

# Package ‘PCBS’

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**Type** Package

**Title** Principal Component BiSulfite

**Version** 0.1.1

**Description** A system for fast, accurate, and flexible whole genome bisulfite sequencing (WGBS) data analysis of two-condition comparisons. Principal Component BiSulfite, ‘PCBS’, assigns methylated loci eigenvector values from the treatment-delineating principal component in lieu of running millions of pairwise statistical tests, which dramatically increases analysis flexibility and reduces computational requirements. Methods: <[https://katlande.github.io/PCBS/articles/Differential\\_Methylation.html](https://katlande.github.io/PCBS/articles/Differential_Methylation.html)>.

**License** GPL-3

**Encoding** UTF-8

**LazyData** true

**Depends** R (>= 2.10)

**URL** <https://github.com/katlande/PCBS>

**BugReports** <https://github.com/katlande/PCBS/issues>

**Imports** ggplot2, tibble, ggrepel, dplyr, data.table

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<b>addRanks</b>	<i>Add ranks to eigenvector scores.</i>
-----------------	---

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## Description

Defines the best principle component to use for downstream analysis.

## Usage

```
addRanks(ranks)
```

## Arguments

ranks	getPCRanks output data frame.
-------	-------------------------------

## Value

The input data.frame with rank order and absolute rank order columns.

## Examples

```
ranks <- getPCRanks(eigen, IDs = c("trt", "ctl"))
ranks <- addRanks(ranks)
```

---

CheckOvercompression *Check if DMR calling seed number is ovcompressed.*

---

## Description

Identifies if seed number to use for DMR calling causes overcompression.

## Usage

```
CheckOvercompression(ranks, CpG_cutoff, values, max.dmr.size, return.plot)
```

## Arguments

ranks	Rank data frame from getPCRanks.
CpG_cutoff	NULL or numeric. If NULL, seed numbers tested will be input of the values argument. If numeric, seed numbers tested will be CpG_cutoff*values argument. Recommended to us rankDist estimate if not null
values	Numeric vector, either seed numbers to test if CpG_cutoff=NULL or multipliers if CpG_cutoff is numeric
max.dmr.size	Automatic=5000. Maximum DMR expansion size in downstream analysis. Note: pipeline is optimized for 5000bp max DMR size, it is not recommended to play with this value.
return.plot	T/F, whether to return a plot or a numeric representing the best seed number for downstream analysis

## Value

If return.plot=T, a grob plotting input seed number vs. compressed seed number is returned. Otherwise, a numeric is returned containing the largest tested input value without detectable overcompression.

## Examples

```
ranks <- getPCRanks(eigen, IDs = c("trt", "ctl"), PC = 1)
CheckOvercompression(ranks, 980)
```

checkRank

*Check rank cut-off values manually.***Description**

Plots a score vs. rank plot with a manually chosen rank cut-off for manual k selection.

**Usage**

```
checkRank(ranks, cutoff)
```

**Arguments**

ranks	getPCRanks output data frame
cutoff	integer, rank value to check

**Value**

Returns a grob plotting the input cutoff on a plot of absolute eigenvector score vs. absolute rank order.

**Examples**

```
ranks <- getPCRanks(eigen, IDs = c("trt", "ctl"), PC = 1)
test_50 <- checkRank(ranks, 50) # set cut-off to 50
test_500 <- checkRank(ranks, 500) # set cut-off to 500
```

chromDict

*Convert a rank object into a chromDict.***Description**

Internal to many functions; creates a chromDict for faster computing times. chromDict can be run separately to speed up functions run iteratively. A chromDict is a list of chromosome-specific data.tables generated from ranks.

**Usage**

```
chromDict(ranks)
```

**Arguments**

ranks	getPCRanks output data frame
-------	------------------------------

**Value**

Returns a list of `data.tables` for each chromosome, for faster analysis. Used internally by many PCBS functions.

**Examples**

```
ranks <- getPCRanks(eigen, IDs = c("trt", "ctl"), PC = 1)
chromDictObj <- chromDict(ranks)
```

---

**chromDictMeth***Create a chromDict of percent methylation difference at all sites.*

---

**Description**

chromDicts are lists of keyed `data.tables` that enable very fast computing times. Much like the primary PCBS function `chromDict()` which makes a chromDict of all locus ranks, the `chromDictMeth()` function makes a list of chromosome-specific `data.tables` containing percent methylation differences at all loci..

**Usage**

```
chromDictMeth(mat, IDs, filter_thresh)
```

**Arguments**

<code>mat</code>	<code>data.frame</code> object containing percent methylation and locus information for all sites, in the format of <code>eigen</code>
<code>IDs</code>	character vector of IDs containing the common names for compared conditions. E.g., for samples trt1 & trt2 vs. ctl1 & ctl2, <code>IDs=c("trt1", "ctl")</code>
<code>filter_thresh</code>	Integer, a coverage threshold for filtering, where CpG coverage of all samples must be larger than this value. Auto=50

**Value**

Returns a list of `data.tables` for each chromosome, for faster analysis.

**Examples**

```
chromDictMethylDiff <- chromDictMeth(eigen, c("trt", "ctl"))
```

---

DefineBestPC	<i>Identify your best principle component.</i>
--------------	--

---

**Description**

Defines the best principle component to use for downstream analysis.

**Usage**

```
DefineBestPC(mat, IDs, filter_thresh, return.plot)
```

**Arguments**

mat	Bismark2Matrix.R output file, or data frame object
IDs	A character vector of IDs containing the common names for compared conditions. E.g., for samples trt1, trt2 vs. ctl1, ctl2, IDs=c("trt", "ctl")
filter_thresh	A coverage threshold for filtering, where CpG coverage of all samples must be larger than this value
return.plot	T/F, whether to return a PCA plot or a numeric representing the best principle component for downstream analysis

**Value**

If `return.plot=T`, a grob plotting a PCA of percent methylation of all samples is returned. Otherwise, a numeric representing the best principal component to use for PCBS analysis is returned.

**Examples**

```
DefineBestPC(eigen, IDs = c("trt", "ctl"))
```

---

eigen	<i>Simulated WGBS data for PCBS vignettes</i>
-------	---

---

**Description**

simulated WGBS data with added DMRs and DMLs

**Usage**

```
eigen
```

**Format**

A data frame with 50000 observations on the following 13 variables.

cpgID a character vector, chrom:locus  
 trt1\_PercMeth a numeric vector, percent methylated of sample trt1  
 trt1\_nCpG a numeric vector, depth of sample trt1  
 trt2\_PercMeth a numeric vector, percent methylated of sample trt2  
 trt2\_nCpG a numeric vector, depth of sample trt2  
 trt3\_PercMeth a numeric vector, percent methylated of sample trt3  
 trt3\_nCpG a numeric vector, depth of sample trt3  
 ctl1\_PercMeth a numeric vector, percent methylated of sample ctl1  
 ctl1\_nCpG a numeric vector, depth of sample ctl1  
 ctl2\_PercMeth a numeric vector, percent methylated of sample ctl2  
 ctl2\_nCpG a numeric vector, depth of sample ctl2  
 ctl3\_PercMeth a numeric vector, percent methylated of sample ctl3  
 ctl3\_nCpG a numeric vector, depth of sample ctl3

**Source**

generated through simulation

getPCRanks

*Get CpG eigenvector scores from a principle component.*

**Description**

Returns eigenvector scores for input CpG sites.

**Usage**

```
getPCRanks(mat, IDs, PC, filter_thresh)
```

**Arguments**

mat	Bismark2Matrix.R output file, or data frame object
IDs	A character vector of IDs containing the common names for compared conditions. E.g., for samples trt1 & trt2 vs. ctl1 & ctl2, IDs=c("trt1", "ctl")
PC	Integer, which principle component to use. Use to DefineBestPC if unsure.
filter_thresh	Integer, a coverage threshold for filtering, where CpG coverage of all samples must be larger than this value.

**Value**

Returns a `data.frame` of eigenvector scores for all loci.

**Examples**

```
ranks <- getPCRanks(eigen, IDs = c("trt", "ctl"), PC = 1)
```

---

**getRegionScores**

*Calculated methylation significance in a set of regions.*

---

**Description**

Returns p-values and Z-scores for CpGs in a set of regions, compared to a local null background distribution.

**Usage**

```
getRegionScores(ranks, regions, chromDictObj)
```

**Arguments**

<code>ranks</code>	getPCRanks output <code>data.frame</code> , only necessary if <code>chromDictObj=NULL</code>
<code>regions</code>	A three-column <code>dataframe</code> containing a set of regions to test. Columns = <code>chrom</code> , <code>start</code> , <code>end</code> .
<code>chromDictObj</code>	<code>chromDict()</code> output object, recommended input instead of <code>ranks</code> .

**Value**

Returns a `data.frame` with significance scores for all input regions.

**Examples**

```
ranks <- getPCRanks(eigen, IDs = c("trt", "ctl"), PC = 1)
DMLs <- addRanks(ranks)

# data.frame of regions to test:
regions <- data.frame(chr=c("chr3", "chr3", "chr1"),
                       s=c(4920450, 3961576, 300000),
                       e=c(4923267, 3963805, 302900),
                       ID=c("Hypo-DMR", "partial Hyper-DMR", "random"))

getRegionScores(DMLs, regions)
```

---

get\_all\_DMRs

*Nested DMR calling function within Get\_Novel\_DMRs()*

---

**Description**

Expands DMRs from collapsed seeds iteratively.

**Usage**

```
get_all_DMRs(chromDictObj, seeds, res=40, max.dmr.size=3000, min.dmr.cpgs=10,  
min.absZscore, null)
```

**Arguments**

chromDictObj	chromDict() output.
seeds	compressed seeds
res	Get_Novel_DMRs arg DMR_resolution
max.dmr.size	Get_Novel_DMRs arg QueryLimit
min.dmr.cpgs	Get_Novel_DMRs arg minCpGs
min.absZscore	Get_Novel_DMRs arg minZ
null	null distribution

**Value**

Returns a list with two indicies, representing intermediate DMR calls within the Get\_Novel\_DMRs() function and a list of background regions..

---

Get\_Novel\_DMRs

*Call DMRs from WGBS data.*

---

**Description**

DMR Calling.

**Usage**

```
Get_Novel_DMRs(ranks, nSeeds, chromDictObj, DMR_resolution,  
QueryLimit, minCpGs, minZ, perms)
```

**Arguments**

ranks	Rank data frame from getPCRanks.
nSeeds	Integer, number of input seeds for DMR expansion.
chromDictObj	chromDict() output. If null, chromDict() is run internally.
DMR_resolution	Automatic=NULL. Integer, number of bases to increase the DMR by with each expansion. If NULL, QueryLimit/25.
QueryLimit	Automatic=5000. Maximum DMR expansion size (bp)
minCpGs	Automatic=15. Minimum CpGs in a DMR region, regions with fewer CpGs will be discarded.
minZ	Automatic=1. Absolute Z score threshold for DMR calling; internal value. Not recommended to play with this setting.
perms	Automatic=1000. Number of permutations to use when defining the null distribution. Increasing this value largely influences computational time with minimal return

**Value**

Returns a data.frame of all novel DMRs.

**Examples**

```
ranks <- getPCRanks(eigen, IDs = c("trt", "ctl"), PC = 1)
DMRs <- Get_Novel_DMRs(ranks, 2940, minCpGs=10)
```

**lin** *Find y value of linear regression given x.*

**Description**

Find y value of linear regression given x.

**Usage**

```
lin(x, l)
```

**Arguments**

x	x coordinate
l	linear model of lm()

**Value**

Returns a column numeric representing the y coordinate at the input x of the linear model l.

---

lmIntx	<i>PC-Intersect nested function.</i>
--------	--------------------------------------

---

### Description

Finds the intersection point of two linear regression models, lm().

### Usage

```
lmIntx(fit1, fit2, rnd=2)
```

### Arguments

fit1	lm() model 1
fit2	lm() model 2
rnd	number of significant figures

### Value

Returns a 2 column data.frame of one row, containing the x and y coordinates of the intersection point of the input models.

---

MethyDiff_Set	<i>Add a mean methylation column to a data.frame of regions.</i>
---------------	--

---

### Description

Using a chromDictMeth() output object, quickly calculate the mean methylation difference across set of regions.

### Usage

```
MethyDiff_Set(chromDictMeth, regions)
```

### Arguments

chromDictMeth	chromDictMeth() output object
regions	data.frame of regions, where column 1 = chromosome, column 2 = region start, and column 3 = region end

### Value

Returns the regions object with a mean percent methylation column.

## Examples

```
chromDictMethylDiff <- chromDictMeth(eigen, c("trt", "ctl"))
regions <- data.frame(chr=c("chr3", "chr3", "chr1"),
                       s=c(4920450, 3961576, 300000),
                       e=c(4923267, 3963805, 302900),
                       ID=c("Hypo-DMR", "partial Hyper-DMR", "random"))

MethylDiff_Set(chromDictMethylDiff, regions)
```

### MethylDiff

*Get the mean methylation difference across a specified region.*

## Description

Using a chromDictMeth() output object, quickly calculate the mean methylation difference across a user-specified region.

## Usage

```
MethylDiff(chromDictMeth, chrom, start, end)
```

## Arguments

chromDictMeth	chromDictMeth() output object
chrom	character, chromosome
start	integer, region start
end	integer, region end

## Value

Returns a list of data.tables for each chromosome, for faster analysis.

## Examples

```
chromDictMethylDiff <- chromDictMeth(eigen, c("trt", "ctl"))
MethylDiff(chromDictMethylDiff, "chr3", 4920450, 4923267)
```

---

`methylDiff_metagene`     *Make a metagene from mean percent methylation differences.*

---

## Description

Uses mean binned percent methylation differences across a set of regions to draw a metagene.

## Usage

```
methylDiff_metagene(chromDictMethObj, regions, bin, title,
xaxis, yaxis, return.data, linecol, value)
```

## Arguments

<code>chromDictMethObj</code>	chromDictMeth() output object
<code>regions</code>	A three-column data.frame containing a set of regions to test. Columns = chrom, start, end.
<code>bin</code>	integer, number of bins to use in metagenes. Default=100.
<code>title</code>	Output plot title
<code>xaxis</code>	Output plot x-axis title
<code>yaxis</code>	Output plot y-axis title
<code>return.data</code>	T/F, whether to return a plot, or data that can be run with plot_metagene() or multiple_metagenes().
<code>linecol</code>	Colour for line, auto="red"
<code>value</code>	Name of the plotted metric in chromDictMethObj. Only needs to be set explicitly for abnormal use cases where chromDictMethObj contains a non-rank value output by chromDict().

## Value

If `return.data=F`, returns a grob containing a metagene plot. Otherwise, returns a list of two data.frames containing metagene and metagene standard error plotting information.

## Examples

```
chromDictMethylDiff <- chromDictMeth(eigen, c("trt", "ctl"))
regions <- data.frame(chr=c("chr3", "chr3", "chr1"),
                       s=c(4920450, 3961576, 300000),
                       e=c(4923267, 3963805, 302900),
                       ID=c("Hypo-DMR", "partial Hyper-DMR", "random"))

methylDiff_metagene(chromDictMethylDiff, regions)
```

**multiple\_metagenes**      *Plot multiple metagene data objects on a single plot.*

## Description

Plots multiple metagene object using the raw data generated by score\_metagene().

## Usage

```
multiple_metagenes(data_list, set_names, title, xaxis, yaxis, legend.title, col, se_alpha)
```

## Arguments

data_list	List of score_metagene() raw data output
set_names	Character vector of names for score_metagene() object
title	Output plot title
xaxis	Output plot x-axis title
yaxis	Output plot y-axis title
legend.title	T/F, whether to show legend title
col	Vector of colours to use for lines
se_alpha	0-1, alpha value for standard error shading

## Value

Returns a grob containing a plot of the input metagene data.

## Examples

```
ranks <- getPCRanks(eigen, IDs = c("trt", "ctl"), PC = 1)
DMRs <- Get_Novel_DMRs(ranks, 2940, minCpGs=10)

# Select all significantly hypomethylated DMRs:
hypo_DMRs <- DMRs[DMRs$FDR <= 0.05 & DMRs$DMR_Zscore < 0,]
# Select all significantly hypermethylated DMRs:
hyper_DMRs <- DMRs[DMRs$FDR <= 0.05 & DMRs$DMR_Zscore > 0,]

# select chrom, start, and end of all hyper DMRs
regions_hypo <- hypo_DMRs[c(1:3)]
regions_hyper <- hyper_DMRs[c(1:3)]

# return.data = T returns raw data instead of a plot:
hyper_metagene <- score_metagene(ranks, regions_hyper, return.data = TRUE)
hypo_metagene <- score_metagene(ranks, regions_hypo, return.data = TRUE)

# The multiple_metagenes function plots multiple metagenes
# using a list of raw data objects from score_metagene().
multiple_metagenes(data_list = list(hyper_metagene, hypo_metagene),
```

---

```
set_names = c("Hyper DMRs", "Hypo DMRs"),
title="Metagenes of DMR Regions", legend.title = FALSE)
```

---

**Ol\_Reliable***PCBS ggplot theme.***Description**

Custom theme for ggplot used by all PCBS output figures.

**Usage**

```
Ol_Reliable()
```

**Value**

Theme for ggplot objects used by PCBS.

**Examples**

```
df <- data.frame(A=c(1,2,3), B=c(1,2,3))

ggplot2::ggplot(df, ggplot2::aes(x=A, y=B))+
  ggplot2::geom_point()+
  Ol_Reliable()
```

---

**oneSeed**

*Nested DMR calling function within within get\_all\_DMRs(), Get\_Novel\_DMRs()*

---

**Description**

Expands one DMR from a single point.

**Usage**

```
oneSeed(chroms, seed, resolution, max.size, mincpgs, null_list, Zlim=1)
```

**Arguments**

chroms	chromDict() output.
seed	seed value input
resolution	Get_Novel_DMRs arg DMR_resolution
max.size	Get_Novel_DMRs arg QueryLimit
mincpgs	Get_Novel_DMRs arg minCpGs
null_list	get_all_DMRs arg null
Zlim	Get_Novel_DMRs arg minZ

**Value**

returns a list of two indices, containing a `data.frame` with the output from a single compressed seed expansion and a `data.frame` of locus information around the seed expansion, intended for use within the `Get_Novel_DMRs()` function.

**plot\_metagene***Generate a metagene plot from raw metagene data.***Description**

Plots a metagene object using the raw data generated by `score_metagene()`.

**Usage**

```
plot_metagene(data, title, xaxis, yaxis, linecol)
```

**Arguments**

<code>data</code>	list, <code>score_metagene()</code> raw data output
<code>title</code>	Output plot title
<code>xaxis</code>	Output plot x-axis title
<code>yaxis</code>	Output plot y-axis title
<code>linecol</code>	Colour for line, auto="red"

**Value**

Returns a grob containing a plot of the input metagene data.

**Examples**

```
ranks <- getPCRanks(eigen, IDs = c("trt", "ctl"), PC = 1)
DMRs <- Get_Novel_DMRs(ranks, 2940, minCpGs=10)

# Select all significantly hypomethylated DMRs:
hypo_DMRs <- DMRs[DMRs$FDR <= 0.05 & DMRs$DMR_Zscore < 0,]

# select chrom, start, and end of all hyper DMRs
regions_hypo <- hypo_DMRs[c(1:3)]

# return.data = T returns raw data instead of a plot:
hypo_metagene <- score_metagene(ranks, regions_hypo, return.data = TRUE)

plot_metagene(hypo_metagene)
```

---

rankDist	<i>Identify the best rank cut-off for significant CpGs.</i>
----------	---

---

**Description**

Automated rank cut-off estimator for input CpGs.

**Usage**

```
rankDist(ranks, draw_intersects, noise_perc, mode, return.plot)
```

**Arguments**

ranks	getPCRanks output data frame.
draw_intersects	T/F whether to draw intersect lines if return.plot=T
noise_perc	Automatic=0.5, numeric between 0 and 1. Fraction of ranks to use to model the background noise. Not recommended to play with this value. Increasing/decreasing returns a looser/stricter threshold, respectively.
mode	"intersect" or "strict", determine cut-off with "intersect" or "strict" method. "Strict" is recommended for sets with lower variability
return.plot	T/F, whether to return a plot or a numeric

**Value**

If return.plot=T, a grob plotting the estimated cutoff on a plot of absolute eigenvector score vs. absolute rank order is returned. Otherwise, a numeric of the estimated cut-off is returned.

**Examples**

```
ranks <- getPCRanks(eigen, IDs = c("trt", "ctl"), PC = 1)
rankDist(ranks, mode="intersect")
```

---

score_metagene	<i>Make a metagene from PC Scores.</i>
----------------	--

---

**Description**

Uses mean binned PC scores across a set of regions to draw a metagene.

**Usage**

```
score_metagene(ranks, regions, bin, title, xaxis, yaxis,
chromDictObj, return.data, linecol)
```

**Arguments**

ranks	getPCRanks output data.frame
regions	A three-column data.frame containing a set of regions to test. Columns = chrom, start, end.
bin	integer, number of bins to use in metagenes. Default=100.
title	Output plot title
xaxis	Output plot x-axis title
yaxis	Output plot y-axis title
chromDictObj	Optional chromDictObject made from chromDict(), runs internally if set to NULL (default). Scripts that run this function multiple times will be sped up by setting this option.
return.data	T/F, whether to return a plot, or data that can be run with plot_metagene() or multiple_metagenes().
linecol	Colour for line, auto="red"

**Value**

If return.data=F, returns a grob containing a metagene plot. Otherwise, returns a list of two data.frames containing metagene and metagene standard error plotting information.

**Examples**

```
ranks <- getPCRanks(eigen, IDs = c("trt", "ctl"), PC = 1)
DMRs <- Get_Novel_DMRs(ranks, 2940, minCpGs=10)

# select chrom, start, and end of all hyper DMRs:
hyper_DMRs <- DMRs[DMRs$FDR <= 0.05 & DMRs$DMR_Zscore > 0,]
regions_hyper <- hyper_DMRs[c(1:3)]
score_metagene(ranks, regions_hyper)
```

se                   *Standard error of a vector.*

**Description**

Takes the standard error of a vector.

**Usage**

```
se(x)
```

**Arguments**

x	numeric vector.
---	-----------------

**Value**

Returns a numeric, representing the standard error of the input vector.

**Examples**

```
x <- sample(1:100, 20)
se(x)
```

---

**tilt***Component of PCBS ggplot theme.*

---

**Description**

Wrapper to title x-axis text in ggplot objects.

**Usage**

```
tilt()
```

**Value**

Theme for ggplot objects that cleanly rotates x-axis text.

---

**trimDMR***Nested DMR calling function within Get\_Novel\_DMRs()*

---

**Description**

Trims the edges off of DMR expansions.

**Usage**

```
trimDMR(df, region, min.dmr.cpgs, max.dmr.size, null_summary, null_values)
```

**Arguments**

df	DMR expansion output dataframe.
region	get_all_DMRs() region output
min.dmr.cpgs	Get_Novel_DMRs arg minCpGs
max.dmr.size	Get_Novel_DMRs arg QueryLimit
null_summary	null distribution conrtainer
null_values	null distribution conrtainer

**Value**

Returns a data.frame of all trimmed DMRs for use within the Get\_Novel\_DMRs() function.

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