methyl\_array.pkl -s fancyimpute -m KNN -k 15

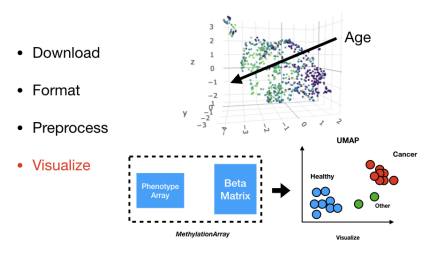
pymethyl-preprocess feature\_select -n 300000

- Download
- Format
- Preprocess
- Visualize



# **Pipeline**

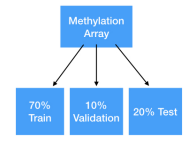
pymethyl-visualize transform\_plot -o visualizations/pre\_vae\_umap.html -c Age -nn 8



# **Pipeline**

pymethyl-utils train\_test\_val\_split -tp .8 -vp .125

- Download
- Format
- Preprocess
- Visualize



## PREPROCESSDATATYPES.PY

Contains datatypes core to downloading IDATs, preprocessing IDATs and samplesheets.

After creating beta, filter out outliers.

Get beta value matrix from minfi after finding RSet.

get\_beta()

```
class pymethylprocess.PreProcessDataTypes.PreProcessIDAT (idat_dir, minfi=None, en-
                                                                              mix=None, base=None,
                                                                              meffil=None)
     Class that will preprocess IDATs using R pipelines.
     idat_dir Location of idats or samplesheet csv.
     minfi Rpy2 importr minfi library, default to None will load through rpy2
     enmix Rpy2 importr enmix library, default to None will load through rpy2
     base Rpy2 importr base library, default to None will load through rpy2
     meffil Rpy2 importr meffil library, default to None will load through rpy2
     export_csv (output_dir)
          Export pheno and beta dataframes to CSVs
          output_dir Where to store csvs.
     export_pickle (output_pickle, disease=")
          Export pheno and beta dataframes to pickle, stored in python dict that can be loaded into MethylationArray
          output_pickle Where to store MethylationArray.
          disease Custom naming scheme for data.
     export_sql (output_db, disease=")
          Export pheno and beta dataframes to SQL
          output_db Where to store data, sqlite db.
          disease Custom naming scheme for data.
     extract_manifest()
          Get manifest from RGSet.
     extract_pheno_data(methylset=False)
          Extract pheno data from MSet or RGSet, minfi.
          methylset If MSet has beenn created, set to True, else extract from original RGSet.
     filter beta()
```

```
get meth()
     Get methylation intensity matrix from MSet
get unmeth()
     Get unmethylated intensity matrix from MSet
load idats()
     For minfi pipeline, load IDATs from specified idat dir.
move_jpg()
     Move jpeg files from current working directory to the idat directory.
output_pheno_beta (meffil=False)
     Get pheno and beta dataframe objects stored as attributes for input to MethylationArray object.
     meffil True if ran meffil pipeline.
plot_original_qc (output_dir)
     Plot QC results from ENmix pipeline and possible minfi. Still experimental.
     output_dir Where to store plots.
plot_qc_metrics(output_dir)
     Plot QC results from ENmix pipeline and possible minfi. Still experimental.
     output_dir Where to store plots.
preprocessENmix (n_cores=6)
     Run ENmix preprocessing pipeline.
     n cores Number of CPUs to use.
preprocessMeffil (n_cores=6,
                                                  qc_report_fname='qc/report.html',
                                                                                       normaliza-
                                     n_pcs=4,
                       tion_report_fname='norm/report.html',
                                                                  pc_plot_fname='qc/pc_plot.pdf',
                       useCache=True, qc_only=True, qc_parameters={'p.beadnum.cpgs':
                       'p.beadnum.samples': 0.1, 'p.detection.cpgs': 0.1, 'p.detection.samples': 0.1},
     Run meffil preprocessing pipeline with functional normalization.
     n cores Number of CPUs to use.
     n_pcs Number of principal components to use for functional normalization, set to -1 to autoselect via
         kneedle algorithm.
     qc_report_fname HTML filename to store QC report.
     normalization report fname HTML filename to store normalization report
     pc_plot_fname PDF file to store principal components plot.
     useCache Use saved QC objects instead of running through QC again.
     qc only Perform QC, then save and quit before normalization.
     qc_parameters Python dictionary with parameters for qc.
     rm_sex Remove non-autosomal cpgs?
preprocessNoob()
     Run minfi preprocessing with Noob normalization
preprocessRAW()
     Run minfi preprocessing with RAW normalization
```

```
preprocess_enmix_pipeline (n_cores=6, pipeline='enmix',
                                                                         noob=False,
                                                                                        ac only=False,
                                         use cache=False)
          Run complete ENmix or minfi preprocessing pipeline.
          n cores Number CPUs.
          pipeline Run enmix or minfi
          noob Noob norm or RAW if minfi running.
          qc only Save and quit after only running QC?
          use cache Load preexisting RGSet instead of running QC again.
     return_beta()
          Return minfi RSet after having created MSet.
     to_methyl_array(disease=")
          Convert results from preprocessing into MethylationArray, and directly return MethylationArray object.
          disease Custom naming scheme for data.
class pymethylprocess.PreProcessDataTypes.PreProcessPhenoData(pheno_sheet,
                                                                                     idat_dir,
                                                                                     header\ line=0)
     Class that will manipute phenotype samplesheet before preprocessing of IDATs.
     pheno_sheet Location of clinical info csv.
     idat dir Location of idats
     header line Where to start reading clinical csv
     concat (other_formatted_sheet)
          Concat multiple PreProcessPhenoData objects, concat their dataframes to accept more than one
          smaplesheet/dataset.
          other_formatted_sheet Other PreProcessPhenoData to concat.
     export (output_sheet_name)
          Export pheno data to csv after done with manipulation.
          output_sheet_name Output csv name.
     format_custom(basename_col, disease_class_column, include_columns={})
          Custom format clinical sheet if user supplied idats.
          basename_col Column name of sample names.
          disease_class_column Disease column of clinical info csv.
          include_columns Dictionary specifying other columns to include, and new names to assign them to.
     format geo(disease class column='methylation class:ch1', include columns={})
          Format clinical sheets if downloaded geo idats.
          disease_class_column Disease column of clinical info csv.
          include_columns Dictionary specifying other columns to include, and new names to assign them to.
     format_tcga (mapping_file='idat_filename_case.txt')
          Format clinical sheets if downloaded tcga idats.
          mapping_file Maps unids to proper tcga sample names, should be downloaded with tcga clinical infor-
```

mation.

```
Print categorical distribution, counts for each unique value in phenotype column.
          key Phenotype Column.
          disease_only Whether to split phenotype column entries by delimiter.
          subtype delimiter Subtype delimiter to split on.
     merge (other_formatted_sheet, use_second_sheet_disease=True, no_disease_merge=False)
          Merge multiple PreProcessPhenoData objects, merge their dataframes to accept more than one
          saplesheet/dataset or add more pheno info.
          other_formatted_sheet Other PreProcessPhenoData to merge.
          use_second_sheet_disease Change disease column to that of second sheet instead of first.
          no_disease_merge Keep both disease columns from both sheets.
     remove_diseases (exclude_disease_list, low_count, disease_only, subtype_delimiter)
          Remove samples with certain diseases from disease column.
          exclude disease list List containing diseases to remove.
          low count Remove samples that have less than x disease occurances in column.
          disease_only Whether to split phenotype column entries by delimiter.
          subtype_delimiter Subtype delimiter to split on.
     split_key (key, subtype_delimiter)
          Split pheno column by key, with subtype delimiter, eg. entry S1,s2 -> S1 with delimiter ",".
          key Pheno column name.
          subtype_delimiter Subtype delimiter to split on.
class pymethylprocess.PreProcessDataTypes.TCGADownloader
     Downloads TCGA and GEO IDAT and clinical data
     download_clinical(output_dir)
          Download TCGA Clinical Data.
          output dir Where to output clinical data csv.
     download_geo (query, output_dir)
          Download GEO IDATs.
          query GEO accession number to query, must be 450k/850k.
          output dir Output directory to store idats and clinical information csv
     download_tcga (output_dir)
          Download TCGA IDATs.
          output_dir Where to output idat files.
```

get\_categorical\_distribution (key, disease\_only=False, subtype\_delimiter=', ')

## **METHYLATIONDATATYPES.PY**

Contains datatypes core to storing beta and phenotype methylation data, and imputation.

class pymethylprocess.MethylationDataTypes.ImputerObject(solver, method, opts={})

Class that stores and accesses different types of imputers. Construct sklearn-like imputer given certain input arguments.

solver Library for imputation, eg. sklearn, fancyimpute.

method Imputation method in library, named.

opts Additional options to assign to imputer.

return\_imputer()

Return initialized sklearn-like imputer.

class pymethylprocess.MethylationDataTypes.MethylationArray ( $pheno\_df$ ,  $beta\_df$ , name=")

Stores beta and phenotype information and performs various operations. Initialize MethylationArray object by inputting dataframe of phenotypes and dataframe of beta values with samples as index.

**pheno df** Phenotype dataframe (samples x covariates)

**beta\_df** Beta Values Dataframe (samples x cpgs)

bin\_column (col, n\_bins)

Turn continuous variable/covariate into categorical bins. Returns name of new column and updates phenotype matrix to reflect this change.

col Continuous column of phenotype array to bin.

**n\_bins** Number of bins to create.

#### categorical\_breakdown (key)

Print categorical distribution, counts for each unique value in phenotype column.

key Phenotype Column.

feature\_select (n\_top\_cpgs, feature\_selection\_method='mad', metric='correlation', nn=10)

Perform unsupervised feature selection on MethylationArray.

**n\_top\_cpgs** Number of CpGs to retain.

**feature\_selection\_method** Method to perform selection.

metric If considering structural feature selection like SPEC, use this distance metric.

nn Number of nearest neighbors.

#### classmethod from\_pickle(input\_pickle)

Load MethylationArray stored in pickle.

```
Usage: MethylationArray.from_pickle([input_pickle])
input_pickle Stored MethylationArray pickle.
```

## groupby(key)

Groupby for Methylation Array. Returns generator of methylation arrays grouped by key.

**preprocess\_sample\_df** New phenotype dataframe.

#### impute (imputer)

Perform imputation on NaN beta vaues. Input imputater returned from ImputerObject.

**imputer** Type of imputer object, in sklearn type interface.

### merge\_preprocess\_sheet (preprocess\_sample\_df)

Feed in another phenotype dataframe that will be merged with existing phenotype array.

**preprocess\_sample\_df** New phenotype dataframe.

### overwrite\_pheno\_data(preprocess\_sample\_df)

Feed in another phenotype dataframe that will overwrite overlapping keys of existing phenotype array.

preprocess\_sample\_df New phenotype dataframe.

### remove\_missingness (cpg\_threshold=None, sample\_threshold=None)

Remove samples and CpGs with certain level of missingness..

cpg\_threshold If more than fraction of Samples for this CpG are missing, remove cpg.

**sample\_threshold** If more than fraction of CpGs for this sample are missing, remove sample.

#### remove\_na\_samples (outcome\_cols)

Remove samples of MethylationArray who have missing values in phenotype column.

outcome\_cols Phenotype columns, if any rows contain missing values, samples are removed.

## remove\_whitespace(key)

Remove whitespaces from phenotype column.

key Phenotype column.

#### return\_cpgs()

Return list of cpgs of MethylationArray

#### return\_idx()

Return sample names of MethylationArray.

### return\_raw\_beta\_array()

Return numpy array of methylation beta vaues.

#### return shape()

Return dimensionality and number of samples of beta matrix.

### split\_by\_subtype (disease\_only, subtype\_delimiter)

Split MethylationArray into generator of MethylationArrays by phenotype column. Much akin to groupby. Only splits from disease column.

disease\_only Consider disease superclass.

**subtype\_delimiter** How to break up disease column if using disease\_only.

#### split\_key (key, subtype\_delimiter)

Manipulate an entire phenotype column, splitting each element up by some delimiter.

**key** Phenotype column.

**subtype\_delimiter** How to break up strings in columns. S1,s2 -> S1 for instance.

 $split\_train\_test$  (train\\_p=0.8, stratified=True, disease\_only=False, key='disease', subtype\_delimiter=', ', val\_p=0.0)

Split MethylationArray into training and test sets, with option to stratify by categorical covariate.

**train\_p** Fraction of methylation array to use as training set.

stratified Whether to stratify by categorical variable.

**disease\_only** Consider disease superclass by some delimiter. For instance if disease is S1,s2, superclass would be S1.

key Column to stratify on.

subtype\_delimiter How to split disease column into super/subclass.

val\_p If set greater than 0, will create additional validation set, fraction of which is broken off from training set.

 $\verb|subsample| (key='disease', n\_samples=None, frac=None, categorical=False)|$ 

Subsample MethylationArray, make the set randomly smaller.

key If stratifying, use this column of pheno array.

**n\_samples** Number of samples to consider overall, or per stratum.

**frac** Alternative to n\_samples, where x frac of array or stratum is considered.

categorical Whether to stratify by column.

#### subset\_cpgs (cpgs)

Subset beta matrix by list of Cpgs. Parameters ———— cpgs

Cpgs to subset by.

#### subset index(index)

Subset MethylationArray by samples.

index Sample names to subset by.

## write\_csvs(output\_dir)

Write phenotype data and beta values to csvs.

output\_dir Directory to output csv files.

#### write\_db (conn, disease=")

Store phenotype data and beta values in SQL database.

**conn** SQLite connection.

disease Create new tables in db that are related to disease state by this name.

#### write\_pickle (output\_pickle, disease=")

Store phenotype data and beta values in pickle file. Is default file format for storing MethylationArray objects.

output\_pickle Pickle file to store MethylationArray data.

**class** pymethylprocess.MethylationDataTypes.MethylationArrays (*list\_methylation\_arrays*)
Literally a list of methylation arrays, with methods operate on these arrays that is memory efficient. Initialize with list of methylation arrays. Can optionally leave list empty or with one element.

**list\_methylation\_arrays** List of methylation arrays.

```
combine (array_generator=None)
```

Combine the list of methylation arrays into one array via concatenation of beta matrices and phenotype arrays.

```
array generator Generator of additional methylation arrays for computational memory minimization.
     impute (imputer)
          Impute all methylation arrays.
          imputer Type of imputation, sklearn-like.
     write dbs(conn)
          Write list of methylation arrays to SQL database. Recommend naming MethylationArray.
          conn SQL connection.
     write_pkls(pkl)
          Write list of methylation arrays to single pickle. Recommend naming each MethylationArray.
          pkl Pickle file to write to.
pymethylprocess.MethylationDataTypes.extract_pheno_beta_df_from_folder(folder)
     Return phenotype and beta dataframes from specified folder with csv.
     folder Input folder.
pymethylprocess.MethylationDataTypes.extract_pheno_beta_df_from_pickle_dict(input_dict,
                                                                                                    dis-
                                                                                                    ease=")
     Return phenotype and beta dataframes from specified dictionary storing MethylationArray python dictionary.
     input_dict Python disctionary storing pheno/beta information.
pymethylprocess.MethylationDataTypes.extract_pheno_beta_df_from_sql(conn,
                                                                                         dis-
                                                                                         ease=")
     Return phenotype and beta dataframes from SQL tables storing MethylationArray info.
     conn SQL connection.
```

# MEFFIL\_FUNCTIONS.PY

```
Contains a few R functions that interact with meffil and minfi.
pymethylprocess.meffil_functions.est_cell_counts_IDOL (rgset, library)
     Given RGSet object, estimate cell counts for 450k/850k using reference approach via IDOL library.
     rgset RGSet object stored in python via rpy2
     library What type of CpG library to use.
pymethylprocess.meffil_functions.est_cell_counts_meffil(qc_list,
                                                                          cell_type_reference)
     Given QCObject list R object, estimate cell counts using reference approach via meffil.
     qc_list R list containing qc objects.
     cell_type_reference Reference blood/tissue set.
pymethylprocess.meffil_functions.est_cell_counts_minfi(rgset)
     Given RGSet object, estimate cell counts using reference approach via minfi.
     rgset RGSet object stored in python via rpy2
pymethylprocess.meffil_functions.load_detection_p_values_beadnum(qc\_list,
                                                                                      n cores)
     Return list of detection p-value matrix and bead number matrix.
     qc_list R list containing qc objects.
     n cores Number of cores to use in computation.
pymethylprocess.meffil_functions.r_autosomal_cpgs(array_type='450k')
     Return list of autosomal cpg probes per platform.
     array_type 450k/850k array?
pymethylprocess.meffil_functions.r_snp_cpgs (array_type='450k')
     Return list of SNP cpg probes per platform.
     array_type 450k/850k array?
pymethylprocess.meffil_functions.remove_sex(beta, array_type='450k')
     Remove non-autosomal cpgs from beta matrix.
     array_type 450k/850k array?
pymethylprocess.meffil_functions.set_missing(beta, pval_beadnum, detection_val=1e-06)
     Set missing beta values to NA, taking into account detection values and bead number the sholds.
     pval_beadnum Detection pvalues and number of beads per cpg/samples
     detection_val If threshold to set site to missingness based on p-value detection.
```

PyMethylProcess Documentation, Release 0.
---

# **GENERAL\_MACHINE\_LEARNING.PY**

Contains a machine learning class to perform scikit-learn like operations, along with held-out hyperparameter grid search.

Machine learning class to run sklearn-like pipeline on MethylationArray data. Initialize object with scikit-learn model, and optionally supply a hyperparameter search grid.

model Scikit-learn-like model, classification, regression, dimensionality reduction, clustering etc.

**options** Options to supply model in form of dictionary.

grid Alternatively, supply search grid to search for bets hyperparameters.

labelencode T/F encode string labels.

n\_eval Number of evaluations for randomized grid search, if set to 0, perform exhaustive grid search

```
assign_results_to_pheno_col (methyl_array, new_col, output_pkl)
```

Assign results to new phenotype column.

methyl\_array MethylationArray.

new col New column name.

output\_pkl Output pickle to dump MethylationArray to.

**fit** (train\_methyl\_array, val\_methyl\_array=None, outcome\_cols=None) Fit data to model.

**train\_methyl\_array** Training MethylationArray.

val\_methyl\_array Validation MethylationArray. Can set to None.

outcome\_cols Set to none if not needed, but phenotype column to train on, can be multiple.

 $\label{lem:constrain_methyl_array} \textbf{fit\_predict} \ (\textit{train\_methyl\_array}, \textit{outcome\_cols=None})$ 

Fit and predict training data.

**train\_methyl\_array** Training MethylationArray.

outcome\_cols Set to none if not needed, but phenotype column to train on, can be multiple.

fit\_transform(train\_methyl\_array, outcome\_cols=None)

Fit and transform to training data.

**train\_methyl\_array** Training MethylationArray.

**outcome\_cols** Set to none if not needed, but phenotype column to train on, can be multiple.

```
predict (test_methyl_array)
     Make new predictions on test methylation array.
     test_methyl_array Testing MethylationArray.
return_outcome_metric (methyl_array, outcome_cols, metric, run_bootstrap=False)
     Supply metric to evaluate results.
     methyl_array MethylationArray to evaluate.
     outcome_cols Outcome phenotype columns.
     metric Sklearn evaluation metric.
     run_bootstrap Make 95% CI from 1k bootstraps.
store_results(output_pkl, results_dict={})
     Store results in pickle file.
     output_pkl Output pickle to dump results to.
     results_dict Supply own results dict to be dumped.
transform(test_methyl_array)
     Transform test methylation array.
     test_methyl_array Testing MethylationArray.
transform_results_to_beta (methyl_array, output_pkl)
     Transform beta matrix into reduced beta matrix and store.
     methyl_array MethylationArray.
     output_pkl Output pickle to dump MethylationArray to.
```

**CHAPTER** 

**FIVE** 

## **PYMETHYL-INSTALL**

pymethyl-install [OPTIONS] COMMAND [ARGS]...

## **Options**

#### --version

Show the version and exit.

# 5.1 change\_gcc\_path

Change GCC and G++ paths if don't have version 7.2.0. [Experimental]

pymethyl-install change\_gcc\_path [OPTIONS]

# 5.2 install\_bioconductor

Installs bioconductor.

pymethyl-install install\_bioconductor [OPTIONS]

# 5.3 install\_custom

Installs bioconductor packages.

pymethyl-install install\_custom [OPTIONS]

## **Options**

-p, --package <package>

Custom packages. [default: ENmix]

-m, --manager

Use BiocManager (recommended).

# 5.4 install\_meffil

Installs meffil (update!).

pymethyl-install install\_meffil [OPTIONS]

# 5.5 install\_minfi\_others

Installs minfi and other dependencies.

pymethyl-install install\_minfi\_others [OPTIONS]

# 5.6 install\_r\_packages

Installs r packages.

pymethyl-install install\_r\_packages [OPTIONS]

## **Options**

-p, --package <package>
 Custom packages. [default: ]

# 5.7 install\_some\_deps

Installs bioconductor, minfi, enmix, tcga biolinks, and meffil.

pymethyl-install install\_some\_deps [OPTIONS]

# 5.8 install\_tcga\_biolinks

Installs tega biolinks.

pymethyl-install install\_tcga\_biolinks [OPTIONS]

**CHAPTER** 

SIX

## **PYMETHYL-VISUALIZE**

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

## **Options**

#### --version

Show the version and exit.

# 6.1 plot\_cell\_type\_results

Plot csv containing cell type results into side by side boxplots.

```
pymethyl-visualize plot_cell_type_results [OPTIONS]
```

## **Options**

```
-i, --input_csv <input_csv>
        Input csv. [default: cell_type_estimates.csv]
-o, --outfilename <outfilename>
        Output png. [default: visualizations/cell_type_results.png]
-cols, --plot_cols <plot_cols>
        Plot columns. [default: Gran, CD4T, CD8T, Bcell, Mono, NK, gMDSC]
-fs, --font_scale <font_scale>
        Font scaling [default: 1.0]
```

# 6.2 plot\_heatmap

Plot heatmap from CSV file.

```
pymethyl-visualize plot_heatmap [OPTIONS]
```

### **Options**

```
-i, --input_csv <input_csv>
     Input csv. [default: ]
-o, --outfilename <outfilename>
     Output png. [default: output.png]
-idx, --index_col <index_col>
     Index load dataframe [default: 0]
-fs, --font_scale <font_scale>
     Font scaling [default: 1.0]
-min, --min_val <min_val>
     Min heat val [default: 0.0]
-max, --max_val <max_val>
     Max heat val, if -1, defaults to None [default: 1.0]
-a, --annot
     Annotate heatmap [default: False]
-n, --norm
     Normalize matrix data [default: False]
-c, --cluster
     Cluster matrix data [default: False]
-m, --matrix_type <matrix_type>
     Type of matrix supplied [default: none]
-x, --xticks
     Show x ticks [default: False]
-y, --yticks
     Show y ticks [default: False]
-t, --transpose
     Transpose matrix data [default: False]
-col, --color_column <color_column>
     Color column. [default: color]
```

# 6.3 transform\_plot

Dimensionality reduce VAE or original beta values using UMAP and plot using plotly.

```
pymethyl-visualize transform_plot [OPTIONS]
```

## **Options**

- -o, --output\_file <output\_file>
   Output visualization. [default: ./visualization.html]
- -nn, --n\_neighbors <n\_neighbors>
   Number of neighbors UMAP. [default: 5]
- -a, --axes\_off
  Whether to turn axes on or off.
- -s, --supervised
  Supervise umap embedding.
- -d, --min\_dist <min\_dist>
   UMAP min distance. [default: 0.1]
- -m, --metric <metric>
   Reduction metric. [default: euclidean]
- -cc, --case\_control\_override

  Add controls from case\_control column and override current disease for classification tasks. [default: False]

6.3. transform\_plot 21

<b>PyMethylProces</b>	s Documentation	, Release 0.1
-----------------------	-----------------	---------------

**CHAPTER** 

SEVEN

## **PYMETHYL-PREPROCESS**

pymethyl-preprocess [OPTIONS] COMMAND [ARGS]...

## **Options**

#### --version

Show the version and exit.

# 7.1 batch\_deploy\_preprocess

Deploy multiple preprocessing jobs in series or parallel.

pymethyl-preprocess batch\_deploy\_preprocess [OPTIONS]

## **Options**

#### -n, --n\_cores <n\_cores>

Number cores to use for preprocessing. [default: 6]

-i, --subtype\_output\_dir <subtype\_output\_dir>
 Output subtypes pheno csv. [default: ./preprocess\_outputs/]

#### -m, --meffil

Preprocess using meffil.

#### -t, --torque

Job submission torque.

## -r, --run

Actually run local job or just print out command.

#### -s, --series

Run commands in series.

#### -p, --pc\_qc\_parameters\_csv <pc\_qc\_parameters\_csv>

For meffil, qc parameters and pcs for final qc and functional normalization. [default: ./preprocess\_outputs/pc\_qc\_parameters.csv]

## -u, --use\_cache

If this is selected, loads qc results rather than running qc again. Only works for meffil selection.

## -qc, --qc\_only

Only perform QC for meffil pipeline, caches results into rds file for loading again, only works if use\_cache is false.

-c, --chunk\_size <chunk\_size>

If not series, chunk up and run these number of commands at once.. -1 means all commands at once.

## 7.2 combine\_methylation\_arrays

If split MethylationArrays by subtype for either preprocessing or imputation, can use to recombine data for down-stream step.

```
pymethyl-preprocess combine_methylation_arrays [OPTIONS]
```

## **Options**

- -i, --input\_pkls <input\_pkls>
  Input pickles for beta and phenotype data. [default: ./preprocess\_outputs/methyl\_array.pkl]
- -d, --optional\_input\_pkl\_dir <optional\_input\_pkl\_dir>
   Auto grab input pkls. [default: ]
- -o, --output\_pkl <output\_pkl>
  Output database for beta and phenotype data. [default: ./combined\_outputs/methyl\_array.pkl]
- -e, --exclude <exclude>
   If -d selected, these diseases will be excluded from study. [default: ]

# 7.3 concat sample sheets

Concat two sample files for more fields for minfi+ input, adds more samples.

```
pymethyl-preprocess concat_sample_sheets [OPTIONS]
```

## **Options**

# 7.4 create\_sample\_sheet

Create sample sheet for input to minfi, meffil, or enmix.

pymethyl-preprocess create\_sample\_sheet [OPTIONS]

## **Options**

- -is, --input\_sample\_sheet <input\_sample\_sheet>
   Clinical information downloaded from tcga/geo/custom. [default: ./tcga\_idats/clinical\_info.csv]
- -s, --source\_type <source\_type>
  Source type of data. [default: tcga]

- -1, --header\_line <header\_line>
   Line to begin reading csv/xlsx. [default: 0]
- -d, --disease\_class\_column <disease\_class\_column>
   Disease classification column, for custom and geo datasets. [default: methylation class:ch1]
- -b, --basename\_col <basename\_col>
  Basename classification column, for custom datasets. [default: Sentrix ID (.idat)]
- -c, --include\_columns\_file <include\_columns\_file>
  Custom columns file containing columns to keep, separated by n. Add a tab for each line if you wish to rename columns: original\_name t new\_column\_name [default:]

# 7.5 download clinical

Download all TCGA 450k clinical info.

```
pymethyl-preprocess download_clinical [OPTIONS]
```

#### **Options**

-o, --output\_dir <output\_dir>
 Output directory for exported idats. [default: ./tcga\_idats/]

# 7.6 download\_geo

Download geo methylation study idats and clinical info.

pymethyl-preprocess download\_geo [OPTIONS]

### **Options**

```
    -g, --geo_query <geo_query>
        GEO study to query. [default: ]
    -o, --output_dir <output_dir>
        Output directory for exported idats. [default: ./geo_idats/]
```

# 7.7 download\_tcga

Download all tega 450k data.

```
pymethyl-preprocess download_tcga [OPTIONS]
```

### **Options**

```
-o, --output_dir <output_dir>
   Output directory for exported idats. [default: ./tcga_idats/]
```

# 7.8 feature\_select

Filter CpGs by taking x top CpGs with highest mean absolute deviation scores or via spectral feature selection.

```
pymethyl-preprocess feature_select [OPTIONS]
```

#### **Options**

```
    -i, --input_pkl <input_pkl>
        Input database for beta and phenotype data. [default: ./imputed_outputs/methyl_array.pkl]
    -o, --output_pkl <output_pkl>
        Output database for beta and phenotype data. [default: ./final_preprocessed/methyl_array.pkl]
    -n, --n_top_cpgs <n_top_cpgs>
        Number cpgs to include with highest variance across population. [default: 300000]
    -f, --feature_selection_method <feature_selection_method>
        -mm, --metric <metric>
```

```
-nn, --n_neighbors <n_neighbors>
```

Number neighbors for feature selection, default enacts rbf kernel. [default: 0]

```
-m, --mad_top_cpgs <mad_top_cpgs>
```

Number cpgs to apply mad filtering first before more sophisticated feature selection. If 0 or primary feature selection is mad, no mad pre-filtering. [default: 0]

# 7.9 get\_categorical\_distribution

Get categorical distribution of columns of sample sheet.

pymethyl-preprocess get\_categorical\_distribution [OPTIONS]

## **Options**

-is, --formatted\_sample\_sheet <formatted\_sample\_sheet>

Clinical information downloaded from tcga/geo/custom, formatted using create\_sample\_sheet. [default: ./tcga\_idats/minfiSheet.csv]

**-k, --key** <key>

Column of csv to print statistics for. [default: disease]

-d, --disease\_only

Only look at disease, or text before subtype\_delimiter.

-sd, --subtype\_delimiter <subtype\_delimiter>
 Delimiter for disease extraction. [default: ,]

# 7.10 imputation\_pipeline

Imputation of subtype or no subtype using various imputation methods.

pymethyl-preprocess imputation\_pipeline [OPTIONS]

## **Options**

- -i, --input\_pkl <input\_pkl>
  - Input database for beta and phenotype data. [default: ./combined\_outputs/methyl\_array.pkl]
- -ss, --split\_by\_subtype

Imputes CpGs by subtype before combining again.

-m, --method <method>

Method of imputation. [default: KNN]

-s, --solver <solver>

Imputation library. [default: fancyimpute]

-k, --n\_neighbors <n\_neighbors>

Number neighbors for imputation if using KNN. [default: 5]

-r, --orientation <orientation>

Impute CpGs or samples. [default: Samples]

-o, --output\_pkl <output\_pkl>

Output database for beta and phenotype data. [default: ./imputed\_outputs/methyl\_array.pkl]

-n, --n\_top\_cpgs <n\_top\_cpgs>

Number cpgs to include with highest variance across population. Greater than 0 allows for mad filtering during imputation to skip mad step. [default: 0]

- -f, --feature selection method <feature selection method>
- -mm, --metric <metric>
- -nfs, --n\_neighbors\_fs <n\_neighbors\_fs>

Number neighbors for feature selection, default enacts rbf kernel. [default: 0]

#### -d, --disease\_only

Only look at disease, or text before subtype\_delimiter.

-sd, --subtype\_delimiter <subtype\_delimiter>

Delimiter for disease extraction. [default: ,]

-st, --sample\_threshold <sample\_threshold>

Value between 0 and 1 for NaN removal. If samples has sample\_threshold proportion of cpgs missing, then remove sample. Set to -1 to not remove samples. [default: -1.0]

-ct, --cpg\_threshold <cpg\_threshold>

Value between 0 and 1 for NaN removal. If cpgs has cpg\_threshold proportion of samples missing, then remove cpg. Set to -1 to not remove samples. [default: -1.0]

## 7.11 meffil encode

Reformat file for meffil input.

pymethyl-preprocess meffil\_encode [OPTIONS]

## **Options**

# 7.12 merge sample sheets

Merge two sample files for more fields for minfi+ input.

pymethyl-preprocess merge\_sample\_sheets [OPTIONS]

## **Options**

-s1, --sample\_sheet1 <sample\_sheet1>

Clinical information downloaded from tcga/geo/custom, formatted using create\_sample\_sheet. [default: ./tcga\_idats/clinical\_info1.csv]

-s2, --sample\_sheet2 <sample\_sheet2>

Clinical information downloaded from tcga/geo/custom, formatted using create\_sample\_sheet. [default: ./tcga\_idats/clinical\_info2.csv]

-d, --second\_sheet\_disease

Use second sheet's disease column.

-nd, --no\_disease\_merge

Don't merge disease columns.

# 7.13 na\_report

Print proportion of missing values throughout dataset.

```
pymethyl-preprocess na_report [OPTIONS]
```

## **Options**

- -o, --output\_dir <output\_dir>
   Output database for na report. [default: ./na\_report/]
- -r, --head\_directory

-i option becomes directory, and searches there for multiple input pickles.

## 7.14 preprocess pipeline

Perform preprocessing of idats using enmix or meffil.

```
pymethyl-preprocess preprocess_pipeline [OPTIONS]
```

## **Options**

-i, --idat\_dir <idat\_dir>

Idat dir for one sample sheet, alternatively can be your phenotype sample sheet. [default: ./tcga\_idats/]

-n, --n\_cores <n\_cores>

Number cores to use for preprocessing. [default: 6]

-o, --output\_pkl <output\_pkl>

Output database for beta and phenotype data. [default: ./preprocess\_outputs/methyl\_array.pkl]

-m, --meffil

Preprocess using meffil.

-pc, --n pcs <n pcs>

For meffil, number of principal components for functional normalization. If set to -1, then PCs are selected using elbow method. [default: -1]

-p, --pipeline <pipeline>

If not meffil, preprocess using minfi or enmix. [default: enmix]

-noob, --noob norm

Run noob normalization of minfi selected.

-u, --use\_cache

If this is selected, loads qc results rather than running qc again and update with new qc parameters. Only works for meffil selection. Minfi and enmix just loads RG Set.

-qc, --qc\_only

Only perform QC for meffil pipeline, caches results into rds file for loading again, only works if use\_cache is false. Minfi and enmix just saves the RGSet before preprocessing.

7.13. na\_report 29

-bns, --p\_beadnum\_samples <p\_beadnum\_samples>

From meffil documentation, "fraction of probes that failed the threshold of 3 beads". [default: 0.05]

-pds, --p\_detection\_samples <p\_detection\_samples>

From meffil documentation, "fraction of probes that failed a detection.pvalue threshold of 0.01". [default: 0.05]

-bnc, --p\_beadnum\_cpgs <p\_beadnum\_cpgs>

From meffil documentation, "fraction of samples that failed the threshold of 3 beads". [default: 0.05]

-pdc, --p\_detection\_cpgs <p\_detection\_cpgs>

From meffil documentation, "fraction of samples that failed a detection.pvalue threshold of 0.01". [default: 0.05]

-sc, --sex\_cutoff <sex\_cutoff>

From meffil documentation, "difference of total median intensity for Y chromosome probes and X chromosome probes". [default: -2]

-sd, --sex\_sd <sex\_sd>

From meffil documentation, "sex detection outliers if outside this range". [default: 5]

## 7.15 remove diseases

Exclude diseases from study by count number or exclusion list.

pymethyl-preprocess remove\_diseases [OPTIONS]

#### **Options**

-is, --formatted\_sample\_sheet <formatted\_sample\_sheet>

Clinical information downloaded from tcga/geo/custom, formatted using create\_sample\_sheet. [default: ./tcga\_idats/clinical\_info.csv]

-e, --exclude\_disease\_list <exclude\_disease\_list>

List of conditions to exclude, from disease column, comma delimited. [default: ]

-os, --output\_sheet\_name <output\_sheet\_name>

CSV for minfi input. [default: ./tcga\_idats/minfiSheet.csv]

-1, --low\_count <low\_count>

Remove diseases if they are below a certain count, default this is not used. [default: 0]

-d, --disease only

Only look at disease, or text before subtype\_delimiter.

-sd, --subtype\_delimiter <subtype\_delimiter>

Delimiter for disease extraction. [default: ,]

# 7.16 split\_preprocess\_input\_by\_subtype

Split preprocess input samplesheet by disease subtype.

pymethyl-preprocess split\_preprocess\_input\_by\_subtype [OPTIONS]

## **Options**

- -i, --idat\_csv <idat\_csv>
  - Idat csv for one sample sheet, alternatively can be your phenotype sample sheet. [default: ./tcga\_idats/minfiSheet.csv]
- -d, --disease\_only

Only look at disease, or text before subtype\_delimiter.

- -sd, --subtype\_delimiter <subtype\_delimiter>
   Delimiter for disease extraction. [default: ,]
- -o, --subtype\_output\_dir <subtype\_output\_dir>
   Output subtypes pheno csv. [default: ./preprocess\_outputs/]

**CHAPTER** 

### **EIGHT**

### **PYMETHYL-UTILS**

```
pymethyl-utils [OPTIONS] COMMAND [ARGS]...
```

#### **Options**

#### --version

Show the version and exit.

# 8.1 backup\_pkl

Copy methylarray pickle to new location to backup.

```
pymethyl-utils backup_pkl [OPTIONS]
```

#### **Options**

```
-o, --output_pkl <output_pkl>
   Output database for beta and phenotype data. [default: ./backup/methyl_array.pkl]
```

## 8.2 bin\_column

Convert continuous phenotype column into categorical by binning.

```
pymethyl-utils bin_column [OPTIONS]
```

#### **Options**

```
-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_pk1 <output_test_pkl>
    Binned shap pickle for further testing. [default: ./train_val_test_sets/test_methyl_array_shap_binned.pkl]
```

### 8.3 concat csv

Concatenate two csv files together.

```
pymethyl-utils concat_csv [OPTIONS]
```

### **Options**

```
-i1, --input_csv <input_csv>
    Beta csv. [default: ./beta1.csv]

-i2, --input_csv2 <input_csv2>
    Beta/other csv 2. [default: ./cell_estimates.csv]

-o, --output_csv <output_csv>
    Output csv. [default: ./beta.concat.csv]

-a, --axis <axis>
    Axis to merge on. Columns are 0, rows are 1. [default: 1]

-i, --index_col <index_col>
    Index Column. [default: 0]
```

#### 8.4 counts

Return categorical breakdown of phenotype column.

```
pymethyl-utils counts [OPTIONS]
```

#### **Options**

```
    -i, --input_pkl <input_pkl>
        Input database for beta and phenotype data. [default: ./final_preprocessed/methyl_array.pkl]
    -k, --key <key>
        Key to split on. [default: disease]
```

## 8.5 create\_external\_validation\_set

Create external validation set containing same CpGs as training set.

```
pymethyl-utils create_external_validation_set [OPTIONS]
```

#### **Options**

```
-t, --train_pkl <train_pkl>
        Input methyl array. [default: ./train_val_test_sets/train_methyl_array.pkl]
-q, --query_pkl <query_pkl>
        Input methylation array to add/subtract cpgs to. [default: ./final_preprocessed/methyl_array.pkl]
-o, --output_pkl <output_pkl>
        Output methyl array external validation. [default: ./external_validation/methyl_array.pkl]
-c, --cpg_replace_method <cpg_replace_method>
        What to do for missing CpGs. [default: mid]
```

### 8.6 est\_age

Estimate age using cgAgeR

```
pymethyl-utils est_age [OPTIONS]
```

#### **Options**

```
    -i, --input_pkl <input_pkl>
        Input methyl array. [default: ./train_val_test_sets/test_methyl_array.pkl]
    -ac, --age_column <age_column>
            Age column of Methylation Array. Leave blank for no age [default: ]
    -a, --analyses <analyses>
            Analyses to run
    -o, --output_csv <output_csv>
            Output csv [default: age_estimation/output_age_estimations.csv]
```

## 8.7 feature\_select\_train\_val\_test

Filter CpGs by taking x top CpGs with highest mean absolute deviation scores or via spectral feature selection.

```
pymethyl-utils feature_select_train_val_test [OPTIONS]
```

#### **Options**

```
    -i, --input_pkl_dir <input_pkl_dir>
        Input database for beta and phenotype data. [default: ./train_val_test_sets/]
    -o, --output_dir <output_dir>
        Output database for beta and phenotype data. [default: ./train_val_test_sets_fs/]
    -n, --n_top_cpgs <n_top_cpgs>
        Number cpgs to include with highest variance across population. [default: 300000]
    -f, --feature_selection_method <feature_selection_method>
    -mm, --metric <metric>
```

8.6. est age 35

-nn, --n\_neighbors <n\_neighbors>

Number neighbors for feature selection, default enacts rbf kernel. [default: 0]

-m, --mad\_top\_cpgs <mad\_top\_cpgs>

Number cpgs to apply mad filtering first before more sophisticated feature selection. If 0 or primary feature selection is mad, no mad pre-filtering. [default: 0]

### 8.8 fix\_key

Format certain column of phenotype array in MethylationArray.

```
pymethyl-utils fix_key [OPTIONS]
```

#### **Options**

**-k, --key** <key>

Key to split on. [default: disease]

-d, --disease\_only

Only look at disease, or text before subtype\_delimiter.

-sd, --subtype\_delimiter <subtype\_delimiter>

Delimiter for disease extraction. [default: ,]

-o, --output\_pkl <output\_pkl>

Input database for beta and phenotype data. [default: ./fixed\_preprocessed/methyl\_array.pkl]

## 8.9 modify\_pheno\_data

Use another spreadsheet to add more descriptive data to methylarray.

```
pymethyl-utils modify_pheno_data [OPTIONS]
```

#### **Options**

-is, --input\_formatted\_sample\_sheet <input\_formatted\_sample\_sheet>
 Information passed through function create\_sample\_sheet, has Basename and disease fields. [default:
 ./tcga\_idats/minfi\_sheet.csv]

-o, --output\_pkl <output\_pkl>

Output database for beta and phenotype data. [default: ./modified\_processed/methyl\_array.pkl]

### 8.10 move\_jpg

Move preprocessing jpegs to preprocessing output directory.

```
pymethyl-utils move_jpg [OPTIONS]
```

#### **Options**

```
    -i, --input_dir <input_dir>
        Directory containing jpg. [default: ./]
    -o, --output_dir <output_dir>
        Output directory for images. [default: ./preprocess_output_images/]
```

### 8.11 overwrite pheno data

Use another spreadsheet to add more descriptive data to methylarray.

```
pymethyl-utils overwrite_pheno_data [OPTIONS]
```

#### **Options**

```
-is, --input_formatted_sample_sheet <input_formatted_sample_sheet>
    Information passed through function create_sample_sheet, has Basename and disease fields. [default:
    ./tcga_idats/minfi_sheet.csv]
```

```
-o, --output_pkl <output_pkl>
   Output database for beta and phenotype data. [default: ./modified_processed/methyl_array.pkl]
```

## 8.12 pkl\_to\_csv

Output methylarray pickle to csv.

```
pymethyl-utils pkl_to_csv [OPTIONS]
```

#### **Options**

```
    -i, --input_pkl <input_pkl>
        Input database for beta and phenotype data. [default: ./final_preprocessed/methyl_array.pkl]
    -o, --output_dir <output_dir>
        Input database for beta and phenotype data. [default: ./final_preprocessed/]
```

```
-c, --col <col>
    Column to color. [default: ]
```

8.10. move\_jpg 37

### 8.13 print\_number\_sex\_cpgs

Print number of non-autosomal CpGs.

```
pymethyl-utils print_number_sex_cpgs [OPTIONS]
```

#### **Options**

```
-a, --array_type <array_type>
Array Type. [default: 450k]
```

## 8.14 print\_shape

Print dimensions of beta matrix.

```
pymethyl-utils print_shape [OPTIONS]
```

#### **Options**

## 8.15 rate\_regression

```
pymethyl-utils rate_regression [OPTIONS]
```

#### **Options**

```
-i, --input_csv <input_csv>
Results csv. [default: ./results.csv]
```

```
-c1, --pred_col pred_col>
    Prediction column. [default: y_pred]
```

```
-c2, --true_col <true_col>
    True Column. [default: y_true]
```

# 8.16 ref\_estimate\_cell\_counts

Reference based cell type estimates.

```
pymethyl-utils ref_estimate_cell_counts [OPTIONS]
```

#### **Options**

## 8.17 ref\_free\_cell\_deconv

Reference free cell type deconvolution

```
pymethyl-utils ref_free_cell_deconv [OPTIONS]
```

#### **Options**

### 8.18 remove\_sex

Remove non-autosomal CpGs.

```
pymethyl-utils remove_sex [OPTIONS]
```

#### **Options**

```
    -i, --input_pkl <input_pkl>
        Input database for beta and phenotype data. [default: ./preprocess_outputs/methyl_array.pkl]
    -o, --output_pkl <output_pkl>
        Output methyl array autosomal. [default: ./autosomal/methyl_array.pkl]
```

```
-a, --array_type <array_type>
Array Type. [default: 450k]
```

### 8.19 remove\_snps

Remove SNPs from methylation array.

```
pymethyl-utils remove_snps [OPTIONS]
```

#### **Options**

```
-o, --output_pkl <output_pkl>
   Output methyl array autosomal. [default: ./no_snp/methyl_array.pkl]
```

```
-a, --array_type <array_type>
Array Type. [default: 450k]
```

### 8.20 set part array background

Set subset of CpGs from beta matrix to background values.

```
pymethyl-utils set_part_array_background [OPTIONS]
```

#### **Options**

```
-c, --cpg_pkl <cpg_pkl>
Pickled numpy array for subsetting. [default: ./subset_cpgs.pkl]
```

```
-o, --output_pkl <output_pkl>
   Output methyl array external validation. [default: ./removal/methyl_array.pkl]
```

## 8.21 stratify

Split methylation array by key and store.

```
pymethyl-utils stratify [OPTIONS]
```

#### **Options**

```
-k, --key <key>
Key to split on. [default: disease]
```

-o, --output\_dir <output\_dir>
 Output directory for stratified. [default: ./stratified/]

### 8.22 subset array

Only retain certain number of CpGs from methylation array.

```
pymethyl-utils subset_array [OPTIONS]
```

### **Options**

```
-c, --cpg_pkl <cpg_pkl>
Pickled numpy array for subsetting. [default: ./subset_cpgs.pkl]
```

-o, --output\_pkl <output\_pkl>
 Output methyl array external validation. [default: ./subset/methyl\_array.pkl]

### 8.23 train\_test\_val\_split

Split methylation array into train, test, val.

```
pymethyl-utils train_test_val_split [OPTIONS]
```

#### **Options**

```
-i, --input_pkl <input_pkl>
        Input database for beta and phenotype data. [default: ./final_preprocessed/methyl_array.pkl]
-o, --output_dir <output_dir>
        Output directory for training, testing, and validation sets. [default: ./train_val_test_sets/]
-tp, --train_percent <train_percent>
        Percent data training on. [default: 0.8]
-vp, --val_percent <val_percent>
        Percent of training data that comprises validation set. [default: 0.1]
-cat, --categorical
        Multi-class prediction. [default: False]
-do, --disease_only
```

Only look at disease, or text before subtype\_delimiter.

-k, --key <key>

Key to split on. [default: disease]

-sd, --subtype\_delimiter <subtype\_delimiter>
 Delimiter for disease extraction. [default: ,]

8.22. subset\_array 41

# 8.24 write\_cpgs

Write CpGs in methylation array to file.

```
pymethyl-utils write_cpgs [OPTIONS]
```

### **Options**

42

- -c, --cpg\_pkl <cpg\_pkl>
   Pickled numpy array for subsetting. [default: ./subset\_cpgs.pkl]

## CHAPTER

# **NINE**

# **INDICES AND TABLES**

- genindex
- modindex
- search

PyMethylProcess Documentation, Release 0.	PyMethylProcess	Documentation,	Release 0.1
---	-----------------	----------------	-------------

## **PYTHON MODULE INDEX**

### р

46 Python Module Index

## **INDEX**

Symbols		pymethyl-preprocess-preprocess_pipeline
-version		command line option, 30
<pre>pymethyl-install command line   option, 17</pre>	-bn	s, -p_beadnum_samples <p_beadnum_samples></p_beadnum_samples>
pymethyl-preprocess command line option, 23		<pre>pymethyl-preprocess-preprocess_pipeline command line option, 29</pre>
pymethyl-utils command line option, 33	-c,	<pre>-cell_type_columns</pre>
<pre>pymethyl-visualize command line   option, 19</pre>		<pre>pymethyl-utils-ref_free_cell_deconv command line option, 39</pre>
-a, -algorithm <algorithm></algorithm>	-c,	-chunk_size <chunk_size></chunk_size>
pymethyl-utils-ref_estimate_cell_courcemmand line option, 39		<pre>pymethyl-preprocess-batch_deploy_preprocess     command line option, 24</pre>
-a, -analyses <analyses></analyses>	-c,	-cluster
<pre>pymethyl-utils-est_age command     line option, 35</pre>		pymethyl-visualize-plot_heatmap command line option, 20
-a, -analysis <analysis></analysis>	-c,	-col <col/>
<pre>pymethyl-utils-ref_free_cell_deconv     command line option, 39</pre>		<pre>pymethyl-utils-bin_column command   line option, 33</pre>
-a, -annot		pymethyl-utils-pkl_to_csv command
<pre>pymethyl-visualize-plot_heatmap</pre>		line option, 37
command line option, 20	-c,	-column_of_interest
-a, -array_type <array_type></array_type>		<column_of_interest></column_of_interest>
<pre>pymethyl-utils-print_number_sex_cpgs</pre>		<pre>pymethyl-visualize-transform_plot</pre>
command line option, 38		command line option, $20$
pymethyl-utils-remove_sex command	-c,	-cpg_pkl <cpg_pkl></cpg_pkl>
line option, 39		<pre>pymethyl-utils-set_part_array_background</pre>
pymethyl-utils-remove_snps command		command line option, $40$
line option, 40		<pre>pymethyl-utils-subset_array</pre>
-a, -axes_off		command line option, 41
pymethyl-visualize-transform_plot		<pre>pymethyl-utils-write_cpgs command</pre>
command line option, 21		line option, 42
-a, -axis <axis></axis>	-c,	-cpg_replace_method
pymethyl-utils-concat_csv command		<cpg_replace_method></cpg_replace_method>
line option, 34		<pre>pymethyl-utils-create_external_validation_set</pre>
-ac, -age_column <age_column></age_column>		command line option, 35
pymethyl-utils-est_age command	-c,	-include_columns_file
line option, 35		<pre><include_columns_file></include_columns_file></pre>
-b, -basename_col <basename_col></basename_col>		pymethyl-preprocess-create_sample_sheet
pymethyl-preprocess-create_sample_sh	eet	command line option, 25
command line option, 25	-c,	-index_col <index_col></index_col>
-bnc, -p_beadnum_cpgs <p_beadnum_cpgs></p_beadnum_cpgs>		<pre>pymethyl-utils-overwrite_pheno_data</pre>

command line option, 37 <pre></pre>	
pymethyl-utils-rate_regression command line option, 30	es
command line option, 38 -f, -feature_selection_method	
-c2, -true_col <true_col> <feature_selection_method></feature_selection_method></true_col>	
<pre>pymethyl-utils-rate_regression</pre>	5
command line option, 38 command line option, 26	
-cat, -categorical pymethyl-preprocess-imputation_pip	peline
<pre>pymethyl-utils-train_test_val_split command line option, 27</pre>	
command line option, 41 pymethyl-utils-feature_select_tra:	in_val_tes
-cc, -case_control_override command line option, 35	
<pre>pymethyl-visualize-transform_plot -fs, -font_scale <font_scale></font_scale></pre>	
command line option, 21 pymethyl-visualize-plot_cell_type_	_results
-col, -color_column <color_column> command line option, 19</color_column>	
<pre>pymethyl-visualize-plot_heatmap</pre> <pre>pymethyl-visualize-plot_heatmap</pre>	
command line option, 20 command line option, 20	
-cols, -plot_cols <plot_cols> -g, -geo_query <geo_query></geo_query></plot_cols>	
pymethyl-visualize-plot_cell_type_resultpymethyl-preprocess-download_geo	
command line option, 19 command line option, 26	
-ct, -cpg_threshold <cpg_threshold> -i, -idat_csv <idat_csv></idat_csv></cpg_threshold>	
pymethyl-preprocess-imputation_pipeline pymethyl-preprocess-split_preproce	ess_input_
command line option, 28 command line option, 31	_
-d, -disease_class_column -i, -idat_dir <idat_dir></idat_dir>	
<pre><disease_class_column></disease_class_column></pre>	_sheet
pymethyl-preprocess-create_sample_sheet command line option, 25	
command line option, 25 pymethyl-preprocess_preprocess_pip	peline
-d, -disease_only command line option, 29	
pymethyl-preprocess-get_categorical_distributexocol <index_col></index_col>	
command line option, 27 pymethyl-utils-concat_csv command	
pymethyl-preprocess-imputation_pipeline line option, 34	
<pre>pymethyl-preprocess-imputation_pipeline line option, 34</pre>	
pymethyl-preprocess-imputation_pipeline line option, 34	
<pre>pymethyl-preprocess-imputation_pipeline</pre>	_results
<pre>pymethyl-preprocess-imputation_pipeline line option, 34     command line option, 27</pre>	_results
pymethyl-preprocess-imputation_pipeline line option, 34 command line option, 27 -i, -input_csv <input_csv> pymethyl-preprocess-remove_diseases pymethyl-utils-rate_regression command line option, 30 command line option, 38 pymethyl-preprocess-split_preprocess_inpptmethylstbtypealize-plot_cell_type_</input_csv>	_results
pymethyl-preprocess-imputation_pipeline line option, 34 command line option, 27 -i, -input_csv <input_csv> pymethyl-preprocess-remove_diseases pymethyl-utils-rate_regression command line option, 30 command line option, 38 pymethyl-preprocess-split_preprocess_inppymbyhylbtypealize-plot_cell_type_ command line option, 31 command line option, 19</input_csv>	_results
pymethyl-preprocess-imputation_pipeline line option, 34 command line option, 27 -i, -input_csv <input_csv> pymethyl-preprocess-remove_diseases pymethyl-utils-rate_regression command line option, 30 command line option, 38 pymethyl-preprocess-split_preprocess_inpptmbthttpealize-plot_cell_type_ command line option, 31 command line option, 19 pymethyl-utils-fix_key command pymethyl-visualize-plot_heatmap</input_csv>	_results
pymethyl-preprocess-imputation_pipeline line option, 34 command line option, 27 -i, -input_csv <input_csv> pymethyl-preprocess-remove_diseases command line option, 30 command line option, 38 pymethyl-preprocess-split_preprocess_inpptmbthttpealize-plot_cell_type_command line option, 31 command line option, 19 pymethyl-utils-fix_key command line option, 36 pymethyl-visualize-plot_heatmap command line option, 20</input_csv>	_results
pymethyl-preprocess-imputation_pipeline line option, 34     command line option, 27	_results
pymethyl-preprocess-imputation_pipeline	_results
pymethyl-preprocess-imputation_pipeline command line option, 27 -i, -input_csv <input_csv> pymethyl-preprocess-remove_diseases command line option, 30 command line option, 38 pymethyl-preprocess-split_preprocess_inpptmbthttpealize-plot_cell_type_command line option, 31 command line option, 19 pymethyl-utils-fix_key command line option, 36 command line option, 20 -d, -min_dist <min_dist> -i, -input_dir <input_dir> pymethyl-visualize-transform_plot command line option, 21 line option, 37</input_dir></min_dist></input_csv>	
pymethyl-preprocess-imputation_pipeline	
pymethyl-preprocess-imputation_pipeline	<u>-</u>
pymethyl-preprocess-imputation_pipeline	<u>-</u>
pymethyl-preprocess-imputation_pipeline command line option, 27 -i, -input_csv <input_csv> pymethyl-preprocess-remove_diseases command line option, 30 command line option, 38  pymethyl-preprocess-split_preprocess_inppymethylbtylptalize-plot_cell_type_command line option, 31 command line option, 19  pymethyl-utils-fix_key command line option, 36 command line option, 36 command line option, 20  -d, -min_dist <min_dist> pymethyl-visualize-transform_plot command line option, 21 line option, 37  -d, -optional_input_pkl_dir -i, -input_pkl <input_pkl> pymethyl-preprocess-feature_select pymethyl-preprocess-combine_methylation_arracyammand line option, 26 command line option, 24 pymethyl-preprocess-imputation_pipeline line option, 34 cinput_csv <input_csv -i,="" -input_dir="" -input_pkl="" 19="" 20="" 37="" 38="" <input_csv="" <input_pkl="" <input_selection="" command="" line="" option,="" pymethyl-visualize-plot_heatmap=""> pymethyl-preprocess-feature_selection option, 26 pymethyl-preprocess-imputation_pipeline command line option, 24</input_csv></input_pkl></min_dist></input_csv>	<u>-</u>
pymethyl-preprocess-imputation_pipeline	<u>-</u>
pymethyl-preprocess-imputation_pipeline command line option, 27 -i, -input_csv <input_csv> pymethyl-preprocess-remove_diseases command line option, 30 command line option, 38  pymethyl-preprocess-split_preprocess_inppymbyhybbbypealize-plot_cell_type_command line option, 31 command line option, 19  pymethyl-utils-fix_key command line option, 36 command line option, 36 command line option, 20  -d, -min_dist <min_dist> -i, -input_dir <input_dir> pymethyl-visualize-transform_plot command line option, 21 line option, 37  -d, -optional_input_pkl_dir -i, -input_pkl <input_pkl> command line option, 24 pymethyl-preprocess-feature_select pymethyl-preprocess-combine_methylation_arraxymmand line option, 27 pymethyl-preprocess-merge_sample_sheets pymethyl-preprocess-na_report</input_pkl></input_dir></min_dist></input_csv>	<u>-</u>
pymethyl-preprocess-imputation_pipeline	<u>-</u>
pymethyl-preprocess-imputation_pipeline command line option, 27 -i, -input_csv <input_csv> pymethyl-preprocess-remove_diseases command line option, 30 command line option, 38 pymethyl-preprocess-split_preprocess_inpptmbthshbttpealize-plot_cell_type_command line option, 31 command line option, 19 pymethyl-utils-fix_key command line option, 36 command line option, 36 command line option, 20 -d, -min_dist <min_dist> -i, -input_dir <input_dir> pymethyl-visualize-transform_plot command line option, 21 line option, 37 -d, -optional_input_pkl_dir -i, -input_pkl <input_pkl> command line option, 26 command line option, 24 pymethyl-preprocess-feature_select pymethyl-preprocess-merge_sample_sheets command line option, 27 pymethyl-preprocess-merge_sample_sheets pymethyl-utils-backup_pkl command line option, 29 -do, -disease_only</input_pkl></input_dir></min_dist></input_csv>	: peline
pymethyl-preprocess-imputation_pipeline	: peline
pymethyl-preprocess-imputation_pipeline	: peline
pymethyl-preprocess-imputation_pipeline command line option, 27 -i, -input_csv <input_csv> pymethyl-preprocess-remove_diseases command line option, 30 command line option, 38  pymethyl-preprocess-split_preprocess_inppymbyhybybypealize-plot_cell_type_command line option, 31 command line option, 19  pymethyl-utils-fix_key command pymethyl-visualize-plot_heatmap command line option, 36 command line option, 20  -d, -min_dist <min_dist> -i, -input_dir <input_dir> pymethyl-visualize-transform_plot command line option, 21 line option, 37  -d, -optional_input_pkl_dir -i, -input_pkl <input_pkl> pymethyl-preprocess-feature_select pymethyl-preprocess-combine_methylation_arraxymmand line option, 26 command line option, 24 pymethyl-preprocess-marge_sample_sheets command line option, 28  -do, -disease_only pymethyl-utils-backup_pkl command line option, 31 pymethyl-utils-backup_pkl command line option, 33 pymethyl-utils-counts command line option, 34</input_pkl></input_dir></min_dist></input_csv>	: peline

line option, 36		pymethyl-utils-modify_pheno_data
pymethyl-utils-modify_pheno_data		command line option, 36
command line option, 36		pymethyl-utils-overwrite_pheno_data
<pre>pymethyl-utils-overwrite_pheno_data</pre>		command line option, 37
command line option, 37	-is	, -input_sample_sheet
pymethyl-utils-pkl_to_csv command		<pre><input_sample_sheet></input_sample_sheet></pre>
line option, 37		pymethyl-preprocess-create_sample_sheet
pymethyl-utils-print_number_sex_cpgs		command line option, 25
command line option, 38		pymethyl-preprocess-meffil_encode
pymethyl-utils-print_shape command		command line option, 28
line option, 38	-k.	-key <key></key>
pymethyl-utils-remove_sex command	1.,	pymethyl-preprocess-get_categorical_distribution
line option, 39		command line option, 27
pymethyl-utils-remove_snps command		pymethyl-utils-counts command line
line option, 40		option, 34
pymethyl-utils-set_part_array_backgr	oun	
command line option, 40	Ounc	line option, 36
pymethyl-utils-stratify command		pymethyl-utils-stratify command
line option, 40		line option, 40
pymethyl-utils-subset_array		pymethyl-utils-train_test_val_split
command line option, 41	,	command line option, 41
pymethyl-utils-train_test_val_split	-ĸ,	-n_cell_types <n_cell_types></n_cell_types>
command line option, 41		pymethyl-utils-ref_free_cell_deconv
pymethyl-utils-write_cpgs command		command line option, 39
line option, 42	-k,	-n_neighbors <n_neighbors></n_neighbors>
pymethyl-visualize-transform_plot		pymethyl-preprocess-imputation_pipeline
command line option, 20		command line option, 27
<pre>-i, -input_pkl_dir <input_pkl_dir></input_pkl_dir></pre>		-header_line <header_line></header_line>
	_val_	_pgmethyl-preprocess-create_sample_sheet
command line option, 35		command line option, 25
<pre>-i, -input_pkls <input_pkls></input_pkls></pre>		-library <library></library>
	ion_	_pymeybyl-utils-ref_estimate_cell_counts
command line option, $24$		command line option, 39
-i, -subtype_output_dir	-1,	-low_count <low_count></low_count>
<subtype_output_dir></subtype_output_dir>		pymethyl-preprocess-remove_diseases
<pre>pymethyl-preprocess-batch_deploy_pre</pre>		
command line option, 23	-m,	-mad_top_cpgs <mad_top_cpgs></mad_top_cpgs>
-i1, -input_csv <input_csv></input_csv>		<pre>pymethyl-preprocess-feature_select</pre>
<pre>pymethyl-utils-concat_csv command</pre>		command line option, $26$
line option, 34		<pre>pymethyl-utils-feature_select_train_val_test</pre>
-i2, -input_csv2 <input_csv2></input_csv2>		command line option, 36
<pre>pymethyl-utils-concat_csv command</pre>	-m,	-manager
line option, 34		<pre>pymethyl-install_custom</pre>
<pre>-idx, -index_col <index_col></index_col></pre>		command line option, 17
<pre>pymethyl-visualize-plot_heatmap</pre>	-m,	<pre>-mapping_file <mapping_file></mapping_file></pre>
command line option, $20$		pymethyl-preprocess-create_sample_sheet
-is, -formatted_sample_sheet		command line option, 25
<formatted_sample_sheet></formatted_sample_sheet>	-m,	-matrix_type <matrix_type></matrix_type>
<pre>pymethyl-preprocess-get_categorical_</pre>	dist	pyhethyh-visualize-plot_heatmap
command line option, 27		command line option, 20
<pre>pymethyl-preprocess-remove_diseases</pre>	-m,	-meffil
command line option, 30	,	pymethyl-preprocess-batch_deploy_preprocess
-is, -input_formatted_sample_sheet		command line option, 23
<pre> <input formatted="" sample="" sheet=""/></pre>		pymethyl-preprocess-preprocess pipeline

command line option, $29$ -o, -m, -method <method></method>	<pre>-outfilename <outfilename> pymethyl-visualize-plot_cell_type_results</outfilename></pre>
pymethyl-preprocess-imputation_pipeline	command line option, 19
command line option, 27	pymethyl-visualize-plot_heatmap
-m, -metric <metric></metric>	command line option, 20
	-output_csv <output_csv></output_csv>
command line option, 21	pymethyl-utils-concat_csv command
-max, -max_val <max_val></max_val>	line option, 34
pymethyl-visualize-plot_heatmap	pymethyl-utils-est_age command
command line option, 20	line option, 35
-min, -min_val <min_val></min_val>	pymethyl-utils-ref_estimate_cell_counts
pymethyl-visualize-plot_heatmap	command line option, 39
	-output_dir <output_dir></output_dir>
-mm, -metric <metric></metric>	pymethyl-preprocess-download_clinical
pymethyl-preprocess-feature_select	command line option, 25
command line option, 26	pymethyl-preprocess-download_geo
pymethyl-preprocess-imputation_pipeline	command line option, 26
command line option, 27	pymethyl-preprocess-download_tcga
pymethyl-utils-feature_select_train_val_	
command line option, 35	pymethyl-preprocess-na_report
-n, -n_bins <n_bins></n_bins>	command line option, 29
pymethyl-utils-bin_column command	<pre>pymethyl-utils-feature_select_train_val_test</pre>
line option, 33	command line option, 35
-n, -n_cores <n_cores></n_cores>	pymethyl-utils-move_jpg command
<pre>pymethyl-preprocess-batch_deploy_preproc</pre>	
command line option, 23	<pre>pymethyl-utils-pkl_to_csv command</pre>
<pre>pymethyl-preprocess-preprocess_pipeline</pre>	line option, 37
command line option, 29	pymethyl-utils-stratify command
-n, -n_top_cpgs <n_top_cpgs></n_top_cpgs>	line option,41
<pre>pymethyl-preprocess-feature_select</pre>	<pre>pymethyl-utils-train_test_val_split</pre>
command line option, $26$	command line option, 41
pymethyl-preprocess-imputation_pipeline,	
command line option, 27	<pre>pymethyl-visualize-transform_plot</pre>
<pre>pymethyl-utils-feature_select_train_val_</pre>	_testcommand line option, 20
command line option, $35$ -o,	-output_pkl <output_pkl></output_pkl>
-n, -norm	$\verb pymethyl-preprocess-combine_methylation_arrays \\$
<pre>pymethyl-visualize-plot_heatmap</pre>	command line option, 24
command line option, 20	pymethyl-preprocess-feature_select
-nd, -no_disease_merge	command line option, 26
<pre>pymethyl-preprocess-merge_sample_sheets</pre>	pymethyl-preprocess-imputation_pipeline
command line option, 28	command line option, 27
-nfs, -n_neighbors_fs <n_neighbors_fs></n_neighbors_fs>	pymethyl-preprocess-preprocess_pipeline
<pre>pymethyl-preprocess-imputation_pipeline</pre>	command line option, 29
command line option, 27	pymethyl-utils-backup_pkl command
-nn, -n_neighbors <n_neighbors></n_neighbors>	line option, 33
pymethyl-preprocess-feature_select	pymethyl-utils-create_external_validation_set
command line option, 26	command line option, 35
pymethyl-utils-feature_select_train_val_	_bemethyl-utils-fix_key command
command line option, 35	line option, 36
pymethyl-visualize-transform_plot	pymethyl-utils-modify_pheno_data
command line option, 21	command line option, 36
-noob_norm	pymethyl-utils-overwrite_pheno_data
pymethyl-preprocess-preprocess_pipeline	command line option, 37
command line option, 29	pymethyl-utils-remove_sex command

```
line option, 39
                                                command line option, 35
   pymethyl-utils-remove snps command
                                         -qc, -qc_only
      line option, 40
                                            pymethyl-preprocess-batch_deploy_preprocess
                                                command line option, 23
   pymethyl-utils-set_part_array_background
      command line option, 40
                                            pymethyl-preprocess-preprocess_pipeline
   pymethyl-utils-subset array
                                                command line option, 29
      command line option, 41
                                         -r, -head directory
                                            pymethyl-preprocess-na_report
-o, -subtype_output_dir
      <subtype_output_dir>
                                                command line option, 29
   pymethyl-preprocess-split_preprocess_inputobienstabtique <orientation>
      command line option, 31
                                            pymethyl-preprocess-imputation_pipeline
-os, -output_sample_sheet
                                                command line option, 27
                                         -r, -run
      <output_sample_sheet>
   pymethyl-preprocess-concat_sample_sheetspymethyl-preprocess-batch_deploy_preprocess
      command line option, 24
                                                command line option, 23
   pymethyl-preprocess-create_sample_sheemtef, -reference <reference>
      command line option, 25
                                            pymethyl-utils-ref_estimate_cell_counts
   pymethyl-preprocess-meffil_encode
                                                command line option, 39
      command line option, 28
                                         -ro, -input_r_object_dir
   pymethyl-preprocess-merge_sample_sheets
                                                <input r object dir>
      command line option, 28
                                            pymethyl-utils-ref_estimate_cell_counts
-os, -output_sheet_name
                                                command line option, 39
      <output_sheet_name>
                                         -s, -series
   pymethyl-preprocess-remove diseases
                                            pymethyl-preprocess-batch_deploy_preprocess
      command line option, 30
                                                command line option, 23
-ot, -output_test_pkl
                                         -s, -solver <solver>
      <output_test_pkl>
                                            pymethyl-preprocess-imputation_pipeline
                                                command line option, 27
   pymethyl-utils-bin_column command
      line option, 34
                                         -s, -source_type <source_type>
-p, -package <package>
                                            pymethyl-preprocess-create_sample_sheet
   pymethyl-install-install_custom
                                                command line option, 25
      command line option, 17
                                         -s, -supervised
                                            pymethyl-visualize-transform_plot
   pymethyl-install-install_r_packages
                                                command line option, 21
      command line option, 18
-p, -pc_qc_parameters_csv
                                         -s1, -sample_sheet1 <sample_sheet1>
                                            pymethyl-preprocess-concat_sample_sheets
      <pc_qc_parameters_csv>
   pymethyl-preprocess-batch_deploy_preprocess command line option, 24
      command line option, 23
                                            pymethyl-preprocess-merge_sample_sheets
-p, -pipeline <pipeline>
                                                command line option, 28
   pymethyl-preprocess-preprocess_pipelins2, -sample_sheet2 <sample_sheet2>
      command line option, 29
                                            pymethyl-preprocess-concat sample sheets
-pc, -n pcs <n pcs>
                                                command line option, 24
   pymethyl-preprocess-preprocess_pipeline pymethyl-preprocess-merge_sample_sheets
                                                command line option, 28
      command line option, 29
                                         -sc, -sex_cutoff <sex_cutoff>
-pdc, -p_detection_cpgs
      <p_detection_cpqs>
                                            pymethyl-preprocess-preprocess_pipeline
   pymethyl-preprocess_preprocess_pipeline
                                                command line option, 30
                                         -sd, -sex_sd <sex_sd>
      command line option, 30
-pds, -p_detection_samples
                                            pymethyl-preprocess_preprocess_pipeline
      <p_detection_samples>
                                                command line option, 30
   pymethyl-preprocess-preprocess_pipelinsd, -subtype_delimiter
      command line option, 30
                                                <subtype delimiter>
-q, -query_pkl <query_pkl>
                                            pymethyl-preprocess-get_categorical_distributio
   pymethyl-utils-create external validation secommand line option, 27
```

	_
pymethyl-preprocess-imputation_pipel	i <b>A</b> e
command line option, $28$	assign_results_to_pheno_col() (pymethyl-
<pre>pymethyl-preprocess-remove_diseases</pre>	process.general_machine_learning.MachineLearning
command line option, $30$	method), 15
<pre>pymethyl-preprocess-split_preprocess</pre>	_input_by_subtype
command line option, 31	В
<pre>pymethyl-utils-fix_key command</pre>	bin_column() (pymethylpro-
line option, 36	cess.MethylationDataTypes.MethylationArray
<pre>pymethyl-utils-train_test_val_split</pre>	method), 9
command line option,41	
-ss, -split_by_subtype	C
<pre>pymethyl-preprocess-imputation_pipel</pre>	ine categorical_breakdown() (pymethylpro-
Command Time Operon, 27	cess.MethylationDataTypes.MethylationArray
-st, -sample_threshold	method), 9
<sample_threshold></sample_threshold>	combine() (pymethylpro-
<pre>pymethyl-preprocess-imputation_pipel</pre>	ine cess.MethylationDataTypes.MethylationArrays
command line option, 28	method), 11
-t, -test_pkl <test_pkl></test_pkl>	concat () (pymethylpro-
<pre>pymethyl-utils-bin_column command</pre>	cess.PreProcessDataTypes.PreProcessPhenoData
line option, 33	method), 7
-t, -torque	memou), i
<pre>pymethyl-preprocess-batch_deploy_pre</pre>	p <b>p</b> ocess
command line option, 23	download_clinical() (pymethylpro-
-t, -train_pkl <train_pkl></train_pkl>	cass ProProcess Data Types TCCA Downloader
pymethyl-utils-create_external_valid	ation_set_thod) &
command line option, 35	download_geo() (pymethylpro-
-t, -transpose	cess.PreProcessDataTypes.TCGADownloader
<pre>pymethyl-visualize-plot_heatmap</pre>	method), 8
command line option, 20	download_tcga() (pymethylpro-
-te, -test_pkl <test_pkl></test_pkl>	cess.PreProcessDataTypes.TCGADownloader
<pre>pymethyl-utils-ref_free_cell_deconv</pre>	method), 8
command line option, 39	memou), o
-tp, -train_percent <train_percent></train_percent>	E
<pre>pymethyl-utils-train_test_val_split</pre>	est_cell_counts_IDOL() (in module pymethyl-
command line option, 41	process.meffil_functions), 13
-tr, -train_pkl <train_pkl></train_pkl>	est_cell_counts_meffil() (in module pymethyl-
<pre>pymethyl-utils-ref_free_cell_deconv</pre>	process.meffil_functions), 13
command line option, 39	est_cell_counts_minfi() (in module pymethyl-
-u, -use_cache	process meffil functions) 13
pymethyl-preprocess-batch_deploy_pre	process export() (pymethylpro-
command line option, 23	cess.PreProcessDataTypes.PreProcessPhenoData
pymethyl-preprocess-preprocess_pipel	ine method), 7
command line option, 29	export_csv() (pymethylpro-
-vp, -val_percent <val_percent></val_percent>	cess.PreProcessDataTypes.PreProcessIDAT
<pre>pymethyl-utils-train_test_val_split</pre>	method), 5
command line option, 41	export_pickle() (pymethylpro-
-x, -xticks	cess.PreProcessDataTypes.PreProcessIDAT
<pre>pymethyl-visualize-plot_heatmap</pre>	method), 5
command line option, 20	
-y, -yticks	export_sql() (pymethylpro- cess.PreProcessDataTypes.PreProcessIDAT
pymethyl-visualize-plot_heatmap	method), 5
command line option, $\overline{20}$	extract_manifest() (pymethylpro-
	cess.PreProcessDataTypes.PreProcessIDAT
	method), 5

(in module pymethylprocess.MethylationDataTypes, let (in module pymethylprocess.MethylationDataTypes), l2	extrac	t_pheno_beta_df_from_f	folder()		(pymethylpro-
extract_pheno_beta_df_from_pickle_dict()     (in module				· · · · · · · · · · · · · · · · · · ·	ylationArray
(in module pymethylprocess.MethylationDataTypes), 12 extract_pheno_beta_df_from_sql() (in module pymethylprocess.MethylationDataTypes), 12 extract_pheno_data() (pymethylprocess.MethylationDataTypes.MethylationDataTypes.MethylationDataTypes.MethylationDataTypes.MethylationDataTypes.MethylationDataTypes.MethylationDataTypes.MethylationDataTypes.MethylationDataTypes.MethylationDataTypes.MethylationDataTypes.MethylationDataTypes.MethylationDataTypes.MethylationDataTypes.MethylationDataTypes.MethylationDataTypes.PreProcessIDAT method), 5  Ft eature_select() (pymethylprocess.MethylationDataTypes.MethylationArray method), 5  fittp=beta() (pymethylprocess.MethylationDataTypes.PreProcessIDAT method), 5  fit() (pymethylprocess.general_machine_learning.MachineLearning method), 15  fit_transform() (pymethylprocess.general_machine_learning.MachineLearning method), 15  fit_transform() (pymethylprocess.general_machine_learning.MachineLearning method), 15  format_custem() (pymethylprocess.PreProcessDataTypes.PreProcessPhenoData method), 7  format_tega() (pymethylprocess.PreProcessDataTypes.PreProcessPhenoData method), 7  format_tega() (pymethylprocess.MethylationDataTypes.PreProcessPhenoData method), 7  format_tega() (pymethylprocess.MethylationDataTypes.PreProcessIDAT method), 5  get_beta() (pymethylprocess.PreProcessDataTypes.PreProcessIDAT method), 6  get_beta() (pymethylprocess.PreProcessDataTypes.PreProcessIDAT method), 6  get_tameth() (pymethylprocess.PreProcessDataTypes.PreProcessIDAT method), 6  potential_df_from_all_d() (pymethylprocess.PreProcessDataTypes.PreProcessIDAT metho		* * * * * * * * * * * * * * * * * * * *			
cess.MethylationDataTypes, 1:2	extrac			) 1	
extract_pheno_beta_dt_from_sql() (in mod-		•		I	
we pymethylprocess.MethylationDataTypes,   cess.PreProcessDataTypes.PreProcessIDAT   method), 5   method), 12   method), 15   method), 16   method), 17   method), 19   method), 20   method), 30					
extract_pheno_data() (pymethylpro-	extrac	ule pymethylprocess.Methylar			ylationArray
cess. PreProcessDataTypes. PreProcessIDAT method), 5  Feature_select() (pymethylpro- cess. MethylationDataTypes. MethylationArray method), 5  filter_beta() (pymethylpro- cess. PreProcessDataTypes. PreProcessIDAT method), 15  fit_transform() (pymethylpro- cess. general_machine_learning. MachineLearning method), 15  fit_transform() (pymethylpro- cess. general_machine_learning. MachineLearning method), 15  format_gen() (pymethylpro- cess. PreProcessDataTypes. PreProcessPhenoData method), 7  format_den(), 0  get_beta() (pymethylpro- cess. MethylationDataTypes. MethylationArray class method), 9  G  get_beta() (pymethylpro- cess. PreProcessDataTypes. PreProcessDataType				±	
Feature_select() (pymethylprocess.MethylationDataTypes.MethylationArray method), 9  filter_beta() (pymethylprocess.PreProcessIDAT method), 5  fit() (pymethylprocess.general_machine_learning.MachineLearning method), 15  fit_predict() (pymethylprocess.general_machine_learning.MachineLearning method), 15  fit_transform() (pymethylprocess.general_machine_learning.MachineLearning method), 15  format_custom() (pymethylprocess.PreProcessDataTypes.Pr	extrac	cess.PreProcessDataTypes.PreF		method), 12	
Feature_select() (pymethylprocess.MethylationDataTypes.MethylationArray method), 9  filter_beta() (pymethylprocess.DataTypes.PreProcessIDAT method), 5  fit() (pymethylprocess.general_machine_learning.MachineLearning method), 15  fit() (pymethylprocess.general_machine_learning.MachineLearning method), 15  fit_predict() (pymethylprocess.general_machine_learning.MachineLearning method), 15  fit_transform() (pymethylprocess.general_machine_learning.MachineLearning cess.general_machine_learning.MachineLearning method), 15  format_oustom() (pymethylprocess.PreProcessIDAT method), 7  format_geo() (pymethylprocess.PreProcessIDAT method), 7  format_toga() (pymethylprocess.PreProcessIDAT method), 7  format_toga() (pymethylprocess.PreProcessIDAT method), 9  Gget_beta() (pymethylprocess.PreProcessIDAT method), 7  get_meth() (pymethylprocess.PreProcessIDAT method), 7  get_meth() (pymethylprocess.PreProcessIDAT method), 5  get_unmeth() (pymethylprocess.PreProcessIDAT method), 6  plot_original_qc() (pymethylprocess.PreProcessIDA		method), 5			pymethylpro-
feature_select() (pymethylprocess.MethylationDataTypes.MethylationArray method), 9 filter_beta() (pymethylprocess.PreProcessIDAT method), 5 filt() (pymethylprocess.general_machine_learning.MachineLearning method), 15 fit_predict() (pymethylprocess.general_machine_learning.MachineLearning method), 15 fit_transform() (pymethylprocess.general_machine_learning.MachineLearning cess.general_machine_learning.MachineLearning method), 15 format_custom() (pymethylprocess.PreProcessDataTypes.PreProcessPhenoData method), 7 format_geo() (pymethylprocess.PreProcessDataTypes.PreProcessPhenoData method), 7 format_tcga() (pymethylprocess.PreProcessDataTypes.PreProcessPhenoData method), 6  G  G  G  G  G  G  G  G  G  G  G  G  G	F			cess. Methylation Data Types), 9	
cess.MethylationDataTypes.MethylationArray method), 9 filter_beta() (pymethylpro- cess.PreProcessDataTypes.PreProcessIDAT method), 5 fit() (pymethylprocess.general_machine_learning.MachineLearning method), 15 fit_predict() (pymethylpro- cess.general_machine_learning.MachineLearning method), 15 fit_transform() (pymethylpro- cess.general_machine_learning.MachineLearning method), 15 format_custom() (pymethylpro- cess.PreProcessDataTypes.PreProcessPhenoData method), 15 format_geo() (pymethylpro- cess.PreProcessDataTypes.PreProcessPhenoData method), 7 format_tcga() (pymethylpro- cess.PreProcessDataTypes.PreProcessPhenoData method), 7 form_pickle() (pymethylpro- cess.MethylationDataTypes.MethylationArray class method), 9  G  G  get_beta() (pymethylpro- cess.MethylationDataTypes.PreProcessIDAT method), 5 get_categorical_distribution() (pymethylpro- cess.PreProcessDataTypes.PreProcessIDAT method), 6  overwrite_pvalues_beaction_pvalues_beaching() (pymethylpro- cess.PreProcessDataTypes.PreProcessIDAT method), 6  overwrite_pvalues_beaction_prevalues_beaching() (pymethylpro- cess.PreProcessDataTypes.PreProcessIDAT method), 6  overwrite_pheno_beta() (pymethylpro- cess.PreProcessDataTypes.PreProcessIDAT m				1	
method), 9  filter_beta() (pymethylprocess.PreProcessDataTypes.PreProcessIDAT method), 5  fit() (pymethylprocess.general_machine_learning.MachineLearning method), 5  fit() (pymethylprocess.general_machine_learning.MachineLearning method), 15  fit_predict() (pymethylprocess.general_machine_learning.MachineLearning method), 15  fit_transform() (pymethylprocess.general_machine_learning.MachineLearning method), 15  format_custom() (pymethylprocess.PreProcessDataTypes.PreProcessPhenoData method), 7  format_geo() (pymethylprocess.PreProcessDataTypes.PreProcessPhenoData method), 7  format_tcga() (pymethylprocess.PreProcessDataTypes.P	featur				
cess.PreProcessDataTypes.PreProcessIDAT method), 5 fit () (pymethylprocess.general_machine_learning.MachineLearning method), 15 fit_predict() (pymethylprocess.general_machine_learning.MachineLearning method), 15 fit_transform() (pymethylprocess.general_machine_learning.MachineLearning method), 15 fit_transform() (pymethylprocess.general_machine_learning.MachineLearning method), 15 format_custom() (pymethylprocess.PreProcessPhenoData method), 7 format_geo() (pymethylprocess.PreProcessPhenoData method), 7 format_tcga() (pymethylprocess.PreProcessDataTypes.PreProcessPhenoData method), 7 format_tcga() (pymethylprocess.PreProcessDataTypes.PreProcessDataTy		method), 9			
method), 5 fit () (pymethylprocess_general_machine_learning.MachineLearning method), 15  fit_predict () (pymethylprocess_general_machine_learning.MachineLearning method), 15 fit_transform () (pymethylprocess_general_machine_learning.MachineLearning method), 15 fit_transform () (pymethylprocess_general_machine_learning.MachineLearning method), 15 format_custom () (pymethylprocess_general_machine_learning.MachineLearning method), 15 format_custom () (pymethylprocess_general_machine_learning.MachineLearning method), 16 format_custom () (pymethylprocess_general_machine_learning.MachineLearning cess.PreProcessDataTypes.PreProcessPhenoData method), 15 format_custom () (pymethylprocess_general_machine_learning.MachineLearning cess.PreProcessDataTypes.PreProcessPhenoData method), 16 format_custom () (pymethylprocess.PreProcessDataTypes.PreProcessPhenoData method), 7 format_ge () (pymethylprocess.MethylationDataTypes.PreProcessPhenoData method), 7 format_tcga () (pymethylprocess.PreProcessDataTypes.PreProcessPhenoData method), 7 format_tcga () (pymethylprocess.PreProcessDataTypes.PreProcessIDAT method), 6  G get_beta () (pymethylprocess.PreProcessDataTypes.PreProcessDataTypes.PreProcessDataTypes.PreProcessIDAT method), 6  get_categorical_distribution () (pymethylprocess.PreProcessDataTypes.PreProcessIDAT method), 5 get_unmeth () (pymethylprocess.PreProcessIDAT method), 5 get_unmeth () (pymethylprocess.PreProcessIDAT method), 6 plot_genetics () (pymethylprocess.PrePr	filter			<pre>load_idats()</pre>	(pymethylpro-
fit () (pymethylprocess general_machine_learning_MachineLearning		* *	ProcessIDAT		ocessIDAT
fit_predict() (pymethylprocess general_machine_learning.MachineLearning	fit()(		ne_learning.Mach	ineLearning	
cess.general_machine_learning.MachineLearning method), 15  fit_transform() (pymethylpro- cess.general_machine_learning.MachineLearning method), 15  format_oustom() (pymethylpro- cess.PreProcessDataTypes.PreProcessPhenoData method), 7  format_geo() (pymethylpro- cess.PreProcessDataTypes.PreProcessPhenoData method), 7  format_tcga() (pymethylpro- cess.PreProcessDataTypes.PreProcessIDAT method), 6  G  G  G  G  G  G  G  Get_categorical_distribution() (pymethylpro- cess.PreProcessDataTypes.PreProcessIDAT method), 5  get_categorical_distribution() (pymethylpro- cess.PreProcessDataTypes.PreProcessIDAT method), 6  output_pheno_beta() (pymethylpro- cess.PreProcessDataTypes.PreProcessIDAT method), 6  output_pheno_beta() (pymethylpro- cess.PreProces	fit nn	· · · · · · · · · · · · · · · · · · ·	(mymathylnra		
fit_transform() (pymethylprocess. PreProcess Data Types. PreProcess Pheno Data method), 15  format_custom() (pymethylprocess. PreProcess Pheno Data method), 7  format_geo() (pymethylprocess. PreProcess Data Types. PreProcess Pheno Data method), 7  format_tcga() (pymethylprocess. PreProcess Pheno Data method), 7  form_pickle() (pymethyl ation Data Types. PreProcess Pheno Data method), 9  G  G  G  G  G  G  G  G  G  G  G  G  G	TTC_bt	cess.general_machine_learning		g cess.general_machine_learning),	15
cess.general_machine_learning.MachineLearning method), 15  format_custom() (pymethylpro- cess.PreProcessDataTypes.PreProcessPhenoData method), 7  format_geo() (pymethylpro- cess.PreProcessDataTypes.PreProcessPhenoData method), 7  format_tcga() (pymethylpro- cess.PreProcessDataTypes.PreProcessPhenoData method), 7  format_tcga() (pymethylpro- cess.PreProcessDataTypes.PreProcessPhenoData method), 7  from_pickle() (pymethylpro- cess.MethylationDataTypes.PreProcessIDAT method), 9  G  get_beta() (pymethylpro- cess.PreProcessDataTypes.PreProcessIDAT method), 5  get_categorical_distribution() (pymethyl- process.PreProcessDataTypes.PreProcessIDAT method), 7  get_meth() (pymethylpro- cess.PreProcessDataTypes.PreProcessIDAT method), 5	fit tr		(nymethylnro-		
format_custom() (pymethylpro-	110_01	cess.general_machine_learning		g method), 8	
cess.PreProcessDataTypes.PreProcessPhenoData method), 7  format_geo() (pymethylpro- cess.PreProcessDataTypes.PreProcessPhenoData method), 7  format_tcga() (pymethylpro- cess.PreProcessDataTypes.PreProcessPhenoData method), 7  format_tcga() (pymethylpro- cess.PreProcessDataTypes.PreProcessPhenoData method), 7  from_pickle() (pymethylpro- cess.MethylationDataTypes.MethylationArray class method), 9  G  get_beta() (pymethylpro- cess.PreProcessDataTypes.PreProcessIDAT method), 6  output_pheno_beta() (pymethylpro- cess.PreProcessDataTypes.PreProcessIDAT method), 6  overwrite_pheno_data() (pymethylpro- cess.MethylationDataTypes.MethylationArray method), 6  overwrite_pheno_data() (pymethylpro- cess.MethylationDataTypes.MethylationArray method), 10  get_meth() (pymethylpro- cess.PreProcessDataTypes.PreProcessIDAT method), 7  get_meth() (pymethylpro- cess.PreProcessDataTypes.PreProcessIDAT method), 5  get_unmeth() (pymethylpro- cess.PreProcessDataTypes.PreProcessIDAT method), 6  plot_qc_metrics() (pymethylpro- cess.PreProcessIDAT method), 6	format		(pymethylpro-		
format_geo() (pymethylpro-			ProcessPhenoDate	a	yiaii0112 XII a y
cess.PreProcessDataTypes.PreProcessPhenoData method), 7  format_tcga()	_			MethylationArray (class in	pymethylpro-
cess.MethylationDataTypes, 11 format_tcga()	format	_geo()	(pymethylpro-	cess. Methylation Data Types), 9	
cess.PreProcessDataTypes.PreProcessPhenoData method), 7  from_pickle() (pymethylprocess.MethylationDataTypes.MethylationArray class method), 9  G  get_beta() (pymethylprocess.PreProcessDataTypes.PreProcessIDAT method), 5  get_categorical_distribution() (pymethylprocess.PreProcessDataTypes.PreProcessPhenoDataTypes.PreProcessDataTypes.PreProcessIDAT method), 7  get_meth() (pymethylprocess.PreProcessDataTypes.PreProcessIDAT method), 5  get_meth() (pymethylprocess.PreProcessDataTypes.PreProcessIDAT method), 5  get_unmeth() (pymethylprocess.PreProcessDataTypes.PreProcessIDAT method), 6  plot_original_qc() (pymethylprocess.PreProcessIDAT method), 6  plot_original_qc() (pymethylprocess.PreProcessDataTypes.PreProcessIDAT method), 6		methoa), /		aMethylationArrays (class in cess.MethylationDataTypes), 11	pymethylpro-
method), 7 from_pickle() (pymethylprocess.MethylationDataTypes.MethylationArray class method), 9  G get_beta() (pymethylprocess.PreProcessIDAT method), 5 get_categorical_distribution() (pymethylprocess.PreProcessDataTypes.PreProcessPhenoDatamethod), 7 get_meth() (pymethylprocess.PreProcessDataTypes.PreProcessIDAT method), 5 get_meth() (pymethylprocess.PreProcessDataTypes.PreProcessIDAT method), 5 get_meth() (pymethylprocess.PreProcessDataTypes.PreProcessIDAT method), 5 get_unmeth() (pymethylprocess.PreProcessDataTypes.PreProcessIDAT method), 6 plot_original_qc() (pymethylprocess.PreProcessDataTypes.PreProcessIDAT method), 6 plot_original_qc() (pymethylprocess.PreProcessDataTypes.PreProcessIDAT method), 6 plot_original_qc() (pymethylprocess.PreProcessIDAT method), 6 plot_original_qc() (pymethylpr	format				
from_pickle() (pymethylpro-		· -		cess.i rei rocessDaiaiypes.i rei ro	ocessidai
cess.MethylationDataTypes.MethylationArray class method), 9  G  get_beta() (pymethylprocess.PreProcessIDAT method), 5  get_categorical_distribution() (pymethylpromethod), 7  get_meth() (pymethylprocess.PreProcessIDAT method), 5  get_meth() (pymethylprocess.PreProcessIDAT method), 5  get_meth() (pymethylprocess.PreProcessIDAT method), 5  get_unmeth() (pymethylprocess.PreProcessIDAT method), 6  get_unmeth() (pymethylprocess.PreProcessIDAT method), 6  get_unmeth() (pymethylprocess.PreProcessIDAT method), 6  get_unmeth() (pymethylprocess.PreProcessIDAT method), 6  plot_qc_metrics() (pymethylprocess.PreProcessIDAT method), 6  plot_qc_metrics() (pymethylprocess.PreProcessIDAT method), 6	from_p		(pymethylpro-	memoa), o	
class method), 9  Quitput_pheno_beta() (pymethylpro-cess.PreProcessIDAT method), 6  Quitput_pheno_beta() (pymethylpro-cess.PreProcessIDAT method), 10  Quitput_pheno_beta() (pymethylpro-c				$\circ$	
cess.PreProcessDataTypes.PreProcessIDAT method), 6  get_beta()		· · · · · · · · · · · · · · · · · · ·		•	/ .1 1
<pre>get_beta()</pre>	G			cess. Pre Process Data Types. Pre Process Data Types. Pre Process Data Types Data Data Data Data Data Data Data Dat	
cess.PreProcessDataTypes.PreProcessIDAT method), 5  get_categorical_distribution() (pymethyl- process.PreProcessDataTypes.PreProcessPhenoDpa method), 7  get_meth() (pymethylpro- cess.PreProcessDataTypes.PreProcessIDAT method), 5  get_unmeth() (pymethylpro- cess PreProcessDataTypes.PreProcessIDAT method), 6  plot_qc_metrics() (pymethylpro- cess.PreProcessDataTypes.PreProcessIDAT method), 6  plot_qc_metrics() (pymethylpro- cess.PreProcessDataTypes.PreProcessIDAT method), 6	ret he	t a ()	(nymethylpro-		
get_categorical_distribution() (pymethyl-	900_00	cess.PreProcessDataTypes.PreF		cess. Methylation Data Types. Methylogistal Methylogista	
method), 7 get_meth()	get_ca	tegorical_distribution		**	
get_meth() (pymethylpro- cess.PreProcessDataTypes.PreProcessIDAT method), 5  get_unmeth() (pymethylpro- cess.PreProcessDataTypes.PreProcessIDAT method), 6  plot_original_qc() (pymethylpro- cess.PreProcessDataTypes.PreProcessIDAT  method), 6  plot_qc_metrics() (pymethylpro- cess.PreProcessDataTypes.PreProcessIDAT				•	(
cess.PreProcessDataTypes.PreProcessIDAT method), 5  get_unmeth() (pymethylpro- cess.PreProcessDataTypes.PreProcessIDAT  method), 6  plot_qc_metrics() (pymethylpro- cess.PreProcessDataTypes.PreProcessIDAT  cess.PreProcessDataTypes.PreProcessIDAT	get_me		(pymethylpro-		
get_unmeth() (pymethylpro- cess PreProcessDataTypes PreProcessIDAT  cess PreProcessDataTypes PreProcessIDAT	_	cess.PreProcessDataTypes.PreF			ocessiDA1
get_unifieth() (pymeinyipro- cess PreProcessDataTypes PreProcessIDAT cess.PreProcessDataTypes.PreProcessIDAT	ao+ :::-		(mmathylana	<pre>plot_qc_metrics()</pre>	(pymethylpro-
method), 6	get_un	cess.PreProcessDataTypes.PreF			ocessIDAT

```
(pymethylpro-
predict()
                                               -o, -output_pkl <output_pkl>, 24
       cess.general_machine_learning.MachineLearningpymethyl-preprocess-concat_sample_sheets
       method), 15
                                                   command line option
preprocess_enmix_pipeline() (pymethylpro-
                                               -os, -output_sample_sheet
       cess. PreProcessDataTypes. PreProcessIDAT
                                                   <output_sample_sheet>, 24
       method), 6
                                               -s1, -sample sheet1
preprocessENmix()
                               (pymethylpro-
                                                   <sample sheet1>,24
       cess.PreProcessDataTypes.PreProcessIDAT
                                               -s2, -sample_sheet2
       method), 6
                                                   <sample_sheet2>, 24
PreProcessIDAT
                                pymethylpro- pymethyl-preprocess-create_sample_sheet
                   (class
                           in
       cess.PreProcessDataTypes), 5
                                                   command line option
                                               -b, -basename_col <basename_col>, 25
preprocessMeffil()
                               (pymethylpro-
       cess.PreProcessDataTypes.PreProcessIDAT
                                               -c, -include_columns_file
       method), 6
                                                   <include_columns_file>, 25
preprocessNoob()
                               (pymethylpro-
                                               -d, -disease_class_column
       cess. Pre Process Data Types. Pre Process IDAT\\
                                                   <disease_class_column>, 25
       method), 6
                                               -i, -idat_dir <idat_dir>,25
PreProcessPhenoData (class in pymethylpro-
                                               -is, -input_sample_sheet
       cess.PreProcessDataTypes), 7
                                                   <input_sample_sheet>, 25
preprocessRAW()
                                               -1, -header_line <header_line>, 25
                               (pymethylpro-
       cess.PreProcessDataTypes.PreProcessIDAT
                                               -m, -mapping_file <mapping_file>, 25
                                               -os, -output_sample_sheet
                                                   <output_sample_sheet>, 25
pymethyl-install command line option
   -version, 17
                                               -s, -source_type <source_type>, 25
pymethyl-install-install_custom
                                           pymethyl-preprocess-download_clinical
       command line option
                                                  command line option
   -m, -manager, 17
                                               -o, -output_dir <output_dir>, 25
   -p, -package <package>, 17
                                           pymethyl-preprocess-download_geo
pymethyl-install-install_r_packages
                                                  command line option
       command line option
                                               -g, -geo_query <geo_query>, 26
   -p, -package <package>, 18
                                               -o, -output_dir <output_dir>, 26
pymethyl-preprocess command line
                                           pymethyl-preprocess-download_tcga
       option
                                                  command line option
   -version, 23
                                               -o, -output_dir <output_dir>, 26
pymethyl-preprocess-batch_deploy_preprocessmethyl-preprocess-feature_select
       command line option
                                                  command line option
   -c, -chunk size <chunk size>, 24
                                               -f, -feature selection method
   -i, -subtype_output_dir
                                                   <feature_selection_method>, 26
       <subtype_output_dir>, 23
                                               -i, -input_pkl <input_pkl>, 26
   -m, -meffil, 23
                                               -m, -mad_top_cpgs <mad_top_cpgs>, 26
   -n, -n_cores <n_cores>, 23
                                               -mm, -metric <metric>, 26
   -p, -pc_qc_parameters_csv
                                               -n, -n_top_cpgs <n_top_cpgs>, 26
                                               -nn, -n_neighbors <n_neighbors>, 26
       <pc_qc_parameters_csv>, 23
                                               -o, -output_pkl <output_pkl>, 26
   -qc, -qc_only, 23
   -r, -run, 23
                                           pymethyl-preprocess-get_categorical_distribution
   -s, -series, 23
                                                  command line option
   -t, -torque, 23
                                               -d, -disease_only, 27
   -u, -use_cache, 23
                                               -is, -formatted_sample_sheet
pymethyl-preprocess-combine_methylation_arrays <formatted_sample_sheet>, 27
       command line option
                                               -k, -key < key >, 27
   -d, -optional_input_pkl_dir
                                               -sd, -subtype_delimiter
       <optional_input_pkl_dir>, 24
                                                   <subtype delimiter>, 27
   -e, -exclude <exclude>, 24
                                           pymethyl-preprocess-imputation_pipeline
   -i, -input_pkls <input_pkls>, 24
                                                  command line option
```

```
-ct, -cpg_threshold
                                              -pdc, -p_detection_cpgs
       <cpg_threshold>, 28
                                                 <p_detection_cpgs>, 30
   -d, -disease only, 27
                                              -pds, -p_detection_samples
   -f, -feature_selection_method
                                                 <p_detection_samples>, 30
       <feature selection method>, 27
                                              -qc, -qc_only, 29
   -i, -input pkl <input pkl>, 27
                                              -sc, -sex cutoff <sex cutoff>, 30
   -k, -n neighbors <n neighbors>, 27
                                              -sd, -sex sd < sex sd > ,30
   -m, -method <method>, 27
                                              -u, -use cache, 29
   -mm, -metric <metric>, 27
                                          pymethyl-preprocess-remove_diseases
   -n, -n_top_cpgs <n_top_cpgs>, 27
                                                 command line option
   -nfs, -n_neighbors_fs
                                              -d, -disease_only, 30
       <n_neighbors_fs>, 27
                                              -e, -exclude_disease_list
   -o, -output_pkl <output_pkl>, 27
                                                 <exclude_disease_list>, 30
   -r, -orientation <orientation>, 27
                                              -is, -formatted_sample_sheet
   -s, -solver < solver >, 27
                                                 <formatted_sample_sheet>, 30
   -sd, -subtype_delimiter
                                              -1, -low_count <low_count>, 30
       <subtype_delimiter>, 28
                                              -os, -output_sheet_name
   -ss, -split_by_subtype, 27
                                                 <output sheet name>, 30
   -st, -sample_threshold
                                              -sd, -subtype_delimiter
       <sample_threshold>, 28
                                                 <subtype delimiter>, 30
pymethyl-preprocess-meffil_encode
                                          pymethyl-preprocess-split_preprocess_input_by_subt
       command line option
                                                 command line option
   -is, -input_sample_sheet
                                              -d, -disease_only, 31
                                              -i, -idat_csv <idat_csv>,31
       <input sample sheet>, 28
   -os, -output_sample_sheet
                                              -o, -subtype_output_dir
       <output_sample_sheet>, 28
                                                 <subtype_output_dir>, 31
pymethyl-preprocess-merge_sample_sheets
                                              -sd, -subtype_delimiter
       command line option
                                                 <subtype_delimiter>, 31
   -d, -second_sheet_disease, 28
                                          pymethyl-utils command line option
   -nd, -no_disease_merge, 28
                                              -version, 33
   -os, -output_sample_sheet
                                          pymethyl-utils-backup_pkl command line
       <output_sample_sheet>, 28
                                                 option
   -s1, -sample_sheet1
                                              -i, -input_pkl <input_pkl>, 33
       <sample_sheet1>, 28
                                              -o, -output_pkl <output_pkl>,33
                                          pymethyl-utils-bin_column command line
   -s2, -sample_sheet2
       <sample_sheet2>, 28
                                                 option
pymethyl-preprocess-na_report command
                                              -c, -col < col >, 33
       line option
                                              -n, -n_bins < n_bins>, 33
   -i, -input_pkl <input_pkl>, 29
                                              -ot, -output_test_pkl
   -o, -output_dir <output_dir>, 29
                                                 <output_test_pkl>, 34
   -r, -head directory, 29
                                              -t, -test pkl <test pkl>,33
pymethyl-preprocess-preprocess_pipeline pymethyl-utils-concat_csv command line
       command line option
                                                 option
   -bnc, -p_beadnum_cpgs
                                              -a, -axis < axis > ,34
      <p_beadnum_cpgs>, 30
                                              -i, -index_col <index_col>, 34
                                              -i1, -input_csv <input_csv>, 34
   -bns, -p_beadnum_samples
                                              -i2, -input_csv2 <input_csv2>,34
       <p_beadnum_samples>, 29
   -i, -idat_dir <idat_dir>,29
                                              -o, -output_csv <output_csv>, 34
   -m, -meffil, 29
                                          pymethyl-utils-counts command line
   -n, -n_cores < n_cores>, 29
                                                 option
                                              -i, -input_pkl <input_pkl>,34
   -noob, -noob_norm, 29
                                              -k, -key < key >, 34
   -o, -output pkl <output pkl>, 29
   -p, -pipeline <pipeline>, 29
                                          pymethyl-utils-create_external_validation_set
                                                 command line option
   -pc, -n_pcs < n_pcs >, 29
```

```
-c, -cpq_replace_method
                                         pymethyl-utils-print_number_sex_cpgs
       <cpg_replace_method>, 35
                                                command line option
   -o, -output_pkl <output_pkl>, 35
                                             -a, -array_type <array_type>, 38
   -q, -query_pkl <query_pkl>,35
                                             -i, -input_pkl <input_pkl>,38
   -t, -train_pkl <train_pkl>,35
                                         pymethyl-utils-print_shape command
pymethyl-utils-est age command line
                                                line option
                                             -i, -input pkl <input pkl>,38
      option
   -a, -analyses <analyses>,35
                                         pymethyl-utils-rate_regression command
   -ac, -age_column <age_column>,35
                                                line option
   -i, -input_pkl <input_pkl>, 35
                                             -c1, -pred_col <pred_col>,38
   -o, -output_csv <output_csv>, 35
                                             -c2, -true_col <true_col>,38
pymethyl-utils-feature_select_train_val_testi, -input_csv <input_csv>,38
      command line option
                                         pymethyl-utils-ref_estimate_cell_counts
   -f, -feature_selection_method
                                                command line option
      <feature_selection_method>, 35
                                             -a, -algorithm <algorithm>,39
   -i, -input_pkl_dir <input_pkl_dir>,
                                             -1, -library <library>, 39
                                             -o, -output_csv <output_csv>, 39
   -m, -mad_top_cpgs <mad_top_cpgs>, 36
                                             -ref, -reference <reference>, 39
   -mm, -metric <metric>, 35
                                             -ro, -input_r_object_dir
   -n, -n_top_cpgs <n_top_cpgs>, 35
                                                <input_r_object_dir>,39
   -nn, -n_neighbors <n_neighbors>, 35
                                         pymethyl-utils-ref_free_cell_deconv
   -o, -output_dir <output_dir>,35
                                                command line option
pymethyl-utils-fix_key command line
                                             -a, -analysis <analysis>, 39
                                             -c, -cell_type_columns
      option
   -d, -disease only, 36
                                                <cell_type_columns>, 39
   -i, -input_pkl <input_pkl>,36
                                             -k, -n_cell_types <n_cell_types>, 39
   -k, -key < key >, 36
                                             -te, -test_pkl <test_pkl>,39
   -o, -output_pkl <output_pkl>,36
                                             -tr, -train_pkl <train_pkl>,39
   -sd, -subtype_delimiter
                                         pymethyl-utils-remove_sex command line
       <subtype_delimiter>,36
                                                option
                                             -a, -array_type <array_type>, 39
pymethyl-utils-modify_pheno_data
      command line option
                                             -i, -input_pkl <input_pkl>,39
   -i, -input_pkl <input_pkl>,36
                                             -o, -output_pkl <output_pkl>, 39
   -is, -input_formatted_sample_sheet
                                         pymethyl-utils-remove_snps command
       <input_formatted_sample_sheet>,
                                                line option
                                             -a, -array_type <array_type>, 40
   -o, -output pkl <output pkl>, 36
                                             -i, -input pkl <input pkl>, 40
pymethyl-utils-move_jpg command line
                                             -o, -output_pkl <output_pkl>, 40
      option
                                         pymethyl-utils-set_part_array_background
   -i, -input_dir <input_dir>,37
                                                command line option
                                             -c, -cpg_pkl < cpg_pkl >, 40
   -o, -output dir <output dir>,37
pymethyl-utils-overwrite_pheno_data
                                             -i, -input_pkl <input_pkl>,40
      command line option
                                             -o, -output_pkl <output_pkl>, 40
   -c, -index_col <index_col>,37
                                         pymethyl-utils-stratify command line
   -i, -input_pkl <input_pkl>,37
                                                option
   -is, -input_formatted_sample_sheet
                                             -i, -input_pkl <input_pkl>,40
       <input_formatted_sample_sheet>,
                                             -k, -key < key >, 40
                                             -o, -output_dir <output_dir>,41
   -o, -output_pkl <output_pkl>, 37
                                         pymethyl-utils-subset_array command
pymethyl-utils-pkl_to_csv command line
                                                line option
                                             -c, -cpg_pkl < cpg_pkl > ,41
      option
   -c, -col <col>, 37
                                             -i, -input pkl <input pkl>,41
   -i, -input_pkl <input_pkl>,37
                                             -o, -output_pkl <output_pkl>,41
```

-o, -output\_dir <output\_dir>,37

<pre>pymethyl-utils-train_test_val_split     command line option</pre>	<pre>pymethylprocess.general_machine_learning</pre>
-cat, -categorical,41	pymethylprocess.meffil_functions (mod-
-do, -disease_only,41	<i>ule</i> ), 12
-i, -input_pkl <input_pkl>,41</input_pkl>	pymethylprocess.MethylationDataTypes
-k, -key <key>,41</key>	(module), 8
-o, -output_dir <output_dir>,41</output_dir>	pymethylprocess.PreProcessDataTypes
-sd, -subtype_delimiter	(module), 3
	(module), 5
<pre><subtype_delimiter>,41</subtype_delimiter></pre>	R
-tp, -train_percent	
<pre><train_percent>,41</train_percent></pre>	r_autosomal_cpgs() (in module pymethylpro-
-vp, -val_percent <val_percent>,41</val_percent>	cess.meffil_functions), 13
<pre>pymethyl-utils-write_cpgs command line</pre>	r_snp_cpgs() (in module pymethylpro-
option	cess.meffil_functions), 13
-c, -cpg_pkl <cpg_pkl>,42</cpg_pkl>	remove_diseases() (pymethylpro-
-i, -input_pkl <input_pkl>,42</input_pkl>	cess. Pre Process Data Types. Pre Process Pheno Data
pymethyl-visualize command line option	method), 8
-version, 19	remove_missingness() (pymethylpro-
<pre>pymethyl-visualize-plot_cell_type_result</pre>	cess.MethylationDataTypes.MethylationArray
command line option	method), 10
-cols, -plot_cols <plot_cols>, 19</plot_cols>	remove_na_samples() (pymethylpro-
-fs, -font_scale <font_scale>, 19</font_scale>	cess.MethylationDataTypes.MethylationArray
-i, -input_csv <input_csv>,19</input_csv>	method), 10
-o, -outfilename <outfilename>, 19</outfilename>	remove_sex() (in module pymethylpro-
<pre>pymethyl-visualize-plot_heatmap</pre>	cess.meffil_functions), 13
command line option	remove_whitespace() (pymethylpro-
-a, -annot, 20	cess.MethylationDataTypes.MethylationArray
-c, -cluster,20	method), 10
-col, -color_column <color_column>,</color_column>	return_beta() (pymethylpro-
20	cess.PreProcessDataTypes.PreProcessIDAT
-fs, -font_scale <font_scale>, 20</font_scale>	method), 7
-i, -input_csv <input_csv>, 20</input_csv>	return_cpgs() (pymethylpro-
-idx, -index_col <index_col>, 20</index_col>	cess.MethylationDataTypes.MethylationArray
-m, -matrix_type <matrix_type>, 20</matrix_type>	method), 10
-max, -max_val <max_val>,20</max_val>	return_idx() (pymethylpro-
-min, -min_val <min_val>, 20</min_val>	cess.MethylationDataTypes.MethylationArray
-n, -norm, 20	method), 10
-o, -outfilename <outfilename>, 20</outfilename>	
-t, -transpose, 20	return_imputer() (pymethylpro-
-x, -xticks, 20	cess.MethylationDataTypes.ImputerObject
-y, -yticks, 20	method), 9
	return_outcome_metric() (pymethylpro-
pymethyl-visualize-transform_plot	cess.general_machine_learning.MachineLearning
command line option	method), 16
-a, -axes_off,21	return_raw_beta_array() (pymethylpro-
-c, -column_of_interest	cess. Methylation Data Types. Methylation Array
<pre><column_of_interest>, 20</column_of_interest></pre>	method), 10
-cc, -case_control_override, 21	return_shape() (pymethylpro-
-d, -min_dist <min_dist>,21</min_dist>	cess. Methylation Data Types. Methylation Array
-i, -input_pkl <input_pkl>,20</input_pkl>	method), 10
-m, -metric <metric>,21</metric>	C
-nn, -n_neighbors <n_neighbors>,21</n_neighbors>	S
-o, -output_file <output_file>, 20</output_file>	<pre>set_missing() (in module pymethylpro-</pre>
-s, -supervised,21	cess.meffil_functions), 13

```
split_by_subtype()
                                       (pymethylpro-
         cess. Methylation Data Types. Methylation Array
        method), 10
                                       (pymethylpro-
split_key()
        cess. Methylation Data Types. Methylation Array
        method), 10
                                       (pymethylpro-
split_key()
         cess. PreProcess Data Types. PreProcess Pheno Data
        method), 8
                                       (pymethylpro-
split_train_test()
        cess. Methylation Data Types. Methylation Array
        method), 10
                                       (pymethylpro-
store_results()
        cess.general_machine_learning.MachineLearning
        method), 16
subsample()
                                       (pymethylpro-
        cess. Methylation Data Types. Methylation Array
        method), 11
subset_cpgs()
                                       (pymethylpro-
         cess. Methylation Data Types. Methylation Array\\
        method), 11
subset index()
                                       (pymethylpro-
        cess. Methylation Data Types. Methylation Array
        method), 11
TCGADownloader
                       (class
                                  in
                                        pymethylpro-
        cess.PreProcessDataTypes), 8
to_methyl_array()
                                       (pymethylpro-
        cess.PreProcessDataTypes.PreProcessIDAT
        method), 7
transform()
                                       (pymethylpro-
         cess.general machine learning.MachineLearning
        method), 16
transform_results_to_beta()
                                       (pymethylpro-
        cess.general_machine_learning.MachineLearning
        method), 16
W
write_csvs()
                                       (pymethylpro-
        cess. Methylation Data Types. Methylation Array
        method), 11
                                       (pymethylpro-
write db()
        cess. Methylation Data Types. Methylation Array
        method), 11
write_dbs()
                                       (pymethylpro-
        cess. Methylation Data Types. Methylation Arrays
        method), 12
write_pickle()
                                       (pymethylpro-
        cess. Methylation Data Types. Methylation Array
         method), 11
                                       (pymethylpro-
write_pkls()
        cess. Methylation Data Types. Methylation Arrays
        method), 12
```