# methylize Documentation

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# CHAPTER 1

# **Getting Started**

methylize is a python package for analyzing output from Illumina methylation arrays. It complements methylprep and methylcheck and provides methods for computing differentially methylated probes and regions, and annotating these regions with the UCSC Genome Browser. View on ReadTheDocs.

# 1.1 methylize is part of the methylsuite

methylize is part of the methylsuite of python packages that provide functions to analyze DNA methylation data from Illumina's Infinium arrays (27k, 450k, and EPIC, as well as mouse arrays). This package is focused on analysis of processed methylation data, such as EWAS using Manhattan and Volcano plots. methylize functions are designed to work with a minimum of knowledge and specification required. But you can always override the "smart" defaults with custom settings if the default settings don't work for your data. The entire methylsuite is designed in this format: to offer ease of use while still maintaining flexibility for customization as needed.

# 1.2 Methylsuite package components

You should install all three components, as they work together. The parts include:

- methylprep: for processing idat files or downloading GEO datasets from NIH. Processing steps include
  - infer type-I channel switch
  - NOOB (normal-exponential convolution on out-of-band probe data)
  - poobah (p-value with out-of-band array hybridization, for filtering low signal-to-noise probes)
  - qualityMask (to exclude historically less reliable probes)
  - nonlinear dye bias correction (AKA signal quantile normalization between red/green channels across a sample)
  - calculate beta-value, m-value, or copy-number matrix
  - large batch memory management, by splitting it up into smaller batches during processing
- methylcheck: (this package) for quality control (QC) and analysis, including
  - functions for filtering out unreliable probes, based on the published literature
    - \* Note that methylprep process will exclude a set of unreliable probes by default. You can disable that using the -no\_quality\_mask option from CLI.
  - sample outlier detection
  - array level QC plots, based on Genome Studio functions
  - a python clone of Illumina's Bead Array Controls Reporter software (QC)
  - data visualization functions based on seaborn and matplotlib graphic libraries.
  - predict sex of human samples from probes
  - interactive method for assigning samples to groups, based on array data, in a Jupyter notebook
- methylize provides more analysis and interpretation functions
  - differentially methylated probe statistics (between treatment and control samples)
  - volcano plots (which probes are the most different?)
  - manhattan plots (where in genome are the differences?)

### 1.3 Table of Contents

- Differentially methylated position (DMP) regression, detection and visualation
  - Logistic Regression
  - Linear Regression
  - Manhattan Plot
  - Volcano plot
- Differentially methylated regions
  - Gene annotation with the UCSC Human Genome Browser

# 1.4 Installation

```
pip3 install methylize
```

Installation will also install the other parts of the methylsuite (methylprep and methylcheck) if they are not already installed.

If progress bar is missing: If you don't see a progress bar in your jupyterlab notebook, try this:

```
- conda install -c conda-forge nodejs
- jupyter labextension install @jupyter-widgets/jupyterlab-manager
```

# 1.5 Methylize Package

The methylize package contains both high-level APIs for processing data from local files and low-level functionality allowing you to analyze your data AFTER running methylprep and methylcheck. For greatest usability, import methylize into a Jupyer Notebook along with your processed sample data (a DataFrame of beta values or m-values and a separate DataFrame containing meta data about the samples).

Methylize allows you to run linear or logistic regression on all probes and identify points of interest in the methylome where DNA is differentially modified. Then you can use these regression results to create *volcano plots* and *manhattan plots*.

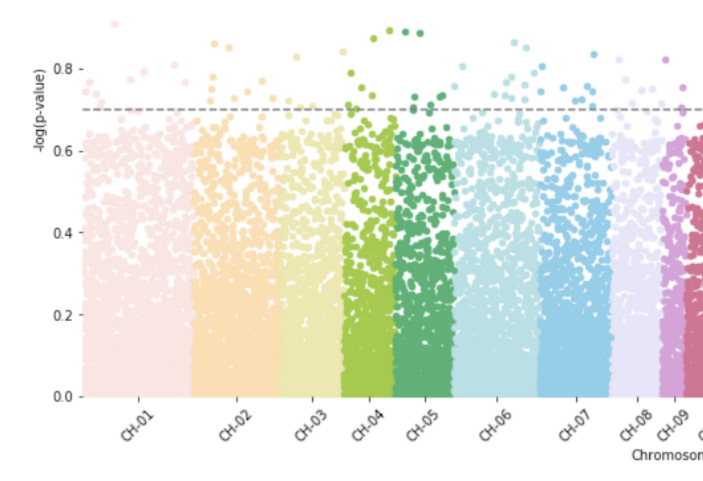
1.4. Installation 3

# 1.5.1 Sample Manhattan Plot

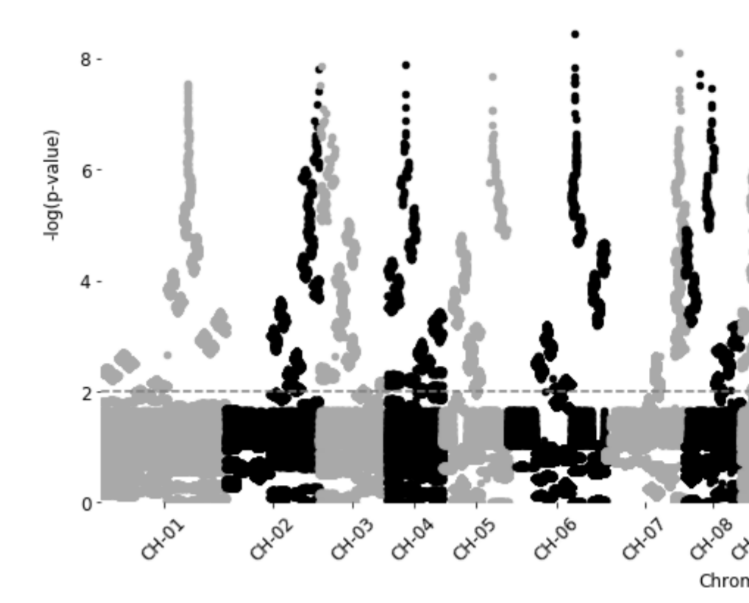
1.4 -

1.2 -

1.0 -

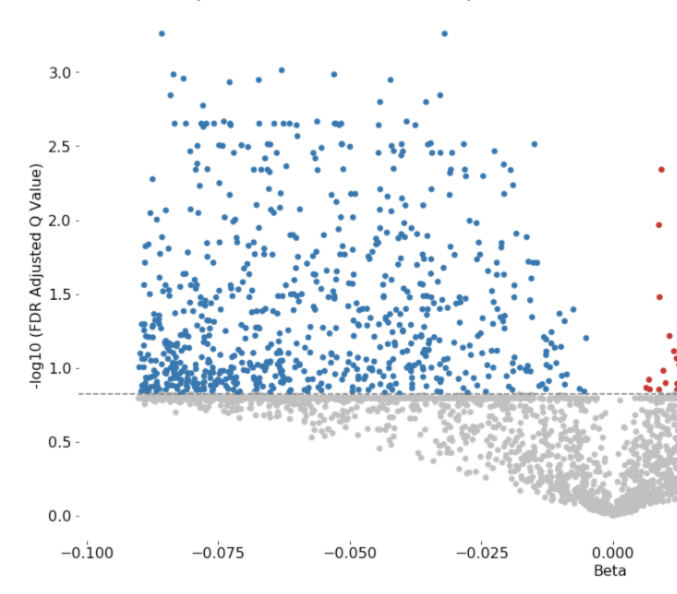


10 -



# 1.5.2 Sample Volcano Plot

Excluded 1128 probes outside of the specified beta coef



Customizable: Plot size, color palette, and cutoff p-value lines can be set by the user. Exporting: You can export all probe statistics, or just the significant probes as CSV or python pickled DataFrame.

# 1.6 Differentially methylated position/probe (DMP) detection

The diff\_meth\_pos(meth\_data, phenotypes) function searches for individual differentially methylated positions/probes (DMPs) by regressing methylation beta values or M-values for each sample at a given genomic location against the phenotype data for those samples.

### 1.6.1 Phenotypes

#### Can be provided as

```
- a list of strings,
- integer binary data,
- numeric continuous data
- pandas Series, DataFrame or numpy array
```

Only 2 phenotypes are allowed with logistic regression. Use Linear regression with numeric (measurement) phenotypes like age or time. You can pass in the GEO meta DataFrame associated with a dataset along with column=<columnname> kwarg. The order of the items in the phenotype should match the order of samples in the beta values or M-values.

Covariates are also supported for logistic (but not linear) regression. Pass in covariates=True to treat all other columns in a phenotype DataFrame as covariates, or pass in a list of column names to specify specific parts of the DataFrame. Note that supplying too many covariates for small sample sizes will lead to most probes failing with Linear Algebra or Perfect Separation errors.

For details on all of the other adjustable input parameters, refer to the API for diff\_meth\_pos()

#### 1.6.2 Returns

A pandas dataframe of regression statistics with one row for each probe and these columns:

```
- `Coefficient`: regression coefficient
- `StandardError`: standard error
- `95%CI_lower`: lower limit of the coefficient's 95% confidence interval
- `95%CI_upper`: upper limit of the coefficient's 95% confidence interval
- `PValue`: p-value: phenotype group A vs B - likelihood that the difference is_
- significant for this probe/location
- `Rsquared`: proportion (0 to 1) of probe variance explained by your phenotype.
- Linear Regression Only.
- `FDR_QValue`: p-values corrected for multiple comparisons using the Benjamini-
- Hochberg FDR method. The False Discovery Rate (FDR) corrected p-values will remain_
- comparable, regardless of the number of additional comparisons (probes) you include.

If a `q_cutoff` is specified, only probes with significant q-values less than the_
- cutoff will be returned in the DataFrame.
```

# 1.7 Differentially methylated regions (DMR)

Pass in your diff\_meth\_pos (DMP) stats results DataFrame as input, and it will calculate and annotate differentially methylated regions (DMR) using the combined-pvalues pipeline. This function returns list of output files.

```
    calculates auto-correlation
    combines adjacent p-values
    performs false discovery rate (FDR) adjustment
    finds regions of enrichment (i.e. series of adjacent low p-values)
    assigns significance to those regions
    annotates significant regions with possibly relevant nearby Genes, using the UCSC Genome Browser Database
    annotates candidate genes with expression levels for the sample tissue type,
```

(continues on next page)

```
if user specifies the sample tissue type.
- returns everything in a CSV that can be imported into other Genomic analysis_
→packages.
```

For more details on customizing the inputs and outputs, see API for the diff\_meth\_regions(stats, array\_type) function.

# 1.8 Loading processed data

Assuming you previously used methylprep to process a data set like this:

```
python -m methylprep -v process -d GSE130030 --betas
```

This creates two files, beta\_values.pkl and sample\_sheet\_meta\_data.pkl. You can load both in methylize like this:

Navigate to the folder where methylrep saved its processed files, and start a python interpreter:

```
>>>import methylcheck
>>>data, meta = methylcheck.load_both()
INFO:methylize.helpers:loaded data (485512, 14) from 1 pickled files (0.159s)
INFO:methylize.helpers:meta.Sample_IDs match data.index (OK)
```

Or if you are running in a notebook, specify the path:

```
import methylcheck
data, meta = methylcheck.load_both('<path_to...>/GSE105018')
```

This also validates both files, and ensures that the Sample\_ID column in meta DataFrame aligns with the column names in the data DataFrame.

# 1.8.1 Differentially methylated regions (DMR) in methylize

Adopted from the "combined-pvalues" package by Brend Pedersen et al, 2013: Comb-p: software for combining, analyzing, grouping and correcting spatially correlated P-values doi: 10.1093/bioinformatics/bts545

### What are Differentially methylated regions (DMRs)?

Genomic regions where DNA methylation levels differ between two groups of samples. DNA methylation is associated with cell differentiation, regulation, and proliferation, so these regions indicate that nearby genes may be involved in transcription regulation. There are several different types of DMRs. These include:

- tissue-specific DMR (tDMR),
- cancer-specific DMR (cDMR),
- development stages (dDMRs),
- reprogramming-specific DMR (rDMR),
- allele-specific DMR (AMR),
- aging-specific DMR (aDMR).

#### How to run DMR

First, assuming you have processed data using methylprep, use methylize to convert a dataframe of beta or M-values into differentially-methylated-probe (DMP) statistics, using methylize.diff\_meth\_pos. You will need to provide the data along with a list of sample labels for how to separate the data into two (treatment / control) groups or more groups (levels of a phenotypic characteristic, such as age or BMI):

```
meth_data = pd.read_pickle('beta_values.pkl')
phenotypes = ["fetal", "fetal", "adult", "adult", "adult"]
test_results = methylize.diff_meth_pos(meth_data, phenotypes)
```

phenotypes can be a list, numpy array, or pandas Series (Anything list-like). The results will be a dataframe with p-values, a measure of the likelihood that each probe is significantly different between the phenotypic groups. The lower the p-value, the more likely it is that groups differ:

IlmnID Coef	ficient	StandardError	PValue	95%CT lower	95
→%CI_upper Rsquared FDR_QValue		50411441451101	1,4140	70 001 <u>-</u> 10.101	30
cg04680738_II_R_C_rep1_EPIC -		0.001627	1.171409e-06	-0.992267	
→-0.992168 0.984496 0.009706					
cg18340948_II_R_C_rep1_EPIC	0.10475	0.005297	1.084911e-06	0.992256	
→ 0.992570 0.984887 0.009706					_
cg03681905_II_F_C_rep1_EPIC -	0.04850	0.002141	4.843320e-07	-0.994254	
→-0.994157 0.988444 0.009706					
cg01815889_II_F_C_rep1_EPIC -	0.05075	0.002735	1.579625e-06	-0.991492	
→-0.991308 0.982875 0.009816					
cg05995891_II_R_C_rep1_EPIC -	0.04050	0.002407	2.812417e-06	-0.989670	<u>.</u>
→-0.989474 0.979254 0.013981					
•••					ш
··· · · · · · · · · · · · · · · · · ·					
cg05855048_II_R_C_rep1_EPIC		0.016362	9.883052e-01	-0.025827	ш
→ 0.038289 0.000039 0.999077					
cg23130711_II_R_C_rep1_EPIC		0.016530	9.884235e-01	-0.026217	ш
→ 0.038553 0.000038 0.999156					
cg10163088_II_F_C_rep1_EPIC -		0.017168	9.888530e-01	-0.039573	ш
→ 0.027696 0.000035 0.999550					
cg04079257_II_F_C_rep1_EPIC -		0.017430	9.890214e-01	-0.039997	ш
→ 0.028300 0.000034 0.999679					
cg24902557_II_F_C_rep1_EPIC		0.017533	9.890857e-01	-0.028535	ш
→ 0.040163 0.000034 0.999704					

The FDR\_QValue is the key result. FDR\_Q is the "False Discovery Rate Q-value": The adjustment corrects individual probe p-values for the number of repeated tests (once for each probe on the array).

Next, you take this whole dataframe output from diff\_meth\_pos and feed it into the DMR function, diff\_meth\_regions, along with the type of methylation array you are using:

This will run a while. It compares all of the probes and clusters CpG probes that show a difference together if they are close to each other in the genome sequence. There are a lot of adjustable parameters you can play with in this function. Refer to the docs for more details.

When it completes, it returns a list of files that have been saved to disk:

```
docs/research_notebooks/dmr.acf.txt
docs/research_notebooks/dmr.args.txt
```

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```
docs/research_notebooks/dmr.fdr.bed.gz
docs/research_notebooks/dmr.manhattan.png
docs/research_notebooks/dmr.regions-p.bed.gz
docs/research_notebooks/dmr.slk.bed.gz
docs/research_notebooks/dmr_regions.csv
docs/research_notebooks/dmr_regions_genes.csv
docs/research_notebooks/dmr_stats.csv
docs/research_notebooks/dmr_stats.csv
docs/research_notebooks/stats.bed
```

Most of these are intermediate processing files that might be useful when imported into other tools, but the main summary file is the one that ends in regions genes.csv:

		art chromEnd descriptions	min_p	z_sidak_	p name	genes_	
21	2535657	2535711	6.662000e-12 .	3.874000e	e-08 cg06415891	VLDLR-AS1	u
$\hookrightarrow$	3 Homo	sapiens VLDI	R antisense RNA	1 (VLDLR-AS1	.)		
22	2535842	2535892	2.816000e-04 .	8.913000e	e-01 cg12443001	HCG22	ш
$\hookrightarrow$	16 Homo	sapiens HLA	complex group 22	2 (HCG22),	•		
25	2577833	2577883	2.048000e-11 .	1.347000e	e-07 cg22948959	HLA-C	
$\hookrightarrow$	33 Homo	sapiens majo	r histocompatib	ility compl			
1	876868	876918	0.00303	1.0	cg05475702		
1	1514374	1514424	0.003309	1.0	cg00088251		

This will reveal clusters of CpG probes that were significantly different and annotate these clusters with one or more nearby genes using the UCSC Genome Browser database. Note the CSV file column headings: genes, distances, descriptions. In some cases, a single diff-meth-region can align with multiple genes. In that case you will be a comma separated list for these fields. The descriptions will be separated by a pipe "|" symbol. The distances are the number of base-pairs of separation between the start of the CpG probe and the gene coding start sequence. Only rows with significant region p-values will have annotation.

If you pass in the tissue=<your tissue type> argument into the diff\_meth\_regions function, and that tissue type is one of the 54 that are part of the GTEx (genome expression levels in humans by tissue dataset), this file will also include a column showing the expression levels for any genes that match, so that you can further narrow down the search for relevant genomic interactions within each experiment.

There are a lot of additional corrections that researchers make at this stage, and many of them are beyond the scope of methylsuite, but this function should get you started. And you can export the output from this function into more sophisticated analysis tools.

#### Gene annotation with UCSC Genome Browser

University of California Santa Cruz maintains a large database of every version of the human genome and its meta data at https://genome.ucsc.edu/cgi-bin/hgTables. You can browse these database tables.

If you are using the latest genome build (hg38), diff\_meth\_regions will annotate your database using the refGene table. It also (partially) supports the knownGene and ncbiRefSeq tables, if you want to use those. This is useful for identifying genes that are nearby your regions of interest, and noting the tissue specificity of those genes, in exploring your data.

### 1.8.2 Methylize Walkthrough

```
[1]: #Install joblib module for parallelization
    import sys
    !conda install --yes --prefix {sys.prefix} joblib
    Collecting package metadata (current_repodata.json): done
    Solving environment: done
    # All requested packages already installed.
[2]: import methylize
    import methylcheck
    import numpy as np
    import pandas as pd
    import matplotlib.pyplot as plt
    from statsmodels.stats.multitest import multipletests
```

#### **Differentially Methylated Position Analysis with Binary Phenotypes**

DMPs are **probes** where methylation levels differ between two groups of samples, while DMRs are **genomic regions** where DNA methylation levels differ between two groups of samples.

Beta or m-values can be used, but the format needs to be samples as rows and probes as columns.

```
[3]: betas, meta = methylcheck.load_both('../data/asthma') #load in the beta values and
    ⊶metadata
    betas = betas.T
    print (meta.shape)
    meta.head()
    INFO:methylcheck.load_processed:Found several meta_data files; attempting to match_
    →each with its respective beta_values files in same folders.
    WARNING: methylcheck.load_processed: Columns in sample sheet meta data files does not_
    →match for these files and cannot be combined:['../data/asthma/sample_sheet_meta_
    data.pkl', '.../data/asthma/GPL13534/GSE157651_GPL13534_meta_data.pkl']
    INFO:methylcheck.load_processed:Multiple meta_data found. Only loading the first file.
    INFO:methylcheck.load_processed:Loading 40 samples.
    Files: 100%|| 1/1 [00:00<00:00, 8.94it/s]
    INFO:methylcheck.load_processed:loaded data (485512, 40) from 1 pickled files (0.118s)
    INFO:methylcheck.load_processed:Transposed data and reordered meta_data so sample_
    ⇔ordering matches.
    INFO:methylcheck.load_processed:meta.Sample_IDs match data.index (OK)
    (40, 19)
[3]:
                       tissue disease smoking_status passage Sex age
    2.0
             Airway Fibroblast Healthy
                                                           4 M 65
                                                   Ex
    38 Parenchymal Fibroblast Asthmatic
                                                             4 F 19
                                                   Non
                                                             4 F 14
    37 Parenchymal Fibroblast Asthmatic
                                                   Non
    32 Parenchymal Fibroblast Asthmatic
                                                             4 F 15
                                              Current
             Airway Fibroblast Asthmatic
                                                             4 F
    2.3
                                                   Non
                                               Sample_ID Sentrix_ID
                            description
    20 fun2norm normalized average beta 9976861129_R03C01 9976861129
    38 fun2norm normalized average beta 9976861129_R04C02 9976861129
    37 fun2norm normalized average beta 9976861129_R05C01 9976861129
    32 fun2norm normalized average beta 9976861129_R06C02 9976861129
```

(continues on next page)

```
fun2norm normalized average beta 9976861137_R06C01 9976861137
  Sentrix_Position Sample_Group
                                 Sample_Name Sample_Plate Sample_Type
20
            R03C01
                                      HAF091
                                                    None
                          None
                                                             Unknown
                                  7294_P4_LG
38
            R04C02
                          None
                                                    None
                                                             Unknown
37
            R05C01
                          None
                                  7291_lg_p4
                                                    None
                                                             Unknown
32
            R06C02
                          None
                               7188_lg_p4_R
                                                    None
                                                             Unknown
23
            R06C01
                          None
                                  7016_BR_P4
                                                    None
                                                             Unknown
  Sub_Type Sample_Well Pool_ID
                                   GSM_ID Control
20
                 None
                         None GSM4772065 False
      None
38
      None
                  None
                         None GSM4772083 False
37
      None
                 None None GSM4772082 False
32
      None
                 None None GSM4772077 False
23
      None
                 None None GSM4772068 False
```

### Get binary phenotype

The phenotypes can be anything list-like (i.e. a list, numpy array, or Pandas series).

When using Logistic Regression DMP, use phenotype data that is binary (has only 2 classes). In this example, we will take disease state as our phenotype because there are only 2 classes, Asthmatic and Healthy. The DMP step will automatically convert your string phenotype to 0 and 1 if there are not in that format already.

#### **DMP Function**

This function searches for individual differentially methylated positions/probes (DMPs) by regressing the methylation beta or M-value for each sample at a given genomic location against the phenotype data for those samples. In this first example, we are using the argument regression\_method="logistic" because we are using binary phenotypes.

impute argument:

- Default: 'delete' probes if any samples have missing data for that probe.
- True or 'auto': if <30 samples, deletes rows; if >=30 samples, uses average.
- False: don't impute and throw an error if NaNs present 'average' use the average of probe values in this batch 'delete' drop probes if NaNs are present in any sample
- 'fast' use adjacent sample probe value instead of average (much faster but less precise)

Note: if you want to analyze every probe in your betas dataframe, remove the .sample(N). What this does is randomly samples N probes from your dataframe and decreases the time it takes to complete this analysis.

```
[5]: test_results = methylize.diff_meth_pos(betas.sample(20000, axis=1), pheno_data,...
    →regression_method="logistic", export=False)
    print(test_results.shape)
    test_results.head()
    WARNING:methylize.diff_meth_pos:Dropped 2902 probes with missing values from all_
    All samples with the phenotype (Asthmatic) were assigned a value of 0 and all samples_
    →with the phenotype (Healthy) were assigned a value of 1 for the logistic regression.
    ⇒analysis.
      0%|
                  | 0/17098 [00:00<?, ?it/s]
    (17098, 6)
             Coefficient StandardError PValue 95%CI_lower 95%CI_upper
[5]:
    cg22048228 -0.320912 0.654584 0.623955 -0.962050 1.603874
    cg19366543
                0.048680
                               1.217084 0.968095
                                                   -2.434121
                                                                  2.336761
                                1.835602 0.969570
               -0.070025
    cg12408293
                                                    -3.527689
                                                                 3.667739
    cg08640790 -0.043054
                               2.412119 0.985759
                                                     -4.684612
                                                                  4.770720
                                                     -3.675343
    cg02835494
                 0.027930
                                1.860959 0.988025
                                                                  3.619483
              FDR_QValue
    cg22048228 0.999978
    cg19366543 0.999978
    cg12408293 0.999978
    cg08640790 0.999978
    cq02835494 0.999978
```

#### Differentially Methylated Positions Analysis with Continuous Numeric Phenotypes

#### **Get Continuous Numeric Phenotype**

In order to find DMPs against a numeric continuous phenotype, the linear regression method must be used. In this example, we are using age as the continuous numeric phenotype.

```
[6]: pheno_data = meta.age.astype(int)
     print (pheno_data.shape)
     (pheno_data.sort_values()).hist()
     plt.xlabel('Age (years)'); plt.ylabel('Frequency')
     (40,)
[6]: Text(0, 0.5, 'Frequency')
        8
        6
      Frequency
4
                                        50
                                              60
                                                           80
              10
                    20
                          30
                                 40
                                                     70
                                  Age (years)
```

#### **DMP Function**

In this second example, we are using the argument regression\_method="linear" because we are using continuous numeric phenotypes.

```
[7]: test_results2 = methylize.diff_meth_pos(betas, pheno_data, regression_method="linear",
     → export=False)
    print(test_results2.shape)
    test results2.head()
    WARNING: methylize.diff_meth_pos:Dropped 71139 probes with missing values from all_
    →samples
      0%|
                   | 0/414373 [00:00<?, ?it/s]
    (414373, 7)
[7]:
                Coefficient StandardError
                                                 PValue 95%CI_lower \
    cg03555227
                 0.004583 0.000380 1.453304e-14
                                                           0.890321
    cg16867657
                  0.008122
                                 0.000718 9.888059e-14
                                                           0.877810
    cg11169102
                 -0.004757
                                 0.000460 1.323437e-12
                                                          -0.859258
                 0.009223
                                 0.000903 1.871221e-12
                                                           0.855763
    cq16909962
    cg23045594
                   0.005227
                                 0.000511 1.856802e-12
                                                           0.856031
                95%CI_upper Rsquared
                                       FDR_QValue
                  0.890629 0.792946 6.022099e-09
    cg03555227
    cg16867657
                  0.878454 0.771117
                                      2.048672e-08
                -0.858785 0.737918 1.550767e-07
    cq11169102
    cg16909962
                  0.856707 0.733140 1.550767e-07
    cq23045594
                  0.856566 0.733247 1.550767e-07
```

#### To BED

This function converts the stats dataframe from the DMP function into BED format. For more information regarding this function and its arguments, please view the API Reference.

```
[8]: bed = methylize.to_BED(test_results2, manifest_or_array_type='450k', save=False,_
    →filename='test_bed.bed')
    bed.head()
    INFO:methylprep.files.manifests:Reading manifest file: HumanMethylation450k_15017482_
[8]:
      chrom chromStart chromEnd
                                    pvalue
                                                  name
          1
               69591.0
                        69641.0 0.060267 cg21870274
               864703.0 864753.0 0.164362 cg08258224
    1
          1
    2
               870161.0 870211.0 0.473560 cg16619049
          1
               877159.0 877209.0 0.267582 cg18147296
    3
          1
               898803.0 898853.0 0.283629 cg13938959
    4
          1
```

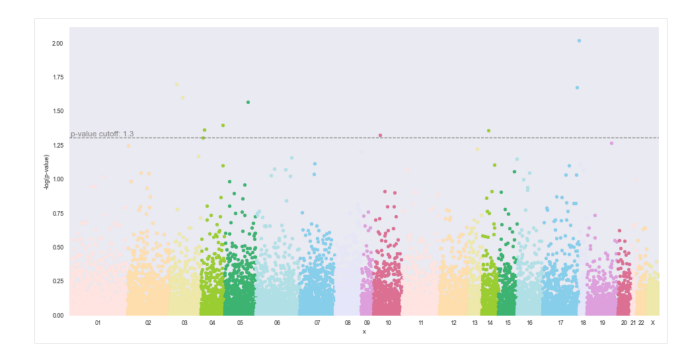
#### **Manhattan Plots**

After we run the DMP analysis, we can move on to visualizing the results. One way to do this is a Manhattan plot.

A Manhattan plot has probe position on the x-axis (grouped and colored by chromosome) and  $-\log(p-value)$  on the y-axis. Because of the scale on the y-axis, the greater the probe is on the y-axis, the more significant that probe is to the phenotype that was used. It is common to have a cutoff on Manhattan plots where we set our  $\alpha$ , which commonly is 0.05. Other common  $\alpha$  are 0.01 and 0.001. The lower your  $\alpha$  is, the more significant probes are that are above that cutoff on the Manhattan plot. The cutoff is for the p-values, meaning the probe is statistically significant if the p-value <  $\alpha$ .

Because there are a high number of probes, a p-value correction is needed to prevent false positives. The Manhattan plot function automatically applies a Bonferroni correction to the cutoff to lower the  $\alpha$  (increase the dotted cutoff line on the plot) to a more conservative value. The post-test correction is controlled by the adjust argument, where "bonferroni" is the default, and other options are "fdr" for an FDR correction, and None for no post-test correction to keep the cutoff the same as is entered in the cutoff argument.

#### **Manhattan Plot for Binary Phenotypes**



### **Find Significant Probes**

To see which probes are significant (above the cutoff line in the Manhattan plot), filter the results dataframe by the p-value. Because we set post\_test=None in the arguments, this means the p-value cutoff that was set is correct, and there was no correction to this value. The next DMR example will have a p-value correction.

interesting interesting	_	st_results[test_	results['F	Val	ue'] <= 0.	05]	
	Coefficient	StandardError	PValue	95	%CI_lower	95%CI_upper	\
cg11703729	-1.403577	0.708667	0.047638		0.014614	2.792539	
cg01096199	-1.518270	0.754115	0.044082		0.040232	2.996308	
cg14543255	-1.834690	0.796496	0.021254		0.273586	3.395794	
cg26494916	1.621125	0.733396	0.027075		-3.058554	-0.183696	
cg27196131	1.991505	0.985877	0.043380		-3.923787	-0.059222	
cg07351758	1.814658	0.810237	0.025113		-3.402694	-0.226623	
cg23103043	1.830804	0.707215	0.009632		-3.216920	-0.444688	
cg25314266	1.448918	0.706773	0.040360		-2.834168	-0.063668	
cg07538039	1.761092	0.897471	0.049730		-3.520103	-0.002081	
cg21664636	-1.624543	0.698487	0.020029		0.255533	2.993553	
	FDR_QValue	minuslog10pvalu	e chromoso	ome	M	APINFO	
cg11703729	0.999978	1.32204	9	10	CHR-1380	9615.0	
cg01096199	0.999978	1.35574	3	14	CHR-8972	4701.0	
cg14543255	0.999978	1.67256	9	17	CHR-900	2458.0	
cg26494916	0.999978	1.56743	4	05	CHR-14945	3321.0	
cg27196131	0.999978	1.36271	2	04	CHR-18717	6529.0	
cg07351758	0.999978	1.60010	5	03	CHR-5976	6624.0	
cg23103043	0.999978	2.01626	4	17	CHR-7165	0582.0	
cg25314266	0.999978	1.39405	0	04	CHR-7618	8042.0	
cg07538039	0.999978	1.30338	5	04	CHR-1537	4511.0	
cq21664636	0.999978	1.69833	8	03	CHR-7038	3513.0	

#### **Manhattan Plot for Continuous Numeric Phenotypes**

For this example, there will be a Benferoni post-test correction. The adjusted p-value cutoff is displayed on the horizontal cutoff line as well as printed above the plot. Please note that this value is the <code>-log(p-value)</code> of the p-value cutoff.

```
[11]: methylize.manhattan_plot(test_results2, cutoff=0.05, palette='default', save=False,...
      →array_type='450k', verbose=True)
      INFO:methylprep.files.manifests:Reading manifest file: HumanMethylation450k_15017482_
      -v3.csv
      Total probes to plot: 414373
      01 40528 | 02 30059 | 03 21877 | 04 17328 | 05 20851 | 06 30620 | 07 24564 | 08 17814.
      → | 09 8501 | 10 20863 | 11 25272 | 12 21229 | 13 10370 | 14 12956 | 15 12921 | 16.
      →18404 | 17 24069 | 18 5215 | 19 21756 | 20 9259 | 21 3329 | 22 7167 | X 9419 | Y 2
      p-value cutoff adjusted: 1.3010299956639813 ==[ Bonferoni ]==> 6.9184214452654755
          p-value cutoff, controlling for FDF
                                  05
                                                     09
                                                        10
                                                                      13
                                                                        14
                                                                                            20 21 22 X Y
```

### **Find Significant Probes**

For this Manhattan plot, there has been a p-value cuttoff correction, meaning that the cutoff line is at a more conservative value (lower p-value, higher cutoff line) to account for false positives. Because of this correction, when filtering the results dataframe to find significant probes, you need to filter by the Bonferoni (or FDR) adjust p-value to find the probes above the cutoff line on the Manhattan plot.

```
[12]: adjusted = multipletests(test_results2.PValue, alpha=0.05)
     pvalue_cutoff_y = -np.log10(adjusted[3])
     interesting_probes2 = test_results2[test_results2['minuslog10pvalue'] >= pvalue_
      →cutoff_y] #bonferoni correction for cutoff
     interesting_probes2
                                                    PValue 95%CI_lower
[12]:
                 Coefficient StandardError
     cg03555227
                    0.004583
                                  0.000380 1.453304e-14
                                                               0.890321
     cq16867657
                    0.008122
                                    0.000718 9.888059e-14
                                                               0.877810
                                                                                (continues on next page)
```

					(continued fro	m previous page)
cg11169102	-0.004757	0.00	00460 1.32343	7e-12 -0.859258		
cg16909962	0.009223	0.00	00903 1.871223	Le-12 0.855763		
cg23045594	0.005227	0.00	00511 1.856802	2e-12 0.856031		
cg04091914	-0.003923	0.00	00604 1.185223	Be-07 -0.726027		
cg25874108	-0.002309		00355 1.190858			
cg01235463	-0.002696		00415 1.188893			
cg11126313	-0.001694		00261 1.199705			
cg04161137	-0.003023	0.00	00465 1.199026	Se-07 -0.725704		
	95%CI_upper	Rsquared	FDR_QValue	minuslog10pvalue	ah romo aomo	\
cg03555227	0.890629	0.792946	6.022099e-09	13.837644	05	\
cg16867657	0.878454	0.771117	2.048672e-08	13.004889		
cg11169102	-0.858785	0.737918	1.550767e-07	11.878297		
cg16909962	0.856707	0.733140	1.550767e-07	11.727875		
cg23045594	0.856566	0.733247	1.550767e-07	11.731235	02	
cg04091914	-0.724906	0.526302	6.947979e-05	6.926200	02	
cg25874108	-0.725057	0.526186	6.947979e-05	6.924140	17	
cg01235463	-0.725029	0.526227	6.947979e-05	6.924857	10	
cg11126313	-0.725020	0.526006	6.972307e-05	6.920925	03	
cg04161137	-0.724839	0.526020	6.972307e-05	6.921172	01	
	MAPI	NEO				
cg03555227	CHR-17086206					
cg16867657	CHR-17080200					
cg11169102	CHR-17085883					
cg16909962	CHR-22927096					
cg23045594	CHR-7127675					
cg04091914	CHR-22768682	2.0				
cg25874108	CHR-7489681					
cg01235463	CHR-9115401					
cg11126313	CHR-4624277	1.0				
cg04161137	CHR-2264768	7.0				

#### **Volcano Plot**

[713 rows x 10 columns]

Logistic Regression Volcano Plots are under construction

Below shows how to create a Volcano plot using the linear regression DMR analysis.

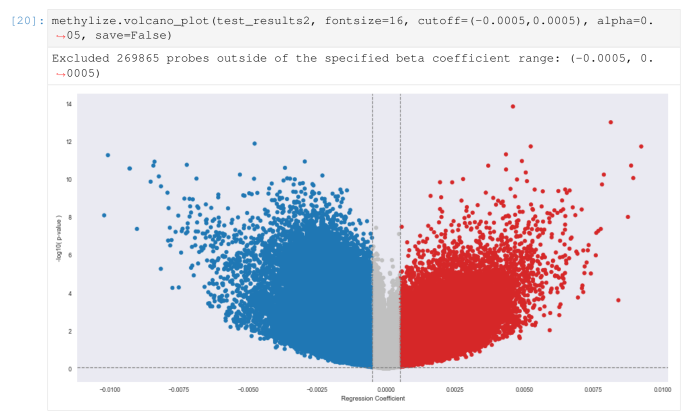
Positive correlation (hypermethylated) in red. Negative correctation (hypomethylated) in blue

A higher | Regression (beta) coefficient | means that that specific probe is more significant in predicting if the probe is methylated (positive) vs not methylated (negative). The non-gray probes are the probes that are statistically significant, and have a higher absolute value of the regression coefficient as the cutoff. The red probes further to the right on the plot are hypermethylated with increase in age, and the opposite is true for the blue probes.

#### **Useful Arguments:**

- alpha: Default: 0.05 alpha level The significance level that will be used to highlight the most significant adjusted p-values (FDR Q-values) on the plot. This is the horizontal cutoff line you see on the plot.
- cutoff: Default: No cutoff format: a list or tuple with two numbers for (min, max) If specified in kwargs, will exclude values within this range of regression coefficients from being "significant" and put dotted vertical lines

on chart. These are the vertical cutoff lines you see on the plot. These cutoffs are dependent on the study and up to the researcher in choosing which cutoff is the most desirable.



### **Find Significant Probes**

```
[21]: interesting_probes3 = test_results2[(test_results2['FDR_QValue'] <= 0.05) & (np.
      →abs(test_results2['Coefficient']) > 0.0005)]
     interesting_probes3
                 Coefficient StandardError
                                                   PValue 95%CI lower \
[21]:
     cq03555227
                    0.004583
                              0.000380 1.453304e-14
                                                              0.890321
     cg16867657
                    0.008122
                                   0.000718 9.888059e-14
                                                              0.877810
     cg11169102
                   -0.004757
                                   0.000460 1.323437e-12
                                                             -0.859258
     cq16909962
                    0.009223
                                   0.000903 1.871221e-12
                                                              0.855763
                    0.005227
                                   0.000511 1.856802e-12
                                                              0.856031
     cg23045594
     cg18227216
                   -0.000972
                                   0.000331
                                                             -0.430024
                                            5.680062e-03
     cg16297435
                    0.001548
                                   0.000528 5.681080e-03
                                                              0.428642
     cq05515570
                   -0.000716
                                   0.000244
                                             5.681038e-03
                                                             -0.429877
     cg13346820
                   -0.001955
                                   0.000667
                                             5.680922e-03
                                                             -0.430553
     cg12773197
                   -0.001329
                                   0.000453 5.681300e-03
                                                             -0.430209
                 95%CI_upper Rsquared
                                         FDR_QValue minuslog10pvalue chromosome
     cg03555227
                    0.890629 0.792946 6.022099e-09
                                                             13.837644
                                                                               0.5
                                                                               06
     cg16867657
                    0.878454 0.771117 2.048672e-08
                                                             13.004889
     cg11169102
                   -0.858785 0.737918 1.550767e-07
                                                             11.878297
                                                                               05
     cg16909962
                    0.856707 0.733140 1.550767e-07
                                                             11.727875
                                                                               01
     cg23045594
                    0.856566 0.733247 1.550767e-07
                                                             11.731235
                                                                               02
                                                                              (continues on next page)
```

```
cg18227216 -0.428965 0.184466 4.997593e-02
                                                      2.245647
                                                                      17
cq16297435
            0.430330 0.184459
                                 4.997954e-02
                                                      2.245569
                                                                      06
            -0.429096 0.184459 4.997954e-02
                                                                      10
cg05515570
                                                      2.245572
             -0.428421 0.184460
                                                                      05
cg13346820
                                 4.997954e-02
                                                      2.245581
cg12773197
             -0.428760 0.184457
                                 4.998041e-02
                                                      2.245552
                                                                      13
                  MAPINFO
cg03555227 CHR-170862066.0
cg16867657 CHR-11044644.0
cg11169102 CHR-170858836.0
cg16909962 CHR-229270964.0
cg23045594 CHR-71276753.0
cq18227216 CHR-10225251.0
cg16297435 CHR-43060558.0
cg05515570 CHR-28125056.0
cg13346820 CHR-865203.0
cg12773197 CHR-111586326.0
[43781 rows x 10 columns]
```

#### Differentiated Methylized Regions (DMR) Analysis

DMRs are **genomic regions** where DNA methylation levels differ between two groups of samples, while DMPs are **probes** where methylation levels differ between two groups of samples. DMR looks at clusters or adjacent probes, and if the whole cluster shows the same direction of effect between groups, then it is more significant (likely to be meaningful).

We have already ran methylize.diff\_meth\_pos with the continuous numeric phenotypes, and the statistics for that fuction is stored in the dataframe test\_restuls2. This is what we will use in the DMR function below.

This following step should create these files: \* dmr.acf.txt

- · dmr.args.txt
- · dmr.fdr.bed.gz
- · dmr.manhattan.png
- · dmr.regions-p.bed.gz
- · dmr.slk.bed.gz
- · dmr\_regions.csv
- dmr\_regions\_genes.csv
- dmr\_stats.csv
- · stats.bed

This could run for a while. This function compares all of the probes and clusters CpG probes that show a difference together if they are close to each other in the genomic sequence.

There are many adjustible parameters for the following function, and you can refer to the API Reference for more information.

```
[15]: files_created = methylize.diff_meth_regions(test_results2, '450k', prefix='../data/
→asthma/dmr/')
```

```
INFO:methylprep.files.manifests:Reading manifest file: HumanMethylation450k_15017482_
⇒v3.csv
INFO:methylprep.files.manifests:Reading manifest file: HumanMethylation450k_15017482_
→v3.csv
/Users/jaredmeyers/opt/anaconda3/lib/python3.8/site-packages/methylize/diff_meth_
→regions.py:150: DtypeWarning: Columns (0) have mixed types.Specify dtype option on_
→import or set low_memory=False.
 results = _pipeline(kw['col_num'], kw['step'], kw['dist'],
Calculating ACF out to: 453
with 17 lags: [1, 31, 61, 91, 121, 151, 181, 211, 241, 271, 301, 331, 361, 391, 421,...

→451, 481]

18581032 bases used as coverage for sidak correction
INFO: methylize.diff_meth_regions: wrote: ../data/asthma/dmr/.regions-p.bed.gz,...
\hookrightarrow (regions with corrected-p < 0.05: 29)
/var/folders/3k/vrkjxj8s7l1gq4x6n7ymqncw0000gn/T/ipykernel_10952/1491901786.py:1:..
→DtypeWarning: Columns (0) have mixed types. Specify dtype option on import or set,
→low_memory=False.
 files_created = methylize.diff_meth_regions(test_results2, '450k', prefix='../data/
→asthma/dmr/')
INFO:methylize.genome_browser:Loaded 4802 CpG regions from ../data/asthma/dmr/_
⇒regions.csv.
INFO:methylize.genome_browser:Using cached `refGene`: /Users/jaredmeyers/opt/
→anaconda3/lib/python3.8/site-packages/methylize/data/refGene.pkl with (135634) genes
Mapping genes: 100%|| 135634/135634 [00:30<00:00, 4381.01it/s]
INFO:methylize.genome_browser:Wrote ../data/asthma/dmr/_regions_genes.csv
```

Many of the files created by this function can be useful when using other tools. The file with the main summary of what was just done is found in dmr\_regions\_genes.csv.

```
[16]: regions_genes = pd.read_csv('../data/asthma/dmr_regions_genes.csv')
     regions_genes.head()
        Unnamed: 0 chrom chromStart
[16]:
                                      chromEnd
                                                 min_p n_probes
                                                                      z_p \
     0
                0
                     1
                           22289407
                                      22289457 0.03571
                                                        1 0.000094
                1
                       1
                           34757646
                                      34757696 0.03110
     1
                                                              1
                                                                 0.000054
                                      34761156 0.03401
                                                             1 0.000081
     2
                2
                      1
                           34761106
                                                             1 0.000038
     3
                         202858152 202858202 0.03110
                3
                      1
     4
                4
                      10
                           22334619
                                     22334669 0.03110
                                                             1 0.000038
        z_sidak_p
                       name genes distances
     0
           0.7979 cg02053477
                             NaN
     1
           0.6018 cg13170235 SNRPC
                                         142
     2
           0.7500 cg15750705
                             NaN
                                         NaN
          0.4790 cg15569630
     3
                               NaN
                                         NaN
          0.4790 cg13327545
     4
                               NaN
                                         NaN
                                           descriptions
     1
        Homo sapiens small nuclear ribonucleoprotein p...
     2.
     3
                                                   NaN
     4
                                                   NaN
```

This shows clusters of CpG probes that were significantly different and annotes these clusters with mne or more nearby genes using the UCSV Genome Browswer database.

Distances are the number of base-pairs that separate the start of the CpG probe and the start of the coding sequence of the gene.

Only the rows with signicicant p-values will have an annotation.

Adopted from the combined-pvalues package by Brend Pedersen et al, 2013: Comb-p: software for combining, analyzing, grouping and correcting spatially correlated P-values doi: 10.1093/bioinformatics/bts545

More information on DMR on this page

#### **Fetching Genes**

This function annotates the DMR region output file using the UCSC Genome Browser database as a reference to what genes are nearby. For more information on this function and the specific argument, please refer to the API Reference

This function can now accept either the filepath or the DMR dataframe output.

```
[17]: reg = pd.read_csv('../data/asthma/dmr_regions.csv')
     genes = methylize.fetch_genes('../data/asthma/dmr_regions.csv')
     genes.head(12)
     INFO:methylize.genome_browser:Loaded 60 CpG regions from ../data/asthma/dmr_regions.
     INFO:methylize.genome_browser:Using cached `refGene`: /Users/jaredmeyers/opt/
     →anaconda3/lib/python3.8/site-packages/methylize/data/refGene.pkl with (135634) genes
     Mapping genes: 100%|| 135634/135634 [00:28<00:00, 4768.33it/s]
     INFO:methylize.genome_browser:Wrote ../data/asthma/dmr_regions_genes.csv
[17]:
         chrom chromStart chromEnd
                                       min_p n_probes
                                                            z_p z_sidak_p \
     0
                            22289457 0.03571
           1
                 22289407
                                                    1 0.000094
                                                                   0.7979
                          34757696 0.03110
     1
            1
                 34757646
                                                    1
                                                       0.000054
                                                                    0.6018
                          34761156 0.03401
     2
            1
                34761106
                                                    1
                                                       0.000081
                                                                    0.7500
     3
            1
                202858152 202858202 0.03110
                                                    1
                                                       0.000038
                                                                   0.4790
                                                    1 0.000038
     4
           10
               22334619 22334669 0.03110
                                                                   0.4790
     5
           11
                 2170376 2170426 0.03110
                                                    1 0.000038
                                                                   0.4790
     6
           11
                 2170376 2170426 0.03110
                                                    1 0.000038
                                                                   0.4790
     7
           11
                 2346840 2346890 0.04207
                                                    1 0.000138
                                                                  0.9041
     8
           11
                61755269 61755319 0.04150
                                                   1 0.000133
                                                                  0.8968
     9
           11
                69403865 69403915 0.03110
                                                   1 0.000068
                                                                  0.6847
     10
           11 114063572 114063622 0.03110
                                                   1 0.000054
                                                                  0.6018
           11
               117861576 117861626 0.03110
                                                   1 0.000038
                                                                   0.4790
     11
              name genes distances \
     0
        cg02053477
         cg13170235 SNRPC
                                142
     1
     2
        cq15750705
     3
        cg15569630
     4
        cg13327545
        cg11911653 VARS2 -165,-135
     5
     6
        cg14714364 VARS2 -165,-135
     7
        cg05532869
        cg11179625 MYRF
                                -108
        cq05582286 TACR2
                                -37
     10 cg10096321
     11 cg14848289
                                            descriptions
     0
     1
         Homo sapiens small nuclear ribonucleoprotein p...
     2
     3
     4
                                                                          (continues on next page)
```

```
Homo sapiens valyl-tRNA synthetase 2, mitochon...

Homo sapiens valyl-tRNA synthetase 2, mitochon...

Homo sapiens myelin regulatory factor (MYRF), ...

Homo sapiens tachykinin receptor 2 (TACR2), mR...
```

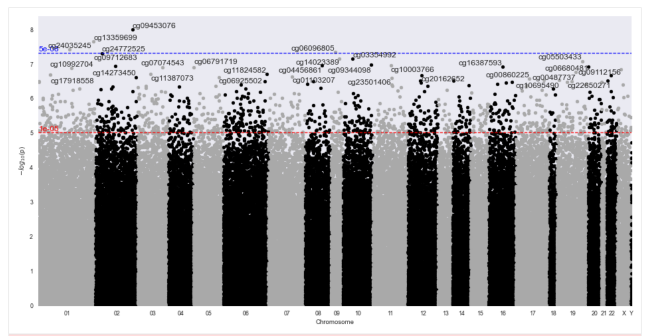
## 1.8.3 Example: Differentially methylated regions – GSE69852

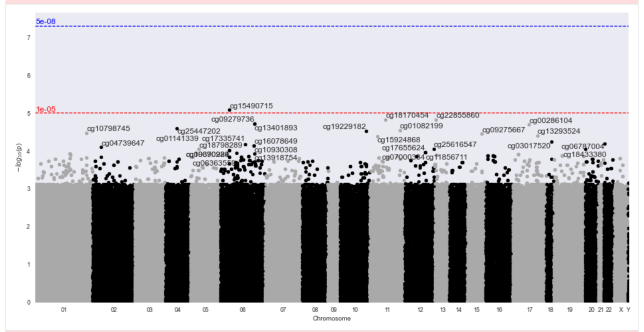
- Using GEO dataset GSE69852 (6 samples, 450k)
- Title: Patterns of DNA methylation in human fetal and adult liver
- https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE69852
- We demonstrate a basic (beta\_values + meta\_data) -> DMP -> DMR calculation.

Claims in the paper: - Nearly half (42%) of the CpGs in human liver show a significant difference in methylation comparing fetal and adult samples. - 69% of the significant sites differed in their mean methylation beta value by 0.2.

Note on phenotype: The original meta data had ages of samples in different units (e.g. 55yr, 21wk) so we manually converted these to a single measurement (converted\_age, float, years) so that linear regression could run. Alternatively, you could treat the two groups (55-56 years vs 20-22 weeks) as a phenotype with two groups (1 and 0) and apply logistic regression.

8 NaNs dropped





INFO:methylize.genome\_browser:Loaded 24371 CpG regions from data\_regions.csv. INFO:methylize.genome\_browser:Using cached `refGene`: /Users/mmaxmeister/methylize/

-methylize/data/refGene.pkl with (135634) genes

(continues on next page)

```
Mapping genes: 100%|| 135634/135634 [00:59<00:00, 2277.89it/s] INFO:methylize.genome_browser:Wrote data_regions_genes.csv
```

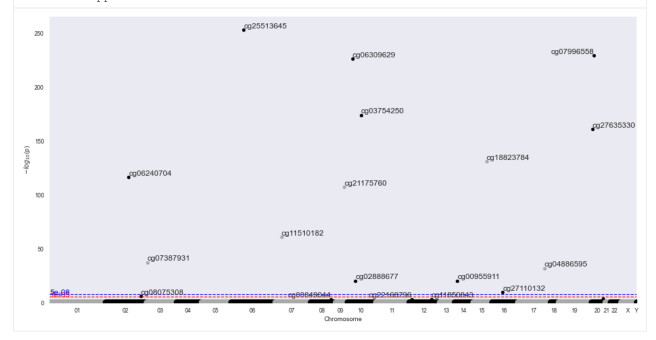
One would interpret the DMR plot to indicate that there were no significantly different methylated regions. Note, this example only ran 100000 of the 485,512 probes.

### 1.8.4 All data, logistic regression

```
[3]: result = m.diff_meth_pos(df, meta['description'])
    m.manhattan plot(result, '450k')
    bed_files = m.diff_meth_regions(result, '450k', prefix='data', plot=True)
    INFO:methylize.diff_meth_pos:Converted your beta values into M-values; (6, 485512)
    INFO: methylize.diff_meth_pos:Logistic regression: Phenotype (Normal Adult liver) was,
     →assigned to 0 and (Normal Fetal Liver) was assigned to 1.
                            | 0/485512 [00:00<?, ?it/s]
    Probes:
              0%1
    218806 probes failed the logistic regression analysis due to perfect separation and_
     \rightarrowcould not be included in the final results.
    13209 probes failed the logistic regression analysis due to LinearAlgebra error and_
     \rightarrowcould not be included in the final results.
    148 probes failed the logistic regression analysis due to other unexplained reasons
     →and could not be included in the final results.
    Linear Algebra and Perfect Separation errors occur when the dataset is (a) too small.

→to observe

    events with low probabilities, or has (b) too many covariates in the model, leading_
     →to individual cell
    sizes that are too small. Singular Matrix Errors occur when there is no variance in _
     \hookrightarrowthe predictors
     (probe values and covariates).
    27 NaNs dropped
```



```
INFO:methylprep.files.manifests:Reading manifest file: HumanMethylation450k_15017482_
     -v3.csv
    INFO:methylprep.files.manifests:Reading manifest file: HumanMethylation450k_15017482_
     ⇔v3.csv
    Calculating ACF out to: 50
    with 3 lags: [1, 31, 61]
    11192348 bases used as coverage for sidak correction
    INFO: methylize.diff_meth_regions: wrote: data.regions-p.bed.gz, (regions with,
     g25513645 g218907656 g231757666309629
g25513645 g218907656 g23754250 g21850843 g218823784
                                                                                 cg07996558
cg27635330
                  cg06240704
       40
                        _cg07387931
       30
                                                                              cg04886595
       20
                                                                    og00955911
                                                     cg02888677
    INFO:methylize.genome_browser:Loaded 17 CpG regions from data_regions.csv.
    INFO:methylize.genome_browser:Using cached `refGene`: /Users/mmaxmeister/methylize/
     →methylize/data/refGene.pkl with (135634) genes
    Mapping genes: 100%|| 135634/135634 [00:56<00:00, 2381.53it/s]
    INFO:methylize.genome_browser:Wrote data_regions_genes.csv
[4]: bed_files
[4]: ['data.args.txt',
     'data.acf.txt',
     'data.fdr.bed.gz',
     'data.slk.bed.gz',
     'data.regions-p.bed.gz',
     'data_regions.csv',
     'data_stats.csv',
     'data_regions_genes.csv',
     'data_dmp_stats.bed']
[5]: bed = pd.read_csv('data.regions-p.bed.gz', sep='\t')
    bed
[5]:
        #chrom
                   start
                                               min_p n_probes
                                  end
                                                                          z_p \
    0
            10
                  4034204
                             4034254 0.000000e+00
                                                      1 1.000000e-07
                                                             1 1.000000e-07
            10
                 31031409
    1
                             31031459 0.000000e+00
                                                            1 1.392000e-16
    2
            10 75210411
                             75210461 2.351000e-12
    3
                                                             1 1.000000e-07
            12 43840121
                             43840171 0.000000e+00
                                                                                (continues on next page)
```

```
12 124453780 124453830 0.000000e+00
                                                    1 1.000000e-07
5
       14
           37592783
                     37592833 2.598000e-12
                                                    1 1.641000e-16
6
       15
           78810879
                     78810929 0.000000e+00
                                                    1 1.000000e-07
                     29005911 6.351000e-24
7
       17
            29005861
                                                    1 3.510000e-28
8
        2
          200911124 200911174 0.000000e+00
                                                    1 1.000000e-07
                      1470918 0.000000e+00
9
       20
            1470868
                                                    1 1.000000e-07
10
       20
            58445879
                     58445929 0.000000e+00
                                                    1
                                                      1.000000e-07
                                                   1 1.900000e-33
11
        3 141066318 141066368 3.702000e-29
                                                   1 1.000000e-07
12
        6
            35150826
                     35150876 0.000000e+00
                                                  1 1.000000e-07
13
        7
           24854884
                     24854934 0.000000e+00
14
        8 144335305 144335355 0.000000e+00
                                                  1 1.000000e-07
15
        9 133804066 133804116 0.000000e+00
                                                  1 1.000000e-07
      z_sidak_p
   2.214000e-02
1
   2.214000e-02
   2.485000e-11
3
   2.214000e-02
4
   2.214000e-02
5
   2.485000e-11
6
   2.214000e-02
7
   7.857000e-23
8
   2.214000e-02
9
   2.214000e-02
10 2.214000e-02
11 4.253000e-28
12 2.214000e-02
13 2.214000e-02
14 2.214000e-02
15 2.214000e-02
```

#### 1.8.5 API Reference

methylize.diff_meth_pos(meth_data,	This function searches for individual differentially
pheno_data)	methylated positions/probes (DMPs) by regressing the
	methylation M-value for each sample at a given ge-
	nomic location against the phenotype data for those
	samples.
<pre>methylize.diff_meth_regions(stats,)</pre>	Calculates and annotates diffentially methylated regions
	(DMR) using the combined-pvalues pipeline and returns
	list of output files.
methylize.fetch_genes([dmr_regions_file,	find genes that are adjacent to significantly different
])	CpG regions provided.
<pre>methylize.manhattan_plot(stats_results,)</pre>	In EWAS Manhattan plots, epigenomic probe locations
	are displayed along the X-axis, with the negative log-
	arithm of the association P-value for each single nu-
	cleotide polymorphism (SNP) displayed on the Y-axis,
	meaning that each dot on the Manhattan plot signifies a
	SNP.
methylize.volcano_plot(stats_results,	This function writes the pandas DataFrame output of
**kwargs)	diff_meth_pos() to a CSV file named by the user.
	Continued on next page

#### Table 1 – continued from previous page

methylize.helpers.to\_BED(stats, ...[, save, Converts & exports manifest and probe p-value ...]) dataframe to BED format.

#### differentially methylated positions

This function searches for individual differentially methylated positions/probes (DMPs) by regressing the methylation M-value for each sample at a given genomic location against the phenotype data for those samples.

Phenotypes can be provided as a list of string-based or integer binary data or as numeric continuous data.

#### **Input Parameters:**

**meth\_data:** A pandas dataframe of methylation array data (as M-values) where each column corresponds to a CpG site probe and each row corresponds to a sample. IF a dataframe of beta-values is supplied instead, this function will detect this and convert to M-values before proceeding.

**pheno\_data:** A list or one dimensional numpy array of phenotypes for each sample row in meth\_data. Binary phenotypes can be presented as a list/array of zeroes and ones or as a list/array of strings made up of two unique words (i.e. "control" and "cancer").

- linear regression requires a measurement for the phenotype
- note: methylprep creates a *sample\_sheet\_meta\_data.pkl* file containing the phenotype

data for this input. You just need to load it and specify which *column* to be used as the pheno\_data. - If pheno\_data is a pandas DataFrame, you must specify a *column* and may optionally specify *covariates* from the other columns in the DataFrame. Numpy matrices are NOT supported.

#### column:

• If pheno\_data is a DataFrame, column='label' will select one series to be used as the phenotype data.

#### covariates: (default None) use to define a more complex model for logistic regression.

- if pheno\_data is not a DataFrame, covariates will be ignored.
- if pheno\_data is a DataFrame, and *column* is specified, and covariates is None, it will ignore other columns in pheno\_data.
- if covariates is a list of strings, only these column names will be used as the covariate data from pheno data.
- If covariates is set to True, then all other columns will be treated as covariates in logistic regression.

#### regression\_method: (logistic | linear)

• Either the string "logistic" or the string "linear"

depending on the phenotype data available. - Phenotypes with only two options (e.g. "control" and "cancer") can be analyzed with a logistic regression - Continuous numeric phenotypes (e.g. age) are required to run a linear regression analysis. - Default: auto-determine, using the data types found in *pheno\_data*. If it is numeric with more than 2 different labels, use "linear."

**impute:** Because methylprep output contains missing values by default, this function requires user to either delete or impute missing values.

- Default: 'delete' probes if ANY samples have missing data for that probe.
- True or 'auto': if <30 samples, deletes rows; if >=30 samples, uses average.
- False: don't impute and throw an error if NaNs present
- 'average' use the average of probe values in this batch
- 'delete' drop probes if NaNs are present in any sample
- 'fast' use adjacent sample probe value instead of average (much faster but less precise)

**alpha:** float Default is 0.05 for all tests where it applies.

**fwer: float** Set the familywise error rate (FWER). Default is 0.05, meaning that we expect 5% of all significant differences to be false positives. 0.1 (10%) is a more conservative convention for FWER.

#### q\_cutoff:

• Select a cutoff value (< 1.0) to return filtered results.

Only those DMPs that meet a particular significance threshold (e.g. 0.1) will be retained. Reported q-values are p-values corrected according to the model's false discovery rate (FDR). - Default: 1 – returns all DMPs regardless of significance.

### export:

- · default: False
- if True or 'csv', saves a csv file with data
- if 'pkl', saves a pickle file of the results as a dataframe.
- Use q\_cutoff to limit what gets saved to only significant results.

By default,  $q\_cutoff == 1$  and this means everything is saved/reported/exported.

#### filename:

• specify a filename for the exported file.

By default, if not specified, filename will be *DMP\_<number of probes in file>\_<number of samples processed>\_<current\_date>.<pkl|csv>* 

max\_workers: (=INT) By default, this will parallelize probe processing, using all available cores. During testing, or when running in a virtual environment like circleci or docker or lambda, the number of available cores is fewer than the system's reported CPU cores, and it breaks. Use this to limit the available cores to some arbitrary number for testing or containerized-usage.

**debug:** Default: False – True turns on extra messages and runs the 'solver' in serial mode, disabling parallel processing of probes. This is slower but provides more detail for debugging code.

**solver:** You can force it to use a different implementation of regression, for debugging. Options include:

- 'statsmodels\_OLS'
- 'linregress' # from scipy
- [logit\_DMP() is the default, based on statsmodels.api.Logit()]
- 'statsmodels\_GLM' # for logistic regression

### Returns:

A pandas dataframe of regression statistics with a row for each probe analyzed and columns listing the individual probe's regression statistics of:

- · regression coefficient
- lower limit of the coefficient's 95% confidence interval
- upper limit of the coefficient's 95% confidence interval
- · standard error
- p-value (phenotype group A vs B likelihood that the difference is significant for this probe/location)
- FDR\_QValue: p-values corrected for multiple comparisons using the Benjamini-Hochberg FDR method

The rows are sorted by q-value in ascending order to list the most significant probes first. If q\_cutoff is specified, only probes with significant q-values less than the cutoff will be returned in the dataframe.

#### Note:

If you get a *RuntimeError: main thread is not in main loop* error running *diff\_meth\_pos*, add *debug=True* to your function inputs. This error occurs in some command line environments when running this function repeatedly, and the parallel processing steps do not all close. Using debug mode forces it to analyze the probes serially, not parallel. It is a little slower but will always run without error under these conditions.

**If Progress Bar Missing:** if you don't see a progress bar in your jupyterlab notebook, try this: - conda install -c conda-forge nodejs - jupyter labextension install @jupyter-widgets/jupyterlab-manager

**Todo:** *shrink\_var* feature: variance shrinkage / squeeze variance using Bayes posterior means. Variance shrinkage is recommended when analyzing small datasets (n < 10). Paper ref: http://www.uvm.edu/~rsingle/JournalClub/papers/Smyth-SAGMB-2004\_eBayes+microarray.pdf [NOT IMPLEMENTED YET]

methylize.diff\_meth\_pos.legacy\_OLS (probe\_data, phenotypes, alpha=0.05) to use this, specify "statsmodels\_OLS" in kwargs to diff\_meth\_pos() – this method gives the same result as the scipy.linregress method when tested in version 1.0.0

methylize.diff\_meth\_pos.linear\_DMP\_regression (probe\_data, phenotypes, alpha=0.05)

This function performs a linear regression on a single probe's worth of methylation data (in the form of beta or M-values). It is called by the diff\_meth\_pos().

Inputs and Parameters:

**probe\_data:** A pandas Series for a single probe with a methylation M-value/beta-value for each sample in the analysis. The Series name corresponds to the probe ID, and the Series is extracted from the meth\_data DataFrame through a parallellized loop in diff\_meth\_pos().

phenotypes: A numpy array of numeric phenotypes with one phenotype per sample (so it must be the same length as probe\_data). This is the same object as the pheno\_data input to diff\_meth\_pos() after it has been checked for data type and converted to the numpy array pheno\_data\_array.

#### Returns:

A pandas Series of regression statistics for the single probe analyzed. The columns of regression statistics are as follows:

- regression coefficient (linregress pearson's 'r')
- lower limit of the coefficient's 95% confidence interval
- upper limit of the coefficient's 95% confidence interval

- · standard error
- p-value

methylize.diff\_meth\_pos.logistic\_DMP\_regression(probe\_data, phenotypes, covariate\_data=None, debug=False)

- · Runs parallelized.
- TESTED, and gives almost the same values as logit\_DMP() (2022-02-25)
- This function performs a logistic regression on a single probe's worth of methylation

data (in the form of beta/M-values). It is called by the diff\_meth\_pos().

Inputs and Parameters:

probe\_data: A pandas Series for a single probe with a methylation M-value or beta\_value for each sample in the analysis. The Series name corresponds to the probe ID, and the Series is extracted from the meth\_data DataFrame through a parallellized loop in diff\_meth\_pos().

**phenotypes:** A numpy array of binary phenotypes with one phenotype per sample (so it must be the same length as probe\_data). This is the same object as the pheno\_data input to diff\_meth\_pos() after it has been checked for data type and converted to the numpy array pheno\_data\_binary.

#### Returns:

A pandas Series of regression statistics for the single probe analyzed. The columns of regression statistics are as follows:

- · regression coefficient
- lower limit of the coefficient's 95% confidence interval
- upper limit of the coefficient's 95% confidence interval
- · standard error
- p-value

If the logistic regression was unsuccessful in fitting to the data due to a Perfect Separation Error (as may be the case with small sample sizes) or a Linear Algebra Error, the exception will be caught and the probe\_stats\_row output will contain dummy values to flag the error. Perfect Separation Errors are coded with the value -999 and Linear Algebra Errors are coded with value -995. These rows are processed and removed in the next step of diff\_meth\_pos() to prevent them from interfering with the final analysis and p-value correction while printing a list of the unsuccessful probes to alert the user to the issues.

```
methylize.diff_meth_pos.logit_DMP(probe_data, phenotypes, covariate_data=None, de-
bug=False)
```

DEFAULT method, because tested and works. uses statsmodels.api.Logit pass in a Series for probe data without the constant added

 $fold\_change\ is\ log2(\ (pheno1.mean\ /\ pheno0.mean)\ )$ 

each covariate will be normalized (to 0...1 range) before running DMP.

```
methylize.diff_meth_pos.manhattan_plot(stats_results, array_type, **kwargs)
```

In EWAS Manhattan plots, epigenomic probe locations are displayed along the X-axis, with the negative logarithm of the association P-value for each single nucleotide polymorphism (SNP) displayed on the Y-axis, meaning that each dot on the Manhattan plot signifies a SNP. Because the strongest associations have the smallest P-values (e.g., 1015), their negative logarithms will be the greatest (e.g., 15).

genomic coordinates along chromosomes vs epigenetic probe locations along chromosomes

• p-values are for the probe value associations, using linear or logistic regression,

between phenotype A and B.

Hints of hidden heritability in GWAS. Nature 2010. (https://www.ncbi.nlm.nih.gov/pubmed/20581876)

**stats\_results:** a pandas DataFrame containing the stats\_results from the linear/logistic regression run on m\_values or beta\_values and a pair of sample phenotypes. The DataFrame must contain A "PValue" column. the default output of diff\_meth\_pos() will work.

**array\_type:** specify the type of array [450k, epic, epic+, mouse, 27k], so that probes can be mapped to chromosomes.

save: specify that it export an image in png format. By default, the function only displays a plot.

**filename:** specify an export filename. The default is f"manhattan\_<stats>\_<timestamp>.png".

- verbose (True/False) default is True, verbose messages, if omitted.
- fwer (default 0.1) familywise error rate (fwer) is used to set p-value threshold and the FDR threshold line.
- genome\_build NEW or OLD. Default is NEWest genome\_build.
- *label-prefix* how to refer to chromosomes. By default, it shows numbers like 1 ... 22, and X, Y. pass in 'CHR-' to add a prefix to plot labels, or rename with 'c' like: c01 ... c22.

There are some preset "override" options for threshold lines on plots:

- *explore*: (default False) include all FOUR significance threshold lines on plot (suggestive, significant, bonferroni, and false discovery rate). Useful for data exploration.
- no\_thresholds: (default False) set to True to hide all FOUR threshold lines from plot.
- plain: (default False) hide all lines AND don't label significant probes.
- *statsmode*: (default False) show the FDR and Bonferroni thresholds, hide the suggestive and genomic significant lines.

These allow you to toggle lines/labels on or off:

- fdr: (default False) draw a threshold line on plot corresponding to the adjusted false discovery rate = 0.05
- bonferroni: (default False) draw the Bonferroni threshold, correcting for multiple comparisons.
- suggestive: (default 1e-5) draw the consensus "suggestive significance" theshold at p < 1e-5, or set to False to hide.
- significant: (default 5e-8) draw the consensus "genomic significance" the shold at p < 5e-8, or set to False to hide.
- plot\_cutoff\_label (default True) label to each dotted line on the plot
- *label\_sig\_probes* (default True) labels significant probes showing the greatest difference between groups.

#### Chart options:

- ymax default: 50. Set to avoid plotting extremely high -10log(p) values.
- width figure width default is 16
- height figure height default is 8

- fontsize figure font size default 16
- border plot border default is OFF
- palette specify one of a dozen options for colors of chromosome regions on plot: ['default', 'Gray', 'Pastel1', 'Pastel2', 'Paired', 'Accent', 'Dark2', 'Set1', 'Set2', 'Set3', 'tab10', 'tab20', 'tab20c', 'Gray2', 'Gray3']

methylize.diff\_meth\_pos.probe\_corr\_plot(stats, group='sig', colorby='pval')

- group='sig' is default (using PValue < 0.05)
- group='chromosome' also kinda works.
- colorby= pval or FDR; what to use to color the significant probes, if group='sig'

```
methylize.diff_meth_pos.volcano_plot(stats_results, **kwargs)
```

This function writes the pandas DataFrame output of diff\_meth\_pos() to a CSV file named by the user. The DataFrame has a row for every successfully tested probe and columns with different regression statistics as follows:

- · regression coefficient
- lower limit of the coefficient's 95% confidence interval
- upper limit of the coefficient's 95% confidence interval
- · standard error
- p-value
- FDR q-value (p-values corrected for multiple testing using the Benjamini-Hochberg FDR method)

Inputs and Parameters:

**stats\_results (required):** A pandas DataFrame output by the function diff\_meth\_pos().

- **'alpha':** Default: 0.05, The significance level that will be used to highlight the most significant p-values (or adjusted FDR Q-values) on the plot.
- 'cutoff': the beta-coefficient cutoff | Default: None format: a list or tuple with two numbers for (min, max) or 'auto'. If specified in kwargs, will exclude values within this range of regression coefficients OR fold-change range from being "significant" and put dotted vertical lines on chart. 'auto' will select a beta coefficient range that excludes 95% of results from appearing significant.
- **'adjust':** (default False) if this will adjust the p-value cutoff line for false discovery rate (Benjamini-Hochberg). Use 'fwer' to set the target rate.
- 'fwer': family-wise error rate (default is 0.1) specify a probability [0 to 1.0] for false discovery rate
- 'data\_type\_label': What to put on X-axis. Either 'Fold Change' (default) or 'Regression Coefficient'.

# visualization kwargs:

- palette color pattern for plot default is [blue, red, grey] other palettes: ['default', 'Gray', 'Pastell', 'Pastell', 'Paired', 'Accent', 'Dark2', 'Set1', 'Set2', 'Set3', 'tab10', 'tab20', 'tab20b', 'tab20c', 'Gray2', 'Gray3']
- width figure width default is 16
- *height* figure height default is 8
- fontsize figure font size default 16

- dotsize figure dot size on chart default 30
- border plot border default is OFF
- data\_type\_label (e.g. Beta Values, M Values) default is 'Beta'
- plot\_cutoff\_label add label to dotted line on plot default False

save: specify that it export an image in png format. By default, the function only displays a plot.

**filename:** specify an export filename. default is *volcano <current date>.png*.

#### Returns:

Displays a plot, but does not directly return an object. The data is color coded and displayed as follows:

- the negative log of adjusted p-values is plotted on the y-axis
- the regression coefficient beta value is plotted on the x-axis
- the significance cutoff level appears as a horizontal gray dashed line
- · non-significant points appear in light gray
- significant points with positive correlations (hypermethylated probes) appear in red
- significant points with negative correlations (hypomethylated probes) appear in blue

# differentially methylated regions

methylize.diff\_meth\_regions.diff\_meth\_regions (stats, manifest\_or\_array\_type, \*\*kwargs)

Calculates and annotates diffentially methylated regions (DMR) using the combined-pvalues pipeline and returns list of output files.

comb-p is a command-line tool and a python library that manipulates BED files of possibly irregularly spaced P-values and

- (1) calculates auto-correlation,
- (2) combines adjacent P-values,
- (3) performs false discovery adjustment,
- (4) finds regions of enrichment (i.e. series of adjacent low P-values) and
- (5) assigns significance to those regions.

ref: https://www.ncbi.nlm.nih.gov/pmc/articles/PMC3496335/

# Input Parameters:

**stats:** dataframe dataframe output from diff\_meth\_pos()

manifest\_or\_array\_type: class instance or string pass in the manifest, or the name of the array

**filename:** filename to prepend to the .BED file created.

creates a <filename>.bed output from diff\_meth\_pos and to\_BED. genome\_build: default 'NEW'

by default, it uses the NEWer genome build. Each manifest contains two genome builds, marked "NEW" and "OLD". To use the OLD build, se this to "OLD".

# Computational Parameters:

dist: int maximum distance from each probe to scan for adjacent peaks (80)

acf-dist: int window-size for smoothing. Default is 1/3 of peak-dist (dist kwarg),

**step:** int step size for bins in the ACF calculation (50)

threshold: foat Extend regions after seeding if pvalue exceeds this value (default: same as seed)

seed: float minimum size of a genomic region (0.05)

no fdr: bool don't use FDR

**genome\_control:** bool correct input pvalues for genome control (inflation factor). This reduces the confounding effects of population stratitication in EWAS data.

Display/output Paramters:

verbose: bool – default False Display additional processing information on screen.

**plot: bool – default False** False will suppress the manhattan plot step.

prefix: str prefix that gets appended to all output files (e.g. 'dmr' becomes 'dmr\_regions.csv')

region\_filter\_p: max adjusted region-level p-value in final output.

region\_filter\_n: req at least this many probes in a region

annotate: bool annotate with fetch\_genes() function that uses UCSC refGene database to add "nearby" genes to differentially methylated regions in the output CSV. If you want to fine-tune the reference database, the tolerance of what "nearby" means, and other parameters, set this to false and call *methylize.fetch\_genes* as a separate step on the '...\_regions.csv' output file.

**tissue: str** if specified, adds additional columns to the annotation output with the expression levels for identified genes in any/all tissue(s) that match the keyword. (e.g. if your methylation samples are whole blood, specify *tissue=blood*) For all 54 tissues, use *tissue=all* 

#### Returns:

list A list of files created.

#### **Fetch Genes**

find genes that are adjacent to significantly different CpG regions provided.

# Summary:

fetch\_genes() annotates the DMR region output file, using the UCSC Genome Browser database as a reference as to what genes are nearby. This is an exploratory tool, as there are many versions of the human genome that map genes to slightly different locations.

fetch genes() is an EXPLORATORY tool and makes a number of simplicifications:

- the DMR regions file saves one CpG probe name and location, even though clusters of probes may map to that nearby area.
- it measures the distance from the start position of the one representative probe per region to any nearby genes, using the 'tol'erance parameter as the cutoff. Tolerance is the max number of base pairs of separation between the probe sequence start and the gene sequence start for it to be considered as a match.
- The default 'tol'erance is 250, but that is arbitrary. Increase it to expand the search area, or decrease it to be more conservative. Remember that Illumina CpG probe sequences are 50 base pairs long, so 100 is nearly overlapping. 300 or 500 would also be reasonable.
- "Adjacent" in the linear sequence may not necessarily mean that the CpG island is FUNCTIONALLY coupled
  to the regulatory or coding region of the nearby protein. DNA superstructure can position regulatory elements
  near to a coding region that are far upstream or downstream from the mapped position, and there is no easy way
  to identify "adjacent" in this sense.

- Changing the 'tol'erance, or the reference database will result major differences in the output, and thus one's interpretation of the same data.
- Before interpreting these "associations" you should also consider filtering candidate genes by specific cell types
  where they are expressed. You should know the tissue from which your samples originated. And filter candidate
  genes to exclude those that are only expressed in your tissue during development, if your samples are from
  adults, and vice versa.

### Arguments:

- **dmr\_regions\_file:** pass in the output file DataFrame or FILEPATH from DMR function. Omit if you specify the *sql* kwarg instead.
- **ref: default is** *refGene* use one of possible\_tables for lookup: 'refGene' 88,819 genes default table used in comb-b and cruzdb packages. 'knownGene' 232,184 genes pseudo genes too (the "WHere TranscriptType == 'coding\_protein'" clause would work, but these fields are missing from the data returned.) 'ncbiRefSeq' 173,733 genes this table won't have gene descriptions, because it cannot be joined with the 'kgXref' (no shared key). Additionally, 'gtexGeneV8' is used for tissue-expression levels. Pseudogenes are ommitted using the "WHERE score > 0" clause in the SQL.
- tol: default 250 +/- this many base pairs consistutes a gene "related" to a CpG region provided.
- **tissue: str** if specified, adds additional columns to output with the expression levels for identified genes in any/all tissue(s) that match the keyword. (e.g. if your methylation samples are whole blood, specify *tissue=blood*) For all 54 tissues, use *tissue=all*
- **genome\_build:** (None, NEW, OLD) Only the default human genome build, hg38, is currently supported. Even though many other builds are available in the UCSC database, most tables do not join together in the same way.
- **use\_cached:** If True, the first time it downloads a dataset from UCSC Genome Browser, it will save to disk and use that local copy thereafter. To force it to use the online copy, set to False.
- **no\_sync:** methylize ships with a copy of the relevant UCSC gene browser tables, and will auto-update these every month. If you want to run this function without accessing this database, you can avoid updating using the *no\_sync=True* kwarg.
- **host, user, password, db:** Internal database connections for UCSC server. You would only need to mess with these of the server domain changes from the current hardcoded value {HOST}. Necessary for tables to be updated and for *tissue* annotation.
- **sql:** a DEBUG mode that bypasses the function and directly queries the database for any information the user wants. Be sure to specify the complete SQL statement, including the ref-table (e.g. refGene or ncbiRefSeq).

**Note:** This method flushes cache periodically. After 30 days, it deletes cached reference gene tables and redownloads.

#### **Manhattan Plot**

In EWAS Manhattan plots, epigenomic probe locations are displayed along the X-axis, with the negative logarithm of the association P-value for each single nucleotide polymorphism (SNP) displayed on the Y-axis, meaning that each dot on the Manhattan plot signifies a SNP. Because the strongest associations have the smallest P-values (e.g., 1015), their negative logarithms will be the greatest (e.g., 15).

# **GWAS vs EWAS**

- genomic coordinates along chromosomes vs epigenetic probe locations along chromosomes
- p-values are for the probe value associations, using linear or logistic regression,

between phenotype A and B.

#### Ref

Hints of hidden heritability in GWAS. Nature 2010. (https://www.ncbi.nlm.nih.gov/pubmed/20581876)

# **Required Inputs**

**stats\_results:** a pandas DataFrame containing the stats\_results from the linear/logistic regression run on m\_values or beta\_values and a pair of sample phenotypes. The DataFrame must contain A "PValue" column. the default output of diff\_meth\_pos() will work.

**array\_type:** specify the type of array [450k, epic, epic+, mouse, 27k], so that probes can be mapped to chromosomes.

# output kwargs

save: specify that it export an image in png format. By default, the function only displays a plot.

**filename:** specify an export filename. The default is f"manhattan <stats> <ti>stats> <timestamp>.png".

#### visualization kwargs

- verbose (True/False) default is True, verbose messages, if omitted.
- fwer (default 0.1) familywise error rate (fwer) is used to set p-value threshold and the FDR threshold line.
- genome\_build NEW or OLD. Default is NEWest genome\_build.
- *label-prefix* how to refer to chromosomes. By default, it shows numbers like 1 ... 22, and X, Y. pass in 'CHR-' to add a prefix to plot labels, or rename with 'c' like: c01 ... c22.

There are some preset "override" options for threshold lines on plots:

- explore: (default False) include all FOUR significance threshold lines on plot (suggestive, significant, bonferroni, and false discovery rate). Useful for data exploration.
- no\_thresholds: (default False) set to True to hide all FOUR threshold lines from plot.
- plain: (default False) hide all lines AND don't label significant probes.
- *statsmode*: (default False) show the FDR and Bonferroni thresholds, hide the suggestive and genomic significant lines.

These allow you to toggle lines/labels on or off:

- fdr: (default False) draw a threshold line on plot corresponding to the adjusted false discovery rate = 0.05
- bonferroni: (default False) draw the Bonferroni threshold, correcting for multiple comparisons.

- suggestive: (default 1e-5) draw the consensus "suggestive significance" the shold at p < 1e-5, or set to False to hide.
- significant: (default 5e-8) draw the consensus "genomic significance" theshold at p < 5e-8, or set to False to hide.
- plot\_cutoff\_label (default True) label to each dotted line on the plot
- *label\_sig\_probes* (default True) labels significant probes showing the greatest difference between groups.

### Chart options:

- ymax default: 50. Set to avoid plotting extremely high -10log(p) values.
- width figure width default is 16
- height figure height default is 8
- fontsize figure font size default 16
- border plot border default is OFF
- palette specify one of a dozen options for colors of chromosome regions on plot: ['default', 'Gray', 'Pastel1', 'Pastel2', 'Paired', 'Accent', 'Dark2', 'Set1', 'Set2', 'Set3', 'tab10', 'tab20', 'tab20c', 'Gray2', 'Gray3']

#### Volcano Plot

This function writes the pandas DataFrame output of diff\_meth\_pos() to a CSV file named by the user. The DataFrame has a row for every successfully tested probe and columns with different regression statistics as follows:

- · regression coefficient
- lower limit of the coefficient's 95% confidence interval
- upper limit of the coefficient's 95% confidence interval
- · standard error
- p-value
- FDR q-value (p-values corrected for multiple testing using the Benjamini-Hochberg FDR method)

# Inputs and Parameters:

stats\_results (required): A pandas DataFrame output by the function diff\_meth\_pos().

- **'alpha':** Default: 0.05, The significance level that will be used to highlight the most significant p-values (or adjusted FDR Q-values) on the plot.
- **'cutoff':** the beta-coefficient cutoff | Default: None format: a list or tuple with two numbers for (min, max) or 'auto'. If specified in kwargs, will exclude values within this range of regression coefficients OR fold-change range from being "significant" and put dotted vertical lines on chart. 'auto' will select a beta coefficient range that excludes 95% of results from appearing significant.
- **'adjust':** (default False) if this will adjust the p-value cutoff line for false discovery rate (Benjamini-Hochberg). Use 'fwer' to set the target rate.
- 'fwer': family-wise error rate (default is 0.1) specify a probability [0 to 1.0] for false discovery rate
- 'data\_type\_label': What to put on X-axis. Either 'Fold Change' (default) or 'Regression Coefficient'. visualization kwargs:

- palette color pattern for plot default is [blue, red, grey] other palettes: ['default', 'Gray', 'Pastel1', 'Pastel2', 'Paired', 'Accent', 'Dark2', 'Set1', 'Set2', 'Set3', 'tab10', 'tab20', 'tab20b', 'tab20c', 'Gray2', 'Gray3']
- width figure width default is 16
- height figure height default is 8
- fontsize figure font size default 16
- dotsize figure dot size on chart default 30
- border plot border default is OFF
- data\_type\_label (e.g. Beta Values, M Values) default is 'Beta'
- plot\_cutoff\_label add label to dotted line on plot default False

save: specify that it export an image in png format. By default, the function only displays a plot.

**filename:** specify an export filename. default is *volcano\_<current\_date>.png*.

#### Returns:

Displays a plot, but does not directly return an object. The data is color coded and displayed as follows:

- the negative log of adjusted p-values is plotted on the y-axis
- the regression coefficient beta value is plotted on the x-axis
- the significance cutoff level appears as a horizontal gray dashed line
- non-significant points appear in light gray
- significant points with positive correlations (hypermethylated probes) appear in red
- significant points with negative correlations (hypomethylated probes) appear in blue

# To BED

Converts & exports manifest and probe p-value dataframe to BED format. \* https://en.wikipedia.org/wiki/BED\_(file\_format)

• BED format: [ chromosome number | start position | end position | p-values]

Where p-values are the output from diff\_meth\_pos() comparing probes across two or more groups of samples for genomic differences in methylation.

This output is required for combined-pvalues library to read and annotate manhattan plots with the nearest Gene(s) for each significant CpG cluster.

manifest\_or\_array\_type: either pass in a Manifest instance from methylprep, or a string that defines which manifest to load. One of {'27k', '450k', 'epic', 'epic+', 'mouse'}.

genome build: pass in 'OLD' to use the older genome build for each respective manifest array type.

note: if manifest has probes that aren't mapped to genome, they are omitted in BED file.

TODO: incorporate STRAND and OLD\_STRAND in calculations.

returns a BED formatted dataframe if save is False, or the saved filename if save is True.

# 1.8.6 Release History

#### v1.1.0

- We found that diff\_meth\_pos results were not accurate in prior versions and have fixed the regression optimization.
- · diff\_meth\_pos function kwargs changed to provide more flexibility in how the model is optimized.
  - Added support for COVARIATES in logistic regression. Provide a dataframe with both the phenotype and covariates, and specify which columns are phenotype or covariates. It will rearrange and normalize to ensure the model works best.
  - Use the new 'solver' kwarg in diff\_meth\_pos to specify which form of linear or logistic regression to run. There are two flavors of each, and both give nearly identical results.
  - Auto-detects logistic or linear based on input: if non-numeric inputs in phenotype of exactly two values, it assumes logistic.
- Upgraded manhattan and volcano plots with many more options. Default settings should mirror most R EWAS packages now, with a "suggestive" and "significant" threshold line on manhattan plots.
- Unit test coverage added.

#### v1.0.1

- Fixed option to use Differentially methylated regions (DMR) via cached local copy of UCSC database (via fetch\_genes) without using the internet. Previously, it would still contact the internet database even if user told it not to.
- · Added testing via github actions, and increased speed
- · updated documentation

#### v1.0.0

• fixed bug in fetch\_genes() from UCSC browser; function will now accept either the filepath or the DMR dataframe output.

# v0.9.9

- Added a differentially methylated regions (DMR) functions that takes the output of the diff\_meth\_pos (DMP) function.
  - DMP maps differences to chromosomes; DMR maps differences to specific genomic locii, and requires more processing.
  - upgraded methylprep manifests to support both old and new genomic build mappings for all array types.
     In general, you can supply a keyword argument (genome\_build='OLD') to change from the new build back to the old one.
  - Illumina 27k arrays are still not supported, but mouse, epic, epic+, and 450k ARE supported. (Genome annotation won't work with mouse array, only human builds.)
  - DMP integrates the combined-pvalues package (https://pubmed.ncbi.nlm.nih.gov/22954632/)
  - DMP integrates with UCSC Genome (refGene) and annotates the genes near CpG regions.

- Annotation includes column(s) showing the tissue specific expression levels of relevant genes (e.g. filter=blood) this function is also available with extended options as methylize. filter\_genes()
- provides output BED and CSV files for each export into other genomic analysis tools
- methylize.to\_BED will convert the diff\_meth\_pos() stats output into a standard BED file (a tab separated CSV format with standardized, ordered column names)

#### v0.9.8

- fixed methylize diff\_meth\_pos linear regression. upgraded features too
  - Fixed bug in diff\_meth\_pos using linear regression was not calculating p-values correctly. Switched from statsmodels OLS to scipy lineareses to fix, but you can use either one with kwargs. They appear to give exactly the same results now after testing.
  - The "CHR-" prefix is omitted from manhattan plots by default now
  - dotted manhattan sig line is Bonferoni corrected (pass in post\_test=None to leave uncorrected)
  - added a probe\_corr\_plot() undocumented function, a scatterplot of probe confidence intervals vs pvalue
  - sorts probes by MAPINFO (chromosome location) instead of FDR\_QValue on manhattan plots now
- Support for including/excluding sex chromosomes from DMP (probe2chr map)

#### v0.9.5

- Added imputation to diff\_meth\_pos() function, because methylprep output contains missing values by default and cannot be used in this function.
  - This can be disabled, and it will throw a warning if NaNs present.
  - Default is to delete probes that have any missing values before running analysis.
  - if 'auto': If there are less than 30 samples, it will delete the missing rows.
  - if 'auto': If there are >= 30 samples in the batch analyzed, it will replace NaNs with the average value for that probe across all samples.
  - User may override the default using: True ('auto'), 'delete', 'average', and False (disable)
  - diff\_meth\_pos() now support mouse array, with multiple copies of the same probe names.

#### v0.9.4

- Fixed bug where methylize could not find a data file in path, causing ImportError
- Improved diff\_meth\_pos() function and added support for all array types. Now user must specify the array\_type when calling the function, as the input data are stats, not probe betas, so it cannot infer the array type from this information.

# CHAPTER 2

Indices and tables

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