# **ember Documentation**

Release 0.1.0

**Pachter Lab** 

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# **GENERATING ENTROPY METRICS**

ember.light\_ember(h5ad\_dir, partition\_label, save\_dir, sampling=True, sample\_id\_col=None, category\_col=None, condition\_col=None, num\_draws=100, save\_draws=False, seed=42, partition\_pvals=True, block\_pvals=False, block\_label=None, n\_pval\_iterations=1000, n\_cpus=1)

Runs the ember entropy metrics and p-value generation workflow on an AnnData object.

This function loads an AnnData .h5ad file, optionally performs balanced sampling across replicates, computes entropy metrics for the specified partition, and generates p-values for Psi and Zeta and optionally Psi\_block for a block of choice.

#### **Entropy metrics generated:**

- Psi: Fraction of infromation explained by partition of choice
- Psi\_block : Specificity of infromation to a block
- Zeta: Speicifcty to a partition/ distance of Psi\_blocks distribution from uniform

#### **Parameters**

- **h5ad\_dir** (*str*, *Required*) Path to the *.h5ad* file to process. Data should be log1p and depth normalized before running ember. Remove genes with less than 100 reads before running ember.
- partition\_label (str, Required) Column in .obs used to partition cells for entropy calculations (e.g., "celltype", "Genotype", "Age"). Required to run process. If performing calculation on interaction term, first create a column in .obs concatnating the two columns of interested with a semicolon (:).
- **save\_dir** (*str*, *Required*) Path to directory where results will be saved. Required to run process.
- **sampling** (bool, default=True) Whether to perform balanced sampling across replicates before entropy calculation. If True, sample\_id\_col, category\_col, and condition\_col must be provided. Sampling should only be False if fast intermediate results are desired or if there are no replicates to sample over. If sampling is set to False but either partition\_pvals or block\_pvals are set to True then the sampling=False will be overridden as pval generation requires sampling.
- **sample\_id\_col** (*str*, *default* = *None*) The column in .*obs* with unique identifiers for each sample or replicate (e.g., 'sample\_id', 'mouse\_id').
- **category\_col** (*str*, *default* = *None*) The column in .*obs* defining the primary group to balance across in order to generate a balanced sample of the experiment. (e.g., 'disease\_status', 'mouse\_strain'). Refer to readme for further explanation on how to select category and condition columns. category\_col and condition\_col are interchangable. If balanc-

ing across more than 2 variables, generate interaction terms, create a column in .obs concatnating the two (or more) columns of interested with a semicolon (:). This way balancing can be done across as many variables as desired.

- **condition\_col** (*str*, *default* = *None*) The column in .*obs* containing the conditions to balance within each categoryto generate a balanced sample of the experiment. (e.g., 'sex', 'treatment'). Refer to readme for further explanation on how to select category and condition columns. category\_col and condition\_col are interchangable. If balancing across more than 2 variables, generate interaction terms, create a column in .*obs* concatnating the two (or more) columns of interested with a semicolon (:). This way balancing can be done across as many variables as desired.
- num\_draws (int, default = 100) The number of balanced subsets to generate, by default 100.
- save\_draws (bool, default=False) Whether to save intermediate draws to save\_dir.
- **seed** (*int*, *default* = 42) The random seed for reproducible draws, by default 42.
- partition\_pvals (bool, default=True) Whether to compute permutation-based p-values for the partition\_label. P-values are generated by sampling. If sampling = False and partition\_pvals = True, the sampling=False will be overwritten. Calls generate\_pavls, which can be called manually after metric generation as well.
- **block\_pvals** (*bool*, *default=False*) Whether to compute permutation-based p-values for the *block\_label*. P-values are generated by sampling. If sampling = False and block\_pvals = True, the sampling=False will be overwritten. Calls generate\_pavls, which can be called manually after metric generation as well.
- **block\_label** (str, default = None) One value in the .obs column for partition\_label to use for block-based permutation tests. Required if block\_pvals=True.
- n\_pval\_iterations (int, default=1000) Number of permutations to use for p-value calculation.
- **n\_cpus** (*int*, *default=1*) Number of CPU cores to use for parallel permutation testing. For this script, performance is I/O-bound and may not improve beyond 4-8 cores.'

#### Return type

None

#### **Notes**

- Results are saved to save\_dir as CSV files.
- one csv file with all entropy metrics
- one csv file in a new Psi\_block\_df folder with psi block values for all blocks in a partition
- Separate file for pvals
- Separate files for each partition
- Alternate file names depending on sampling on or off.

#### What to expect inside 'entropy\_metrics.csv':

- gene\_name: All genes in .var
- Psi\_mean: Psi scores averaged over n draws (between 0 and 1) corresponding to the selected partition for each gene in .var.

- Psi\_std: Standard deviation of Psi scores across n draws corresponding to the selected partition for each gene in .var.
- Psi\_valid\_counts: Number of valid Psi scores observed across n draws. Only use genes for downstream analysis that have valid counts=num\_draws. If valid counts is not close to num\_draws, increase threshold for filtering genes with low reads beforehand(recommended <100 reads, increase as needed).
- Zeta\_mean: Zeta scores averaged over n draws (between 0 and 1) corresponding to the selected partition for each gene in .var.
- Zeta\_std: Standard deviation of Zeta scores across n draws corresponding to the selected partition for each gene in .var.
- Zeta\_valid\_counts: Number of valid Psi scores observed across n draws. Only use genes for downstream analysis that have valid counts=num\_draws. If valid counts is not close to num\_draws, increase threshold for filtering genes with low reads beforehand (recommended <100 reads, increase as needed).

#### What to expect inside 'Psi\_block\_df/':

- mean\_Psi\_block\_df.csv : A dataframe of mean Psi\_block scores (between 0 and 1) corresponding to the selected partition for each gene in .var. Scores are caluclated for all blocks, each column of the dataframe corresponds to one block.
- std\_Psi\_block\_df.csv : A dataframe of standard deviations for Psi\_block scores corresponding to the selected partition for each gene in .var.Scores are caluclated for all blocks, each column of the dataframe corresponds to one block.

#### What to expect inside 'pvals entropy metrics.csv':

- gene name: All genes in .var
- Psi: Psi scores averaged over n draws (between 0 and 1) generated by light\_ember for each gene in .var.
- Psi p-value: Permutation based empirical p-values for observed Psi scores for each gene in .var.
- Zeta: Zeta scores averaged over n draws (between 0 and 1) generated by light\_ember for each gene in .var.
- Zeta p-value: Permutation based empirical p-values for observed Zeta scores for each gene in .var.
- Psi FDR: Multiple testing corrected q-values for Psi scores.
- Zeta FDR: Multiple testing corrected q-values for Zeta scores. Correction perfromed to include all p-values generated in a single file (Psi and Zeta).

#### If block pvals = True and a single block label is given:

- psi\_block: psi\_block scores (between 0 and 1) generated by light\_ember for each gene in .var.
- psi\_block p-value: Permutation based empirical p-values for observed psi\_block scores for each gene in
   .var.
- psi\_block FDR: Multiple testing corrected q-values for psi\_block scores. Correction perfromed to include all p-values generated in a single file (Psi, psi\_block and Zeta).

# **GENERATING P-VALUES**

ember.generate\_pvals.generate\_pvals(h5ad\_dir, partition\_label, entropy\_metrics\_dir, save\_dir, sample\_id\_col, category\_col, condition\_col, block\_label=None, seed=42, n\_iterations=1000, n\_cpus=1, Psi\_real=None, Psi\_block\_df\_real=None, Zeta\_real=None)

Calculate empirical p-values for entropy metrics from permutation test results. This function can be called manually or accessed through light\_ember with partition\_pvals = True or block\_pvals = True.

Manual access useful if wanting to generate p-values for multiple blocks and partitions of interest after initial investigation using entropy metrics.

Integrated access with light\_ember is easier if investigating only a partition or a block in a partition.

#### **Entropy metrics generated:**

- Psi: Fraction of infromation explained by partition of choice
- Psi\_block : Specificity of infromation to a block
- Zeta: Speicifcty to a partition/distance of Psi\_blocks distribution from uniform

#### **Parameters**

- **h5ad\_dir** (*str*, *Required*) Path to the *.h5ad* file to process. Data should be log1p and depth normalized before running ember. Remove genes with less than 100 reads before running ember.
- partition\_label (str, Required) Column in .obs used to partition cells for entropy calculations (e.g., "celltype", "Genotype", "Age"). Required to run process. If performing calculation on interaction term, first create a column in .obs concatnating the two columns of interested with a semicolon (:).
- entropy\_metrics\_dir (str, Required) Path to csv with entropy metrics to use for generating pvals.
- **save\_dir** (*str*, *Required*) Path to directory where results will be saved.
- **sample\_id\_col** (*str*, *Required*) The column in *.obs* with unique identifiers for each sample or replicate (e.g., 'sample\_id', 'mouse\_id').
- category\_col (str, Required) The column in .obs defining the primary group to balance across in order to generate a balanced sample of the experiment. (e.g., 'disease\_status', 'mouse\_strain'). Refer to readme for further explanation on how to select category and condition columns. category\_col and condition\_col are interchangable. If balancing across more than 2 variables, generate interaction terms, create a column in .obs concatnating the two (or more) columns of interested with a semicolon (:). This way balancing can be done across as many variables as desired.

- **condition\_col** (*str*, *Required*) The column in .*obs* containing the conditions to balance within each category to generate a balanced sample of the experiment. (e.g., 'sex', 'treatment'). Refer to readme for further explanation on how to select category and condition columns. category\_col and condition\_col are interchangable. If balancing across more than 2 variables, generate interaction terms, create a column in .*obs* concatnating the two (or more) columns of interested with a semicolon (:). This way balancing can be done across as many variables as desired.
- **block\_label** (*str*, *default=None*) Block in partition to calucate p-values for. Default set to None, program will continue generating p-values for only Psi and Zeta.
- **seed** (*int*, *default=42*) The random seed for reproducible draws, by default 42.
- n\_iterations (int, default = 1000) Number of iterations to calulate p-vales. Default set to 1000. Note that doing fewer than 1000 iterations is a good choice to get first pass p-values but for reliable p-values 1000 iterations is recommended. Larger than 1000 will give you more reliable p-values but will increase runtime significantly.
- n\_cpus (int, default=1) Number of cpus to use to perfrom p-value calculation. Default set to 1 assuming no parallel compute power on local machine. User can input -1 to use all available cpus but one.
- **Psi\_real** (*pd.Series*, *default=None*) Observed Psi values for each gene. Used by light\_ember, not necessary for user use.
- Psi\_block\_df\_real (pd.Dataframe, default = None) Observed Psi\_block values for all blocks in chosen partition. Used by light\_ember, not necessary for user use.
- **Zeta\_real** (*pd. Series*, *default=None*) Observed Zeta values for each gene. Used by light\_ember, not necessary for user use.

#### Return type

None

#### **Notes**

#### What to expect inside 'pvals\_entropy\_metrics.csv':

- gene\_name: All genes in .var
- Psi: Psi scores averaged over n draws (between 0 and 1) generated by light\_ember for each gene in .var.
- Psi p-value: Permutation based empirical p-values for observed Psi scores for each gene in .var.
- Zeta: Zeta scores averaged over n draws (between 0 and 1) generated by light\_ember for each gene in .var.
- Zeta p-value: Permutation based empirical p-values for observed Zeta scores for each gene in .var.
- Psi FDR: Multiple testing corrected q-values for Psi scores.
- Zeta FDR: Multiple testing corrected q-values for Zeta scores. Correction perfromed to include all p-values generated in a single file (Psi and Zeta).

if block\_pvals = True and a single block\_label is given:

- psi\_block: psi\_block scores (between 0 and 1) generated by light\_ember for each gene in .var.
- psi\_block p-value: Permutation based empirical p-values for observed psi\_block scores for each gene in .var.
- psi\_block FDR: Multiple testing corrected q-values for psi\_block scores. Correction perfromed to include all p-values generated in a single file (Psi, psi\_block and Zeta).

### PLOTTING FUNCTIONS

# 3.1 Psi vs. Zeta scatter plots

Generate a Zeta vs. Psi scatter plot to visualize partition-specific genes.

This function reads p-value data, colors genes based on their statistical significance for Psi and Zeta scores, and highlights top "marker" and "housekeeping" genes. Allows for custom highlighting of a user-provided gene list. Fontsize and color pallette can be customized.

#### **Parameters**

- partition\_label (str, Required.) The label for the partition being plotted, used in the plot title.
- **pvals\_dir** (*str*, *Required*.) Path to the input CSV file containing p-values and scores (Psi, Zeta, FDRs). The CSV must have gene names as its index column.
- **save\_dir** (*str*, *Required*.) Path where the output plot image will be saved.
- highlight\_genes (list[str], default=None.) A list of gene names to highlight and annotate on the plot, by default None.
- fontsize (int, default=18.) The base font size for plot labels and text, by default 18.
- **custom\_palette** (*list[str]*, *default=None*.) A list of 7 hex color codes to customize the plot's color scheme. If None, a default palette is used. Please provide list in this order ['significant by psi', 'significant by zeta', 'highlight genes', 'significant by both', 'cirlce markers', 'circle housekeeping genes', 'significant by neither']

#### Return type

None

# 3.2 Psi vs. psi\_block scatter plots

Generate a psi\_block vs. Psi scatter plot to visualize block-specific genes.

This function reads p-value data, colors genes based on their statistical significance for Psi and psi\_block scores, and highlights the top genes significant in both metrics. Allows for custom highlighting of a user-provided gene list. Fontsize and color pallette can be customized.

#### **Parameters**

- partition\_label (str, Required.) The label for the partition, used in the plot title.
- **block\_label** (*str*, *Required*.) The label for the block variable (e.g., a cell type or condition).
- **pvals\_dir**(*str*, *Required*.) Path to the input CSV file containing p-values and scores. The CSV must have gene names as its index column.
- **save\_dir** (*str*, *Required*.) Path where the output plot image will be saved.
- highlight\_genes (list[str], default=None.) A list of gene names to highlight and annotate on the plot, by default None.
- **fontsize** (*int*, *default* = 18.) The base font size for plot labels and text, by default 18.
- **custom\_palette** (*list[str]*, *default=None*.) A list of 6 hex color codes to customize the plot's color scheme. If None, a default palette is used. Provide list of colors int his order: ['significant by psi', 'significant by psi\_block', 'highlight genes', 'significant by both', 'circle markers', 'circle housekeeping genes', 'significant by neither']

#### Return type

None

# 3.3 Descriptive bar plot of sample counts

Generate a bar plot showing the number of unique individuals per category and condition.

This function reads an AnnData object from an .h5ad file in backed mode, calculates the number of unique individuals for each combination of a given category and condition, and visualizes these counts as a grouped bar plot. Fontsize can be customized.

### **Parameters**

- **h5ad\_dir** (*str*, *Required*) Path to the input AnnData (.h5ad) file.
- **save\_dir** (*str*, *Required*) Path to directory to save the output plot image.
- **sample\_id\_col** (*str*, *Required*) The column name in adata.obs that contains unique sample IDs.
- **category\_col** (*str*, *Required*) The column name to use for the primary categories on the x-axis.
- condition\_col (str, Required) The column name to use for grouping the bars (hue).
- **fontsize** (*int*, *default* = 18.) The base font size for plot labels and text, by default 18.

#### Return type

None

# 3.4 psi\_blocks bar plots with error bars

ember.plots.plot\_psi\_blocks(gene\_name, partition\_label, psi\_block\_df\_dir, save\_dir, fontsize=18)

Generates and saves a bar plot of mean psi block values with error bars.

This function reads two CSV files from a specified directory: one for mean psi block values and one for standard deviations. It plots the mean values for a specific gene as a bar plot with corresponding standard deviation error bars. Fontsize can be customized.

#### **Parameters**

- **gene\_name** (*str*, *Required*) The name of the gene (row) to select and plot from the CSV files.
- partition\_label (*str*, *Required*) The partition label used to find the correct files (e.g., 'Genotype').
- **psi\_block\_df\_dir** (*str*, *Required*) Path to the directory containing the mean and std CSV files. Files must be named 'mean\_Psi\_block\_df\_{partition\_label}.csv' and 'std\_Psi\_block\_df\_{partition\_label}.csv'.
- **save\_dir** (*str*, *Required*) Path to directory to save the output plot image.
- **fontsize** (*int*, *default=18*.) The base font size for plot labels and text, by default 18.

#### **Return type**

None

**CHAPTER** 

**FOUR** 

# **COMMAND-LINE INTERFACE (CLI)**

The ember also has a command-line interface (CLI). This allows you to run workflows and plotting directly from the terminal.

A command-line toolkit for ember: Entropy Metrics for Biological ExploRation.

# 4.1 Positional Arguments

command

Possible choices: light\_ember, generate\_pvals, plot\_partition\_specificity, plot\_block\_specificity, plot\_sample\_counts, plot\_psi\_blocks

Available sub-commands

### 4.2 Sub-commands

# 4.2.1 light\_ember

Runs the ember entropy metrics and p-value generation workflow on an AnnData object.

This function loads an AnnData .h5ad file, optionally performs balanced sampling across replicates, computes entropy metrics for the specified partition, and generates p-values for Psi and Zeta and optionally Psi block for a block of choice.

### **Entropy metrics generated:**

- Psi: Fraction of information explained by partition of choice
- Psi\_block : Specificity of information to a block
- Zeta: Specificity to a partition / distance of Psi\_blocks distribution from uniform

#### Notes:

- Results are saved to save\_dir as CSV files.
- One CSV file with all entropy metrics.
- One CSV file in a new Psi\_block\_df folder with Psi\_block values for all blocks in a partition.
- Separate file for p-values.

- Separate files for each partition.
- Alternate file names depending on sampling on or off.

#### **Positional Arguments**

**h5ad\_dir** Path to the .h5ad file to process. Data should be log1p and depth normalized

before running ember. Remove genes with <100 reads before running ember.

partition\_label Column in .obs used to partition cells for entropy calculations (e.g., 'celltype',

'Genotype', 'Age'). For interaction terms, create a new column concatenating

multiple .obs columns with a semicolon (:).

**save\_dir** Path to directory where results will be saved.

**Sampling Parameters** 

--no\_sampling Disable balanced sampling. Default: True. Note: If partition\_pvals or

block\_pvals are enabled, sampling will be re-enabled.

Default: True

**--sample\_id\_col** Column in .obs with unique identifiers for each sample or replicate (e.g., 'sam-

ple id', 'mouse id').

--category\_col Column in .obs defining the primary group to balance across (e.g., 'dis-

ease status', 'mouse strain'). Interchangeable with condition col. For >2 vari-

ables, create interaction terms by concatenating columns with:.

**--condition\_col** Secondary column in .obs to balance sampling across (e.g., 'sex', 'treatment').

Interchangeable with category\_col. Supports interaction terms.

**--num\_draws** Number of balanced subsets to generate (default: 100).

Default: 100

**--save\_draws** Save intermediate sampled draws to save\_dir (default: False).

Default: False

**--seed** Random seed for reproducible draws (default: 42).

Default: 42

### **P-value Parameters**

**--no\_partition\_pvals** Disable permutation p-value calculation for the main partition. Default: True.

Default: True

**--block pvals** Enable permutation p-value calculation for a specific block. Default: False.

Default: False

**--block\_label** Specific value in 'partition\_label' for block p-values. Required if -block\_pvals is

set.

**--n\_pval\_iterations** Number of permutations for p-value calculation (default: 1000).

Default: 1000

#### **Performance Parameters**

**--n\_cpus** Number of CPU cores to use for parallel processing (default: 1). Performance is

I/O-bound and may not improve beyond 4–8 cores.

Default: 1

#### **Example:**

ember light\_ember ~/ember\_test/test\_adata\_cwc22.h5ad Genotype ~/ember\_test/ -sample\_id\_col Mouse\_ID -category\_col Genotype -condition\_col Sex -num\_draws 50 -no\_partition\_pvals -n\_cpus 4

### 4.2.2 generate pvals

Calculate empirical p-values for entropy metrics from permutation test results.

#### **Entropy metrics generated:**

- Psi: Fraction of information explained by partition of choice
- Psi block: Specificity of information to a block
- Zeta: Specificity to a partition / distance of Psi\_blocks distribution from uniform

```
ember generate_pvals [-h] [--block_label BLOCK_LABEL] [--seed SEED]

[--n_iterations N_ITERATIONS] [--n_cpus N_CPUS]

[--Psi_real PSI_REAL]

[--Psi_block_df_real PSI_BLOCK_DF_REAL]

[--Zeta_real ZETA_REAL]

h5ad_dir partition_label entropy_metrics_dir save_dir

sample_id_col category_col condition_col
```

#### **Positional Arguments**

**h5ad dir** Path to the .h5ad file to process. Data should be log1p and depth normalized

before running ember. Remove genes with <100 reads before running ember.

partition\_label Column in .obs used to partition cells for entropy calculations (e.g., 'celltype',

'Genotype', 'Age'). For interaction terms, create a new column concatenating

multiple .obs columns with a semicolon (:).

**entropy\_metrics\_dir** Path to CSV with entropy metrics to use for generating p-values.

**save\_dir** Path to directory where results will be saved.

**sample\_id\_col** Column in .obs with unique identifiers for each sample or replicate (e.g., 'sam-

ple\_id', 'mouse\_id').

category\_col Column in .obs defining the primary group to balance across (e.g., 'dis-

ease\_status', 'mouse\_strain'). Interchangeable with condition\_col. For >2 vari-

ables, create interaction terms by concatenating columns with:.

**condition\_col** Column in .obs containing the conditions to balance within each category (e.g.,

'sex', 'treatment'). Interchangeable with category\_col. Supports interaction

terms.

4.2. Sub-commands

### **Named Arguments**

**--block label** Block in partition to calculate p-values for. Default: None (Psi and Zeta only).

#### **Performance Parameters**

**--seed** Random seed for reproducible draws (default: 42).

Default: 42

**--n\_iterations** Number of iterations to calculate p-values (default: 1000). Use fewer for quick

runs, more for reliable results.

Default: 1000

**--n\_cpus** Number of CPUs to use for p-value calculation (default: 1). Set to -1 to use all

available cores but one.

Default: 1

# Internal Arguments (used by light\_ember)

**--Psi\_real** Observed Psi values for each gene (pd.Series). Not required for user runs.

--Psi\_block\_df\_real Observed Psi\_block values for all blocks in chosen partition (pd.DataFrame). Not

required for user runs.

**--Zeta\_real** Observed Zeta values for each gene (pd.Series). Not required for user runs.

#### **Example:**

ember generate\_pvals test\_adata\_cwc22.h5ad Genotype ~/ember\_test/ ~/ember\_test/output Mouse\_ID Genotype Sex -block\_label WSBJ -n\_cpus 4

# 4.2.3 plot partition specificity

Generate a Zeta vs. Psi scatter plot to visualize partition-specific genes.

This function reads p-value data, colors genes based on their statistical significance for Psi and Zeta scores, and highlights top "marker" and "housekeeping" genes. Allows for custom highlighting of a user-provided gene list. Font size and color palette can be customized.

#### **Positional Arguments**

**partition\_label** Label for the partition being plotted, used in the plot title.

**pvals\_dir** Path to input CSV containing p-values and scores (Psi, Zeta, FDRs). CSV must

have gene names as its index.

**save dir** Path where the output plot image will be saved.

#### **Named Arguments**

**--highlight\_genes** List of gene names to highlight and annotate on the plot (default: None).

**--fontsize** Base font size for plot labels and text (default: 18).

Default: 18

**--custom\_palette** List of 7 hex color codes to customize the color scheme. Order: ['significant by

psi', 'significant by zeta', 'highlight genes', 'significant by both', 'circle markers', 'circle housekeeping genes', 'significant by neither']. Default: None (uses built-in

palette).

#### **Example:**

ember plot\_partition\_specificity Genotype pvals\_entropy\_metrics\_Genotype\_WSBJ.csv output/ -highlight\_genes Cwc22 -fontsize 25

# 4.2.4 plot\_block\_specificity

Generate a psi\_block vs. Psi scatter plot to visualize block-specific genes.

This function reads p-value data, colors genes based on their statistical significance for Psi and psi\_block scores, and highlights the top genes significant in both metrics. Allows for custom highlighting of a user-provided gene list. Font size and color palette can be customized.

#### **Positional Arguments**

**partition\_label** Label for the partition, used in the plot title.

**block\_label** Label for the block variable (e.g., a cell type or condition).

**pvals\_dir** Path to input CSV containing p-values and scores. CSV must have gene names as

its index.

**save\_dir** Path where the output plot image will be saved.

#### **Named Arguments**

**--highlight\_genes** List of gene names to highlight and annotate on the plot (default: None).

**--fontsize** Base font size for plot labels and text (default: 18).

Default: 18

**--custom palette** List of 6 hex color codes to customize the color scheme. Order: ['significant

by psi', 'significant by psi\_block', 'highlight genes', 'significant by both', 'circle markers', 'circle housekeeping genes', 'significant by neither']. Default: None

(uses built-in palette).

#### **Example:**

ember plot\_block\_specificity Genotype WSBJ pvals\_entropy\_metrics\_Genotype\_WSBJ.csv output/-highlight\_genes Cwc22 –fontsize 25

4.2. Sub-commands

### 4.2.5 plot sample counts

Generate a bar plot showing the number of unique individuals per category and condition.

This function reads an AnnData object from an .h5ad file in backed mode, calculates the number of unique individuals for each combination of a given category and condition, and visualizes these counts as a grouped bar plot. Font size can be customized.

ember plot\_sample\_counts [-h] [--fontsize FONTSIZE]

h5ad\_dir save\_dir sample\_id\_col category\_col

condition\_col

#### **Positional Arguments**

**h5ad\_dir** Path to the input AnnData (.h5ad) file.

**save\_dir** Path to directory to save the output plot image.

**sample\_id\_col** Column name in .obs that contains unique sample IDs.

**category\_col** Column name to use for the primary categories on the x-axis.

**condition\_col** Column name to use for grouping the bars (hue).

#### **Named Arguments**

**--fontsize** Base font size for plot labels and text (default: 18).

Default: 18

#### **Example:**

ember plot\_sample\_counts test\_adata\_cwc22.h5ad ~/ember\_test/output Mouse\_ID Genotype Sex –fontsize 20

# 4.2.6 plot psi blocks

Generates and saves a bar plot of mean psi block values with error bars.

This function reads two CSV files from a specified directory: one for mean psi block values and one for standard deviations. It plots the mean values for a specific gene as a bar plot with corresponding standard deviation error bars. Font size can be customized.

```
ember plot_psi_blocks [-h] [--fontsize FONTSIZE]
```

gene\_name partition\_label psi\_block\_df\_dir save\_dir

# **Positional Arguments**

**gene\_name** Name of the gene (row) to select and plot from the CSV files.

**partition\_label** Partition label used to find the correct files (e.g., 'Genotype').

psi\_block\_df\_dir Directory containing the mean and std CSV files. Files

must be named 'mean\_Psi\_block\_df\_{partition\_label}.csv' and

'std\_Psi\_block\_df\_{partition\_label}.csv'.

**save\_dir** Path to directory to save the output plot image.

# **Named Arguments**

**--fontsize** Base font size for plot labels and text (default: 18).

Default: 18

**Example:** 

ember plot\_psi\_blocks Cwc22 Genotype ~/ember\_test/output/Psi\_block\_df/ ~/em-

ber\_test/output/figs –fontsize 30

4.2. Sub-commands

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